JOURNAL OF THE CHEMICAL SOCIETY, DALTON TRANSACTIONS

Instructions for Authors (1988)

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1.0 General Policy

The Journal of the Chemical Society is a medium for reporting selected original and significant contributions to new chemical knowledge. Articles which do not present original work (e.g. reviews) will not normally be considered for publication in the Journal.

All contributions are judged on the criteria of (i) originality and quality of scientific content and (ii) appropriateness of the length to content of new science. Thus, papers reporting results which would be routinely predicted or result from application of standard procedures or techniques are unlikely to prove acceptable in the absence of other attributes which themselves make publication desirable.

Although short articles are acceptable, the Society strongly discourages fragmentation of a substantial body of work into a number of short publications. Unnecessary fragmentation will be a valid reason for rejection of manuscripts.

The *Journal* is published in six sections, of which five are termed *Transactions*; these are distinguished by their subject matter, as follows:

Dalton Transactions (Inorganic Chemistry). All aspects of the chemistry of inorganic and organometallic compounds, including bioinorganic chemistry and solid-state inorganic chemistry; the application of physicochemical techniques to the study of their structures, properties, and reactions, including kinetics and mechanisms; new or improved experimental techniques and syntheses.

Faraday Transactions I (Physical Chemistry in Condensed Phases). Colloid and interface science, surface science, physisorption and chromatographic science, chemisorption and heterogeneous catalysis, zeolites and ion-exchange phenomena. Electrode processes, liquids and solutions: experimental and theoretical. Solid-state chemistry: microstructures and dynamics. Reactions in condensed phases, physical chemistry of macromolecules and polymers. Biophysical chemistry and radiation chemistry.

Faraday Transactions II (Molecular and Chemical Physics). Gas-phase reaction kinetics and dynamics: experimental and theoretical; molecular beam kinetics and spectroscopy. Photochemistry and photophysics, energy transfer and relaxation processes. Laser-induced chemistry. Spectroscopies of molecules, molecular ensembles, and gas-phase complexes. Quantum theory of molecular structure. Statistical mechanics of gaseous molecules and complexes; statistical mechanics and quantum theory of the condensed phase. Computational chemistry; molecular dynamics.

Perkin Transactions I (Organic Chemistry). All aspects of organic and bio-organic chemistry. These include synthetic organic chemistry of all types, organometallic chemistry, chemistry and biosynthesis of natural products, the relationship between molecular structure and biological activity, the chemistry of polymers and biological macromolecules, and medicinal and agricultural chemistry where there is originality in the science.

Perkin Transactions II (Physical Organic Chemistry). Physicochemical aspects of organic, organometallic and bioorganic chemistry including kinetic, mechanistic, structural, spectroscopic, and theoretical studies. Such topics include structure—activity relationships and physical aspects of biological processes and of the study of polymers and biological macromolecules.

Authors are requested to indicate, at the time they submit a typescript, the journal for which it is intended. Should this seem unsuitable, the Editor will inform the author.

The sixth section of the *Journal of the Chemical Society* is Chemical Communications, which is intended as a forum for preliminary accounts of original and significant work, in any area of chemistry that is likely to prove of wide general appeal or exceptional specialist interest. Such preliminary reports should be followed up eventually by full papers in other journals (e.g. the five *Transactions*) providing detailed accounts of the work.

In addition to full papers, *Dalton Transactions* also publishes Notes and Letters (see section 1.2).

1.1 Conditions Governing Acceptance

Contributions which have appeared or have been accepted for publication with essentially the same content in another journal or which incorporate freely available printed work will not be published in the *Journal* except by permission of the Council. This restriction does not apply to results previously published in materially abbreviated form, as a paper presented at a symposium, as a preliminary communication (e.g. to Chemical Communications), as a letter to the Editor of some other periodical, or as a patent. However, the acceptance of a contribution for Chemical Communications does not guarantee that the corresponding full paper will be accepted for the *Journal*; although publication of a full account is strongly encouraged, its acceptability will depend on whether or not it contains significant new details, new interpretations, or new results.

Contributions are accepted by the Society on the understanding that the authors (a) have obtained any necessary authority for publication, and (b) will, if requested, execute a formal licence granting the Society exclusive licence under any copyright therein.

Authors are solely responsible for the factual accuracy of their contributions.

Since the Society reserves the right to retain all typescripts sent to it, authors are advised to keep copies. When contributions have been submitted for publication the authors are not at liberty, save by permission of the Society, to withdraw or delay them or to publish them elsewhere until after publication by the Society.*

* Attention is drawn to the following extract from the Society's By-Laws:

1.2 Notes and Letters

Notes are intended for the description of essentially complete pieces of work which are not of the length to justify a full paper. They are not preliminary communications, nor in any way an alternative to *Chemical Communications* for which there are additional criteria of novelty and urgency.

The normal length of a Note should not exceed 2 printed pages (corresponding to about 8 pages of typescript, including Figures, Tables, *etc.*). It should comprise a short abstract and Discussion, but adequate experimental details are required.

The quality of material contained in a Note should be the same as that in a full paper. Investigations arising out of some larger project but not prosecuted to the same degree are particularly appropriate.

Letters are a medium for the expression of scientific opinions and views normally concerning material published in the Society's journals. The Letter section is for scientific discussion, and is not intended to compete with media for the publication of more general matters such as *Chemistry in Britain*.

Only rarely should a Letter exceed one printed column in length (about 1—2 pages of typescript). Where a Letter is polemical in nature, and if it is accepted, a Reply will be solicited from other parties implicated for publication alongside the original Letter.

1.3 Submission of Articles

Typescripts should be addressed to: The Manager, Journals, The Royal Society of Chemistry, Burlington House, Piccadilly, London WIV 0BN.

Three copies of the typescript (a top copy and two good quality carbon or Xerox copies) are required. Copies of any related, relevant, unpublished material should also be provided.

Rapid publication is aided by careful preparation of text and illustrations and strict adherence to the format and conventions of individual *Transactions* as laid down in these Instructions for Authors.

Particular attention is drawn to the use of (i) SI units and associated conventions, (ii) IUPAC nomenclature for compounds, and (iii) standard methods of literature citation.

Owing to the non-availability of many referees and editorial staff in July-August each year, authors are advised to limit their submissions during this period in order to minimize delays.

2.0 Administration and Publication Procedure

Receipt of a contribution for consideration will be acknowledged immediately by the Editorial Office. The acknowledgement will indicate the paper reference number assigned to the contribution. Authors are particularly asked to quote this number on all subsequent correspondence.

The paper is sent simultaneously to at least two referees, whose names are not disclosed to the authors. On the basis of the referees' reports, the Editor decides whether the paper is suitable for publication, either unchanged or after appropriate revision. This decision and relevant comments of the referees are communicated to the author. Differences of opinion are mediated by the Editor, possibly after consultation with further referees, or, in the last resort, by the Editorial Board.

When rejection of a paper is recommended, the Editor informs the author, and returns the top copy of the manuscript. Authors have the right to appeal to the Editorial Board if they regard a decision to reject as unfair.

Acceptance of a paper is confirmed when the edited manuscript is sent to the printer. The author receives two copies of proofs, together with the edited manuscript and reprint order

^{91. (}iii) Every member who submits a paper or other communication with a view to its publication by the Society shall by so doing undertake: (a) that his communication has not been published and that he will not permit its publication before it is accepted or declined by the Society, and

⁽b) that if it is accepted for publication the Society shall thereupon become entitled to an exclusive licence under any copyright therein (which shall include the right to sublicence) and that he will, if then called upon to do so, execute a formal licence to the Society of the said copyright, including the sole right to publish in any form in any language and in any part of the world, the whole or any part of his communication. The Council shall not refuse any reasonable request from an author to reproduce his own work elsewhere in whole or in part.

⁽iv) The Society shall have the right to retain manuscripts and illustrative drawings sent to the Society for consideration for publication.

⁽v) The attention of every member who submits any paper or other communication with a view to its publication shall be drawn to paragraph (iii) of this By-Law above, and any person other than a member shall be required to sign an undertaking in the terms set out therein.

form. The Society supplies 50 reprints free of charge, and further copies can be purchased.

One corrected proof and the manuscript, the reprint order form, and payment (if any) should be sent to the Editor. Checking of proofs is the author's responsibility (although the Editor will carry out a further check before publication), and particular attention should be paid to numerical data both in tables and in the text, references, structural formulae, and diagrams.

An author may be required to pay the cost of any extensive changes made by him at proof stage (other than the correction of printer's errors). So far as possible, essential changes should be made without altering the length of the text, or at the end of a paragraph. The standard signs for proof correction set out in British Standard BS 5261: Part 2 (1976) may be used: these are conveniently summarised in the pamphlet 'Authors' alterations cost money and cause delay...' which can be purchased from the British Printing Industries Federation (11 Bedford Row, London WC1R 4DX). However, the author may prefer simply to put a line through the incorrect characters and write the correct version in the margin. Corrections should be made in ink, clearly and without ambiguity, and any queries from the printer or editorial staff on the manuscript or proof should be answered fully.

3.0 Presentation of Papers

Every latitude, consistent with brevity, in the form and style of papers is permitted, and no rigid pattern for either is prescribed. Nevertheless, adherence to the methods outlined in this section is recommended unless there is good reason for deviation. For the format of Notes and Letters, see section 1.2.

3.1 Organization of Material

3.1.1 Title.—The choice of a title for a paper is of the greatest importance, since it is from the title that the important key-words used in information retrieval are taken. Not only should the title clearly and accurately indicate the content of that paper but also it should be as specific as the content and emphasis of the work permit. Brevity in a title, though desirable, should be balanced against its accuracy and usefulness.

The use of abbreviations and symbols in a title is discouraged; terms should be written out in full unless they are extremely cumbersome. However the use of linear formulae to represent complex structures is permitted; in such cases the editor may insert a systematic name in a footnote.

The preceding part of a series must be referred to (as reference 1) in the title in the form:

Chemistry of the Metal Carbonyls, Part 81. Homonuclear Di- and Tri-metal Carbonyl Complexes Derived from Dicarbonyl(pentamethylcyclopentadienyl)rhodium; X-Ray Crystal Structure of [MnRh(μ -CO)₂(η -C₅H₅)(η -C₅Me₅)]. The corresponding reference should be in the form:

1 Part 80, L. J. Farrugia, J. A. K. Howard, P. Mitrprachachon, F. G. A. Stone, and P. Woodward, *J. Chem.* Soc., Dalton Trans., 1981, 1274.

When the preceding part has been submitted to the Society but is not yet published, the paper reference number should be given.

3.1.2 Summary.—Every paper for the Journal (including Notes) must be accompanied by a summary (50—250 words) setting out briefly and clearly the main objects and results of the work; it should give a reader a clear idea of what has been achieved. The summary should be essentially independent of the main text; however, names, partial names, or linear formulae

of compounds may be accompanied by the numbers referring to the corresponding displayed formulae in the body of the text.

Examples:

Reaction of $[Rh_2(C_5Me_5)_2Cl_4]$ with $PhC\equiv CH$ in acetonitrile in the presence of Na_2CO_3 gives two complexes (3) and (4) as well as some isomers of triphenylbenzene and acetophenone. Complexes (3) and (4) were characterised by ^{13}C n.m.r. spectroscopy and by single-crystal X-ray structure determinations. Complex (3) has the rhodium π -bonded η^5 to a C_5Me_5 ring and η^4 to the C_4 ring of a tetraphenylbenzocyclobutene. The benzocyclobutene is very close to planar and only a little distorted upon coordination; the structure of the ligand is best understood in terms of a '1,2-divinylcyclobutadiene' type of bonding. Complex (4) has the rhodium π -bonded η^5 to a C_5Me_5 ring and η^4 to a cyclobutadiene; this cyclobutadiene carries two phenyl substituents (1,3-) and an (uncoordinated) 6-(1,3,6-triphenylfulvenyl) substituent. Possible routes by which (3) and (4) could be formed are discussed.

Water-soluble manganese(III) porphyrins are oxidised in alkaline aqueous solution to the corresponding manganese(IV) porphyrins which, from magnetic moment measurements, appear to exist in solution as μ-oxo-dimers. Midpoint potentials and rate constants for oxidation of the manganese(III) porpyhrins have been measured for a series of oxidants and throughout the range 9 < pH < 14, but the overall electronic charge on the metalloporphyrin had little effect upon either parameter. The midpoint potentials for the MnIIIIV couple are strongly dependent upon pH and increase with decreasing pH. Although manganese(IV) porphyrins are mild oxidants at pH 14, they should be capable of oxidising water to molecular oxygen in neutral solution. With hypochlorite as oxidant, a second oxidation step is possible and the final product is believed to be a manganese(v) oxoporphyrin.

No summary is required for Letters to Dalton Transactions.

- 3.1.3 *Introduction.*—This should give clearly and briefly, with relevant references, both the nature of the problem under investigation and its background.
- 3.1.4 Results and Discussion.—It is usual for the results to be presented first, followed by a discussion of their significance. Only strictly relevant results should be presented and figures, tables, and equations should be used for purposes of clarity and brevity. The use of flow diagrams and reaction schemes is encouraged. Data must not be reproduced in more than one form, e.g. in both figures and tables, without good reason.
- 3.1.5 Experimental Section.—Descriptions of experiments should be given in detail sufficient to enable experienced experimental workers to repeat them; the degree of purity of materials should be given, as should the relative quantities used. Descriptions of established procedures are unnecessary. Standard techniques and methods used throughout the work should be stated at the beginning of the section. Apparatus should be described only if it is non-standard; commercially available instruments are referred to by their stock numbers (e.g. Perkin-Elmer 457 or Varian HA-100 spectrometers). The accuracy of primary measurements should be stated. Unexpected hazards encountered during the experimental work should be noted. In general there is no need to report unsuccessful experiments.

- 3.1.6 Acknowledgements.—Contributors other than coauthors may be acknowledged in a separate paragraph at the end of the paper; acknowledgements should be as brief as possible. Titles, Mr., Mrs., Miss, Dr., Professor, etc., should be given but not degrees.
- 3.1.7 Bibliographic References.—These should be given on a separate sheet at the end of the manuscript; for details see section 3.7.

3.2 Brevity

For reasons of economy, brevity in the presentation of papers is essential. Authors should note that the following practices are likely grounds for rejection of a manuscript, or acceptance only after substantial revision.

- (a) Unnecessary division of work into separate parts of a series of papers.
- (b) Submission of fragmentary work which can be included in a larger communication.
 - (c) Undue elaboration of hypotheses.
 - (d) Over-detailed and verbose exposition of ideas.
- (e) Excessive use of diagrams; for example, a straight-line plot can be adequately expressed as an equation together with, if necessary, a table of deviations.
 - (f) Duplication of data in text, tables, and figures, etc.
- (g) Descriptions of slight variations of essentially the same technique.

3.3 Linguistic and Typographical Conventions

- 3.3.1 Grammar and Spelling.—Standard English spelling is used (Oxford English Dictionary). Latitude with respect to alternative spellings is allowed, but consistency should be maintained within a paper. Difficult grammatical points may be elucidated by reference to Fowler's Modern English Usage.
- 3.3.2 Abbreviations.—The following common initial letter abbreviations may be used without definition: b.p., c.d., e.s.r., g.l.c., i.r., m.p., n.m.r., o.r.d., t.l.c., u.v., v/v, w/w. Other such abbreviations should be defined at first mention, as should abbreviations for ligands, reagents, etc.
- 3.3.3 Punctuation.—Punctuation follows standard English practice; the following conventions are observed:
- (a) A comma is placed before 'and' or 'or' in a series such as 'oxygen, sulphur, and selenium' or ' λ_{max} . 237, 295, and 343 nm.'
- (b) The 'nesting' order for parentheses, square brackets, and braces is $\{ [()] \}$.
- (c) Punctuation follows, rather than precedes parentheses, e.g. 'm.p. 234 °C (decomp.),' and not 'm.p. 234 °C, (decomp.)'.
- (d) A colon is used to separate a ratio as in 1:20—not a solidus 1/20.
- (e) Parenthetical expressions of the same physical quantity in different units are separated by comma, e.g. (3.9 g, 0.1 mol), (30 ml, 1 mol).
- 3.3.4 Use of Italics.—(a) Foreign words and phrases and Latin abbreviations are given in italics: e.g., in toto, in vivo, ca., cf., i.e., etc.
- (b) In the names of chemical compounds or radicals italics are used for prefixes (other than numerals or symbols) when they define the positions of named substituents, or when they define stereoisomers: other prefixes are printed in roman. (Note: Initial capital letters are not to be used with italic prefixes or single-

letter prefixes: full stops are not to be associated with letter prefixes.)

Examples:

- o-, m-, and p-nitrotoluenes, but ortho-, meta-, and paracompounds (o-, m-, and p- are used only with specific names; ortho-, meta-, and para- are used with classes), N,N-dimethylaniline, trans- and cis-bis(glycinato)platinum(II), gem- and vic-diols, benzil anti-oxime.
- (c) The names of periodicals or their abbreviations are set in italics

Note: Greek letters are not italicised.

- 3.3.5 *Headings.*—(a) Main sections (Experimental, Discussion, etc.): side-heading, bold, no final fullstop.
- (b) Main side-heading: italics, initial capital letter for each noun and adjective, final fullstop and dash.
- (c) Subsidiary side-heading: italics, first initial capital only, final fullstop but no dash.
- (d) Further subdivision: by italic (a), (b), etc. (no following fullstop), and finally (i), (ii), etc. If (a), (b), etc. are used in front of a subsidiary side-heading, then for contrast these letters are not italicized.

Letters and prefixes which are ordinarily printed in italics are transferred for contrast into roman type in italicised phrases (see example below, where *N,N*-dimethyl) becomes *N,N*-dimethyl).

Physicochemical symbols, however, remain in their prescribed form, and structural formulae, numerals, and Greek letters are not italicised.

Examples:

Experimental

Preparation of the Thiolate Complexes.—(a) Bis(dimethyldithiocarbamato)[N,N-dimethylhydrazido(2—)] bis-(benzenethiolato)molybdenum(vi) (10). (a) Benzenethiol (1 g) was added to . . .

Action of 2-Benzylaminopyridine on $[Os_3(CO)_{10}-(C_8H_{14})_2]$ at 40 °C.—A solution of the bis(cyclo-octene) complex . . .

3.4 Formulae and Figures

The purpose of all illustrative matter in a paper is to clarify the arguments and descriptions rather than to duplicate them. The Society strongly encourages the use of displayed formulae, particularly in the form of schemes where the details of a reaction sequence are often more easily understood when illustrated than when described in the text.

All formulae and figures should be clearly drawn, and in the case of figures, provided with captions; the latter should be typed on a separate sheet. Since all formulae carry key numbers by which they are identified, unless they form part of the running text or unless they are part of a scheme which itself has a caption, they are not generally further described. Blocks of formulae do not need captions.

- 3.4.1 Structural Formulae.—(a) Only those formulae which are displayed outside the text should be given key numbers. In other cases compounds should be referred to by a name or a linear formula.
- (b) Formulae should be numbered with bold arabic numerals in parentheses [(1), (2), and (3) etc.] in the order in which they are displayed and not in the order of mention in text.
 - (c) In complex reaction schemes formulae should be

numbered serially following the reaction sequence. Nonsequential numbering in a collection of formulae can render it hard to locate an individual number.

- (d) Structural or displayed formulae must be carefully and accurately drawn or typed on a separate sheet, rather than inserted into the text, although a marginal indication of where they are to go in the text is desirable.
- (e) Formulae inserted into the body of the text (as distinct from those displayed separately) should be written on one line if possible, e.g.

$$[Cu_2(\mu\text{-}Cl)_2(OR)_4]$$
 and $[NiR(PPh_3)_2(\overrightarrow{CH=CHCH_2CH_2O})]$

rather than

- (f) Guidelines for writing linear formulae of complexes are given in IUPAC Nomenclature of Inorganic Chemistry (see Section 3.8 of these Instructions). Authors having particular reasons for wishing to deviate from these guidelines should inform the editor when the paper is submitted.
- (g) In formulae of organic ligands the abbreviations Me, Et, Prⁿ, Prⁱ, Buⁿ, Buⁱ Bu^s, Bu^t, and Ph may be used. Other special symbols, if used, require an explanatory footnote. The carboxygroup is written CO₂H (not COOH); similarly CO₂R.
- (h) One variable univalent substituent is indicated by R; when more than one independently variable general substituent is present, R^1 , R^2 , and R^3 should be used not R, R^1 , R^2 , R^3 ; or R_1 , R_2 , and R_3 , which indicate $1 \times R$, $2 \times R$, etc. A variable metal may be indicated by M, variable ligands by L^1 , L^2 , etc., and a variable halogen or chalcogen by X.
- (i) Often it is desirable to use one formula to represent a number of related compounds (or classes of compounds) by the use of one or more independently variable substituents. It is preferable to give each compound thus represented a separate key number rather than subdivide individual key numbers of alphabetical suffixes [i.e. (1a), (1b), (1c) etc.]. The use of more

Examples:

than four independently variable substituents or atoms on one generalized formula is discouraged.

- (j) Once a formula has been displayed it is permissible to employ its key number in later reaction schemes or equations rather than to re-display the formula.
- (k) Displayed formulae may be included in tables provided that they can be typed on one line [see point (e) above]; otherwise they should be displayed elsewhere and referred to by number only in the table itself.
- (1) The key number for a compound may be used in the cursive text to avoid repetition of long chemical names; this device must not be used to excess. In general it is preferred if the key number is qualified by a partial name as in the following example:

'When the iridium complex (1) was stirred with an excess of iodomethane at room temperature, the adduct (7) was obtained in high yield. The rhodium complexes (4)—(6) react with iodomethane under similar conditions to give the acetyl complexes (8)—(10) respectively, formed by isomerization of the first-formed methyl complexes (11)—(13).'

- (m) Reference to compounds in the summary by key number alone is discouraged, since a summary should be comprehensible without reference to the body of the paper.
- 3.4.2 Figures.—(a) Figures must bear on the back the names of the authors, the title of the paper (abbreviated if necessary), and the number of the figure.
- (b) Figures must be in black ink, on board, white smooth cartridge paper, tracing linen, plastic film (it is essential that the special plastic ink developed for this is used), or graph paper with faint blue lines (red or brown lines must not be present as they may be reproduced by the photographic process employed). Since lines must be black and sharp, photostats or similar prints are often not suitable. If paper is used, it must be strong enough to withstand repeated handling.
- (c) Lettering and numerals must be in blue pencil (not red or black pencil or ink) clearly legible but not so heavily scored as to make a permanent impression on the paper or board.
- (d) When the figures are large (more than 20×25 cm), smaller copies (which may be rough, as long as they are clear) should be supplied for submission to the referees; editing will not be undertaken, however, before the final figures are received.
- (e) Figures should be drawn about three times the required size, with lines thick enough to withstand photoreduction.
- (f) Five-cm margins should be left all round figures. Lettering for insertion at margins should be placed well clear of the ordinate or abscissa line so that it can be copied before erasure.

Lettering and touching-up are done by the Society and clarity of instructions is essential. When there is much lettering, or complicated lettering, and always when tracing linen or plastic film is used, a rough tracing should be provided with the lettering shown in ink.

- (g) Since, for printing, the size is reduced, lines should not be too thin. Given lines must be of even thickness, angles neat, and curves smooth. Particular care should be taken with pairs of crystal structure diagrams for stereoscopic viewing: for good reproduction an adequate line thickness is essential.
- (h) Graphs should have only the requisite minimum of scale divisions (not less than three points) marked by numerals, and the scale lines should not normally be continued into the body of the figure.
- (i) Graphs in any one paper should be drawn to the same scale when convenient, and scale markings should be identical when possible so that the graphs may be placed adjacent on the page. Two curves drawn to different scales can be shown on one graph by having the appropriate scales on the left-hand and the right-hand side. The use of both right- and left-hand axes and top and bottom axes on figures which have quantitative significance is encouraged.
- (j) The expression used to define the numerical values of a physical quantity plotted on a graph should be dimensionless, e.g. $\ln (p/atm)$, $10^3 (T/K)^{-1}$.
- (k) Experimental points must be shown sufficiently large to be distinguishable when reduced in size. Whenever possible, they should be confined to open and closed circles, crosses, squares, and triangles. Partly black circles and similar signs frequently become indistinguishable in print.
- (1) Curves may be distinguished as full lines (—), broken (---) or dotted lines ($\cdots \cdot$), and dot-dash lines ($--\cdot ---$).
- (m) For reference in legends, it is preferable to mark curves A,B,C, etc. rather than to reproduce the type of line in print.
- (n) There must be no unnecessary waste space, e.g. around curves; ordinates and abscissae should start at zero only if the curve extends to that range. Enlargements of parts of a figure can occasionally be placed on a corner of the complete figure.
- (o) It is not advisable to insert much or complicated lettering on curves or in blank spaces; mistakes (in copying by the artist) can rarely be rectified once the block is made. It is better to label the curves A,B,C, etc. and to use explanatory legends.
- (p) Large solid objects should be represented by hatching rather than by black surfaces, otherwise the ink may smear on printing.
- (q) Photographs are reproduced by a half-tone process. The prints supplied must be very clear and of good contrast, as considerable definition may be lost in reproduction.
- (r) Captions and explanatory legends to be set by the printer should be typed on a separate page attached to the manuscript, and not given on the figure itself.
- (s) Figures are numbered consecutively Figure 1, Figure 2, etc. (in arabic numerals).

3.5 Presentation of Experimental Data

3.5.1 Tables.—If there is extensive reference to any particular data in the text, presentation of the data in tabular form is preferred. It is difficult to give general rules for the economical layout of Tables but authors will find it helpful to consult recent issues of the Journal for examples. A layout taking up the full width of the printed page, with repetition of column headings if necessary, is normally preferred to a lengthy half-page-width presentation. Columns containing very few entries are wasteful of space, and better replaced by footnotes.

When Tables (and Figures) are reproduced in the journal,

they will be positioned at the top or the bottom of a printed page, as near as possible to their first mention.

Column headings should be brief, as their width, rather than that of the entries beneath them, often determines the number of columns that can be accommodated.

Column headings should be in accord with the conventions associated with SI; thus the expression at the head of a column of numerical values of a physical quantity should be dimensionless, *i.e.* the quotient of the symbol for the physical quantity and the symbol for the unit used, *e.g.* p/atm, or the symbol for a dimensionless physical quantity, *e.g.* p/G, or some mathematical function of such a number, *e.g.* $1n(p^m/atm)$.

Example:

$$\theta/^{\circ}C$$
 T/K 10^{3} K/T p/atm 1 n(p/atm) $V_{\rm m}^{\rm G}$ /cm³ mol⁻¹ $pV_{\rm m}^{\rm G}$ /RT -51.60 216.55 4.6179 5.112 1.6316 3 177.6 0.9142

Space requirements may favour the use of a horizontal rule, e.g. $\frac{V_{\rm m}^{\rm G}}{{\rm cm}^3 {\rm mol}^{-1}}$ rather than an oblique stroke.

If possible, tables should be arranged so as not to require printing sideways on the page ('landscape') unless their depth is such that the page will be filled; otherwise division into two tables is preferred.

3.5.2 Physical Characteristics of Compounds.—Data associated with particular compounds should be listed after the name of the compound concerned, following the description of its preparation, or else presented in tabular form.

The following is suggested as the order in which the most commonly encountered data for a new compound should be cited: yield, melting point, optical rotation, refractive index, elemental analysis, u.v. absorptions, i.r. absorptions, n.m.r., spectrum, mass spectrum. Appropriate formats for the citation of each are as follows.

Yield. In parentheses after the compound name (or its equivalent). Weight and percentage are separated by a comma, e.g. 'the carbonyl complex (7.1 g, 56%)'.

Melting point. In the form 'm.p. 75 °C (from EtOH)', i.e. the crystallisation solvent in parentheses. If an identical mixed melting point is to be recorded, the form 'm.p. and mixed m.p. 75 °C' is appropriate.

Refractive index. Given in the form n_D^{22} 1.653.

Elemental analysis. In the presentation of elemental analyses, a distinction is made between 'new' and 'known' compounds (see section 3.6).

New compounds should be indicated by underlining the name (for italics) at its first mention (excluding headings) in the Experimental section only, and by giving analytical results in the form: (Found: C, 56.4; H, 4.00. $C_{12}H_{10}CrO_3$ requires C, 56.7; H, 3.95%). If analytical results for compounds which have been adequately described in the literature are to be included, they should be given in the form: (Found: C, 56.5; H, 4.00. Calc. for $C_{12}H_{10}CrO_3$: C, 56.7; H, 3.95%). Analyses are normally quoted to the nearest 0.1%, but a '5' in the second place of decimals is retained.

If a molecular weight is to be included, the appropriate form is: [Found: C, 56.5; H, 4.00%; M (mass spectrum), 254 (or simply M^+ , 254). $C_{12}H_{10}CrO_3$ requires C, 56.7; H, 3.95%; M, 254].

U.v. absorptions. These are given in the form λ_{max} (EtOH) 228 (ε 40 900 dm³ mol⁻¹ cm⁻¹), 262 (19 200), and 302 nm (11 500). Inflections and shoulders are specified as 228infl or 262sh. Alternatively the following form may be used: λ_{max} (EtOH) 228, 262, and 302 nm (ε 40 900, 19 200, and 11 500 dm³ mol⁻¹ cm⁻¹). Log ε may be quoted instead of ε.

Table 0. Analytical and physical data for the platinum-tungsten complexes

	>		37:-136		Analys	sis (%)
Complex	$M.p.^b$ $(\theta/^{\circ}C)$	Colour	Yield ' (%)	$v(CO)^d/cm^{-1}$	C	Н
(1) $[PtW(\mu-CC_6H_4Me-4)(CO)_3(PMe_3)(\eta-C_5H_5)]$	110—112	Pink	18	2 033vs, 1 939s, 1 865m,sh, 1 845m	32.3 (32.3)	3.0 (3.2)
(2) $[PtW(\mu-CC_6H_4Me-4)(CO)_3(PMe_2Ph)(\eta-C_5H_5)]$		Red	10	2 016s, 1 939m, 1 843m,br	37.6 (37.5)	3.2 (3.0)
(4) $[(PtW(\mu-CC_6H_4Me-4)(CO)_3(PPh_3)(\eta-C_5H_5)]^e$		Red	40	2 027s, 1 938s, 1 857m,br		
(6) $[Pt_2W(\mu_3\text{-}CC_6H_4Me\text{-}4)(CO)_4(PMe_3)_2(\eta\text{-}C_5H_5)]$	128—134	Orange	40	2 003s, 1 989s, 1 857m,br, 1 763m,br	26.9 (27.4)	3.4 (3.0)
(7) $[Pt_2W(\mu_3-CC_6H_4Me-4)(CO)_4(PMe_2Ph)_2(\eta-C_5H_5)]^f$	136—140	Orange	41	1 993s, 1 980s, 1 729m	34.3 (34.3)	(3.0)
(8) $[Pt_2W(\mu_3-CC_6H_4Me-4)(CO)_4(PMePh_2)_2(\eta-C_5H_5)]^f$	141—145	Dark orange	43	1 999vs, 1 839s,br, 1 746m,br ^g	40.5 (40.3)	3.2 (3.0)
(9) $[Pt_2W(\mu_3-CC_6H_4Me-4)(CO)_4(PPh_3)_2(\eta-C_5H_5)]^h$	112—115	Red	27	2 003s, 1 983s, 1 967s, 1 923s,	48.2 (47.6)	3.6 (3.2)
(10) $[Pt_2W(\mu_3\text{-CC}_6H_4\text{Me-4})(CO)_4(PEt_3)_2(\eta\text{-C}_5H_5)]$	160—169	Red	10	1 864m,br, 1 784m,br 1 990s, 1 829m, 1 732m	32.1 (31.9)	3.6 (3.2)

^a Required values are given in parentheses. ^b With decomposition. ^c Based on tungsten. ^d In methylcyclohexane, unless otherwise stated. ^e Compound not obtained analytically pure due to contamination with PPh₃. ^f Crystallised with 0.2 molecules CH₂Cl₂. ^g In dichloromethane. ^h Crystallised with ½ molecule PhMe.

I.r. absorptions. Shown as follows: v_{max} . 2 029 and 1 955 (CO), and 1 714 cm⁻¹ (NO). The type of signal (s, w, vs, br) can be indicated by appended letters (e.g. 1 760vs).

N.m.r. data. For all spectra δ values should be used, with the nucleus indicated by subscript if necessary (e.g. $\delta_{\rm H}$, $\delta_{\rm C}$). Instrument frequency, solvent, and standard should be specified. For example: $\delta_{\rm H}$ (100 MHz; solvent CDCl₃; standard Me₄Si) 5.28 [4 H, m, $J({\rm PtH})$ 72 Hz, 4 CH], 2.04 (8 H, s, 4 CH₂), and 1.80 (30 H, s, 2 C₅Me₅). A broad signal may be denoted by 'br', e.g. 2.43 (1 H, br s, NH). Order of citation in parentheses: (i) number of equivalent nuclei (by integration), (ii) multiplicity (s, d, t, q), (iii) coupling constant, e.g. $J({\rm RhP})$ 15 Hz, $J({\rm PH})$ 4 Hz, (iv) assignment; underlining for italics can be used to specify the nuclei concerned (e.g. CH₃CH₂).

Mass spectrum. Given in the form: m/z 183 (M^+ , 41%), 168 (38), 154 (9), 138 (31) etc. The molecular ion may be specified as shown if desired. Relative intensities in parentheses (% only included once). Other assignments may be included in the form m/z 152 (33, $M - \text{CH}_3\text{CONH}_2$). Metastable peaks may be listed as: m^* 160 (189 \longrightarrow 174), 147 (176 \longrightarrow 161), etc. The type of spectrum (field desorption, electron impact, etc.) should be indicated.

Literature citations. If comparison is to be made with literature values, these should be quoted in parentheses, e.g. m.p. 157 $^{\circ}$ C (from chloroform) (lit., ¹⁹ 156 $^{\circ}$ C), or v_{max} . 2 020 and 1 592 cm⁻¹ (lit., ²⁴ 2 015 and 1 600 cm⁻¹).

Example of a typical experimental section format. The following paragraph exemplifies many of the points made in the preceding paragraphs. Authors should note in particular the specification of quantities in parentheses after the names of reagents, and the use of the past tense.

Synthesis of the Rhodium-Manganese Complex (1).— Tricarbonyl(η -cyclopentadienyl)manganese (0.365 g, 1.79 mmol) in tetrahydrofuran (50 cm³) was irradiated (u.v.) for 2 h at 0 °C under argon. The solution was then treated with $[Rh(CO)_2(\eta-C_5Me_5)]$ (0.259 g, 0.88 mmol) and the mixture stirred at room temperature for 18 h. Removal of solvent and chromatography afforded unchanged [Rh(CO)₂(η -C₅Me₅] and [Mn(CO)₂(η -C₅H₅)] followed by yellow-brown crystals of [MnRh(μ -CO)₂(CO)₂(η -C₅H₅)(η -C₅Me₅)] (1) (0.327 g, 79%), m.p. 158—160 °C (Found: C, 48.7; H, 4.5%; M^+ , 470. C₁₉H₂₀MnO₄Rh requires C, 48.5; H, 4.3%; M, 470); v_{max} (CO) 1 997sh, 1 991s, 1 921vs, 1 809w, and 1 788vs cm⁻¹ (Nujol); 1 983s, 1 935vs, 1 818w,sh, and 1 807s cm⁻¹ (hexane); δ_{H} (CDCl₃) 1.84 (15 H, s, C₅Me₅) and 4.62 (5 H, s, C₅H₅); δ_{C} (CD₂Cl₂-CH₂Cl₂) 252.2 [2 C, d, J(RhC) 22 Hz, μ -CO], 227.6 (1 C, s, MnCO), 188.3 [1 C, d, J(RhC) 84 Hz, RhCO], 105.0 (C_5 Me₅), 86.5 (C₅H₅), and 8.9 (C₅Me₅); m/z 470 (M^+), 442 (M – CO), 414 (M – 2CO), 386 (M – 2CO), and 358 (M – 4CO).

An example of an alternative, tabular presentation of data is also shown (Table 0).

3.6 Authentication of New Compounds

It is the responsibility of authors to provide fully convincing evidence for the homogeneity and identity of all compounds they claim as new. Evidence of both purity and identity is required to establish that the properties and constants reported are those of the compound with the new structure claimed.

A compound is considered as new (a) if it has not been prepared before, (b) if it has been prepared before but not adequately purified, (c) if it has been purified but not adequately characterised, (d) if, earlier, it has been assigned an erroneous consitution, or (e) if it is a natural product isolated or synthesised for the first time. In preliminary communications compounds are often recorded with limited characterising data; in spite of (c) above later preparations of such compounds are not considered as new if the properties previously reported are confirmed; the same applies to patents.

Referees will assess, as a whole, the evidence in support of the homogeneity and structure of all new compounds. No hard and

fast rules can be laid down to cover all types of compounds, but evidence for the unequivocal identification of new compounds should wherever possible include good elemental analytical data; an accurate mass measurement of a molecular ion does not provide evidence of purity of a compound and must be accompanied by independent evidence of homogeneity. Low-resolution mass spectrometry must be treated with even more reserve in the absence of firm evidence to distinguish between alternative molecular formulae. Where elemental analytical data are not available, appropriate evidence which is convincing to an expert in the field may be acceptable, but authors should include, for the referees, a brief explanation of the special nature of their problem.

Spectroscopic information necessary to the assignment of structure should normally be given. Just how complete this information should be must depend upon the circumstances; the structure of a compound obtained from an unusual reaction or isolated from a natural source needs much stronger supporting

evidence than one derived by a standard reaction from a precursor of undisputed structure.

3.7 Bibliographic References and Footnotes

A clear distinction is made between bibliographic references and footnotes. The latter are used to present material which, if included in the body of the text, would disrupt the flow of the argument but which is, nevertheless, of importance in qualifying or amplifying the textual material. Such footnotes are referred to with the following symbols: *,†,‡,§,¶,||, etc. [Note: Since an asterisk is used to indicate the author to whom correspondence should be addressed, its use early on in a paper is not advised; a dagger (†) is preferred.]

Bibliographic reference to the source of statements in the text is made by use of *superior numerals* at the appropriate place. The references themselves are given at the end of the final printed text. It is essential that they are numbered in the order in which they are cited in the text.

Journal Abbreviations

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Acc. Chem. Res.
Acta Acad. Abo, Ser. B
Acta Biochim. Biophys. Acad. Sci. Hung.
Acta Biochim. Iran.
                                                                                                                                             Annu. Rep. Prog. Chem., Sect. C, Phys. Chem.
Annu. Rev. Biochem.
Annu. Rev. Ind. Eng. Chem.
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Chem. Eng. Progr.
Chem. Eng. (Rugby, Engl.)
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Environ. Sci. Technol.
Erdoel Kohle, Erdgas, Petrochem., Brennst.
                                                                                                                                             Annu. Rev. Phys. Chem.
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Chem. Erde
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Appl. Spectrosc.
Arch. Pharm. Chemi, Sci. Ed.
Arch. Pharm. (Weinheim, Ger.)
 Acta Biochim, Pol.
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Chem. Heterocycl. Compd. (Engl. Transl.)
Chem. Ind. (Duesseldorf)
Chem. Ind. Int. (Engl. Transl.)
Chem. Ind. (London)
Chem. Ind. - Tech.
Chem. Ind. - Tech.
Acta Chem. Scand., Ser. A
Acta Chem. Scand., Ser. B
Acta Chim. Acad. Sci. Hung.
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Acta Crystallogr., Sect. A
Acta Crystallogr., Sect. B
Acta Crystallogr., Sect. C
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                                                                                                                                                                                                                                                                                         Chem. Ing. - Iech.
Chem. Listy
Chem. Nat. Compd. (Engl. Transl.)
Chem. NZ.
Chem. Pharm. Bull.
Chem. Phys.
Carbon
Chem. Phys. Carbon
Chem. Phys. Lipids
Chem. Prum.
Chem. Prum.
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                                                                                                                                             At. Absorbt. Newsl.
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Faraday Symp. Chem. Soc.
FEBS Lett.
Fermenin, Spirt. Prom.
Fette, Seifen. Anstrichm.
Finn Chem. Lett.
Fiz.-Khim. Mekh. Mater.
Fiz. Met. Metalloved.
                                                                                                                                             Aust. J. Biol. Sci.
Aust. J. Chem.
Aust. J. Phys.
 Acta Metall.
Acta Phys. Acad. Sci. Hung.
Acta Phys. Chem.
Acta Vitaminol. Enzymol.
                                                                                                                                             Azerb. Khim. Zh
Acta Vitaminol. Enzymol.
Adv. Act. Anal.
Adv. Alicyclic Chem.
Adv. Anal. Chem. Instrumen.
Adv. Carbohydr. Chem. Biochem.
Adv. Catol.
Adv. Chem. Phys.
Adv. Chromatogr.
Adv. Colloid Interface Sci.
Adv. Enzymol. Relat. Areas Mol. Biol.
Adv. Free-Radical Chem.
Adv. Herocycl. Chem.
                                                                                                                                             Ber. Bunsenges. Phys. Chem.
                                                                                                                                             Biochem. Biophys. Res. Commun.
Biochem. Educ.
Biochemistry
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Fluorine Chem. Rev.
Food Manuf.
Fortschr. Chem. Org. Naturst.
Fortschr. Hochpolym.-Forsch.
Fresenius' Z. Anal. Chem.
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ChemSA
                                                                                                                                             Biochemistry (Eng. Transl.)
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Chem. Soc. Rev.
Chem. Stosow.
Chem. Tech. (Leipzig)
Chem. Technol.
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Biochem. J.
Biochem. Pharmacol.
Biochem. Prep.
Biochem. Soc. Trans.
Biochim. Biophys. Acta
Biochimie
  Adv. Heterocycl. Chem.
Adv. Heterocycl. Chem.
Adv. Inorg. Bioinorg. Mech.
Adv. Lipid Res.
Adv. Lipid Res.
Adv. Macromol. Chem.
Adv. Mollen Salt Chem.
Adv. Mollen Salt Chem.
Adv. Organomet. Chem.
Adv. Org. Chem.
Adv. Org. Chem.
Adv. Org. Chem.
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Gen. Cytochem. Methods
Geokhimiya
Ger. Chem. Eng. (Engl. Transl.)
Gidrokhim. Mat.
Glas. Hem. Drus., Beograd
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Chem. Weekbl
                                                                                                                                             Biofizika
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Bioinorg, Chem.
Biokhimiya
Bioorg, Chem.
Bioorg, Khim.
Biopolymers
Biotechnol. Bioeng.
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Chem. Zvesti
Chim. Acta Turc.
Chim. Actual.
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God. Vissh. Khim.-Tekhnol. Inst., Sofia
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Clin. Biochem.
                                                                                                                                             Bochu-Kagaku
Boll. Soc. Ital. Biol. Sper.
Br. Corros. J.
Br. J. Pharmacol.
  Adv. Phys. Org. Chem.
Adv. Protein Chem.
Adv. Quantum Chem.
Adv. Struct. Res. Diffr. Methods
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Clin. Chem. (Winston-Salem, N.C.)
Clin. Chim. Acta
Collect. Czech. Chem. Commun.
Colloid J. USSR (Engl. Transl.)
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High Energy Chem. (Engl. Transl.)
Hist. Stud. Phys. Sci.
Hoppe-Seyler's Z. Physiol. Chem.
Hua Hsueh
                                                                                                                                             Br. Polym. J.
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Bul. Inst. Politeh. Iasi.
Bull. Acad. Sci. USSR, Div. Chem. Sci.
Bull. Chem. Soc. Jpn.
Bull. Inst. Chem. Res., Kyoto Univ.
Bull. Pol. Acad. Sci., Chem.
Bull. Sci., Cons. Acad. Sci. Arts RSF Yougosl.,
  Aub. Struct. Res. D
Afinidad
Agric. Biol. Chem.
Agrokem. Talajtan
AIChE J.
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Colloid Polym. Sci.
Combust. Flame
Commun. Fac. Sci. Univ. Ankara
  AICHE J.
Ambix
Am. J. Pharm.
Am. J. Sci.
An. Acad. Bras. Cienc.
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Commun. R. Soc. Edinburgh, Phys. Sci.
Comput. Chem.
Coord. Chem. Rev.
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Huaxue Tongbao
Huaxue Xuebao
Hung, J. Ind. Chem.
Hwahak Kwa Kongop Ui Chinbo
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Bull. Soc. Chim. Belg.
Bull. Soc. Chim. Fr.
Bunseki Kagaku
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  An. Acaa. Bras. C
Anal. Biochem.
Anal. Chem.
Anal. Chim. Acta
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CRC Crit. Rev. Biochem.
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Crit. Rev. Anal. Chem.
Croat. Chem. Acta
C.R. Seances Soc. Biol. Ses Fil.
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  Anal. Lett.
Analusis
Analyst (London)
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Ind. Eng. Chem., Process. Des. Dev.
Ind. Eng. Chem., Prod. Res. Dev.
Indian J. Agric. Chem.
Indian J. Biochem. Biophys.
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Can. J. Chem.
Can. J. Chem. Eng.
Can. J. Pharm. Sci.
Can. J. Phys.
Can. J. Spectrosc.
Carbohydr. Res.
                                                                                                                                                                                                                                                                                                                                                                                                                                        Indian J. Agric. Chem.
Indian J. Biochem. Biophys.
Indian J. Chem., Sect. A
Indian J. Chem., Sect. A
Indian J. Chem., Sect. B
Indian J. Pure Appl. Phys.
Ind. Lab. (Engl. Transl.)
Inorg. Chem.
Inorg. Chim. Acta
Inorg. Mater. (Engl. Transl.)
Inorg. Synth.
Int. Chem. Eng.
Int. Flavours Food Addit.
Int. J. Appl. Radiat. Isot.
Int. J. Appl. Radiat. Isot.
Int. J. Mass Spectrum. Ion Processes
Int. J. Chem. Kinet.
Int. J. J. Wasnum Chem.
Int. J. Quantum Chem.
Int. J. Sulfur Chem.
Int. J. Sulfur Chem.
Int. J. Sulfur Chem.
Int. J. Vitam. Nutr. Res.
   An Bromatol
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  An. Bromatol.
Angew. Chem.
Angew. Chem., Int. Ed. Engl.
Angew. Makromol. Chem.
Ann. Acad. Sci. Fenn., Ser. A2
Ann. Chim. (Paris)
Ann. Chim. (Rome)
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Denki Kagaku yobi Kogyo Butsuri Kagaku
Dokl. Akad. Nauk Arm. SSR
Dokl. Bolg. Akad. Nauk SSSR
Dokl. Bolg. Akad. Nauk
Dokl. Chem. (Engl. Transl.)
Dokl. Chem. Technol. (Engl. Transl.)
Dokl. Phys. Chem. (Engl. Transl.)
Dopov. Akad. Nauk Ukr. RSR. Ser. B
Dauble-Liaison
Disch. Lehonm - Rundsch
                                                                                                                                               Carbon
                                                                                                                                               Catal. Rev.
                                                                                                                                              Cellul. Chem. Technol.
Cereal Chem.
Cesk. Farm.
   Ann Endocrinol
  Ann. N.Y. Acad. Sci.
Ann. Pharm. Fr.
Ann. Soc. Sci. Bruxelles, Ser. 2
                                                                                                                                              Chelates Anal. Chem.
Chem. Age (London)
Chem. Anal. (Warsaw)
  Ann. Univ. Mariae Curie-Sklodowska, Sect.
                                                                                                                                                                                                                                                                                         Dtsch. Lebensm.-Rundsch.
Dyn. Mass Spectrom.
 AA
Annu. Rep. Anal. At. Spectrosc.
Annu. Rep. Med. Chem.
Annu. Rep. N.M.R. Spectrosc.
Annu. Rep. Prog. Chem., Sect. A, Inorg.
                                                                                                                                               Chem. Ber.
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Chem. Br.
Chem. Chron.
Chem. Econ. Eng. Rev.
Chem. Eng. Commun.
Chem. Eng. J. (Lausanne)
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                                                                                                                                                                                                                                                                                          Egypt. J. Chem.
Egypt. J. Chem.
Electroanal. Chem.
Electrochim. Acta
  Annu. Rep. Prog. Chem., Sect. B, Org. Chem.
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Journal Abbreviations (continued)

Intra-Sci. Chem. Rep.	J. Phys. E.	Pak. J. Sci. Ind. Res.	Sci. Rep. Tohoku Univ., Ser. 1
Inz. Chem.	J. Polym. Sci., Polym. Chem. Ed.	Pak J. Sci. Res.	Sci. Sinica
Ion Exch. Solvent Extr.	J. Polym. Sci., Polym. Phys. Ed.	Periodia Polytech., Chem. Eng.	Sel. Annu. Rev. Anal. Sci.
Isr. J. Chem.	J. Polym. Sci., Polym. Symp.	Pestic. Sci.	Semicond. Insul.
Istanbul Univ. Fen Fak. Mecm., Seri C	J. Prakt. Chem.	Philos. Mag.	Sep. Purif. Methods
Ital. J. Biochem.	J. Protein Chem.	Philos. Trans. R. Soc. London, A	Sep. Sci. Technol.
Itsuu Kenkyusho Nempo	J. Quant. Spectrosc. Radiat. Transfer	Phosphorus Sulfur	Soap. Cosmet., Chem. Spec.
Izv. Akad. Nauk Kaz. SSR, Ser. Khim.	J. Radioanal. Nucl. Chem.	Photochem. Photobiol.	Sov. Electrochem. (Engl. Transl.)
Izv. Akad. Nauk SSSR, Neorg. Mater.	J. Raman Spectrosc.	Phys. Chem. Glasses	Sov. J. Bioorg. Chem. (Engl. Transl.)
Izv. Akad. Nauk SSSR, Ser. Khim. Izv. Akad. Nauk Turkm. SSR, Ser. Fiz	J. Res. Inst. Catal., Hokkaido Univ. J. Res. Nat. Bur. Stand., Sect A	Phys. Rev.	Sov. J. Coord. Chem. (Engl. Transl.)
Tekh., Khim. Geol. Nauk	J. Res. Nat. Bur. Stana., Sect A J. Sci. Food Agric.	Phys. Rev. Lett.	Sov. PhysCrystallogr. (Engl. Transl.) Sov. Radiochem. (Engl. Transl.)
Izv. Sib. Otd. Akad. Nauk SSSR, Ser.	J. Sci. Hiroshima Univ., Ser. A2	Phys. Scr.	Spec. Publ. Chem. Soc.
Khim. Nauk	J. Sci. Ind. Res.	Phytochemistry	Spectrochim. Acta, Part A
1	J. Soc. Dyers Colour.	Pigm. Resin Technol. Pis'ma Zh. Eksp. Teor. Fiz.	Spectrochim. Acta, Part B
J. Agric. Food Chem.	J. Soc. Leather Technol. Chem.	Plast. Polym.	Spectrosc. Lett.
J. Am. Chem. Soc.	J. Solid State Chem.	Pol. J. Chem.	Steroids
J. Am. Leather Chem. Assoc.	J. Solution Chem.	Polyhedron	Steroids Lipids Res.
J. Am. Oil Chem. Soc.	J. Steroid Biochem.	Polym. Age	Struct. Bonding (Berlin)
J. Anal. At. Spectrom.	J. Struct. Chem. (Engl. Transl.)	Polym. Commun.	Stud. Univ. Babes-Bolyai, Chem.
J. Anal. Chem. USSR (Engl. Transl.)	J. Text. Inst.	Polymer	Sub-Cell. Biochem.
J. Appl. Chem. Biotechnol.	J. Therm. Anal.	Polym. Sci. USSR (Engl. Transl.)	Surf. Colloid Sci.
J. Appl. Chem. USSR (Engl. Transl.)		Postepy Biochem.	Surf. Sci.
J. Appl. Crystallogr.	Kagaku Kogaku	Prikl. Biokhim. Mikrobiol.	Synth. Commun.
J. Appl. Polym. Sci.	Kanazawa Daigaku Yakugakubu Kenkyu	Priroda (Moscow)	Synthesis
J. Assoc. Off. Anal. Chem. J. Assoc. Public Anal.	Nempo	Proc. Am. Soc. Brew. Chem.	Synth. React. Inorg. Metal-Org. Chem.
J. Biochem. (Tokyo)	KemKemi	Process Biochem.	Total and House I also Chi
J. Biol. Chem.	Kem. Tidskr. Khim. Geterotsikl. Soedin.	Processing	Taehan Hwahakhoe Chi Talanta
J. Carbohydr. Chem.	Khim. Ind. (Sofia)	Proc. Indian Acad. Sci., Sect. A	Technol. Rep. Osaka Univ.
J. Catal.	Khim. Neft. Mashinostr.	Proc. Indian Acad. Sci., Sect. B	Teor. Eksp. Khim.
J. Chem. Educ.	Khim. Prir. Soedin.	Proc., K. Ned. Akad. Wet., Ser. B	Teor. Osn. Khim. Tekhnol.
J. Chem. Eng. Datu	Khim. Promst. (Moscow)	Proc., K. Ned. Akad. Wet., Ser. C Proc. Natl. Acad. Sci., India, Sect. A	Tetrahedron
J. Chem. Inf. Comput. Sci.	Khim. Volokna	Proc. Natl. Acad. Sci., India, Seci. A Proc. Natl. Acad. Sci. USA	Tetrahedron Lett.
J. Chem. Phys.	Khim. Vys. Energ.	Proc. R. Soc. Edinburgh, Sect. A	Text. Res. J.
J. Chem. Res. (M)	Kinet. Catal. (Engl. Transl.)	Proc. R. Soc. London, A	Theor. Exp. Chem. (Engl. Transl.)
J. Chem. Res. (S)	Kinet. Katal.	Proc. R. Soc. London, B	Thermochim. Acta
J. Chem. Soc., Chem. Commun.	Kjemi	Proc. Soc. Exp. Biol. Med.	Tin Its Uses
J. Chem. Soc., Dalton Trans.	Kobunshi Kagaku	Proc. Soc. Exp. Biol. Med. Prog. Bioorg. Chem. Prog. Colloid Polym. Sci.	Top. Curr. Chem.
J. Chem. Soc., Faraday Trans. 1	Kogyo Kagaku Zasshi	Prog. Colloid Polym. Sci.	Top. Stereochem.
J. Chem. Soc., Faraday Trans. 2	Kolloidn. Zh.	Prog. Inorg. Chem.	Trans. Inst. Met. Finish.
J. Chem. Soc., Perkin Trans. 1	Koord. Khim.	Prog. Inorg. Chem. Prog. Med. Chem.	Transition Met. Chem. (Weinheim, Ger.)
J. Chem. Soc., Perkin Trans. 2	Kristallografiya	Prog. Nucleic Acid Res. Mol. Biol.	Trant. J. Br. Ceram. Soc.
J. Chem. Thermodyn.	KunststPlast. (Solothurn, Switz.)	Prog. Nucl. Magn. Reson. Spectrosc.	Trends Anal. Chem.
J. Chim. Phys. PhysChim. Biol.		Prog. Phys. Org. Chem.	Trends Biochem. Sci.
J. Chin. Chem. Soc. (Taipei)	Lab. Pract.	Prog. React. Kinet.	Tr. Inst. Elektrokhim., Ural. Nauchn. Tsentr.
J. Chromatogr.	Langmuir	Prog. Solid State Chem.	Akad. Nauk SSSR
J. Chromatogr. Sci. J. Colloid Interface Sci.	Laser Chem.	Prog. Stereochem.	## b:-1.1: 71
J. Coord. Chem.	Latv. PSR Zinat. Akad. Vestis, Kim. Ser.	Prog. Surf. Membr. Sci.	Ukr. Biokhim. Zh.
J. Cryst. Mol. Struct.	Liebigs Ann. Chem.	Prog. Surf. Sci.	Ukr. Khim. Zh. (Russ. Ed.)
J. Doc.	Lipids	Prog. Thin-Layer Chromatogr. Relat.	Usp. Khim. UV Spectrum. Group Bull.
J. Electroanal. Chem. Interfacial	Macromolecules	Methods	Uzb. Khim. Zh.
Electrochem.	Macromole Cules Macromol. Synth.	Przem. Chem.	UZU. KIIIII. ZII.
J. Electrochem. Soc.	Magy. Kem. Foly.	Pure Appl. Chem.	Vestn. Leningr. Univ., Fiz., Khim.
J. Electrochem. Soc. India	Magy. Kem. Lapja	Pyrethrum Post	Vestn. Mosk. Univ., Ser. 2: Khim.
J. Fac. Sci. Univ. Tokyo	Makromol. Chem.	Quad. Ing. Chim. Ital.	Vestn. Slov. Kem. Drus.
J. Fluorine Chem.	Manuf. Chem. Aerosol News	Quim. Nova	Vestsi Akad. Navuk BSSR. Ser. Khim.
J. Food Sci.	Meded. Vlaam. Chem. Ver.	Quim. Nota	Navuk
J. Franklin Inst.	Mekh. Polim.	Radiat. Phys. Chem.	Veszpremi Vegyip, Egy. Kozl.
J. Gen. Chem. USSR (Engl. Transl.)	Mem. Fac. Sci. Kyushu Univ., Ser. C	Radiat. Res.	Vitam. Horm. (N.Y.)
J. Hazard. Mater.	Mem. Inst. Protein Res., Osaka Univ.	Radiochim. Acta	Vopr. Med. Khim.
J. Heterocycl. Chem.	Mem. Inst. Sci. Ind. Res., Osaka Univ.	Radiokhimiya	Vysokomol. Soedin., Ser. A
J. Histochem. Cytochem. J. Indian Chem. Soc.	Mendeleev Chem. J. (Engl. Transl.)	React. Kinet. Catal. Lett.	Vysokomol. Soedin., Ser. B
	Methods Biochem. Anal.	Recent Dev. Chem. Nat. Carbon Compd.	w tra
J. Indian Inst. Sci. J. Inorg. Chem. USSR (Engl. Transl.)	Methods Free-Radical Chem.	Recent Prog. Horm. Res.	Xenobiotica
J. Inst. Brew.	Microchem. J. Mikrochim. Acta	Recherches	Yakugaku Zasshi
J. Inst. Chem. (India)	Mikrochim. Acta Mol. Cell. Biochem.	Recl. Trav. Chim. Pays-Bas	Yuki Gosei Kagaku Kyokaishi
J. Inst. Fuel	Mol. Cryst. Liq. Cryst.	Rend. Accad. Sci. Fis. Mat., Naples	I MAI GUSEL RUBUKU KYUKUISIII
J. Labelled Compd. Radiopharm.	Mol. Phys.	Rep. Prog. Appl. Chem. Residue Rev.	Z. Anorg. Allg. Chem.
J. Less-Common Met.	Monatsh. Chem.	Rev. Anal. Chem.	Zavod. Lab.
J. Lipid Res.		Rev. Asoc. Bioquim. Argent.	Zb. Pr. Chemickotechnol. Fac. SVST
J. Liq. Chromatogr.	Nahrung	Rev. Chim. (Bucharest)	Z. Chem.
J. Lumin.	Nat. Prod. Rep.	Rev. Phys. Chem. Jpn.	Zentralbl. Pharm., Pharmakother. Labora-
J. Macromol. Sci., Chem.	Nature (London)	Rev. Port. Quim.	toriumsdiagn.
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J. Mol. Biol. J. Mol. Catal.	Nippon Nogei Kagaku Kaishi	Rev. Soc. Quim. Mex.	Zh. Fiz. Khim.
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J. Mol. Spectrosc. J. Mol. Struct.	Nucleosides, Nucleotides	Rubber Chem. Technol.	Zh. Neorg. Khim.
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J. Nonmet. Semiconduct.	Online (Weston, Conn.)	Russ. J. Inorg. Chem. (Engl. Transl.)	Zh. Org. Khim. Zh. Prikl. Khim. (Leningrad)
J. Oil Colour Chem. Assoc.	Orbital	Russ. J. Phys. Chem. (Engl. Transl.)	Zh. Prikl. Knim. (Leningraa) Zh. Prikl. Spektrosk.
J. Organomet. Chem.	Organometallics	S Afr. I Cham	Zh. Priki. Spekirosk. Zh. Struki. Khim.
J. Org. Chem.	Org. Magn. Reson.	S. Afr. J. Chem.	Zh. Strukt. Knim. Zh. Vses. Khim. Ova im D.I. Mendeleeva
J. Org. Chem. USSR (Engl. Transl.)	Org. Mass. Spectrom.	Sankyo Kenkyusho Nempo Sb. Ved. Pr., Vys. Sk. Chemickotechnol.,	Z. Kristallogr.
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J. Pharm. Sci.	Org. React. Mech.	Sci. Cult.	Z. Naturforsch., C
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J. Phys. Chem. Solids	Pak. J. Sci.	Sci. Rep. Res. Inst., Tohoku Univ.	Z. Wiss. Photogr., Photophys., Photochem.
***************************************	Tuk. V. Sci.	Sci. Rep. Res. Inst., Tonoka Oniv.	

The position of the superior numeral should be chosen with care, particularly when it does not follow an author's name. If placed adjacent to punctuation, the numeral should normally be placed after the punctuation mark, e.g. 'This compound was shown to be the dienone,³ which...'.

Particular care is necessary where a reference number is likely to be confused with a superscript numeral indicating a power index: '... which gave a value of 2.3 cm³...' should be written as '... which gave a value of 2.3 cm' or '... which gave a value of 2.3 cm (ref. 3).'

Since it is difficult to predict the final position of a table in the text, references cited only in the table should be incorporated into the printed footnotes to the table. References which are also cited elsewhere in the text should be referred to in the footnotes by the numbers used in the text citations, e.g. a Ref. 15.

Journals. The style of journal abbreviations to be used in the Society's publications is that defined in Chemical Abstracts Service Source Index (CASSI). The abbreviations listed in CASSI are based upon internationally recognised systems. The list of CASSI-style abbreviations on pages xii—xiii covers most of the journals received in the library of the Royal Society of Chemistry. It is not, of course, a full list; CASSI plus its quarterly supplements run to more than 2 000 pages.

If you cannot locate an authoritative abbreviation for a journal, and if it is not obvious how the title should be abbreviated, please cite the full title.

Bibliographic details should be cited in the order: year, volume, page.

Books. Titles of books are cited in quotation marks, in upright letters, and the authors(s), title, publisher, town, date (or edition, if more than one has been published), and page number (if required) must be given in that order:

D. Brown, 'Halides of the Lanthanides and Actinides,' Wiley, London, 1968, p. 50.

H. A. O. Hill in 'New Trends in Bioinorganic Chemistry,' ed. R. J. P. Williams and J. R. F. Dasilva, Academic Press, London, 1978, p. 85.

Patents. Patents should be indicated in the form: B.P. 357 450, 367 455—7. U.S.P. 1 171 230. G.P. 436 112—4, Jap.P. 20 101. Dates are indicated thus: B.P.666 776/1956. Patents which are applied for must always be given a year, e.g. B.P. Appl. 102/1982.

Reports and Bulletins, etc.

R. A. Allen, D. B. Smith, and J. E. Hiscott, 'Radioisotope Data,' UKAEA Research Group Report AERE-R 2938, H.M.S.O., London, 1961.

G. M. Sheldrick, SHELX-76, Program for Crystal Structure Determinations, University of Cambridge, 1976.

Material presented at meetings.

- R. G. Kidd and H. G. Spinney, presented at the 5th International Conference on Non-Aqueous Solutions, Leeds, 1976
- H. C. Freeman, Proceedings of the 21st International Conference on Coordination Chemistry, Toulouse, 1980.

 Theses.
 - A. D. Mount, Ph.D. Thesis, University of London, 1977.

Reference to unpublished material. For material presented at a meeting, congress, or before a Society, etc., but not published, the following form is used:

1 A. R. Jones, presented in part at the 28th Congress of the International Union of Pure and Applied Chemistry, Vancouver, August, 1981.

For material accepted for publication, but not yet published, the following form is used:

2 A. R. Jones, J. Chem. Soc., Dalton Trans., in the press.

If the paper has been submitted to the Society, the paper number should be given:

3 A. R. Jones, J. Chem. Soc., Dalton Trans., in the press (2/556).

For material submitted for publication but not yet accepted the following form is used:

- 4 A. R. Jones, submitted for publication in *Angew. Chem.* For personal communications the following is used:
 - 5 G. B. Ball, personal communication. (*Note:* the form G. B. Ball, private communication, is inappropriate.)

If material is to be published but has yet to be submitted the following form is used:

6 G. B. Ball, unpublished work.

Reference to unpublished work should not be made without the permission of those by whom the work was performed.

Names. The names and initials of all authors are always given in the reference; they must not be replaced by the phrase et al. This does not prevent some, or all, of the names being mentioned at their first citation in the cursive text: initials are not necessary in the text.

For Chinese and Spanish authors all names should be given as in the original, since the patronymic is not always given last in these languages. If co-authors are to be collectively cited, as in 'Smith and co-workers' or 'Smith et al.,' the latter form is inappropriate unless the individual name 'Smith' appears first among the authors named in the original.

Composite references. Whenever possible, composite references should be used rather than a series of individual references. The style for composite references is as follows:

- 1 A. B. Jones, J. Chem. Soc., Dalton Trans., 1975, 234.
- 2 A. B. Jones, J. Chem. Soc., Dalton Trans., 1977, 123; 1978, 234.
- 3 A. B. Jones, J. Chem. Soc., Dalton Trans., 1977, 123; J. Am. Chem. Soc., 1956, 78, 1234.
- 4 A. B. Jones, J. Chem. Soc., 1956, 234; A. B. Jones and C. D. Brown, J. Chem. Soc. B, 1967, 234, 1077; 1968, 599.
- 5 A. B. Jones, J. Am. Chem. Soc., 1956, 78, 1234; A. B. Jones and C. D. Brown, *ibid.*, 1957, 79, 567; A. B. Jones and E. F. Green, *ibid.*, p. 999.

If only one paper from a composite reference is required for citation later, then two numbers may be assigned to the first citation (e.g. Jones ^{1,2}); alternatively, long composite references may be divided by letters, e.g.:

(a) A. B. Jones, J. Chem. Soc., Dalton Trans., 1978, 467; (b) A. B. Jones and C. D. Brown, J. Chem. Soc., Perkin Trans. 2, 1979, 234.

A. B. Jones, J. Chem. Soc. A, (a) 1967, 267; (b) 1968, 1742; (c) etc.

A composite reference may cite a previous reference in the form:

12 A. B. Jones, J. Chem. Soc., 1956, 234; C. D. Brown, ref. 5. (Note: ibid. is used only within a given reference and not to refer from one reference number to another: the abbreviated title for the journal should be repeated for separate reference numbers.)

Idem, loc. cit., and op. cit are not used in references.

3.8 Nomenclature

For many years the Society has actively encouraged the use of standard IUPAC nomenclature and symbolism in its publications as an aid to the accurate and unambiguous communication of chemical information between authors and readers.

In order to encourage authors to use IUPAC nomenclature

rules when drafting papers, attention is drawn to the following publications in which both the rules themselves and guidance on their use are given:

Nomenclature of Organic Chemistry, Sections A, B, C, D, E, F, and H, Pergamon, Oxford, 1979 edn.

Nomenclature of Inorganic Chemistry, Butterworths, London, 1971 (now published by Pergamon).

Biochemical Nomenclature and Related Documents, The Biochemical Society, London, 1978.

Compendium of Chemical Terminology: IUPAC Recommendations, Blackwell, Oxford, 1987.

A listing of all relevant IUPAC nomenclature publications appears as an appendix to these Instructions.

It is recommended that where there are no IUPAC rules for the naming of particular compounds or authors find difficulty in applying the existing rules, they should seek the advice of the Society's editorial staff.

3.9 Units and Symbols

The Symbols Committee of The Royal Society, of which The Royal Society of Chemistry is a participating member, has produced a set of recommendations in a pamphlet 'Quantities, Units, and Symbols,' 1975 (copies of this pamphlet and further details can be obtained from the Manager, Journals, The Royal Society of Chemistry, Burlington House, Piccadilly, London, W1V 0BN). These recommendations are applied by The Royal Society of Chemistry in all its publications. Their basis is the 'Système International d'Unités' (SI).

A more detailed treatment of units and symbols with specific application to chemistry is given in the IUPAC Manual of Symbols and Terminology for Physicochemical Quantities and Units (Pergamon, Oxford, 1979).

Guidelines for the publications of the Society. An author will not be denied any reasonable usage, but if non-SI units are used for critical data or for quantities measured to a high order of accuracy (as opposed to the rough physical conditions of an experiment), the definitive values will be expressed in SI units as well.

The following will be the guidelines used:

- (a) A metric system will always be used in preference to a nonmetric one.
 - (b) SI will be the standard usage.
- (c) The units used to record the definitive values of 'critical data' or quantities measured to a high degree of accuracy will be
- (d) When non-SI units are used they must be adequately explained unless their definition is obvious (e.g. degree Celsius, mmHg, g, h). The derivation of derived non-SI units will be indicated.
- (e) Equations involving electrical quantities should normally be those appropriate for use with SI (rationalized m.k.s) units. If authors wish to use equations suitable for e.s.u. or e.m.u. the

Table 1. Base units

Physical quantity	Name of base-unit	Symbol for unit
length	metre	m
mass	kilogram	kg
time	second	S
electrical current	ampere	Α
thermodynamic		
temperature	kelvin	K
luminous intensity	candela	cd
amount of substance	mole	mol

lack of consistency with SI units must be explicitly noted.

- (1) Base-units. The SI base-units are given in Table 1.
- (2) Supplementary units. The SI also includes two 'supplementary' dimensionless units as follows:

Physical quantity	Name of unit	Symbol for unit
plane angle	radian	rad
solid angle	steradian	sr

(3) Multiples and sub-multiples. In the SI there is one and only one basic unit for each physical quantity. Decimal fractions and multiples of these basic units may, however, be constructed by use of certain prefixes (see Table 2). They may also be used with derived SI units.

The combination of a prefix and a unit symbol constitutes a new single unit symbol; compounding of prefixes is not permitted.

Although it will not always be possible, particularly in Tables, the general principle should be to choose a unit (i.e. including multiple or sub-multiple) such that the resulting numerical value is between 0.1 and 1 000.

- (4) Derived units. Some derived units have special names and symbols, and these are given in Table 3. Others do not (Table 4).
- (5) Symbol. The symbol for a unit will be printed in roman (upright) type, remains unaltered in the plural and does not take a full point, i.e. 5 cm not 5 cm. or 5 cms or 5 cms.

The symbol will be separated from the numerical value by a thin space.

(6) Decimal fractions and multiples of SI units having special names. These names are not part of the SI, but for the time being their use in the Society's publications may continue. The list given in Table 5 is not exhaustive.

Table 2. Prefixes

Fraction	Prefix	Symbol	Multiple	Prefix	Symbol,
10^{-1}	deci	d	10	deka	da
10^{-2}	centi	c	10 ²	hecto	h
10^{-3}	milli	m	10^{3}	kilo	k
10 ⁻⁶	micro	μ	10 ⁶	mega	M
10^{-9}	nano	n	10 ⁹	giga	G
10^{-12}	pico	p	10^{12}	tera	T
10^{-15}	femto	f			
10^{-18}	atto	a			

Table 3. Derived units with special names and symbols

Physical quantity	Name of SI unit	Symbol for SI unit	Definition of SI unit
energy	joule	J	$kg m^2 s^{-2}$
force	newton	N	$kg \ m \ s^{-2} = J \ m^{-1}$
power	watt	W	$kg m^2 s^{-3} = J s^{-1}$
electric charge	coulomb	C	As
electric potential			
difference	volt	V	$kg m^2 s^{-3} A^{-1} = J A^{-1} s^{-1}$
electric resistance	ohm	Ω	$kg m^2 s^{-3} A^{-2} = V A^{-1}$
electric capacitance	farad	F	$A^2 s^4 kg^{-1} m^{-2} = A s V^{-1}$
magnetic flux	weber	Wb	$kg m^2 s^{-2} A^{-1} = V s$
inductance	henry	Н	$kg m^2 s^{-2} A^{-2} = V A^{-1} s$
magnetic flux			
density	tesla	T	$kg s^{-2} A^{-1} = V s m^{-1}$
luminous flux	lumen	lm	cd sr
illumination	lux	lx	cd sr m ⁻²
frequency	hertz	Hz	s ⁻¹

(7) Units defined in terms of the best available experimental values of certain physical constants. These units are not part of the SI. The factors for conversion of these units to SI units are subject to change in the light of new experimental measurements of the constants involved. Their use outside the restricted contexts to which they are appropriate should be discouraged. The following list is not exhaustive.

Physical quantity	Name of unit	Symbol for unit	Conversion factor
quantity	ume	ioi uiiit	Conversion factor
energy	electron volt	eV	$eV = 1.6021 \times 10^{-19} J$
mass	unified atomic	u	$u = 1.660 41 \times 10^{-27} \mathrm{kg}$
	mass unit		

(8) Other units now exactly defined in terms of the SI units. These units are not part of the SI. It is recognized that their use may be continued for some time but it is recommended that except in special circumstances they should be progressively abandoned in conformity with international recommendations.

Table 4. Derived units with no special names or symbols

Physical quantity	SI unit	Symbol for SI unit
area	square metre	m²
volume	cubic metre	m ³
density	kilogram per cubic metre	k m ⁻³
velocity	metre per second	$m s^{-1}$
angular velocity	radian per second	rad s ⁻¹
acceleration	metre per second squared	m s ⁻²
pressure	newton per square metre	$N m^{-2}$
kinematic viscosity, diffusion coefficient	square metre per second	$m^2 \ s^{-1}$
dynamic viscosity	newton second per square metre	$N s m^{-2}$
electric field strength	volt per metre	$V m^{-1}$
magnetic field strength	ampere per metre	$A m^{-1}$
luminance	candela per square metre	cd m ⁻²

The list given in Table 6 is by no means exhaustive. Each of the definitions given in the fourth column is exact.

3.10 Notes for Typists

Manuscripts must be typed in double-line spacing, single sided on A4 paper, with margins at top, bottom, and left-hand side of at least 4 cm.

The first page should be set out as follows (see appended example on p. xvii):

- (i) Name and address for proofs.
- (ii) Title of paper, with capitals for first letter of each noun or adjective only.
- (iii) Authors' names; an asterisk should follow the name of the author who is to receive any correspondence.
- (iv) The address where the work was carried out; if this is different from the present address of the asterisked author, a footnote indicating this present address should be included. Present addresses of other authors are not normally given.
- (v) Summary, preceded and followed by a horizontal line, and typed in double-line spacing.
 - (vi) Main text.

Table 5. Fractions and multiples of units with special names

	•	•	
Physical quantity	Name of unit	Symbol for unit	Definition of unit
length	ångström	Å	$10^{-10} \mathrm{m} = 10^{-1} \mathrm{nm}$
length	micron	μm	10 ⁻⁶ m
area	barn	ь	10 ⁻²⁸ m ²
volume	litre	1	$10^{-3} \text{ m}^3 = \text{dm}^3$
mass	tonne	t	$10^3 \text{ kg} = \text{Mg}$
force	dyne	dyn	10 ⁻⁵ N
pressure	bar	bar	10^5 N m^{-2}
pressure	pascal	Pa	$N m^{-2}$
energy	erg	erg	10 ⁻⁷ J
kinematic viscosity,			
diffusion coefficient	stokes	St	$10^{-4} \text{ m}^2 \text{ s}^{-1}$
dynamic viscosity	poise	P	10 ⁻¹ kg m ⁻¹ s ⁻¹
magnetic flux	maxwell	Mx	10 ⁻⁸ Wb
magnetic flux density			
(magnetic induction)	gauss	G	10 ⁻⁴ T
conductance	sîemens	S	Ω^{-1}

Table 6. Units defined in terms of SI units

Physical quantity	Name of unit	Symbol for unit	Definition of unit
length	inch	m	$2.54 \times 10^{-2} \text{ m}$
mass	pound (avoirdupois)	lb	0.453 592 37 kg
time *	minute	min	60 s
time *	hour	h	3 600 s
force	kilogram-force	kgf	9.806 65 N
force	pound-force	lbf	9.806 65 × 0.453 592 37 N
pressure	atmosphere	atm	101 325 N m ²
pressure	conventional millimetre of mercury	mmHg	$13.5951 \times 9.806 65 \text{ N m}^{-2}$
pressure	torr	Torr	(101 325/760) N m ⁻²
pressure	pound-force per square inch	lbf in ⁻²	$\frac{9.806\ 65\ +\ 4\ 535.9237}{6.4516}\ \text{N m}^{-2}$
energy	kilowatt hour	kW h	$3.6 \times 10^4 \text{ J}$
energy	thermochemical calorie	cal(thermochem.)	4.184 J
energy	I.T. calorie	cal _{it}	4.1868 J
thermodynamic temperature	degree Rankine	°R	(5/9) K
radioactivity	curie	Ci	$3.7 \times 10^{16} \mathrm{s}^{-1}$

^{*} Use of other common units (min, h, day) may continue in normal expressions of intervals of time.

Specimen first page of typescript

Proofs to: J. Dalton,
Royal Society of Chemistry,
Burlington House,
Piccadilly,
LONDON
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Inclusion Properties of Structures of the Type (RSC)_n

John Dalton,* Michael Faraday, and William H. Perkin Royal Society of Chemistry, Burlington House, Piccadilly, LONDON, W1V OBN

The clathrating ability of various structures of the type $(RSC)_n$ has been studied. Hexakis(alkylthio)benzenes, $(RSC)_6$, exhibit considerable activity as host molecules, and exceptional stability is conferred by SAc substituents.

Recent reports of inclusion behaviour by hexakis(phenylthio)-benzene, $(PhSC)_6$, ¹ and some initial studies by R.S. Cahn, ² led us to investigate the host-guest chemistry of further structures of the type $(RSC)_n$. We were particularly interested in the benzene derivatives $(RSC)_6$, which were expected to possess an attractive divisional structure.

The multi-step synthesis of (RSC)₆, carried out over a period of several years, was achieved......

Tables and captions for Figures should be typed on separate sheets at the end of the manuscript.

For typing of headings see section 3.3.5.

No underlining (for italics) is called for, as this will be done by the editor.

4.0 Deposition of Data: Supplementary Publications Scheme

Bulk information (such as primary kinetic data, computer programs, and output, evidence for amino-acid sequences, spectra, etc.), which accompanies papers published in the Journal of the Chemical Society may be deposited, free of charge, with the Society's Supplementary Publications Scheme, either at the request of the author and with the approval of the referees or on the recommendation of referees and with the approval of the author.

Under this scheme, authors should submit articles and the supplementary material to the *Journal* simultaneously in the normal way, and both will be refereed. If the paper is accepted

for publication the supplementary material will be sent by the Society to the British Library Document Supply Centre (Boston Spa) (BLDSC), where it will be stored. Copies will be obtainable by individuals both in the U.K. and abroad on quoting a supplementary publication number that will appear in the parent article.

4.1 Preparation of Material

Authors will be responsible for the preparation of cameraready copy according to the following specifications (although the Society will be prepared to help in case of difficulty).

- (a) Optimum page size for text or tables in typescript: up to $30 \text{ cm} \times 21 \text{ cm}$.
- (b) Limiting page size for text or tables in typescript: 33 cm \times 24 cm.
- (c) Limiting size for diagrams, graphs, spectra, etc.: 39 cm \times 28.5 cm.
- (d) Tabular matter should be headed descriptively on the first page, with column headings recurring on each page.

(e) Pages should be clearly numbered.

It is recommended that all material which is to be deposited should be accompanied by some prefatory text. Normally this will be the summary from the parent paper and authors will greatly aid the deposition of the material if a duplicate copy of the summary is provided.

4.2 Deposition

The Society will be responsible for the deposition of the material with the BLDSC. The BLDSC will not receive material direct from authors since the Library wishes to ensure that the material has been properly and adequately refereed.

4.3 Action by the Society

The Society will receive a manuscript for publication together with any supplementary material for deposition and will circulate all this to referees in the normal way. When the edited manuscript is sent to the printers the supplementary material will be sent for deposition to the BLDSC. The Society will add to the paper a footnote indicating what material has been deposited in the Supplementary Publications Scheme, and the supplementary publication number.

4.4 Availability

Copies of Supplementary Publications may be obtained from the BLDSC on demand by organizations which are registered borrowers. They should use the normal forms and coupons for such requests addressing them as follows:

> Special Acquisitions, British Library Document Supply Centre, Boston Spa, Wetherby, West Yorkshire, LS23 7BQ, U.K.

Non-registered users may also obtain copies of Supplementary Publications but should first apply for price quotations. These are available from the Loans Office at the above address.

5.0 Publication of X-Ray Crystallographic Work

Crystallographic work will be assessed mainly for its chemical interest. Thus crystallographic work carried out as part of a wider chemical study should not normally be submitted for publication separately from the results of that study. However, papers reporting only the results of crystal structure determinations may be accepted for publication provided that these results are considered to possess specifically chemical significance.

The description of a crystallographic structure determination should be as brief as possible, consistent with the following guidelines, and should be included at the end of a paper (or at the end of the Experimental section, if this precedes the Discussion). If the procedures for data collection and structure analysis were routine, their description should be particularly concise.

5.1 Title and Summary

The title of a paper reporting a crystallographic structure determination should normally contain the words 'Crystal Structure of...'

The summary need not contain cell dimensions and other crystal data, but should make clear that a crystal structure analysis has been performed.

5.2 Presentation of Crystal Data

The following sequence should be employed (although not all items will be required in all cases): (1) chemical formula and formula weight (M); (2) crystal system; (3) unit cell dimensions (Å or pm, degrees) and volume, with estimated standard deviations, method of determination, and temperature (if not ambient); (4) type and wavelength of radiation used; (5) space group symbol (if non-standard give related standard setting); (6) measured density (D_m) , no. of molecules in unit cell (Z), and calculated density (D_x) (with any implications for 'molecular symmetry'); (7) colour, size, and shape of crystal used for data collection; (8) linear absorption coefficient (μ) .

5.3 Data Collection and Processing

The description of data collection should contain sufficient information for the reliability of the determination to be assessed. The following should normally be included:

- (1) Type of diffractometer, measuring mode, and temperature if not ambient (with reference if possible to an earlier full description).
 - (2) Theta range and reciprocal lattice segment.
- (3) Number of data measured, number observed, and $I/\sigma(I)$ criterion used (if more than one asymmetric unit is recorded, the merging R value should be given).
 - (4) Absorption correction and method used (with reference).
 - (5) Stability of intensity controls and any action taken.

5.4 Structure Analysis and Refinement

The information provided should fulfil the same criteria as for data collection (section 5.3). This would normally include:

- (1) Method of solution (direct, heavy atom, or combination) and development.
- (2) Mode of refinement (full matrix, blocked, block diagonal, anisotropic, isotropic, or mixed), treatment of hydrogen atoms, and weighting scheme.
- (3) Final values of R and R_w (and their definitions), with a statement of whether unobserved reflections were included (and if so, how), and if possible some estimate of the accuracy of the F_o data.
- (4) Programs or packages and computers used (with references) and source of f data (also f', f'').
 - (5) Range of standard deviations for bond lengths and angles.
- (6) Any procedures used for definition of chirality (quotation of a single Hamilton ratio without specifying the R values from which the ratio was derived, and the conditions under which these were obtained, will not normally be acceptable).

When the analysis has not been of a routine nature, authors should explain concisely all procedures used.

5.5 Example of Presentation

The following example demonstrates the application of the recommendations in the preceding sections (5.2 to 5.4):

Experimental

Crystals were prepared as described in ref. 11 and sealed under nitrogen in Lindemann capillaries.

Crystal Data.— $C_{24}H_{26}Cl_3Re_3Si_6$, $M=1\,028.0$. Monoclinic, a=12.021(3), b=20.489(5), c=18.194(5) Å, $\beta=91.03(3)^\circ$, $V=4\,480$ Å³ (by least-squares refinement on diffractometer angles for 15 automatically centred reflections, $\lambda=0.710\,69$ Å), space group $P2_1/n$ (alt. $P2_1/c$, No. 14), Z=4, $D_x=1.76$ g cm⁻³. Dark blue, air-sensitive tablets. Crystal dimensions (distance to faces from centre):

 $0.313(211,\overline{2}\overline{1}\overline{1}) \times 0.163(010,\overline{0}\overline{1}\overline{0}) \times 0.063(10t,\overline{1}\overline{0}\overline{1})$ mm, $\mu(\text{Mo-K}_{\pi}) = 85.12 \text{ cm}^{-1}$.

Data Collection and Processing. ¹²—CAD4 diffractometer, $\omega/2\theta$ mode with ω scan width = 0.85 + 0.35 tan θ , ω scan speed 1.3—6.8 deg min⁻¹, graphite-monochromated Mo- K_{α} radiation; 6 383 reflections measured (1.5 $\leq \theta \leq 25^{\circ}$, +h,k,l), 5 797 unique [merging R=0.024 after absorption correction (max., min. transmission factors = 0.37, 0.10)], giving 4 170 with $I > 2\sigma(I)$. Linear and approx. isotropic crystal decay, ca. 37%, corrected during processing.

Structure Analysis and Refinement.—Direct methods (Re and Cl atoms) followed by normal heavy atom procedures. Full-matrix least squares refinement (in two blocks for final, anisotropic cycles) with all non-hydrogen atoms anisotropic and hydrogens in calculated positions with one, overall, refined $\mu_{\rm iso}$ [= 0.08(2) Å²). The weighting scheme $w = 1/[\sigma^2(F_{\rm o}) + 0.0078F_{\rm o}^2]$, with $\sigma(F_{\rm o})$ from counting statistics ¹² gave satisfactory agreement analyses. Final R and $R_{\rm w}$ values are 0.062, 0.064. Programs and computers used and sources of scattering factor data are given in ref. 12.

References

- 11 Ref. to prep of compound.
- 12 For full details of experimental techniques used see ref. to previous paper giving full details of data collection procedures *etc.* used in author's laboratory.

5.6 Presentation of Tables and Diagrams

A clear distinction should be made between material to appear in print and material for deposition.

- 5.6.1 For Publication in the Journal.—(1) A table of final fractional atomic co-ordinates (labelled x, y, z) must be included. If the origin chosen differs from that adopted in International Tables this should be mentioned and justified. Where the asymmetric unit consists of a discrete molecule (or molecules) co-ordinates should refer to atoms which are all in the same molecule. Hydrogen atom co-ordinates should be included only when experimentally determined or refined; when they have been determined only by calculation, deposition is appropriate.
- (2) A table of selected bond lengths and angles, with estimated standard deviations. This should be restricted to significant dimensions only (for example it is rarely necessary to include data for phenyl rings). Average values may be given (with a range of e.s.d.s) for chemically equivalent groups or for similar bonds. As an alternative to tabular presentation it is often clearer to give important dimensions in a structural diagram. Differences from expected norms should be noted.
- (3) A conventional *line drawing* of the structure must be included except in the simplest cases and one *perspective diagram* (or *stereo pair*) if appropriate. Packing diagrams should not be included unless required to illustrate a specific chemical point. The *atom numbering system* should be clearly shown in one of the diagrams. Any differences from that required by standard rules of chemical nomenclature should be pointed out. Each atom of the asymmetric unit should be assigned an arabic numeral in parentheses following the chemical symbol: C(2), O(1"), etc.; it is often convenient to associate a particular number of primes with a particular

asymmetric unit. Alternatively, roman numeral superscripts can be employed: $C(2^I) \cdot \cdot \cdot C(2^{IV})$.

- 5.6.2 For the Referees and/or for Deposition.—(1) Any calculated co-ordinates (e.g. hydrogen).
- (2) A full list of bond lengths and angles with estimated standard deviations.
- (3) A full list of thermal parameters in the form B_{ij} or U_{ij} (in \mathring{A}^2 or pm²).
- (4) Details of any *least squares planes* used to provide parameters for the paper.
- (5) A legible table of structure factors (F_0, F_c) (for the referees only; not for deposition).*
- 5.6.3 Deposition of Material at the Cambridge Crystallographic Data Centre.—The table of final fractional atomic coordinates, the full list of bond lengths and angles, and the list of thermal parameters will be deposited at the Cambridge Crystallographic Data Centre, University Chemical Laboratory, Lensfield Road, Cambridge CB2 1EW (or at the Institut für Anorganische Chemie, Universität, Gerhard-Domagk-Str. 1. D-5300 Bonn, via Prof. Dr. G. Bergerhoff, for molecules not containing 'organic' carbon). All Tables should be of publication quality. Computer printout is acceptable provided that it is clearly legible. The print program used should yield concise tables of atomic and other derived parameters. The content of the tables should be non-redundant, and their interpretation immediately obvious. The thermal parameters should be submitted as separate tables, with clear definition of the units used. Any request to Cambridge or to Bonn for deposited material should be accompanied by the full literature citation for the paper concerned.

5.7 Reference to Crystallographic Work Published in Chemical Communications

It is permissible to regard a fully refined crystal structure determination published in *Chem. Comm.* as archival material. If an author does not wish to discuss the structure again at any length in the corresponding full paper, his purpose will be served by a simple reference back to the original communication, and he need not re-present the associated data for publication or for the referees. However if these conditions are not fulfilled, the data should be re-presented and will be re-published if considered necessary.

5.8 Reference to Unpublished Crystallographic Work

There may be cases (other than that just described) when an author wishes to publish a paper in which the result of a crystal structure determination is discussed, but where he/she does not wish to include details or extensive discussion. He/she may not even wish to include the crystallographer as co-author (for example when the determination is carried out by a commercial company). If the author is able to show the referees that this procedure is appropriate, it will be allowed provided that it does not lead to unnecessary fragmentation. However the author must provide, as supplementary information, sufficient data relating to the crystal structure determination to allow a referee to make sure that the point made is correct, and co-ordinates etc. will be deposited at the Cambridge Crystallographic Data Centre (or Bonn). The brief published description of the

^{*} Authors should preserve copies of structure factor tables in their own

determination should be supplemented by appropriate reference to 'unpublished work'.

6.0 Publication of Theoretical and Computational Papers

Authors should note the following guidelines for the preparation of computational papers, so that the material can be presented concisely and effectively.

- (i) Papers should be submitted to the appropriate journal: a paper containing innovations in theory to Faraday Transactions II, one in which the computations are incidental to the chemistry to Perkin, Dalton, or Faraday Transactions I. Papers concerned mainly with computational details are unlikely to be accepted.
- (ii) The purpose of the paper and the precise objectives of the calculations performed should be clearly stated; the results obtained should be reported only in so far as they relate to those objectives.
 - (iii) Many papers use a routine procedure based on a well

documented method, be it semi-empirical or *ab initio*. It is then sufficient to name the particular variant, referring to key papers in which the method was developed, to cite the computer program used, and to indicate *briefly* any modification made by the author. A review of theoretical background would be out of place, but an author should say why he considers the method adequate for his purposes.

(iv) Extensive tabulation of numerical results, such as the magnitudes of atomic orbital coefficients, electron populations, contour maps of molecular orbitals and electron densities, and peripheral material of a similar nature, is normally unnecessary. Lengthy line-by-line discussion of such material is, as a general rule, unacceptable. Where an author considers that there is a special need to make such material available to other workers, as with highly accurate computations, for example, then this may be deposited with the British Library as a Supplementary Publication. Such material should be submitted with the main paper, clearly distinguished from it, and referred to in the main text.

IUPAC Publications on Nomenclature and Symbolism

1.0 Compilations

- 1.1 Nomenclature of Organic Chemistry, a 550-page hard-cover volume published in 1979, available from Pergamon, Oxford.
 - Section A: Hydrocarbons
 - Section B: Fundamental heterocyclic systems
 - Section C: Characteristic groups containing carbon, hydrogen, oxygen, nitrogen, halogen, sulphur, selenium, and tellurium
 - Section D: Organic compounds containing elements not exclusively those referred to in the title of Section C
 - Section E: Stereochemistry
 - Section F: General principles for the naming of natural products and related compounds
 - Section H: Isotopically modified compounds
- 1.2 Nomenclature of Inorganic Chemistry, a 110-page hardcover volume published in 1970, available from Pergamon, Oxford.
 - Chapter 1: Elements
 - Chapter 2: Formulae and names of compounds in general
 - Chapter 3: Names for ions and radicals
 - Chapter 4: Iso- and hetero-polyanions
 - Chapter 5: Acids
 - Chapter 6: Salts and salt-like compounds
 - Chapter 7: Co-ordination compounds
 - Chapter 8: Addition compounds
 - Chapter 9: Crystalline phases of variable composition
 - Chapter 10: Polymorphism
 - Chapter 11: Boron compounds
- 1.3 Biochemical Nomenclature and Related Documents, a 220-page softcover manual published in 1978 by The Biochemical Society for IUB, and available from the Biochemical Society Book Depot, P.O. Box 32, Commerce Way, Colchester, Essex CO2 8HP. The contents are as follows:

General

Nomenclature of organic chemistry. Section E: Stereochemistry (1974)

Nomenclature of organic chemistry. Section F: Natural products and related compounds (1976)

Nomenclature of organic chemistry. Section H: Isotopically modified compounds (1977)

Isotopically labelled compounds: common biochemical practice

Recommendations for measurement and presentation of biochemical equilibrium data (1976)

Abbreviations and symbols for chemical names of special interest in biological chemistry (1965)

Abbreviations and symbols: a compilation (1976)

Citation of bibliographic references in biochemical journals (1971)

Amino acids, peptides and proteins

Nomenclature of α -amino acids (1974)

Symbols for amino-acid derivatives and peptides (1971)

Rules for naming synthetic modifications of natural peptides (1966)

Abbreviated nomenclature of synthetic polypeptides or polymerized amino acids (1971)

A one-letter notation for amino-acid sequences (1968)

Abbreviations and symbols for the description of the conformation of polypeptide chains (1969)

Nomenclature of peptide hormones (1974)

Recommendations for the nomenclature of human immunoglobulins

Protein data bank. A computer-based archival file for macromolecular structures (1977)

Nomenclature of multiple forms of enzymes (1976)

Nucleotides and nucleic acids

Abbreviations and symbols for nucleic acids, polynucleotides and their constituents (1970)

Lipids

Nomenclature of lipids (1976)

Nomenclature of steroids (1967)

Nomenclature of quinones with isoprenoid side chains (1973)

Tentative rules for the nomenclature of carotenoids (1970). Amendments (1974)

Nomenclature of tocopherols and related compounds (1973)

Carbohydrates, etc.

Tentative rules for carbohydrate nomenclature. Part 1 (1969)

Nomenclature of cyclitols (1973)

Phosphorus-containing compounds

Nomenclature of phosphorus-containing compounds of biochemical importance (1976)

Miscellaneous

Trivial names of miscellaneous compounds of importance in biochemistry (1965)

Nomenclature and symbols for folic acids and related compounds (1965)

Nomenclature for vitamins B-6 and related compounds (1973)

Nomenclature of corrinoids (1973)

- 1.4 Compendium of Analytical Nomenclature, a 222-page volume published in 1978, available in hardcover and softcover from Pergamon, Oxford.
 - Chapter 1: Recommendations for the presentation of the results of chemical analysis
 - Chapter 2: Recommendations for terminology to be used with precision balances
 - Chapter 3: Recommended nomenclature for scales of working in analysis
 - Chapter 4: Recommendations on nomenclature for contamination phenomena in precipitation from aqueous solution
 - Chapter 5: Recommended nomenclature for automatic analysis
 - Chapter 6: Recommendations for nomenclature of thermal analysis
 - Chapter 7: Recommendations for nomenclature of mass spectrometry

Chapter 8: Recommended nomenclature for titrimetric analysis

Chapter 9: Report on the standardization of pH and related technology

Chapter 10: Practical measurements of pH in amphiprotic and mixed solvents

Chapter 11: Recommended symbols for solution equilibria

Chapter 12: Recommended nomenclature for liquidliquid distribution

Chapter 13: Recommendations on nomenclature and presentation of data in gas chromatography

Chapter 14: Recommendations on nomenclature for chromatography

Chapter 15: Recommendations on ion-exchange nomenclature

Chapter 16–18: Nomenclature, symbols, units and their usage in spectrochemical analysis. I, General atomic emission spectroscopy. II, Data interpretation. III, Analytical flame spectroscopy and associated non-flame procedures

Chapter 19: Classification and nomenclature of electroanalytical techniques

Chapter 20: Recommendations for sign conventions and plotting of electrochemical data

Chapter 21: Recommendations for nomenclature of ion-selective electrodes

Appendix: Recommendations on the usage of the terms 'equivalent' and 'normal'

1.5 Compendium of Chemical Terminology: IUPAC Recommendations, a 456-page volume published in 1987, available in hardcover and softcover from Blackwells, Oxford.

2.0 Documents not included in the compilations

2.1 Nomenclature of Elements and Compounds

2.1.1 Amino acids and Peptides

Nomenclature and symbolism for amino acids and peptides (Pure Appl. Chem., 1984, 56, 595; Eur. J. Biochem., 1984, 138, 9). 2.1.2 Analytical Reagents

Guide to trivial names, trade names, and synonyms for substances used in analytical chemistry (*Pure Appl. Chem.*, 1978, **50**, 339).

2.1.3 Boron Compounds

Nomenclature of inorganic boron compounds (*Pure Appl. Chem.*, 1972, **30**, 681).

2.1.4 Carbohydrates

Conformational nomenclature for five- and six-membered ring forms of monosaccharides and their derivatives (provisional) (*Pure Appl. Chem.*, 1981, 53, 1901; *Eur. J. Biochem.*, 1980, 111, 295).

Abbreviated terminology of oligosaccharide chains (provisional) (*Pure Appl. Chem.*, 1982, **54**, 1517; *J. Biol. Chem.*, 1982, **257**, 2347.

Polysaccharide nomenclature (provisional) (*Pure Appl. Chem.*, 1982, **54**, 1523; *J. Biol. Chem.*, 1982, **257**, 3352).

Nomenclature of unsaturated monosaccharides (provisional) (*Pure Appl. Chem.*, 1982, **54**, 207; *Eur. J. Biochem.*, 1981, **119**, 1; errata *Eur. J. Biochem.*, 1982, **125**, 1).

Nomenclature of branched-chain monosaccharides (provisional) (*Pure Appl. Chem.*, 1982, **54**, 211; *Eur. J. Biochem.*, 1981, **119**, 5; errata *Eur. J. Biochem.*, 1982, **125**, 1).

Symbols for specifying the conformation of polysaccharide chains (provisional) (*Pure Appl. Chem.*, 1983, **55**, 1269; *Eur. J. Biochem.*, 1983, **131**, 5).

2.1.5 Elements

Recommendations for the names of elements of atomic number greater than 100 (*Pure Appl. Chem.*, 1979, **51**, 381).

2.1.6 Enzymes

Enzyme Nomenclature (1984), published by Academic Press in hardcover and softcover editions.

2.1.7 Folic Acid

Nomenclature and symbols for folic acid and related compounds (*Pure Appl. Chem.*, 1987, **59**, 833).

2.1.8 Heterocyclic Compounds

Revision of the extended Hantzsch-Widman system of nomenclature for heteromonocycles (*Pure Appl. Chem.*, 1983, 55, 409).

2.1.9 Isotopically Modified Compounds

Nomenclature of inorganic chemistry. Part II. 1. Isotopically modified compounds (*Pure Appl. Chem.*, 1981, **53**, 1887).

2.1.10 Lambda Convention

Treatment of variable valence in organic nomenclature (*Pure Appl. Chem.*, 1984, **56**, 769).

2.1.11 Nitrogen Hydrides

Nomenclature of hydrides of nitrogen and derived cations, anions, and ligands (*Pure Appl. Chem.*, 1982, **54**, 2545).

2.1.12 Nucleotides

Abbreviations and symbols for the description of conformations of polynucleotide chains (provisional) (*Pure Appl. Chem.*, 1983, 55, 1279; *Eur. J. Biochem.*, 1983, 131, 9).

2.1.13 Numerical Terms

Extension of Rules A-1.1 and A-2.5 concerning numerical terms used in organic chemical nomenclature (*Pure Appl. Chem.*, 1986, 58, 1693).

2.1.14 Polymers

Nomenclature of regular single-strand organic polymers (*Pure Appl. Chem.*, 1976, **48**, 373).

Nomenclature for regular single-strand and quasi single-strand inorganic and co-ordination polymers (*Pure Appl. Chem.*, 1985, 57, 149).

Source-based nomenclature for copolymers (*Pure Appl. Chem.*, 1985, **57**, 1427).

Stereochemical definitions and notations relating to polymers (*Pure Appl. Chem.*, 1981, 53, 733).

Use of abbreviations for names of polymeric substances (*Pure Appl. Chem.*, 1987, **59**, 691).

Basic definitions of terms relating to polymers (*Pure Appl. Chem.*, 1975, **40**, 477).

2.1.15 Prenols

Nomenclature of prenols (*Pure Appl. Chem.*, 1987, **59**, 683; *Eur. J. Biochem.*, 1987, **167**, 181).

2.1.16 Retinoids

Nomenclature of retinoids (provisional) (Pure Appl. Chem., 1983, 55, 721; Eur. J. Biochem., 1982, 129, 1).

2.1.17 Tetrapyrroles

Nomenclature of tetrapyrroles (Pure Appl. Chem., 1987, 59, 779).

2.1.18 Tocopherols

Nomenclature of tocopherols and related compounds (*Pure Appl. Chem.*, 1982, **54**, 1507; *Eur. J. Biochem.*, 1982, **123**, 473). 2.1.19 *Vitamins*

Nomenclature of Vitamin D (provisional) (Pure Appl. Chem., 1982, 54, 1511; Eur. J. Biochem., 1982, 124, 223).

2.1.20 Zeolites

Chemical nomenclature and formulation of compositions of synthetic and natural zeolites (*Pure Appl. Chem.*, 1979, **51**, 1091).

2.2 Terminology, Symbols, and Units, and Presentation of Results

2.2.1 General

Glossary of terms used in physical organic chemistry (*Pure Appl. Chem.*, 1983, **55**, 1281).

Manual of symbols and terminology for physicochemical quantities and units (*Pure Appl. Chem.*, 1979, **51**, 1, also available from Pergamon, Oxford, as a 40-page softcover booklet). 2.2.2 *Analytical*

Nomenclature, symbols, units, and their usage in spectrochemical analysis. Part IV, X-Ray emission spectroscopy (*Pure Appl. Chem.*, 1980, **52**, 2543). Part V, Radiation sources (*Pure Appl. Chem.*, 1985, **57**, 1453). Part VI, Molecular luminescence spectroscopy (*Pure Appl. Chem.*, 1984, **55**, 231).

Recommendations for nomenclature, standard procedures, and reporting of experimental data for surface analysis techniques (*Pure Appl. Chem.*, 1979, **51**, 2243).

Glossary of terms used in nuclear analytical chemistry (provisional) (*Pure Appl. Chem.*, 1982, **54**, 1533).

Recommendations for publication of papers on a new analytical method based on ion exchange or ion-exchange chromatography (*Pure Appl. Chem.*, 1980, **52**, 2555).

Recommendations for presentation of data on compleximetric indicators. 1. General (*Pure Appl. Chem.*, 1979, **51**, 1357).

Recommendations for publishing manuscripts on ion-selective electrodes (*Pure Appl. Chem.*, 1981, 53, 1907).

Recommendations on use of the term amplification reactions (*Pure Appl. Chem.*, 1982, **54**, 2553).

Recommendations for the usage of selective, selectivity, and related terms in analytical chemistry (*Pure Appl. Chem.*, 1983, 55, 553).

Proposed terminology and symbols for the transfer of solutes from one solvent to another (*Pure Appl. Chem.*, 1978, **50**, 589). Nomenclature, symbols and units recommended for *in situ* microanalysis (provisional) (*Pure Appl. Chem.*, 1983, **55**, 2023).

Physicochemical quantities and units in clinical chemistry with special emphasis on activities and activity coefficients (*Pure Appl. Chem.*, 1984, **56**, 567).

Quantities and units in clinical chemistry (*Pure Appl. Chem.*, 1979, **51**, 2451).

Quantities and units in clinical chemistry: nebulizer and flame properties in flame emission and absorption spectrometry (*Pure Appl. Chem.*, 1986, **58**, 1737).

List of quantities in clinical chemistry (*Pure Appl. Chem.*, 1979, 51, 2481).

2.2.4 Colloids and Surface Chemistry

Definitions, terminology, and symbols in colloid and surface chemistry. I (*Pure Appl. Chem.*, 1972, 31, 577). II, Heterogeneous catalysis (*Pure Appl. Chem.*, 1976, 46, 71). Part 1.14: Light scattering (provisional) (*Pure Appl. Chem.*, 1983, 55, 931).

Reporting experimental pressure-area data with film balances (Pure Appl. Chem., 1985, 57, 621).

Reporting physisorption data for gas/solid systems with special reference to the determination of surface area and porosity (*Pure Appl. Chem.*, 1985, 57, 603).

Reporting data on adsorption from solution at the solid/solution interface (*Pure Appl. Chem.*, 1986, **58**, 967).

2.2.5 Electrochemistry

Nomenclature for transfer phenomena in electrolytic systems (*Pure Appl. Chem.*, 1981, 53, 1827).

Electrode reaction orders, transfer coefficients, and rate

constants—amplification of definitions and recommendations for publication of parameters (*Pure Appl. Chem.*, 1980, **52**, 233). Recommended terms, symbols, and definitions for electroanalytical chemistry (*Pure Appl. Chem.*, 1985, **57**, 1491).

Classification and nomenclature of electroanalytical techniques (*Pure Appl. Chem.*, 1976, **45**, 81).

Recommendations for sign conventions and plotting of electrochemical data (*Pure Appl. Chem.*, 1976, **45**, 131).

Electrochemical nomenclature (*Pure Appl. Chem.*, 1974, 37, 499). Recommendations on reporting electrode potentials in non-aqueous solvents (*Pure Appl. Chem.*, 1984, 56, 461).

Definition of pH scales, standard reference values, measurement of pH and related terminology (*Pure Appl. Chem.*, 1985, 57, 531).

Interphases in systems of conducting phases (*Pure Appl. Chem.*, 1986, **58**, 437).

The absolute electrode potential: an explanatory note (Pure Appl. Chem., 1986, 58, 955).

2.2.6 Kinetics

Symbolism and terminology in chemical kinetics (provisional) (*Pure Appl. Chem.*, 1981, **53**, 753).

2.2.7 Photochemistry

Recommended standards for reporting photochemical data (*Pure Appl. Chem.*, 1984, **56**, 939).

2.2.8 Quantum Chemistry

Expression of results in quantum chemistry (*Pure Appl. Chem.*, 1978, **50**, 75).

2.2.9 Reactions

Nomenclature for straightforward transformations (provisional) (Pure Appl. Chem., 1981, 53, 306).

2.2.10 Rheological Properties

Selected definitions, terminology, and symbols for rheological properties (*Pure Appl. Chem.*, 1979, **51**, 1215).

2.2.11 Spectroscopy

Recommendations for publication of papers on methods of molecular absorption spectrophotometry in solution (*Pure Appl. Chem.*, 1978, **50**, 237).

Nomenclature and spectral presentation in electron spectroscopy resulting from excitation by photons (*Pure Appl. Chem.*, 1976, **45**, 221).

Recommendations for the presentation of infrared absorption spectra in data collections. A, Condensed phases (*Pure Appl. Chem.*, 1978, **50**, 231).

Definition and symbolism of molecular force constants (*Pure Appl. Chem.*, 1978, **50**, 1709).

Recommendations for symbolism and nomenclature for mass spectrometry (*Pure Appl. Chem.*, 1978, **50**, 65).

Nomenclature and conventions for reporting Mössbauer spectroscopic data (*Pure Appl. Chem.*, 1976, 45, 211).

Recommendations for the presentation of NMR data for publication in chemical journals. A, Proton spectra (*Pure Appl. Chem.*, 1972, 39, 625). B, Spectra from nuclei other than protons (*Pure Appl. Chem.*, 1976, 45, 217).

Presentation of Raman spectra in data collections (*Pure Appl. Chem.*, 1981, **53**, 1879).

Names, symbols, definitions and units of quantities in optical spectroscopy (*Pure Appl. Chem.*, 1985, **57**, 105).

2.2.12 Thermal Analysis

Nomenclature of thermal analysis. I (Pure Appl. Chem., 1974, 37, 439). II, DTA and TG apparatus and technique. III, DTA and TG curves (Pure Appl. Chem., 1980, 52, 2387). IV (Pure Appl. Chem., 1985, 57, 1737).

Calorimetric measurements on cellular systems: recommendations for measurements and presentation of results (provisional) (*Pure Appl. Chem.*, 1982, **54**, 671).

2.2.13 Thermodynamics

A guide to procedures for the publication of thermodynamic data (*Pure Appl. Chem.*, 1972, 39, 395).

Assignment and presentation of uncertainties of the numerical results of thermodynamic measurements (*Pure Appl. Chem.*, 1981, 53, 1805).

Notation for states and processes; significance of the word 'standard' in chemical thermodynamics and remarks on commonly tabulated forms of thermodynamic functions (*Pure Appl. Chem.*, 1982, **54**, 1239).