

INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY

Physical and Biophysical Chemistry Division



# Quantities, Units and Symbols in Physical Chemistry

Third Edition

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## PREFACE

The purpose of this manual is to improve the exchange of scientific information among the readers in different disciplines and across different nations. As the volume of scientific literature expands, each discipline has a tendency to retreat into its own jargon. This book attempts to provide a readable compilation of widely used terms and symbols from many sources together with brief understandable definitions. This Third Edition reflects the experience of the contributors with the previous editions and we are grateful for the many thoughtful comments we have received. Most of the material in this book is “standard”, but a few definitions and symbols are not universally accepted. In such cases, we have attempted to list acceptable alternatives. The references list the reports from IUPAC and other sources in which some of these notational problems are discussed further. IUPAC is the acronym for International Union of Pure and Applied Chemistry.

A spectacular example of the consequences of confusion of units is provided by the loss of the United States NASA satellite, the “Mars Climate Orbiter” (MCO). The Mishap Investigation Board (Phase I Report, November 10, 1999)<sup>1</sup> found that the root cause for the loss of the MCO was “the failure to use metric units in the coding of the ground (based) software file”. The impulse was reported in Imperial units of pounds (force)-seconds (lbf-s) rather than in the metric units of Newton (force)-seconds (N-s). This caused an error of a factor of 4.45 and threw the satellite off course.<sup>2</sup> We urge the users of this book always to define explicitly the terms, the units, and the symbols that they use.

This edition has been compiled in machine-readable form by Martin Quack and Jürgen Stohner. The entire text of the manual will be available on the Internet some time after the publication of the book and will be accessible via the IUPAC web site, <http://www.iupac.org>. Suggestions and comments are welcome and may be addressed in care of the

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Corrections to the manual will be listed periodically.

The book has been systematically brought up to date and new sections have been added. As in previous editions, the first chapter describes the use of quantity calculus for handling physical quantities and the general rules for the symbolism of quantities and units and includes an expanded description on the use of roman and italic fonts in scientific printing. The second chapter lists the symbols for quantities in a wide range of topics used in physical chemistry. New parts of this chapter include a section on surface structure. The third chapter describes the use of the International System of units (SI) and of a few other systems such as atomic units. Chapter 4 outlines mathematical symbols and their use in print. Chapter 5 presents the 2006 revision of the fundamental physical constants, and Chapter 6 the properties of elementary particles, elements and nuclides. Conversion of units follows in Chapter 7, together with the equations of electricity and magnetism in their various forms. Chapter 8 is entirely new and outlines the treatment of uncertainty in physical measurements. Chapter 9 lists abbreviations and acronyms. Chapter 10 provides the references, and Chapter 11, the Greek alphabet. In Chapters 12 and 13, we end with indexes. The IUPAC Periodic Table of the Elements is shown on the inside back cover and the preceding pages list frequently used conversion factors for pressure and energy units.

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<sup>1</sup> The MCO report can be found at [ftp://ftp.hq.nasa.gov/pub/pao/reports/1999/MCO\\_report.pdf](ftp://ftp.hq.nasa.gov/pub/pao/reports/1999/MCO_report.pdf).

<sup>2</sup> Impulse (change of momentum) means here the time-integral of the force.

Many people have contributed to this volume. The people most directly responsible are acknowledged in the Historical Introduction. Many of the members of IUPAC I.1 have continued to make active contributions long after their terms on the Commission expired. We also wish to acknowledge the members of the other Commissions of the Physical Chemistry Division: Chemical Kinetics, Colloid and Surface Chemistry, Electrochemistry, Spectroscopy, and Thermodynamics, who have each contributed to the sections of the book that concern their various interests.

We also thank all those who have contributed whom we have inadvertently missed out of these lists.

*Commission on Physicochemical Symbols,  
Terminology and Units*

Jeremy G. Frey and Herbert L. Strauss

## HISTORICAL INTRODUCTION

The *Manual of Symbols and Terminology for Physicochemical Quantities and Units* [1.a], to which this is a direct successor, was first prepared for publication on behalf of the Physical Chemistry Division of IUPAC by M. L. McGlashan in 1969, when he was chairman of the Commission on Physicochemical Symbols, Terminology and Units (I.1). He made a substantial contribution towards the objective which he described in the preface to that first edition as being “to secure clarity and precision, and wider agreement in the use of symbols, by chemists in different countries, among physicists, chemists and engineers, and by editors of scientific journals”. The second edition of that manual prepared for publication by M. A. Paul in 1973 [1.b], and the third edition prepared by D. H. Whiffen in 1976 [1.c], were revisions to take account of various developments in the *Système International d’Unités* (International System of Units, international abbreviation SI), and other developments in terminology.

The first edition of *Quantities, Units and Symbols in Physical Chemistry* published in 1988 [2.a] was a substantially revised and extended version of the earlier manuals. The decision to embark on this project originally proposed by N. Kallay was taken at the IUPAC General Assembly at Leuven in 1981, when D. R. Lide was chairman of the Commission. The working party was established at the 1983 meeting in Lyngby, when K. Kuchitsu was chairman, and the project has received strong support throughout from all present and past members of the Commission I.1 and other Physical Chemistry Commissions, particularly D. R. Lide, D. H. Whiffen, and N. Sheppard.

The extensions included some of the material previously published in appendices [1.d–1.k]; all the newer resolutions and recommendations on units by the *Conférence Générale des Poids et Mesures* (CGPM); and the recommendations of the International Union of Pure and Applied Physics (IUPAP) of 1978 and of Technical Committee 12 of the International Organization for Standardization, *Quantities, units, symbols, conversion factors* (ISO/TC 12). The tables of physical quantities (Chapter 2) were extended to include defining equations and SI units for each quantity. The style was also slightly changed from being a book of rules towards a manual of advice and assistance for the day-to-day use of practicing scientists. Examples of this are the inclusion of extensive notes and explanatory text inserts in Chapter 2, the introduction to quantity calculus, and the tables of conversion factors between SI and non-SI units and equations in Chapter 7.

The second edition (1993) was a revised and extended version of the previous edition. The revisions were based on the recent resolutions of the CGPM [3]; the new recommendations by IUPAP [4]; the new international standards ISO 31 [5,6]; some recommendations published by other IUPAC commissions; and numerous comments we have received from chemists throughout the world. The revisions in the second edition were mainly carried out by Ian Mills and Tom Cvitaš with substantial input from Robert Alberty, Kozo Kuchitsu, Martin Quack as well as from other members of the IUPAC Commission on Physicochemical Symbols, Terminology and Units.

The manual has found wide acceptance in the chemical community, and various editions have been translated into Russian [2.c], Hungarian [2.d], Japanese [2.e], German [2.f], Romanian [2.g], Spanish [2.h], and Catalan [2.i]. Large parts of it have been reproduced in the 71st and subsequent editions of the *Handbook of Chemistry and Physics* published by CRC Press.

The work on revisions of the second edition started immediately after its publication and between 1995 and 1997 it was discussed to change the title to “Physical-Chemical Quantities, Units and Symbols” and to apply rather complete revisions in various parts. It was emphasized that the book covers as much the field generally called “physical chemistry” as the field called “chemical physics”. Indeed we consider the larger interdisciplinary field where the boundary between physics and chemistry has largely disappeared [10]. At the same time it was decided to produce the whole book as a text file in computer readable form to allow for future access directly by computer, some

time after the printed version would be available. Support for this decision came from the IUPAC secretariat in the Research Triangle Park, NC (USA) (John W. Jost). The practical work on the revisions was carried out at the ETH Zürich, while the major input on this edition came from the group of editors listed now in full on the cover. It fits with the new structure of IUPAC that these are defined as project members and not only through membership in the commission. The basic structure of this edition was finally established at a working meeting of the project members in Engelberg, Switzerland in March 1999, while further revisions were discussed at the Berlin meeting (August 1999) and thereafter. In 2001 it was decided finally to use the old title. In this edition the whole text and all tables have been revised, many chapters substantially. This work was carried out mainly at ETH Zürich, where Jürgen Stohner coordinated the various contributions and corrections from the current project group members and prepared the print-ready electronic document. Larger changes compared to previous editions concern a complete and substantial update of recently available improved constants, sections on uncertainty in physical quantities, dimensionless quantities, mathematical symbols and numerous other sections.

At the end of this historical survey we might refer also to what might be called the tradition of this manual. It is not the aim to present a list of recommendations in form of commandments. Rather we have always followed the principle that this manual should help the user in what may be called “good practice of scientific language”. If there are several well established uses or conventions, these have been mentioned, giving preference to one, when this is useful, but making allowance for variety, if such variety is not harmful to clarity. In a few cases possible improvements of conventions or language are mentioned with appropriate reference, even if uncommon, but without specific recommendation. In those cases where certain common uses are deprecated, there are very strong reasons for this and the reader should follow the corresponding advice.

Zürich, 2007

Martin Quack

The membership of the Commission during the period 1963 to 2006, during which the successive editions of this manual were prepared, was as follows:

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### *National representatives*

Numerous national representatives have served on the commission over many years. We do not provide this long list here.



# 1 PHYSICAL QUANTITIES AND UNITS



## 1.1 PHYSICAL QUANTITIES AND QUANTITY CALCULUS

The value of a *physical quantity*  $Q$  can be expressed as the product of a *numerical value*  $\{Q\}$  and a *unit*  $[Q]$

$$Q = \{Q\} [Q] \quad (1)$$

Neither the name of the physical quantity, nor the symbol used to denote it, implies a particular choice of unit (see footnote <sup>1</sup>, p. 4).

Physical quantities, numerical values, and units may all be manipulated by the ordinary rules of algebra. Thus we may write, for example, for the wavelength  $\lambda$  of one of the yellow sodium lines

$$\lambda = 5.896 \times 10^{-7} \text{ m} = 589.6 \text{ nm} \quad (2)$$

where m is the symbol for the unit of length called the metre (or meter, see Sections 3.2 and 3.3, p. 86 and 87), nm is the symbol for the nanometre, and the units metre and nanometre are related by

$$1 \text{ nm} = 10^{-9} \text{ m} \quad \text{or} \quad \text{nm} = 10^{-9} \text{ m} \quad (3)$$

The equivalence of the two expressions for  $\lambda$  in Equation (2) follows at once when we treat the units by the rules of algebra and recognize the identity of 1 nm and  $10^{-9}$  m in Equation (3). The wavelength may equally well be expressed in the form

$$\lambda/\text{m} = 5.896 \times 10^{-7} \quad (4)$$

or

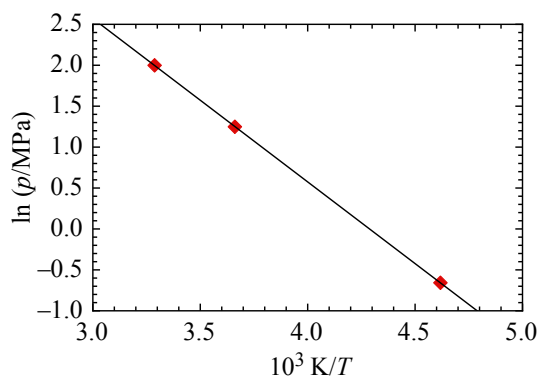
$$\lambda/\text{nm} = 589.6 \quad (5)$$

It can be useful to work with variables that are defined by dividing the quantity by a particular unit. For instance, in tabulating the numerical values of physical quantities or labeling the axes of graphs, it is particularly convenient to use the quotient of a physical quantity and a unit in such a form that the values to be tabulated are numerical values, as in Equations (4) and (5).

*Example*

$$\begin{aligned} \ln(p/\text{MPa}) &= a + b/T = \\ &= a + b'(10^3 \text{ K}/T) \end{aligned} \quad (6)$$

| $T/\text{K}$ | $10^3 \text{ K}/T$ | $p/\text{MPa}$ | $\ln(p/\text{MPa})$ |
|--------------|--------------------|----------------|---------------------|
| 216.55       | 4.6179             | 0.5180         | -0.6578             |
| 273.15       | 3.6610             | 3.4853         | 1.2486              |
| 304.19       | 3.2874             | 7.3815         | 1.9990              |



Algebraically equivalent forms may be used in place of  $10^3 \text{ K}/T$ , such as  $\text{kK}/T$  or  $10^3 (T/\text{K})^{-1}$ . Equations between numerical values depend on the choice of units, whereas equations between quantities have the advantage of being independent of this choice. Therefore the use of equations between quantities should generally be preferred.

The method described here for handling physical quantities and their units is known as *quantity calculus* [11–13]. It is recommended for use throughout science and technology. The use of quantity calculus does not imply any particular choice of units; indeed one of the advantages of quantity calculus is that it makes changes between units particularly easy to follow. Further examples of the use of quantity calculus are given in Section 7.1, p. 131, which is concerned with the problems of transforming from one set of units to another.

## 1.2 BASE QUANTITIES AND DERIVED QUANTITIES

By convention physical quantities are organized in a dimensional system built upon seven *base quantities*, each of which is regarded as having its own dimension. These base quantities in the International System of Quantities (ISQ) on which the International System of units (SI) is based, and the principal symbols used to denote them and their dimensions are as follows:

| <i>Base quantity</i>      | <i>Symbol for quantity</i> | <i>Symbol for dimension</i> |
|---------------------------|----------------------------|-----------------------------|
| length                    | $l$                        | L                           |
| mass                      | $m$                        | M                           |
| time                      | $t$                        | T                           |
| electric current          | $I$                        | I                           |
| thermodynamic temperature | $T$                        | Θ                           |
| amount of substance       | $n$                        | N                           |
| luminous intensity        | $I_v$                      | J                           |

All other quantities are called *derived quantities* and are regarded as having dimensions derived algebraically from the seven base quantities by multiplication and division.

*Example*      dimension of energy is equal to dimension of  $M L^2 T^{-2}$   
This can be written with the symbol dim for dimension (see footnote <sup>1</sup>, below)  
 $\dim(E) = \dim(m \cdot l^2 \cdot t^{-2}) = M L^2 T^{-2}$

The quantity *amount of substance* is of special importance to chemists. Amount of substance is proportional to the number of specified elementary entities of the substance considered. The proportionality factor is the same for all substances; its reciprocal is the *Avogadro constant* (see Section 2.10, p. 47, Section 3.3, p. 88, and Chapter 5, p. 111). The SI unit of amount of substance is the mole, defined in Section 3.3, p. 88. The physical quantity “amount of substance” should no longer be called “number of moles”, just as the physical quantity “mass” should not be called “number of kilograms”. The name “amount of substance”, sometimes also called “chemical amount”, may often be usefully abbreviated to the single word “amount”, particularly in such phrases as “amount concentration” (see footnote <sup>2</sup>, below), and “amount of N<sub>2</sub>”. A possible name for international usage has been suggested: “enplethy” [10] (from Greek, similar to enthalpy and entropy).

The number and choice of base quantities is pure convention. Other quantities could be considered to be more fundamental, such as electric charge  $Q$  instead of electric current  $I$ .

$$Q = \int_{t_1}^{t_2} I \, dt \quad (7)$$

However, in the ISQ, electric current is chosen as base quantity and ampere is the SI base unit. In atomic and molecular physics, the so-called *atomic units* are useful (see Section 3.9, p. 94).

---

<sup>1</sup> The symbol  $[Q]$  was formerly used for *dimension* of  $Q$ , but this symbol is used and preferred for *unit* of  $Q$ .

<sup>2</sup> The Clinical Chemistry Division of IUPAC recommended that “amount-of-substance concentration” be abbreviated “substance concentration” [14].

## 1.3 SYMBOLS FOR PHYSICAL QUANTITIES AND UNITS [5.a]

A clear distinction should be drawn between the names and symbols for physical quantities, and the names and symbols for units. Names and symbols for many quantities are given in Chapter 2, p. 11; the symbols given there are *recommendations*. If other symbols are used they should be clearly defined. Names and symbols for units are given in Chapter 3, p. 83; the symbols for units listed there are quoted from the Bureau International des Poids et Mesures (BIPM) and are *mandatory*.

### 1.3.1 General rules for symbols for quantities

The symbol for a physical quantity should be a single letter (see footnote <sup>1</sup>, below) of the Latin or Greek alphabet (see Section 1.6, p. 7). Capital or lower case letters may both be used. The letter should be printed in italic (sloping) type. When necessary the symbol may be modified by subscripts and superscripts of specified meaning. Subscripts and superscripts that are themselves symbols for physical quantities or for numbers should be printed in italic type; other subscripts and superscripts should be printed in Roman (upright) type.

|                 |                      |                                                         |
|-----------------|----------------------|---------------------------------------------------------|
| <i>Examples</i> | $C_p$                | for heat capacity at constant pressure                  |
|                 | $p_i$                | for partial pressure of the $i$ th substance            |
| but             | $C_B$                | for heat capacity of substance B                        |
|                 | $\mu_B^\alpha$       | for chemical potential of substance B in phase $\alpha$ |
|                 | $E_k$                | for kinetic energy                                      |
|                 | $\mu_r$              | for relative permeability                               |
|                 | $\Delta_r H^\ominus$ | for standard reaction enthalpy                          |
|                 | $V_m$                | for molar volume                                        |
|                 | $A_{10}$             | for decadic absorbance                                  |

The meaning of symbols for physical quantities may be further qualified by the use of one or more subscripts, or by information contained in parentheses.

|                 |                                                                                                                                   |
|-----------------|-----------------------------------------------------------------------------------------------------------------------------------|
| <i>Examples</i> | $\Delta_f S^\ominus(\text{HgCl}_2, \text{cr}, 25^\circ\text{C}) = -154.3 \text{ J K}^{-1} \text{ mol}^{-1}$                       |
|                 | $\mu_i = (\partial G / \partial n_i)_{T,p,\dots,n_j,\dots; j \neq i}$ or $\mu_i = (\partial G / \partial n_i)_{T,p,n_{j \neq i}}$ |

Vectors and matrices may be printed in bold-face italic type, e.g. ***A***, ***a***. Tensors may be printed in bold-face italic sans serif type, e.g. ***S***, ***T***. Vectors may alternatively be characterized by an arrow,  $\vec{A}$ ,  $\vec{a}$  and second-rank tensors by a double arrow,  $\vec{\vec{S}}$ ,  $\vec{\vec{T}}$ .

### 1.3.2 General rules for symbols for units

Symbols for units should be printed in Roman (upright) type. They should remain unaltered in the plural, and should not be followed by a full stop except at the end of a sentence.

*Example*  $r = 10 \text{ cm}$ , not cm. or cms.

---

<sup>1</sup> An exception is made for certain characteristic numbers or “dimensionless quantities” used in the study of transport processes for which the internationally agreed symbols consist of two letters (see Section 2.15.1, p. 82).

*Example* Reynolds number,  $Re$ ; another example is pH (see Sections 2.13 and 2.13.1 (viii), p. 70 and 75).

When such symbols appear as factors in a product, they should be separated from other symbols by a space, multiplication sign, or parentheses.

Symbols for units shall be printed in lower case letters, unless they are derived from a personal name when they shall begin with a capital letter. An exception is the symbol for the litre which may be either L or l, i.e. either capital or lower case (see footnote <sup>2</sup>, below).

*Examples*      m (metre), s (second), but J (joule), Hz (hertz)

Decimal multiples and submultiples of units may be indicated by the use of prefixes as defined in Section 3.6, p. 91.

*Examples*      nm (nanometre), MHz (megahertz), kV (kilovolt)

## 1.4 USE OF THE WORDS “EXTENSIVE”, “INTENSIVE”, “SPECIFIC”, AND “MOLAR”

A quantity that is additive for independent, noninteracting subsystems is called *extensive*; examples are mass  $m$ , volume  $V$ , Gibbs energy  $G$ . A quantity that is independent of the extent of the system is called *intensive*; examples are temperature  $T$ , pressure  $p$ , chemical potential (partial molar Gibbs energy)  $\mu$ .

The adjective *specific* before the name of an extensive quantity is used to mean *divided by mass*. When the symbol for the extensive quantity is a capital letter, the symbol used for the specific quantity is often the corresponding lower case letter.

*Examples*      volume,  $V$ , and  
                  specific volume,  $v = V/m = 1/\rho$  (where  $\rho$  is mass density);  
                  heat capacity at constant pressure,  $C_p$ , and  
                  specific heat capacity at constant pressure,  $c_p = C_p/m$

ISO [5.a] and the Clinical Chemistry Division of IUPAC recommend systematic naming of physical quantities derived by division with mass, volume, area, and length by using the attributes massic or specific, volumic, areic, and lineic, respectively. In addition the Clinical Chemistry Division of IUPAC recommends the use of the attribute entitic for quantities derived by division with the number of entities [14]. Thus, for example, the specific volume could be called massic volume and the surface charge density would be areic charge.

The adjective *molar* before the name of an extensive quantity generally means *divided by amount of substance*. The subscript m on the symbol for the extensive quantity denotes the corresponding molar quantity.

*Examples*      volume,  $V$       molar volume,  $V_m = V/n$  (Section 2.10, p. 47)  
                  enthalpy,  $H$       molar enthalpy,  $H_m = H/n$

If the name enplethy (see Section 1.2, p. 4) is accepted for “amount of substance” one can use enplethic volume instead of molar volume, for instance. The word “molar” violates the principle that the name of the quantity should not be mixed with the name of the unit (mole in this case). The use of enplethic resolves this problem. It is sometimes convenient to divide all extensive quantities by amount of substance, so that all quantities become intensive; the subscript m may then be omitted if this convention is stated and there is no risk of ambiguity. (See also the symbols recommended for partial molar quantities in Section 2.11, p. 57, and in Section 2.11.1 (iii), p. 60.)

There are a few cases where the adjective *molar* has a different meaning, namely *divided by amount-of-substance concentration*.

*Examples*      absorption coefficient,  $a$   
                  molar absorption coefficient,  $\varepsilon = a/c$  (see Section 2.7, note 22, p. 37)  
                  conductivity,  $\kappa$   
                  molar conductivity,  $\Lambda = \kappa/c$  (see Section 2.13, p. 73)

---

<sup>2</sup> However, only the lower case l is used by ISO and the International Electrotechnical Commission (IEC).

## 1.5 PRODUCTS AND QUOTIENTS OF PHYSICAL QUANTITIES AND UNITS

Products of physical quantities may be written in any of the ways

$$a\ b \quad \text{or} \quad ab \quad \text{or} \quad a \cdot b \quad \text{or} \quad a \times b$$

and similarly quotients may be written

$$a/b \quad \text{or} \quad \frac{a}{b} \quad \text{or by writing the product of } a \text{ and } b^{-1}, \text{ e.g. } ab^{-1}$$

*Examples*  $F = ma, \quad p = nRT/V$

Not more than one solidus (/) shall be used in the same expression unless parentheses are used to eliminate ambiguity.

*Example*  $(a/b)/c$  or  $a/(b/c)$  (in general different), not  $a/b/c$

In evaluating combinations of many factors, multiplication written without a multiplication sign takes precedence over division in the sense that  $a/bc$  is interpreted as  $a/(bc)$  and not as  $(a/b)c$ ; however, it is necessary to use parentheses to eliminate ambiguity under all circumstances, thus avoiding expressions of the kind  $a/bcd$  etc. Furthermore,  $a/b + c$  is interpreted as  $(a/b) + c$  and not as  $a/(b + c)$ . Again, the use of parentheses is recommended (required for  $a/(b + c)$ ).

Products and quotients of units may be written in a similar way, except that the cross ( $\times$ ) is not used as a multiplication sign between units. When a product of units is written without any multiplication sign a space shall be left between the unit symbols.

*Example*  $1\ \text{N} = 1\ \text{m kg s}^{-2} = 1\ \text{m} \cdot \text{kg} \cdot \text{s}^{-2} = 1\ \text{m kg/s}^2, \quad \text{not} \quad 1\ \text{mkg s}^{-2}$

## 1.6 THE USE OF ITALIC AND ROMAN FONTS FOR SYMBOLS IN SCIENTIFIC PUBLICATIONS

Scientific manuscripts should follow the accepted conventions concerning the use of italic and Roman fonts for symbols. An italic font is generally used for emphasis in running text, but it has a quite specific meaning when used for symbols in scientific text and equations. The following summary is intended to help in the correct use of italic fonts in preparing manuscript material.

1. The general rules concerning the use of italic (sloping) font or Roman (upright) font are presented in Section 1.3.2, p. 5 and in Section 4.1, p. 103 in relation to mathematical symbols and operators. These rules are also presented in the International Standards ISO 31 (successively being replaced by ISO/IEC 80000) [5], ISO 1000 [6], and in the SI Brochure [3].
2. The overall rule is that symbols representing physical quantities or variables are italic, but symbols representing units, mathematical constants, or labels, are roman. Sometimes there may seem to be doubt as to whether a symbol represents a quantity or has some other meaning (such as label): a good rule is that quantities, or variables, may have a range of numerical values, but labels cannot. Vectors, tensors and matrices are denoted using a bold-face (heavy) font, but they shall be italic since they are quantities.

*Examples* The Planck constant  $h = 6.626\ 068\ 96(33) \times 10^{-34}\ \text{J s}$ .  
The electric field strength  $\mathbf{E}$  has components  $E_x, E_y$ , and  $E_z$ .  
The mass of my pen is  $m = 24\ \text{g} = 0.024\ \text{kg}$ .

3. The above rule applies equally to all letter symbols from both the Greek and the Latin alphabet, although some authors resist putting Greek letters into italic.

*Example* When the symbol  $\mu$  is used to denote a physical quantity (such as permeability or reduced mass) it should be italic, but when it is used as a prefix in a unit such as microgram,  $\mu\text{g}$ , or when it is used as the symbol for the muon,  $\mu$  (see paragraph 5 below), it should be roman.

4. Numbers, and labels, are roman (upright).

*Examples* The ground and first excited electronic state of the  $\text{CH}_2$  molecule are denoted  $\dots(2a_1)^2(1b_2)^2(3a_1)^1(1b_1)^1$ ,  $\tilde{X}^3B_1$ , and  $\dots(2a_1)^2(1b_2)^2(3a_1)^2$ ,  $\tilde{a}^1A_1$ , respectively. The  $\pi$ -electron configuration and symmetry of the benzene molecule in its ground state are denoted:  $\dots(a_{2u})^2(e_{1g})^4$ ,  $\tilde{X}^1A_{1g}$ . All these symbols are labels and are roman.

5. Symbols for elements in the periodic system shall be roman. Similarly the symbols used to represent elementary particles are always roman. (See, however, paragraph 9 below for use of italic font in chemical-compound names.)

*Examples* H, He, Li, Be, B, C, N, O, F, Ne, ... for atoms; e for the electron, p for the proton, n for the neutron,  $\mu$  for the muon,  $\alpha$  for the alpha particle, etc.

6. Symbols for physical quantities are single, or exceptionally two letters of the Latin or Greek alphabet, but they are frequently supplemented with subscripts, superscripts or information in parentheses to specify further the quantity. Further symbols used in this way are either italic or roman depending on what they represent.

*Examples*  $H$  denotes enthalpy, but  $H_m$  denotes molar enthalpy (m is a mnemonic label for molar, and is therefore roman).  $C_p$  and  $C_V$  denote the heat capacity at constant pressure  $p$  and volume  $V$ , respectively (note the roman m but italic  $p$  and  $V$ ). The chemical potential of argon might be denoted  $\mu_{\text{Ar}}$  or  $\mu(\text{Ar})$ , but the chemical potential of the  $i$ th component in a mixture would be denoted  $\mu_i$ , where  $i$  is italic because it is a variable subscript.

7. Symbols for mathematical operators are always roman. This applies to the symbol  $\Delta$  for a difference,  $\delta$  for an infinitesimal variation,  $d$  for an infinitesimal difference (in calculus), and to capital  $\Sigma$  and  $\Pi$  for summation and product signs, respectively. The symbols  $\pi$  (3.141 592...),  $e$  (2.718 281..., base of natural logarithms),  $i$  (square root of minus one), etc. are always roman, as are the symbols for specified functions such as  $\log$  ( $\lg$  for  $\log_{10}$ ,  $\ln$  for  $\log_e$ , or  $\text{lb}$  for  $\log_2$ ),  $\exp$ ,  $\sin$ ,  $\cos$ ,  $\tan$ ,  $\text{erf}$ , ***div***, ***grad***, ***rot***, etc. The particular operators ***grad*** and ***rot*** and the corresponding symbols  $\nabla$  for ***grad***,  $\nabla \times$  for ***rot***, and  $\nabla \cdot$  for ***div*** are printed in bold-face to indicate the vector or tensor character following [5.k]. Some of these letters, e.g.  $e$  for elementary charge, are also sometimes used to represent physical quantities; then of course they shall be italic, to distinguish them from the corresponding mathematical symbol.

*Examples*  $\Delta H = H(\text{final}) - H(\text{initial})$ ;  $(dp/dt)$  used for the rate of change of pressure;  $\delta x$  used to denote an infinitesimal variation of  $x$ . But for a damped linear oscillator the amplitude  $F$  as a function of time  $t$  might be expressed by the equation  $F = F_0 \exp(-\delta t) \sin(\omega t)$  where  $\delta$  is the decay coefficient (SI unit: Np/s) and  $\omega$  is the angular frequency (SI unit: rad/s). Note the use of roman  $\delta$  for the operator in an infinitesimal variation of  $x$ ,  $\delta x$ , but italic  $\delta$  for the decay coefficient in the product  $\delta t$ . Note that the products  $\delta t$  and  $\omega t$  are both dimensionless, but are described as having the unit neper ( $\text{Np} = 1$ ) and radian ( $\text{rad} = 1$ ), respectively.

8. The fundamental physical constants are always regarded as quantities subject to measurement (even though they are not considered to be variables) and they should accordingly always be italic. Sometimes fundamental physical constants are used as though they were units, but they are still given italic symbols. An example is the hartree,  $E_h$  (see Section 3.9.1, p. 95). However, the electronvolt, eV, the dalton, Da, or the unified atomic mass unit, u, and the astronomical unit, ua, have been recognized as units by the Comité International des Poids et Mesures (CIPM) of the BIPM and they are accordingly given Roman symbols.

*Examples*  $c_0$  for the speed of light in vacuum,  $m_e$  for the electron mass,  $h$  for the Planck constant,  $N_A$  or  $L$  for the Avogadro constant,  $e$  for the elementary charge,  $a_0$  for the Bohr radius, etc.

The electronvolt  $1 \text{ eV} = e \cdot 1 \text{ V} = 1.602\,176\,487(40) \times 10^{-19} \text{ J}$ .

9. Greek letters are used in systematic organic, inorganic, macromolecular, and biochemical nomenclature. These should be roman (upright), since they are not symbols for physical quantities. They designate the position of substitution in side chains, ligating-atom attachment and bridging mode in coordination compounds, end groups in structure-based names for macromolecules, and stereochemistry in carbohydrates and natural products. Letter symbols for elements are italic when they are locants in chemical-compound names indicating attachments to heteroatoms, e.g. *O*-, *N*-, *S*-, and *P*-. The italic symbol *H* denotes indicated or added hydrogen (see reference [15]).

*Examples*  $\alpha$ -ethylcyclopentaneacetic acid  
 $\beta$ -methyl-4-propylcyclohexaneethanol  
 tetracarbonyl( $\eta^4$ -2-methylidenepropane-1,3-diyl)chromium  
 $\alpha$ -(trichloromethyl)- $\omega$ -chloropoly(1,4-phenylenemethylene)  
 $\alpha$ -D-glucopyranose  
 5 $\alpha$ -androstan-3 $\beta$ -ol  
*N*-methylbenzamide  
*O*-ethyl hexanethioate  
 3*H*-pyrrole  
 naphthalen-2(1*H*)-one



## 2 TABLES OF PHYSICAL QUANTITIES

The following tables contain the internationally recommended names and symbols for the physical quantities most likely to be used by chemists. Further quantities and symbols may be found in recommendations by IUPAP [4] and ISO [5].

Although authors are free to choose any symbols they wish for the quantities they discuss, provided that they define their notation and conform to the general rules indicated in Chapter 1, it is clearly an aid to scientific communication if we all generally follow a standard notation. The symbols below have been chosen to conform with current usage and to minimize conflict so far as possible. Small variations from the recommended symbols may often be desirable in particular situations, perhaps by adding or modifying subscripts or superscripts, or by the alternative use of upper or lower case. Within a limited subject area it may also be possible to simplify notation, for example by omitting qualifying subscripts or superscripts, without introducing ambiguity. The notation adopted should in any case always be defined. Major deviations from the recommended symbols should be particularly carefully defined.

The tables are arranged by subject. The five columns in each table give the name of the quantity, the recommended symbol(s), a brief definition, the symbol for the coherent SI unit (without multiple or submultiple prefixes, see Section 3.6, p. 91), and note references. When two or more symbols are recommended, commas are used to separate symbols that are equally acceptable, and symbols of second choice are put in parentheses. A semicolon is used to separate symbols of slightly different quantities. The definitions are given primarily for identification purposes and are not necessarily complete; they should be regarded as useful relations rather than formal definitions. For some of the quantities listed in this chapter, the definitions given in various IUPAC documents are collected in [16]. Useful definitions of physical quantities in physical organic chemistry can be found in [17] and those in polymer chemistry in [18]. For dimensionless quantities, a 1 is entered in the SI unit column (see Section 3.10, p. 97). Further information is added in notes, and in text inserts between the tables, as appropriate. Other symbols used are defined within the same table (not necessarily in the order of appearance) and in the notes.



## 2.1 SPACE AND TIME

The names and symbols recommended here are in agreement with those recommended by IUPAP [4] and ISO [5.b].

| <i>Name</i>                                         | <i>Symbols</i>                                                     | <i>Definition</i>                                            | <i>SI unit</i>                        | <i>Notes</i> |
|-----------------------------------------------------|--------------------------------------------------------------------|--------------------------------------------------------------|---------------------------------------|--------------|
| cartesian<br>space coordinates                      | $x; y; z$                                                          |                                                              | m                                     |              |
| cylindrical coordinates                             | $\rho; \vartheta; z$                                               |                                                              | m, 1, m                               |              |
| spherical polar<br>coordinates                      | $r; \vartheta; \varphi$                                            |                                                              | m, 1, 1                               |              |
| generalized coordinates                             | $q, q_i$                                                           |                                                              | (varies)                              |              |
| position vector                                     | $\mathbf{r}$                                                       | $\mathbf{r} = x\mathbf{e}_x + y\mathbf{e}_y + z\mathbf{e}_z$ | m                                     |              |
| length                                              | $l$                                                                |                                                              | m                                     |              |
| special symbols:                                    |                                                                    |                                                              |                                       |              |
| height                                              | $h$                                                                |                                                              |                                       |              |
| breadth                                             | $b$                                                                |                                                              |                                       |              |
| thickness                                           | $d, \delta$                                                        |                                                              |                                       |              |
| distance                                            | $d$                                                                |                                                              |                                       |              |
| radius                                              | $r$                                                                |                                                              |                                       |              |
| diameter                                            | $d$                                                                |                                                              |                                       |              |
| path length                                         | $s$                                                                |                                                              |                                       |              |
| length of arc                                       | $s$                                                                |                                                              |                                       |              |
| area                                                | $A, A_s, S$                                                        |                                                              | m <sup>2</sup>                        | 1            |
| volume                                              | $V, (v)$                                                           |                                                              | m <sup>3</sup>                        |              |
| plane angle                                         | $\alpha, \beta, \gamma, \vartheta, \varphi, \dots$                 | $\alpha = s/r$                                               | rad, 1                                | 2            |
| solid angle                                         | $\Omega, (\omega)$                                                 | $\Omega = A/r^2$                                             | sr, 1                                 | 2            |
| time, duration                                      | $t$                                                                |                                                              | s                                     |              |
| period                                              | $T$                                                                | $T = t/N$                                                    | s                                     | 3            |
| frequency                                           | $\nu, f$                                                           | $\nu = 1/T$                                                  | Hz, s <sup>-1</sup>                   |              |
| angular frequency                                   | $\omega$                                                           | $\omega = 2\pi\nu$                                           | rad s <sup>-1</sup> , s <sup>-1</sup> | 2, 4         |
| characteristic                                      | $\tau, T$                                                          | $\tau =  dt/d \ln x $                                        | s                                     |              |
| time interval,<br>relaxation time,<br>time constant |                                                                    |                                                              |                                       |              |
| angular velocity                                    | $\omega$                                                           | $\omega = d\varphi/dt$                                       | rad s <sup>-1</sup> , s <sup>-1</sup> | 2, 5         |
| velocity                                            | $\mathbf{v}, \mathbf{u}, \mathbf{w}, \mathbf{c}, \dot{\mathbf{r}}$ | $\mathbf{v} = d\mathbf{r}/dt$                                | m s <sup>-1</sup>                     |              |
| speed                                               | $v, u, w, c$                                                       | $v =  \mathbf{v} $                                           | m s <sup>-1</sup>                     | 6            |
| acceleration                                        | $\mathbf{a}$                                                       | $\mathbf{a} = d\mathbf{v}/dt$                                | m s <sup>-2</sup>                     | 7            |

(1) An infinitesimal area may be regarded as a vector  $\mathbf{e}_n dA$ , where  $\mathbf{e}_n$  is the unit vector normal to the plane.

(2) The units radian (rad) and steradian (sr) for plane angle and solid angle are derived. Since they are of dimension one (i.e. dimensionless), they may be included in expressions for derived SI units if appropriate, or omitted if clarity and meaning is not lost thereby.

(3)  $N$  is the number of identical (periodic) events during the time  $t$ .

(4) The unit Hz is not to be used for angular frequency.

(5) Angular velocity can be treated as a vector,  $\boldsymbol{\omega}$ , perpendicular to the plane of rotation defined by  $\mathbf{v} = \boldsymbol{\omega} \times \mathbf{r}$ .

(6) For the speeds of light and sound the symbol  $c$  is customary.

(7) For acceleration of free fall the symbol  $g$  is used.

## 2.2 CLASSICAL MECHANICS

The names and symbols recommended here are in agreement with those recommended by IUPAP [4] and ISO [5.c]. Additional quantities and symbols used in acoustics can be found in [4,5.g].

| <i>Name</i>                | <i>Symbol</i>     | <i>Definition</i>                                | <i>SI unit</i>                        | <i>Notes</i> |
|----------------------------|-------------------|--------------------------------------------------|---------------------------------------|--------------|
| mass                       | $m$               |                                                  | kg                                    |              |
| reduced mass               | $\mu$             | $\mu = m_1 m_2 / (m_1 + m_2)$                    | kg                                    |              |
| density, mass density      | $\rho$            | $\rho = m/V$                                     | kg m <sup>-3</sup>                    |              |
| relative density           | $d$               | $d = \rho/\rho^\ominus$                          | 1                                     | 1            |
| surface density            | $\rho_A, \rho_S$  | $\rho_A = m/A$                                   | kg m <sup>-2</sup>                    |              |
| specific volume            | $v$               | $v = V/m = 1/\rho$                               | m <sup>3</sup> kg <sup>-1</sup>       |              |
| momentum                   | $\mathbf{p}$      | $\mathbf{p} = m\mathbf{v}$                       | kg m s <sup>-1</sup>                  |              |
| angular momentum           | $\mathbf{L}$      | $\mathbf{L} = \mathbf{r} \times \mathbf{p}$      | J s                                   | 2            |
| moment of inertia          | $I, J$            | $I = \sum_i m_i r_i^2$                           | kg m <sup>2</sup>                     | 3            |
| force                      | $\mathbf{F}$      | $\mathbf{F} = d\mathbf{p}/dt = m\mathbf{a}$      | N                                     |              |
| moment of force,<br>torque | $\mathbf{M}, (T)$ | $\mathbf{M} = \mathbf{r} \times \mathbf{F}$      | N m                                   |              |
| energy                     | $E$               |                                                  | J                                     |              |
| potential energy           | $E_p, V, \Phi$    | $E_p = - \int \mathbf{F} \cdot d\mathbf{r}$      | J                                     | 4            |
| kinetic energy             | $E_k, T, K$       | $E_k = (1/2)mv^2$                                | J                                     |              |
| work                       | $W, A, w$         | $W = \int \mathbf{F} \cdot d\mathbf{r}$          | J                                     |              |
| power                      | $P$               | $P = \mathbf{F} \cdot \mathbf{v} = dW/dt$        | W                                     |              |
| generalized coordinate     | $q$               |                                                  | (varies)                              |              |
| generalized momentum       | $p$               |                                                  | (varies)                              |              |
| Lagrange function          | $L$               | $L(q, \dot{q}) = T(q, \dot{q}) - V(q)$           | J                                     |              |
| Hamilton function          | $H$               | $H(q, p) = \sum_i p_i \dot{q}_i - L(q, \dot{q})$ | J                                     |              |
| action                     | $S$               | $S = \int L dt$                                  | J s                                   | 5            |
| pressure                   | $p, (P)$          | $p = F/A$                                        | Pa, N m <sup>-2</sup>                 |              |
| surface tension            | $\gamma, \sigma$  | $\gamma = dW/dA$                                 | N m <sup>-1</sup> , J m <sup>-2</sup> |              |
| weight                     | $G, (W, P)$       | $G = mg$                                         | N                                     |              |
| gravitational constant     | $G$               | $F = Gm_1 m_2 / r^2$                             | N m <sup>2</sup> kg <sup>-2</sup>     |              |

(1) Usually  $\rho^\ominus = \rho(\text{H}_2\text{O}, 4^\circ\text{C})$ .

(2) Other symbols are customary in atomic and molecular spectroscopy (see Section 2.6, p. 25).

(3) In general  $\mathbf{I}$  is a tensor quantity:  $I_{\alpha\alpha} = \sum_i m_i (\beta_i^2 + \gamma_i^2)$ , and  $I_{\alpha\beta} = -\sum_i m_i \alpha_i \beta_i$  if  $\alpha \neq \beta$ , where  $\alpha, \beta, \gamma$  is a permutation of  $x, y, z$ . For a continuous mass distribution the sums are replaced by integrals.

(4) Strictly speaking, only potential energy differences have physical significance, thus the integral is to be interpreted as a definite integral, for instance

$$E_p(r_1, r_2) = - \int_{r_1}^{r_2} \mathbf{F} \cdot d\mathbf{r}$$

or possibly with the upper limit infinity

$$E_p(r) = - \int_r^\infty \mathbf{F} \cdot d\mathbf{r}$$

(5) Action is the time integral over the Lagrange function  $L$ , which is equivalent to  $\int p dq - \int H dt$  (see [19]).

| <i>Name</i>                               | <i>Symbol</i>        | <i>Definition</i>                        | <i>SI unit</i>              | <i>Notes</i> |
|-------------------------------------------|----------------------|------------------------------------------|-----------------------------|--------------|
| normal stress                             | $\sigma$             | $\sigma = F/A$                           | Pa                          | 6            |
| shear stress                              | $\tau$               | $\tau = F/A$                             | Pa                          | 6            |
| linear strain,<br>relative elongation     | $\varepsilon, e$     | $\varepsilon = \Delta l/l$               | 1                           |              |
| modulus of elasticity,<br>Young's modulus | $E$                  | $E = \sigma/\varepsilon$                 | Pa                          | 6            |
| shear strain                              | $\gamma$             | $\gamma = \Delta x/d$                    | 1                           | 6, 7         |
| shear modulus,<br>Coulomb's modulus       | $G$                  | $G = \tau/\gamma$                        | Pa                          | 6            |
| volume strain,<br>bulk strain             | $\vartheta$          | $\vartheta = \Delta V/V_0$               | 1                           | 6            |
| bulk modulus,<br>compression modulus      | $K$                  | $K = -V_0 (dp/dV)$                       | Pa                          | 6            |
| viscosity,<br>dynamic viscosity           | $\eta, (\mu)$        | $\tau_{xz} = \eta (dv_x/dz)$             | Pa s                        |              |
| fluidity                                  | $\varphi$            | $\varphi = 1/\eta$                       | $\text{m kg}^{-1} \text{s}$ |              |
| kinematic viscosity                       | $\nu$                | $\nu = \eta/\rho$                        | $\text{m}^2 \text{s}^{-1}$  |              |
| dynamic friction factor                   | $\mu, (f)$           | $F_{\text{frict}} = \mu F_{\text{norm}}$ | 1                           |              |
| sound energy flux                         | $P, P_a$             | $P = dE/dt$                              | W                           |              |
| acoustic factors,                         |                      |                                          |                             |              |
| reflection                                | $\rho$               | $\rho = P_r/P_0$                         | 1                           | 8            |
| absorption                                | $\alpha_a, (\alpha)$ | $\alpha_a = 1 - \rho$                    | 1                           | 9            |
| transmission                              | $\tau$               | $\tau = P_{\text{tr}}/P_0$               | 1                           | 8            |
| dissipation                               | $\delta$             | $\delta = \alpha_a - \tau$               | 1                           |              |

(6) In general these can be tensor quantities.

(7)  $d$  is the distance between the layers displaced by  $\Delta x$ .

(8)  $P_0$  is the incident sound energy flux,  $P_r$  the reflected flux and  $P_{\text{tr}}$  the transmitted flux.

(9) This definition is special to acoustics and is different from the usage in radiation, where the absorption factor corresponds to the acoustic dissipation factor.

## 2.3 ELECTRICITY AND MAGNETISM

The names and symbols recommended here are in agreement with those recommended by IUPAP [4] and ISO [5.e].

| <i>Name</i>                                                                                 | <i>Symbol</i>                            | <i>Definition</i>                                                 | <i>SI unit</i>                                                                 | <i>Notes</i> |
|---------------------------------------------------------------------------------------------|------------------------------------------|-------------------------------------------------------------------|--------------------------------------------------------------------------------|--------------|
| electric current                                                                            | $I, i$                                   |                                                                   | A                                                                              | 1            |
| electric current density                                                                    | $\mathbf{j}, \mathbf{J}$                 | $I = \int \mathbf{j} \cdot \mathbf{e}_n \, dA$                    | A m <sup>-2</sup>                                                              | 2            |
| electric charge,<br>quantity of electricity                                                 | $Q$                                      | $Q = \int I \, dt$                                                | C                                                                              | 1            |
| charge density                                                                              | $\rho$                                   | $\rho = Q/V$                                                      | C m <sup>-3</sup>                                                              |              |
| surface density of charge                                                                   | $\sigma$                                 | $\sigma = Q/A$                                                    | C m <sup>-2</sup>                                                              |              |
| electric potential                                                                          | $V, \phi$                                | $V = dW/dQ$                                                       | V, J C <sup>-1</sup>                                                           |              |
| electric potential difference,<br>electric tension                                          | $U, \Delta V, \Delta \phi$               | $U = V_2 - V_1$                                                   | V                                                                              |              |
| electromotive force                                                                         | $E,$<br>$(E_{\text{mf}}, E_{\text{MK}})$ | $E = \oint (\mathbf{F}/Q) \cdot d\mathbf{r}$                      | V                                                                              | 3            |
| electric field strength                                                                     | $\mathbf{E}$                             | $\mathbf{E} = \mathbf{F}/Q = -\nabla V$                           | V m <sup>-1</sup>                                                              |              |
| electric flux                                                                               | $\Psi$                                   | $\Psi = \int \mathbf{D} \cdot \mathbf{e}_n \, dA$                 | C                                                                              | 2            |
| electric displacement                                                                       | $\mathbf{D}$                             | $\nabla \cdot \mathbf{D} = \rho$                                  | C m <sup>-2</sup>                                                              |              |
| capacitance                                                                                 | $C$                                      | $C = Q/U$                                                         | F, C V <sup>-1</sup>                                                           |              |
| permittivity                                                                                | $\varepsilon$                            | $\mathbf{D} = \varepsilon \mathbf{E}$                             | F m <sup>-1</sup>                                                              | 4            |
| electric constant,<br>permittivity of vacuum                                                | $\varepsilon_0$                          | $\varepsilon_0 = \mu_0^{-1} c_0^{-2}$                             | F m <sup>-1</sup>                                                              | 5            |
| relative permittivity                                                                       | $\varepsilon_r$                          | $\varepsilon_r = \varepsilon/\varepsilon_0$                       | 1                                                                              | 6            |
| dielectric polarization,<br>electric polarization<br>(electric dipole moment<br>per volume) | $\mathbf{P}$                             | $\mathbf{P} = \mathbf{D} - \varepsilon_0 \mathbf{E}$              | C m <sup>-2</sup>                                                              |              |
| electric susceptibility                                                                     | $\chi_e$                                 | $\chi_e = \varepsilon_r - 1$                                      | 1                                                                              |              |
| 1st hyper-susceptibility                                                                    | $\chi_e^{(2)}$                           | $\chi_e^{(2)} = \varepsilon_0^{-1} (\partial^2 P / \partial E^2)$ | C m J <sup>-1</sup> , m V <sup>-1</sup>                                        | 7            |
| 2nd hyper-susceptibility                                                                    | $\chi_e^{(3)}$                           | $\chi_e^{(3)} = \varepsilon_0^{-1} (\partial^3 P / \partial E^3)$ | C <sup>2</sup> m <sup>2</sup> J <sup>-2</sup> , m <sup>2</sup> V <sup>-2</sup> | 7            |

(1) The electric current  $I$  is a base quantity in ISQ.

(2)  $\mathbf{e}_n dA$  is a vector element of area (see Section 2.1, note 1, p. 13).

(3) The name electromotive force and the symbol emf are no longer recommended, since an electric potential difference is not a force (see Section 2.13, note 14, p. 71).

(4)  $\varepsilon$  can be a second-rank tensor.

(5)  $c_0$  is the speed of light in vacuum.

(6) This quantity was formerly called dielectric constant.

(7) The hyper-susceptibilities are the coefficients of the non-linear terms in the expansion of the magnitude  $P$  of the dielectric polarization  $\mathbf{P}$  in powers of the electric field strength  $\mathbf{E}$ , quite related to the expansion of the dipole moment vector described in Section 2.5, note 17, p. 24. In isotropic media, the expansion of the component  $i$  of the dielectric polarization is given by

$$P_i = \varepsilon_0 [\chi_e^{(1)} E_i + (1/2) \chi_e^{(2)} E_i^2 + (1/6) \chi_e^{(3)} E_i^3 + \dots]$$

where  $E_i$  is the  $i$ -th component of the electric field strength, and  $\chi_e^{(1)}$  is the usual electric susceptibility  $\chi_e$ , equal to  $\varepsilon_r - 1$  in the absence of higher terms. In anisotropic media,  $\chi_e^{(1)}$ ,  $\chi_e^{(2)}$ , and  $\chi_e^{(3)}$  are tensors of rank 2, 3, and 4. For an isotropic medium (such as a liquid), or for a crystal with a centrosymmetric unit cell,  $\chi_e^{(2)}$  is zero by symmetry. These quantities are macroscopic

| <i>Name</i>                                             | <i>Symbol</i>                  | <i>Definition</i>                                 | <i>SI unit</i>                        | <i>Notes</i> |
|---------------------------------------------------------|--------------------------------|---------------------------------------------------|---------------------------------------|--------------|
| electric dipole moment                                  | $\mathbf{p}, \boldsymbol{\mu}$ | $\mathbf{p} = \sum_i Q_i \mathbf{r}_i$            | C m                                   | 8            |
| magnetic flux density,<br>magnetic induction            | $\mathbf{B}$                   | $\mathbf{F} = Q\mathbf{v} \times \mathbf{B}$      | T                                     | 9            |
| magnetic flux                                           | $\Phi$                         | $\Phi = \int \mathbf{B} \cdot \mathbf{e}_n \, dA$ | Wb                                    | 2            |
| magnetic field strength,<br>magnetizing field strength  | $\mathbf{H}$                   | $\nabla \times \mathbf{H} = \mathbf{j}$           | A m <sup>-1</sup>                     |              |
| permeability                                            | $\mu$                          | $\mathbf{B} = \mu \mathbf{H}$                     | N A <sup>-2</sup> , H m <sup>-1</sup> | 10           |
| magnetic constant,<br>permeability of vacuum            | $\mu_0$                        | $\mu_0 = 4\pi \times 10^{-7} \text{ H m}^{-1}$    | H m <sup>-1</sup>                     |              |
| relative permeability                                   | $\mu_r$                        | $\mu_r = \mu/\mu_0$                               | 1                                     |              |
| magnetization<br>(magnetic dipole<br>moment per volume) | $\mathbf{M}$                   | $\mathbf{M} = \mathbf{B}/\mu_0 - \mathbf{H}$      | A m <sup>-1</sup>                     |              |
| magnetic susceptibility                                 | $\chi, \kappa, (\chi_m)$       | $\chi = \mu_r - 1$                                | 1                                     | 11           |
| molar magnetic susceptibility                           | $\chi_m$                       | $\chi_m = V_m \chi$                               | m <sup>3</sup> mol <sup>-1</sup>      |              |
| magnetic dipole moment                                  | $\mathbf{m}, \boldsymbol{\mu}$ | $\mathbf{E} = -\mathbf{m} \cdot \mathbf{B}$       | A m <sup>2</sup> , J T <sup>-1</sup>  |              |
| electric resistance                                     | $R$                            | $R = U/I$                                         | $\Omega$                              | 12           |
| conductance                                             | $G$                            | $G = 1/R$                                         | S                                     | 12           |
| loss angle                                              | $\delta$                       | $\delta = \varphi_U - \varphi_I$                  | rad                                   | 13           |
| reactance                                               | $X$                            | $X = (U/I) \sin \delta$                           | $\Omega$                              |              |
| impedance,<br>(complex impedance)                       | $Z$                            | $Z = R + iX$                                      | $\Omega$                              |              |
| admittance,<br>(complex admittance)                     | $Y$                            | $Y = 1/Z$                                         | S                                     |              |
| susceptance                                             | $B$                            | $Y = G + iB$                                      | S                                     |              |
| resistivity                                             | $\rho$                         | $\mathbf{E} = \rho \mathbf{j}$                    | $\Omega \text{ m}$                    | 14           |
| conductivity                                            | $\kappa, \gamma, \sigma$       | $\mathbf{j} = \boldsymbol{\kappa} \mathbf{E}$     | S m <sup>-1</sup>                     | 14, 15       |
| self-inductance                                         | $L$                            | $E = -L(dI/dt)$                                   | H, V s A <sup>-1</sup>                |              |
| mutual inductance                                       | $M, L_{12}$                    | $E_1 = -L_{12}(dI_2/dt)$                          | H, V s A <sup>-1</sup>                |              |
| magnetic vector potential                               | $\mathbf{A}$                   | $\mathbf{B} = \nabla \times \mathbf{A}$           | Wb m <sup>-1</sup>                    |              |
| Poynting vector                                         | $\mathbf{S}$                   | $\mathbf{S} = \mathbf{E} \times \mathbf{H}$       | W m <sup>-2</sup>                     | 16           |

(7) (continued) properties and characterize a dielectric medium in the same way that the microscopic quantities polarizability ( $\alpha$ ) and hyper-polarizabilities ( $\beta, \gamma$ ) characterize a molecule. For a homogeneous, saturated, isotropic dielectric medium of molar volume  $V_m$  one has  $\alpha_m = \epsilon_0 \chi_e V_m$ , where  $\alpha_m = N_A \alpha$  is the molar polarizability (Section 2.5, note 17, p. 24 and Section 2.7.2, p. 40).

(8) When a dipole is composed of two point charges  $Q$  and  $-Q$  separated by a distance  $r$ , the direction of the dipole vector is taken to be from the negative to the positive charge. The opposite convention is sometimes used, but is to be discouraged. The dipole moment of an ion depends on the choice of the origin.

(9) This quantity should not be called “magnetic field”.

(10)  $\mu$  is a second-rank tensor in anisotropic materials.

(11) The symbol  $\chi_m$  is sometimes used for magnetic susceptibility, but it should be reserved for molar magnetic susceptibility.

(12) In a material with reactance  $R = (U/I) \cos \delta$ , and  $G = R/(R^2 + X^2)$ .

(13)  $\varphi_I$  and  $\varphi_U$  are the phases of current and potential difference.

(14) This quantity is a tensor in anisotropic materials.

(15) ISO gives only  $\gamma$  and  $\sigma$ , but not  $\kappa$ .

(16) This quantity is also called the Poynting-Umov vector.

## 2.4 QUANTUM MECHANICS AND QUANTUM CHEMISTRY

The names and symbols for quantities used in quantum mechanics and recommended here are in agreement with those recommended by IUPAP [4]. The names and symbols for quantities used mainly in the field of quantum chemistry have been chosen on the basis of the current practice in the field. Guidelines for the presentation of methodological choices in publications of computational results have been presented [20]. A list of acronyms used in theoretical chemistry has been published by IUPAC [21]; see also Chapter 9, p. 155.

| <i>Name</i>                                         | <i>Symbol</i>                       | <i>Definition</i>                                                             | <i>SI unit</i>                     | <i>Notes</i> |
|-----------------------------------------------------|-------------------------------------|-------------------------------------------------------------------------------|------------------------------------|--------------|
| momentum operator                                   | $\hat{\mathbf{p}}$                  | $\hat{\mathbf{p}} = -i\hbar \nabla$                                           | J s m <sup>-1</sup>                | 1            |
| kinetic energy operator                             | $\hat{T}$                           | $\hat{T} = -(\hbar^2/2m)\nabla^2$                                             | J                                  | 1            |
| hamiltonian operator,<br>hamiltonian                | $\hat{H}$                           | $\hat{H} = \hat{T} + \hat{V}$                                                 | J                                  | 1            |
| wavefunction,<br>state function                     | $\Psi, \psi, \phi$                  | $\hat{H}\psi = E\psi$                                                         | (m <sup>-3/2</sup> )               | 2, 3         |
| hydrogen-like<br>wavefunction                       | $\psi_{nlm}(r, \theta, \phi)$       | $\psi_{nlm} = R_{nl}(r)Y_{lm}(\theta, \phi)$                                  | (m <sup>-3/2</sup> )               | 3            |
| spherical harmonic<br>function                      | $Y_{lm}(\theta, \phi)$              | $Y_{lm} = N_{l m }P_l^{ m }(\cos \theta)e^{im\phi}$                           | 1                                  | 4            |
| probability density                                 | $P$                                 | $P = \psi^*\psi$                                                              | (m <sup>-3</sup> )                 | 3, 5         |
| charge density<br>of electrons                      | $\rho$                              | $\rho = -eP$                                                                  | (C m <sup>-3</sup> )               | 3, 5, 6      |
| probability current<br>density,<br>probability flux | $\mathbf{S}$                        | $\mathbf{S} = -(i\hbar/2m) \times$<br>$(\psi^*\nabla\psi - \psi\nabla\psi^*)$ | (m <sup>-2</sup> s <sup>-1</sup> ) | 3            |
| electric current density<br>of electrons            | $\mathbf{j}$                        | $\mathbf{j} = -e\mathbf{S}$                                                   | (A m <sup>-2</sup> )               | 3, 6         |
| integration element                                 | d $\tau$                            | d $\tau = dx \, dy \, dz$                                                     | (varies)                           |              |
| matrix element<br>of operator $\hat{A}$             | $A_{ij}, \langle i   A   j \rangle$ | $A_{ij} = \int \psi_i^* \hat{A} \psi_j d\tau$                                 | (varies)                           | 7            |
| expectation value<br>of operator $\hat{A}$          | $\langle A \rangle, \bar{A}$        | $\langle A \rangle = \int \psi^* \hat{A} \psi d\tau$                          | (varies)                           | 7            |

(1) The circumflex (or “hat”),  $\hat{\phantom{x}}$ , serves to distinguish an operator from an algebraic quantity. This definition applies to a coordinate representation, where  $\nabla$  denotes the nabla operator (see Section 4.2, p. 107).

(2) Capital and lower case  $\psi$  are commonly used for the time-dependent function  $\Psi(x, t)$  and the amplitude function  $\psi(x)$  respectively. Thus for a stationary state  $\Psi(x, t) = \psi(x)\exp(-iEt/\hbar)$ .

(3) For the normalized wavefunction of a single particle in three-dimensional space the appropriate SI unit is given in parentheses. Results in quantum chemistry, however, are commonly expressed in terms of atomic units (see Section 3.9.2, p. 95 and Section 7.3 (iv), p. 145; and reference [22]). If distances, energies, angular momenta, charges and masses are all expressed as dimensionless ratios  $r/a_0$ ,  $E/E_h$ , etc., then all quantities are dimensionless.

(4)  $P_l^{|m|}$  denotes the associated Legendre function of degree  $l$  and order  $|m|$ .  $N_{l|m|}$  is a normalization factor.

(5)  $\psi^*$  is the complex conjugate of  $\psi$ . For an anti-symmetrized  $n$  electron wavefunction  $\Psi(r_1, \dots, r_n)$ , the total probability density of electrons is  $\int_2 \dots \int_n \Psi^* \Psi \, d\tau_2 \dots d\tau_n$ , where the integration extends over the coordinates of all electrons but one.

(6)  $-e$  is the charge of an electron.

(7) The unit is the same as for the physical quantity  $A$  that the operator represents.

| <i>Name</i>                                  | <i>Symbol</i>                              | <i>Definition</i>                                        | <i>SI unit</i> | <i>Notes</i> |
|----------------------------------------------|--------------------------------------------|----------------------------------------------------------|----------------|--------------|
| hermitian conjugate<br>of operator $\hat{A}$ | $\hat{A}^\dagger$                          | $(A^\dagger)_{ij} = (A_{ji})^*$                          | (varies)       | 7            |
| commutator<br>of $\hat{A}$ and $\hat{B}$     | $[\hat{A}, \hat{B}], [\hat{A}, \hat{B}]_-$ | $[\hat{A}, \hat{B}] = \hat{A}\hat{B} - \hat{B}\hat{A}$   | (varies)       | 8            |
| anticommutator<br>of $\hat{A}$ and $\hat{B}$ | $[\hat{A}, \hat{B}]_+$                     | $[\hat{A}, \hat{B}]_+ = \hat{A}\hat{B} + \hat{B}\hat{A}$ | (varies)       | 8            |
| angular momentum<br>operators                | see SPECTROSCOPY, Section 2.6.1, p. 30     |                                                          |                |              |
| spin wavefunction                            | $\alpha; \beta$                            |                                                          | 1              | 9            |

### Hückel molecular orbital theory (HMO)

|                                  |                      |                                               |                   |           |
|----------------------------------|----------------------|-----------------------------------------------|-------------------|-----------|
| atomic-orbital basis<br>function | $\chi_r$             |                                               | $\text{m}^{-3/2}$ | 3         |
| molecular orbital                | $\phi_i$             | $\phi_i = \sum_r \chi_r c_{ri}$               | $\text{m}^{-3/2}$ | 3, 10     |
| coulomb integral                 | $H_{rr}, \alpha_r$   | $H_{rr} = \int \chi_r^* \hat{H} \chi_r d\tau$ | J                 | 3, 10, 11 |
| resonance integral               | $H_{rs}, \beta_{rs}$ | $H_{rs} = \int \chi_r^* \hat{H} \chi_s d\tau$ | J                 | 3, 10, 12 |
| energy parameter                 | $x$                  | $-x = (\alpha - E) / \beta$                   | 1                 | 13        |
| overlap integral                 | $S_{rs}, S$          | $S_{rs} = \int \chi_r^* \chi_s d\tau$         | 1                 | 10        |
| charge order                     | $q_r$                | $q_r = \sum_{i=1}^n b_i c_{ri}^2$             | 1                 | 14, 15    |
| bond order                       | $p_{rs}$             | $p_{rs} = \sum_{i=1}^n b_i c_{ri} c_{si}$     | 1                 | 15, 16    |

(8) The unit is the same as for the product of the physical quantities  $A$  and  $B$ .

(9) The spin wavefunctions of a single electron,  $\alpha$  and  $\beta$ , are defined by the matrix elements of the  $z$  component of the spin angular momentum,  $\hat{s}_z$ , by the relations  $\langle \alpha | \hat{s}_z | \alpha \rangle = +(1/2)$ ,  $\langle \beta | \hat{s}_z | \beta \rangle = -(1/2)$ ,  $\langle \beta | \hat{s}_z | \alpha \rangle = \langle \alpha | \hat{s}_z | \beta \rangle = 0$  in units of  $\hbar$ . The total electron spin wavefunctions of an atom with many electrons are denoted by Greek letters  $\alpha, \beta, \gamma$ , etc. according to the value of  $\sum m_s$ , starting from the greatest down to the least.

(10)  $\hat{H}$  is an effective hamiltonian for a single electron,  $i$  and  $j$  label the molecular orbitals, and  $r$  and  $s$  label the atomic orbitals. In Hückel MO theory,  $H_{rs}$  is taken to be non-zero only for bonded pairs of atoms  $r$  and  $s$ , and all  $S_{rs}$  are assumed to be zero for  $r \neq s$ .

(11) Note that the name ‘‘coulomb integral’’ has a different meaning in HMO theory (where it refers to the energy of the orbital  $\chi_r$  in the field of the nuclei) from Hartree-Fock theory discussed below (where it refers to a two-electron repulsion integral).

(12) This expression describes a bonding interaction between atomic orbitals  $r$  and  $s$ . For an anti-bonding interaction, the corresponding resonance integral is given by the negative value of the resonance integral for the bonding interaction.

(13) In the simplest application of Hückel theory to the  $\pi$  electrons of planar conjugated hydrocarbons,  $\alpha$  is taken to be the same for all carbon atoms, and  $\beta$  to be the same for all bonded pairs of carbon atoms; it is then customary to write the Hückel secular determinant in terms of the dimensionless parameter  $x$ .

(14)  $-eq_r$  is the electronic charge on atom  $r$ .  $q_r$  specifies the contribution of all  $n$   $\pi$  electrons to the total charge at center  $r$ , with  $\sum q_r = n$ .

(15)  $b_i$  gives the number of electrons which occupy a given orbital energy level  $\varepsilon_i$ ; for non-degenerate orbitals,  $b_i$  can take the values 0, 1, or 2.

(16)  $p_{rs}$  is the bond order between atoms  $r$  and  $s$ .

### 2.4.1 Ab initio Hartree-Fock self-consistent field theory (ab initio SCF)

Results in quantum chemistry are typically expressed in atomic units (see Section 3.9.1, p. 94 and Section 7.3 (iv), p. 145). In the remaining tables of this section all lengths, energies, masses, charges and angular momenta are expressed as dimensionless ratios to the corresponding atomic units,  $a_0$ ,  $E_h$ ,  $m_e$ ,  $e$  and  $\hbar$  respectively. Thus all quantities become dimensionless, and the SI unit column is therefore omitted.

| Name                                         | Symbol                                               | Definition                                                                                                      | Notes      |
|----------------------------------------------|------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------|------------|
| molecular orbital                            | $\phi_i(\mu)$                                        |                                                                                                                 | 17         |
| molecular spin orbital                       | $\phi_i(\mu)\alpha(\mu);$<br>$\phi_i(\mu)\beta(\mu)$ |                                                                                                                 | 17         |
| total wavefunction                           | $\Psi$                                               | $\Psi = (n!)^{-1/2} \ \phi_i(\mu)\ $                                                                            | 17, 18     |
| core hamiltonian of<br>a single electron     | $\hat{H}_\mu^{\text{core}}$                          | $\hat{H}_\mu = -(1/2)\nabla_\mu^2 - \sum_A Z_A/r_{\mu A}$                                                       | 17, 19     |
| one-electron integrals:                      |                                                      |                                                                                                                 |            |
| expectation value of<br>the core hamiltonian | $H_{ii}$                                             | $H_{ii} = \int \phi_i^*(1)\hat{H}_1^{\text{core}}\phi_i(1)d\tau_1$                                              | 17, 19     |
| two-electron repulsion<br>integrals:         |                                                      |                                                                                                                 |            |
| coulomb integral                             | $J_{ij}$                                             | $J_{ij} = \iint \phi_i^*(1)\phi_j^*(2)\frac{1}{r_{12}}\phi_i(1)\phi_j(2)d\tau_1d\tau_2$                         | 17, 20     |
| exchange integral                            | $K_{ij}$                                             | $K_{ij} = \iint \phi_i^*(1)\phi_j^*(2)\frac{1}{r_{12}}\phi_j(1)\phi_i(2)d\tau_1d\tau_2$                         | 17, 20     |
| one-electron orbital<br>energy               | $\varepsilon_i$                                      | $\varepsilon_i = H_{ii} + \sum_j (2J_{ij} - K_{ij})$                                                            | 17, 21     |
| total electronic energy                      | $E$                                                  | $E = 2 \sum_i H_{ii} + \sum_i \sum_j (2J_{ij} - K_{ij})$<br>$= \sum_i (\varepsilon_i + H_{ii})$                 | 17, 21, 22 |
| coulomb operator                             | $\hat{J}_i$                                          | $\hat{J}_i\phi_j(2) = \left\langle \phi_i(1) \left  \frac{1}{r_{12}} \right  \phi_i(1) \right\rangle \phi_j(2)$ | 17         |
| exchange operator                            | $\hat{K}_i$                                          | $\hat{K}_i\phi_j(2) = \left\langle \phi_i(1) \left  \frac{1}{r_{12}} \right  \phi_j(1) \right\rangle \phi_i(2)$ | 17         |
| Fock operator                                | $\hat{F}$                                            | $\hat{F} = \hat{H}^{\text{core}} + \sum_i \left( 2\hat{J}_i - \hat{K}_i \right)$                                | 17, 21, 23 |

(17) The indices  $i$  and  $j$  label the molecular orbitals, and either  $\mu$  or the numerals 1 and 2 label the electron coordinates.

(18) The double vertical bars denote an anti-symmetrized product of the occupied molecular spin orbitals  $\phi_i\alpha$  and  $\phi_i\beta$  (sometimes denoted  $\phi_i$  and  $\bar{\phi}_i$ ); for a closed-shell system  $\Psi$  would be a normalized Slater determinant.  $(n!)^{-1/2}$  is the normalization factor and  $n$  the number of electrons.

(19)  $Z_A$  is the proton number (charge number) of nucleus A, and  $r_{\mu A}$  is the distance of electron  $\mu$  from nucleus A.  $H_{ii}$  is the energy of an electron in orbital  $\phi_i$  in the field of the core.

(20) The inter-electron repulsion integral is written in various shorthand notations: In  $J_{ij} = \langle ij|ij \rangle$  the first and third indices refer to the index of electron 1 and the second and fourth indices to electron 2. In  $J_{ij} = (i^*i|j^*j)$ , the first two indices refer to electron 1 and the second two indices to electron 2. Usually the functions are real and the stars are omitted. The exchange integral is written in various shorthand notations with the same index convention as described:  $K_{ij} = \langle ij|ji \rangle$  or  $K_{ij} = (i^*j|j^*i)$ .

(21) These relations apply to closed-shell systems only, and the sums extend over the occupied molecular orbitals.

(22) The sum over  $j$  includes the term with  $j = i$ , for which  $J_{ii} = K_{ii}$ , so that this term in the sum simplifies to give  $2J_{ii} - K_{ii} = J_{ii}$ .

### 2.4.2 Hartree-Fock-Roothaan SCF theory, using molecular orbitals expanded as linear combinations of atomic-orbital basis functions (LCAO-MO theory)

| Name                                | Symbol    | Definition                                                                                                    | Notes  |
|-------------------------------------|-----------|---------------------------------------------------------------------------------------------------------------|--------|
| atomic-orbital basis function       | $\chi_r$  |                                                                                                               | 24     |
| molecular orbital                   | $\phi_i$  | $\phi_i = \sum_r \chi_r c_{ri}$                                                                               |        |
| overlap matrix element              | $S_{rs}$  | $S_{rs} = \int \chi_r^* \chi_s d\tau$ , $\sum_{r,s} c_{ri}^* S_{rs} c_{sj} = \delta_{ij}$                     |        |
| density matrix element              | $P_{rs}$  | $P_{rs} = 2 \sum_i^{\text{occ}} c_{ri}^* c_{si}$                                                              | 25     |
| integrals over the basis functions: |           |                                                                                                               |        |
| one-electron integrals              | $H_{rs}$  | $H_{rs} = \int \chi_r^* (1) \hat{H}_1^{\text{core}} \chi_s(1) d\tau_1$                                        |        |
| two-electron integrals              | $(rs tu)$ | $(rs tu) = \iint \chi_r^* (1) \chi_s (1) \frac{1}{r_{12}} \chi_t^* (2) \chi_u (2) d\tau_1 d\tau_2$            | 26, 27 |
| total electronic energy             | $E$       | $E = \sum_r \sum_s P_{rs} H_{rs} + (1/2) \sum_r \sum_s \sum_t \sum_u P_{rs} P_{tu} [(rs tu) - (1/2) (ru ts)]$ | 25, 27 |
| matrix element of the Fock operator | $F_{rs}$  | $F_{rs} = H_{rs} + \sum_t \sum_u P_{tu} [(rs tu) - (1/2) (ru ts)]$                                            | 25, 28 |

(Notes continued)

(23) The Hartree-Fock equations read  $(\hat{F} - \varepsilon_j)\phi_j = 0$ . Note that the definition of the Fock operator involves all of its eigenfunctions  $\phi_i$  through the coulomb and exchange operators,  $\hat{J}_i$  and  $\hat{K}_i$ .

(24) The indices  $r$  and  $s$  label the basis functions. In numerical computations the basis functions are either taken as Slater-type orbitals (STO) or as Gaussian-type orbitals (GTO). An STO basis function in spherical polar coordinates has the general form  $\chi(r, \theta, \phi) = N r^{n-1} \exp(-\zeta_{nl} r) Y_{lm}(\theta, \phi)$ , where  $\zeta_{nl}$  is a shielding parameter representing the effective charge in the state with quantum numbers  $n$  and  $l$ . GTO functions are typically expressed in cartesian space coordinates, in the form  $\chi(x, y, z) = N x^a y^b z^c \exp(-\alpha r^2)$ . Commonly, a linear combination of such functions with varying exponents  $\alpha$  is used in such a way as to model an STO.  $N$  denotes a normalization factor.

(25) For closed-shell species with two electrons per occupied orbital. The sum extends over all occupied molecular orbitals.  $P_{rs}$  may also be called the bond order between atoms  $r$  and  $s$ .

(26) The contracted notation for two-electron integrals over the basis functions,  $(rs|tu)$ , is based on the same convention outlined in note 20.

(27) Here the two-electron integral is expressed in terms of integrals over the spatial atomic-orbital basis functions. The matrix elements  $H_{ii}$ ,  $J_{ij}$ , and  $K_{ij}$  may be similarly expressed in terms of integrals over the spatial atomic-orbital basis functions according to the following equations:

$$\begin{aligned}
 H_{ii} &= \sum_r \sum_s c_{ri}^* c_{si} H_{rs} \\
 J_{ij} &= (i^* i | j^* j) = \sum_r \sum_s \sum_t \sum_u c_{ri}^* c_{si} c_{tj}^* c_{uj} (r^* s | t^* u) \\
 K_{ij} &= (i^* j | j^* i) = \sum_r \sum_s \sum_t \sum_u c_{ri}^* c_{si} c_{tj}^* c_{uj} (r^* u | t^* s)
 \end{aligned}$$

(28) The Hartree-Fock-Roothaan SCF equations, expressed in terms of the matrix elements of the Fock operator  $F_{rs}$ , and the overlap matrix elements  $S_{rs}$ , take the form:

$$\sum_s (F_{rs} - \varepsilon_i S_{rs}) c_{si} = 0$$

## 2.5 ATOMS AND MOLECULES

The names and symbols recommended here are in agreement with those recommended by IUPAP [4] and ISO [5.i]. Additional quantities and symbols used in atomic, nuclear and plasma physics can be found in [4,5.j].

| <i>Name</i>                       | <i>Symbol</i> | <i>Definition</i>                     | <i>SI unit</i>  | <i>Notes</i> |
|-----------------------------------|---------------|---------------------------------------|-----------------|--------------|
| nucleon number, mass number       | $A$           |                                       | 1               |              |
| proton number, atomic number      | $Z$           |                                       | 1               |              |
| neutron number                    | $N$           | $N = A - Z$                           | 1               |              |
| electron mass                     | $m_e$         |                                       | kg              | 1, 2         |
| mass of atom, atomic mass         | $m_a, m$      |                                       | kg              |              |
| atomic mass constant              | $m_u$         | $m_u = m_a(^{12}\text{C})/12$         | kg              | 1, 3         |
| mass excess                       | $\Delta$      | $\Delta = m_a - Am_u$                 | kg              |              |
| elementary charge                 | $e$           | proton charge                         | C               | 2            |
| Planck constant                   | $h$           |                                       | J s             |              |
| Planck constant divided by $2\pi$ | $\hbar$       | $\hbar = h/2\pi$                      | J s             | 2            |
| Bohr radius                       | $a_0$         | $a_0 = 4\pi\epsilon_0\hbar^2/m_e e^2$ | m               | 2            |
| Hartree energy                    | $E_h$         | $E_h = \hbar^2/m_e a_0^2$             | J               | 2            |
| Rydberg constant                  | $R_\infty$    | $R_\infty = E_h/2hc$                  | $\text{m}^{-1}$ |              |
| fine-structure constant           | $\alpha$      | $\alpha = e^2/4\pi\epsilon_0\hbar c$  | 1               |              |
| ionization energy                 | $E_i, I$      |                                       | J               | 4            |
| electron affinity                 | $E_{ea}, A$   |                                       | J               | 4            |
| electronegativity                 | $\chi$        | $\chi = (1/2)(E_i + E_{ea})$          | J               | 5            |
| dissociation energy               | $E_d, D$      |                                       | J               |              |
| from the ground state             | $D_0$         |                                       | J               | 6            |
| from the potential minimum        | $D_e$         |                                       | J               | 6            |

(1) Analogous symbols are used for other particles with subscripts: p for proton, n for neutron, a for atom, N for nucleus, etc.

(2) This quantity is also used as an atomic unit (see Section 3.9.1, p. 94 and Section 7.3 (iv), p. 145).

(3)  $m_u$  is equal to the unified atomic mass unit, with symbol u, i.e.  $m_u = 1 \text{ u}$  (see Section 3.7, p. 92). The name dalton, with symbol Da, is used as an alternative name for the unified atomic mass unit [23].

(4) The ionization energy is frequently called the ionization potential ( $I_p$ ). The electron affinity is the energy needed to detach an electron.

(5) The concept of electronegativity was introduced by L. Pauling as the power of an atom in a molecule to attract electrons to itself. There are several ways of defining this quantity [24]. The one given in the table has a clear physical meaning of energy and is due to R. S. Mulliken. The most frequently used scale, due to Pauling, is based on bond dissociation energies  $E_d$  in eV and it is relative in the sense that the values are dimensionless and that only electronegativity differences are defined. For atoms A and B

$$\chi_{r,A} - \chi_{r,B} = \sqrt{\frac{E_d(\text{AB})}{\text{eV}} - \frac{1}{2} \frac{[E_d(\text{AA}) + E_d(\text{BB})]}{\text{eV}}}$$

where  $\chi_r$  denotes the Pauling relative electronegativity. The scale is chosen so as to make the relative electronegativity of hydrogen  $\chi_{r,H} = 2.1$ . There is a difficulty in choosing the sign of the square root, which determines the sign of  $\chi_{r,A} - \chi_{r,B}$ . Pauling made this choice intuitively.

(6) The symbols  $D_0$  and  $D_e$  are used for dissociation energies of diatomic and polyatomic molecules.

| <i>Name</i>                                      | <i>Symbol</i>                     | <i>Definition</i>                                                                | <i>SI unit</i>                  | <i>Notes</i> |
|--------------------------------------------------|-----------------------------------|----------------------------------------------------------------------------------|---------------------------------|--------------|
| principal quantum number<br>(hydrogen-like atom) | $n$                               | $E = hcZ^2 R_\infty / n^2$                                                       | 1                               | 7            |
| angular momentum<br>quantum numbers              | see SPECTROSCOPY, Section 2.6     |                                                                                  |                                 |              |
| magnetic dipole moment<br>of a molecule          | $\mathbf{m}, \boldsymbol{\mu}$    | $E_p = -\mathbf{m} \cdot \mathbf{B}$                                             | J T <sup>-1</sup>               | 8            |
| magnetizability<br>of a molecule                 | $\xi$                             | $\mathbf{m} = \xi \mathbf{B}$                                                    | J T <sup>-2</sup>               |              |
| Bohr magneton                                    | $\mu_B$                           | $\mu_B = e\hbar/2m_e$                                                            | J T <sup>-1</sup>               |              |
| nuclear magneton                                 | $\mu_N$                           | $\mu_N = e\hbar/2m_p = (m_e/m_p) \mu_B$                                          | J T <sup>-1</sup>               |              |
| gyromagnetic ratio,<br>(magnetogyric ratio)      | $\gamma$                          | $\gamma_e = -g_e \mu_B / \hbar$                                                  | s <sup>-1</sup> T <sup>-1</sup> | 9            |
| $g$ -factor                                      | $g, g_e$                          | $g_e = -\gamma_e (2m_e/e)$                                                       | 1                               | 10           |
| nuclear $g$ -factor                              | $g_N$                             | $g_N = \gamma_N (2m_p/e)$                                                        | 1                               | 10           |
| Larmor angular frequency                         | $\boldsymbol{\omega}_L$           | $\boldsymbol{\omega}_L = -\gamma \mathbf{B}$                                     | s <sup>-1</sup>                 | 11           |
| Larmor frequency                                 | $\nu_L$                           | $\nu_L = \omega_L / 2\pi$                                                        | Hz                              |              |
| relaxation time,<br>longitudinal                 | $T_1$                             |                                                                                  | s                               | 12           |
| transverse                                       | $T_2$                             |                                                                                  | s                               | 12           |
| electric dipole moment<br>of a molecule          | $\mathbf{p}, \boldsymbol{\mu}$    | $E_p = -\mathbf{p} \cdot \mathbf{E}$                                             | C m                             | 13           |
| quadrupole moment<br>of a molecule               | $\mathbf{Q}; \boldsymbol{\Theta}$ | $E_p = (1/2)\mathbf{Q} : \mathbf{V}'' = (1/3)\boldsymbol{\Theta} : \mathbf{V}''$ | C m <sup>2</sup>                | 14           |

(7) For an electron in the central coulomb field of an infinitely heavy nucleus of atomic number  $Z$ .

(8) Magnetic moments of specific particles may be denoted by subscripts, e.g.  $\mu_e, \mu_p, \mu_n$  for an electron, a proton, and a neutron. Tabulated values usually refer to the maximum expectation value of the  $z$  component. Values for stable nuclei are given in Section 6.3, p. 121.

(9) The gyromagnetic ratio for a nucleus is  $\gamma_N = g_N \mu_N / \hbar$ .

(10) For historical reasons,  $g_e > 0$ .  $e$  is the (positive) elementary charge, therefore  $\gamma_e < 0$ . For nuclei,  $\gamma_N$  and  $g_N$  have the same sign. A different sign convention for the electronic  $g$ -factor is discussed in [25].

(11) This is a vector quantity with magnitude  $\omega_L$ . This quantity is sometimes called Larmor circular frequency.

(12) These quantities are used in the context of saturation effects in spectroscopy, particularly spin-resonance spectroscopy (see Section 2.6, p. 27–28).

(13) See Section 2.6, note 9, p. 26.

(14) The quadrupole moment of a molecule may be represented either by the tensor  $\mathbf{Q}$ , defined by an integral over the charge density  $\rho$ :

$$Q_{\alpha\beta} = \int r_\alpha r_\beta \rho \, dV$$

in which  $\alpha$  and  $\beta$  denote  $x, y$  or  $z$ , or by the tensor  $\boldsymbol{\Theta}$  of trace zero defined by

$$\Theta_{\alpha\beta} = (1/2) \int (3r_\alpha r_\beta - \delta_{\alpha\beta} r^2) \rho \, dV = (1/2) [3Q_{\alpha\beta} - \delta_{\alpha\beta} (Q_{xx} + Q_{yy} + Q_{zz})]$$

$\mathbf{V}''$  is the second derivative of the electronic potential:

$$V_{\alpha\beta}'' = -q_{\alpha\beta} = \partial^2 V / \partial \alpha \partial \beta$$

The contribution to the potential energy is then given by

$$E_p = (1/2)\mathbf{Q} : \mathbf{V}'' = (1/2) \sum_\alpha \sum_\beta Q_{\alpha\beta} V_{\alpha\beta}''$$

| <i>Name</i>                                                 | <i>Symbol</i>      | <i>Definition</i>                                                         | <i>SI unit</i>                                | <i>Notes</i> |
|-------------------------------------------------------------|--------------------|---------------------------------------------------------------------------|-----------------------------------------------|--------------|
| quadrupole moment<br>of a nucleus                           | $eQ$               | $eQ = 2 \langle \Theta_{zz} \rangle$                                      | C m <sup>2</sup>                              | 15           |
| electric field gradient<br>tensor                           | $\mathbf{q}$       | $q_{\alpha\beta} = -\partial^2 V / \partial \alpha \partial \beta$        | V m <sup>-2</sup>                             |              |
| quadrupole interaction<br>energy tensor                     | $\chi$             | $\chi_{\alpha\beta} = eQq_{\alpha\beta}$                                  | J                                             | 16           |
| electric polarizability<br>of a molecule                    | $\alpha$           | $\alpha_{ab} = \partial p_a / \partial E_b$                               | C <sup>2</sup> m <sup>2</sup> J <sup>-1</sup> | 17           |
| 1st hyper-polarizability                                    | $\beta$            | $\beta_{abc} = \partial^2 p_a / \partial E_b \partial E_c$                | C <sup>3</sup> m <sup>3</sup> J <sup>-2</sup> | 17           |
| 2nd hyper-polarizability                                    | $\gamma$           | $\gamma_{abcd} = \partial^3 p_a / \partial E_b \partial E_c \partial E_d$ | C <sup>4</sup> m <sup>4</sup> J <sup>-3</sup> | 17           |
| activity (of a radio-<br>active substance)                  | $A$                | $A = -dN_B/dt$                                                            | Bq                                            | 18           |
| decay (rate) constant,<br>disintegration (rate)<br>constant | $\lambda, k$       | $A = \lambda N_B$                                                         | s <sup>-1</sup>                               | 18           |
| half life                                                   | $t_{1/2}, T_{1/2}$ | $N_B(t_{1/2}) = N_B(0) / 2$                                               | s                                             | 18, 19       |
| mean life, lifetime                                         | $\tau$             | $\tau = 1/\lambda$                                                        | s                                             | 19           |
| level width                                                 | $\Gamma$           | $\Gamma = \hbar/\tau$                                                     | J                                             |              |
| disintegration energy                                       | $Q$                |                                                                           | J                                             |              |
| cross section                                               | $\sigma$           |                                                                           | m <sup>2</sup>                                |              |
| electroweak charge of a<br>nucleus                          | $Q_W$              | $Q_W \approx Z(1 - 4 \sin^2 \theta_W) - N$                                | 1                                             | 20           |

(15) Nuclear quadrupole moments are conventionally defined in a different way from molecular quadrupole moments.  $Q$  has the dimension of an area and  $e$  is the elementary charge.  $eQ$  is taken to be twice the maximum expectation value of the  $zz$  tensor element (see note 14). The values of  $Q$  for some nuclei are listed in Section 6.3, p. 121.

(16) The nuclear quadrupole interaction energy tensor  $\chi$  is usually quoted in MHz, corresponding to the value of  $eQq/h$ , although the  $h$  is usually omitted.

(17) The polarizability  $\alpha$  and the hyper-polarizabilities  $\beta, \gamma, \dots$  are the coefficients in the expansion of the dipole moment  $\mathbf{p}$  in powers of the electric field strength  $\mathbf{E}$  (see Section 2.3, note 7, p. 16). The expansion of the component  $a$  is given by

$$p_a = p_a^{(0)} + \sum_b \alpha_{ab} E_b + (1/2) \sum_{bc} \beta_{abc} E_b E_c + (1/6) \sum_{bcd} \gamma_{abcd} E_b E_c E_d + \dots$$

in which  $\alpha_{ab}$ ,  $\beta_{abc}$ , and  $\gamma_{abcd}$  are elements of the tensors  $\alpha$ ,  $\beta$ , and  $\gamma$  of rank 2, 3, and 4, respectively. The components of these tensors are distinguished by the subscript indices  $abc\dots$ , as indicated in the definitions, the first index  $a$  always denoting the component of  $p$ , and the subsequent indices the components of the electric field. The polarizability and the hyper-polarizabilities exhibit symmetry properties. Thus  $\alpha$  is commonly a symmetric tensor, and all components of  $\beta$  are zero for a molecule with a centre of symmetry, etc. Values of the polarizability are commonly quoted as the value  $\alpha/4\pi\epsilon_0$ , which is a volume. The value is commonly expressed in the unit Å<sup>3</sup> (Å should not be used, see Section 2.6, note 11, p. 27) or in the unit  $a_0^3$  (atomic units, see Section 3.9.1, p. 94). Similar comments apply to the hyper-polarizabilities with  $\beta/(4\pi\epsilon_0)^2$  in units of  $a_0^5 e^{-1}$ , and  $\gamma/(4\pi\epsilon_0)^3$  in units of  $a_0^7 e^{-2}$ , etc.

(18)  $N_B$  is the number of decaying entities B (1 Bq = 1 s<sup>-1</sup>, see Section 3.4, p. 89).

(19) Half lives and mean lives are commonly given in years (unit a), see Section 7.2, note 4, p. 137.  $t_{1/2} = \tau \ln 2$  for exponential decays.

(20) The electroweak charge of a nucleus is approximately given by the neutron number  $N$  and the proton number  $Z$  with the weak mixing angle  $\theta_W$  (see Chapter 5, p. 111). It is important in calculations of atomic and molecular properties including the weak nuclear interaction [26].

## 2.6 SPECTROSCOPY

This section has been considerably extended compared with the original Manual [1.a–1.c] and with the corresponding section in the IUPAP document [4]. It is based on the recommendations of the ICSU Joint Commission for Spectroscopy [27, 28] and current practice in the field which is well represented in the books by Herzberg [29–31]. The IUPAC Commission on Molecular Structure and Spectroscopy has also published recommendations which have been taken into account [32–43].

| <i>Name</i>                                      | <i>Symbol</i>                                         | <i>Definition</i>                                               | <i>SI unit</i>  | <i>Notes</i> |
|--------------------------------------------------|-------------------------------------------------------|-----------------------------------------------------------------|-----------------|--------------|
| total term                                       | $T$                                                   | $T = E_{\text{tot}}/hc$                                         | $\text{m}^{-1}$ | 1, 2         |
| transition wavenumber                            | $\tilde{\nu}$                                         | $\tilde{\nu} = T' - T''$                                        | $\text{m}^{-1}$ | 1            |
| transition frequency                             | $\nu$                                                 | $\nu = (E' - E'')/h$                                            | Hz              |              |
| electronic term                                  | $T_e$                                                 | $T_e = E_e/hc$                                                  | $\text{m}^{-1}$ | 1, 2         |
| vibrational term                                 | $G$                                                   | $G = E_{\text{vib}}/hc$                                         | $\text{m}^{-1}$ | 1, 2         |
| rotational term                                  | $F$                                                   | $F = E_{\text{rot}}/hc$                                         | $\text{m}^{-1}$ | 1, 2         |
| spin-orbit coupling constant                     | $A$                                                   | $T_{\text{so}} = A < \hat{\mathbf{L}} \cdot \hat{\mathbf{S}} >$ | $\text{m}^{-1}$ | 1, 3         |
| principal moments<br>of inertia                  | $I_A; I_B; I_C$                                       | $I_A \leq I_B \leq I_C$                                         | $\text{kg m}^2$ |              |
| rotational constants,<br>in wavenumber           | $\tilde{A}; \tilde{B}; \tilde{C}$                     | $\tilde{A} = h/8\pi^2 c I_A$                                    | $\text{m}^{-1}$ | 1, 2         |
| in frequency                                     | $A; B; C$                                             | $A = h/8\pi^2 I_A$                                              | Hz              |              |
| inertial defect                                  | $\Delta$                                              | $\Delta = I_C - I_A - I_B$                                      | $\text{kg m}^2$ |              |
| asymmetry parameter                              | $\kappa$                                              | $\kappa = \frac{2B - A - C}{A - C}$                             | 1               | 4            |
| centrifugal distortion constants,<br>S reduction | $D_J; D_{JK}; D_K; d_1; d_2$                          |                                                                 | $\text{m}^{-1}$ | 5            |
| A reduction                                      | $\Delta_J; \Delta_{JK}; \Delta_K; \delta_J; \delta_K$ |                                                                 | $\text{m}^{-1}$ | 5            |
| harmonic vibration<br>wavenumber                 | $\omega_e; \omega_r$                                  |                                                                 | $\text{m}^{-1}$ | 6            |
| vibrational anharmonicity<br>constant            | $\omega_e x_e; x_{rs}; g_{tt'}$                       |                                                                 | $\text{m}^{-1}$ | 6            |

(1) In spectroscopy the unit  $\text{cm}^{-1}$  is almost always used for the quantity wavenumber, and term values and wavenumbers always refer to the reciprocal wavelength of the equivalent radiation in vacuum. The symbol  $c$  in the definition  $E/hc$  refers to the speed of light in vacuum. Because “wavenumber” is not a number ISO suggests the use of repetency in parallel with wavenumber [5,6]. The use of the word “wavenumber” in place of the unit  $\text{cm}^{-1}$  must be avoided.

(2) Term values and rotational constants are sometimes defined in wavenumber units (e.g.  $T = E/hc$ ), and sometimes in frequency units (e.g.  $T = E/h$ ). When the symbol is otherwise the same, it is convenient to distinguish wavenumber quantities with a tilde (e.g.  $\tilde{\nu}, \tilde{T}, \tilde{A}, \tilde{B}, \tilde{C}$  for quantities defined in wavenumber units), although this is not a universal practice.

(3)  $\hat{\mathbf{L}}$  and  $\hat{\mathbf{S}}$  are electron orbital and electron spin operators, respectively.

(4) The Wang asymmetry parameters are also used: for a near prolate top  $b_p = (C - B) / (2A - B - C)$ , and for a near oblate top  $b_o = (A - B) / (2C - A - B)$ .

(5) S and A stand for the symmetric and asymmetric reductions of the rotational hamiltonian respectively; see [44] for more details on the various possible representations of the centrifugal distortion constants.

(6) For a diatomic molecule:  $G(v) = \omega_e(v + \frac{1}{2}) - \omega_e x_e(v + \frac{1}{2})^2 + \dots$ . For a polyatomic molecule the  $3N - 6$  vibrational modes ( $3N - 5$  if linear) are labeled by the indices  $r, s, t, \dots$ , or  $i, j, k, \dots$ . The index  $r$  is usually assigned to be increasing with descending wavenumber, symmetry species by

| <i>Name</i>                            | <i>Symbol</i>                           | <i>Definition</i>                                   | <i>SI unit</i>  | <i>Notes</i> |
|----------------------------------------|-----------------------------------------|-----------------------------------------------------|-----------------|--------------|
| vibrational quantum numbers            | $v_r; l_t$                              |                                                     | 1               | 6            |
| vibrational fundamental wavenumber     | $\tilde{\nu}_r, \tilde{\nu}_r^0, \nu_r$ | $\tilde{\nu}_r = T(v_r = 1) - T(v_r = 0)$           | $\text{m}^{-1}$ | 6            |
| Coriolis $\zeta$ -constant             | $\zeta_{rs}^\alpha$                     |                                                     | 1               | 7            |
| angular momentum quantum numbers       | see Section 2.6.1, p. 30                |                                                     |                 |              |
| degeneracy, statistical weight         | $g, d, \beta$                           |                                                     | 1               | 8            |
| electric dipole moment of a molecule   | $\mathbf{p}, \boldsymbol{\mu}$          | $E_p = -\mathbf{p} \cdot \mathbf{E}$                | C m             | 9            |
| transition dipole moment of a molecule | $\mathbf{M}, \mathbf{R}$                | $\mathbf{M} = \int \psi^* \mathbf{p} \psi \, d\tau$ | C m             | 9, 10        |

(6) (continued) symmetry species starting with the totally symmetric species. The index  $t$  is kept for degenerate modes. The vibrational term formula is

$$G(v) = \sum_r \omega_r (v_r + d_r/2) + \sum_{r \leq s} x_{rs} (v_r + d_r/2) (v_s + d_s/2) + \sum_{t \leq t'} g_{tt'} l_t l_{t'} + \dots$$

Another common term formula is defined with respect to the vibrational ground state

$$T(v) = \sum_i v_i \tilde{\nu}_i' + \sum_{r \leq s} x_{rs}' v_r v_s + \sum_{t \leq t'} g_{tt}' l_t l_{t'} + \dots$$

these being commonly used as the diagonal elements of effective vibrational hamiltonians.

(7) Frequently the Coriolis coupling constants  $\xi_\alpha^{v'v}$  with units of  $\text{cm}^{-1}$  are used as effective hamiltonian constants ( $\alpha = x, y, z$ ). For two fundamental vibrations with harmonic wavenumbers  $\omega_r$  and  $\omega_s$  these are connected with  $\zeta_{rs}^\alpha$  by the equation ( $v_r = 1$  and  $v_s = 1$ )

$$\xi_\alpha^{v'v} = \tilde{B}_\alpha \zeta_{rs}^\alpha [\sqrt{\omega_s/\omega_r} + \sqrt{\omega_r/\omega_s}]$$

where  $\tilde{B}_\alpha$  is the  $\alpha$  rotational constant. A similar equation applies with  $B_\alpha$  if  $\xi_\alpha^{v'v}$  is defined as a quantity with frequency units.

(8)  $d$  is usually used for vibrational degeneracy, and  $\beta$  for nuclear spin degeneracy.

(9)  $E_p$  is the potential energy of a dipole with electric dipole moment  $\mathbf{p}$  in an electric field of strength  $\mathbf{E}$ . Molecular dipole moments are commonly expressed in the non-SI unit debye, where  $1 \text{ D} \approx 3.335 \, 64 \times 10^{-30} \text{ C m}$ . The SI unit C m is inconvenient for expressing molecular dipole moments, which results in the continued use of the deprecated debye (D). A convenient alternative is to use the atomic unit,  $ea_0$ . Another way of expressing dipole moments is to quote the electric dipole lengths,  $l_p = p/e$ , analogous to the way the nuclear quadrupole areas are quoted (see Section 2.5, notes 14 and 15, p. 23 and 24 and Section 6.3, p. 121). This gives the distance between two elementary charges of the equivalent dipole and conveys a clear picture in relation to molecular dimensions (see also Section 2.3, note 8, p. 17).

| <i>Examples</i>  | Dipole moment          |              |          | Dipole length   |
|------------------|------------------------|--------------|----------|-----------------|
|                  | SI                     |              | a.u.     |                 |
|                  | $p/\text{C m}$         | $p/\text{D}$ | $p/ea_0$ | $l_p/\text{pm}$ |
| HCl              | $3.60 \times 10^{-30}$ | 1.08         | 0.425    | 22.5            |
| H <sub>2</sub> O | $6.23 \times 10^{-30}$ | 1.87         | 0.736    | 38.9            |
| NaCl             | $3.00 \times 10^{-29}$ | 9.00         | 3.54     | 187             |

(10) For quantities describing line and band intensities see Section 2.7, p. 34–41.

| <i>Name</i>                                                                  | <i>Symbol</i>                    | <i>Definition</i>                                                                   | <i>SI unit</i>                        | <i>Notes</i> |
|------------------------------------------------------------------------------|----------------------------------|-------------------------------------------------------------------------------------|---------------------------------------|--------------|
| interatomic distances,                                                       |                                  |                                                                                     |                                       | 11, 12       |
| equilibrium                                                                  | $r_e$                            |                                                                                     | m                                     |              |
| zero-point average                                                           | $r_z$                            |                                                                                     | m                                     |              |
| ground state                                                                 | $r_0$                            |                                                                                     | m                                     |              |
| substitution structure                                                       | $r_s$                            |                                                                                     | m                                     |              |
| vibrational coordinates,                                                     |                                  |                                                                                     |                                       | 11           |
| internal                                                                     | $R_i, r_i, \theta_j$ , etc.      |                                                                                     | (varies)                              |              |
| symmetry                                                                     | $S_j$                            |                                                                                     | (varies)                              |              |
| normal                                                                       |                                  |                                                                                     |                                       |              |
| mass adjusted                                                                | $Q_r$                            |                                                                                     | kg <sup>1/2</sup> m                   |              |
| dimensionless                                                                | $q_r$                            |                                                                                     | 1                                     |              |
| vibrational force constants,                                                 |                                  |                                                                                     |                                       |              |
| diatomic                                                                     | $f, (k)$                         | $f = \partial^2 V / \partial r^2$                                                   | J m <sup>-2</sup> , N m <sup>-1</sup> | 11, 13       |
| polyatomic,                                                                  |                                  |                                                                                     |                                       |              |
| internal coordinates                                                         | $f_{ij}$                         | $f_{ij} = \partial^2 V / \partial r_i \partial r_j$                                 | (varies)                              |              |
| symmetry coordinates                                                         | $F_{ij}$                         | $F_{ij} = \partial^2 V / \partial S_i \partial S_j$                                 | (varies)                              |              |
| dimensionless normal                                                         | $\phi_{rst\dots}, k_{rst\dots},$ |                                                                                     | m <sup>-1</sup>                       | 14           |
| coordinates                                                                  | $C_{rst\dots}$                   |                                                                                     |                                       |              |
| <b>electron spin resonance (ESR), electron paramagnetic resonance (EPR):</b> |                                  |                                                                                     |                                       |              |
| gyromagnetic ratio                                                           | $\gamma$                         | $\gamma = \mu / \hbar \sqrt{S(S+1)}$                                                | s <sup>-1</sup> T <sup>-1</sup>       | 15           |
| <i>g</i> -factor                                                             | $g$                              | $h\nu = g\mu_B B$                                                                   | 1                                     | 16           |
| hyperfine coupling constant                                                  |                                  |                                                                                     |                                       |              |
| in liquids                                                                   | $a, A$                           | $\hat{H}_{\text{hfs}}/h = a \hat{\mathbf{S}} \cdot \hat{\mathbf{I}}$                | Hz                                    | 17           |
| in solids                                                                    | $\mathbf{T}$                     | $\hat{H}_{\text{hfs}}/h = \hat{\mathbf{S}} \cdot \mathbf{T} \cdot \hat{\mathbf{I}}$ | Hz                                    | 17           |

(11) Interatomic (internuclear) distances and vibrational displacements are commonly expressed in the non-SI unit ångström, where 1 Å = 10<sup>-10</sup> m = 0.1 nm = 100 pm. Å should not be used.

(12) The various slightly different ways of representing interatomic distances, distinguished by subscripts, involve different vibrational averaging contributions; they are discussed in [45], where the geometrical structures of many free molecules are listed. Only the equilibrium distance  $r_e$  is isotopically invariant, to a good approximation. The effective distance parameter  $r_0$  is estimated from the rotational constants for the ground vibrational state and has a more complicated physical significance for polyatomic molecules.

(13) Force constants are commonly expressed in mdyn Å<sup>-1</sup> = aJ Å<sup>-2</sup> for stretching coordinates, mdyn Å = aJ for bending coordinates, and mdyn = aJ Å<sup>-1</sup> for stretch-bend interactions. Å should not be used (see note 11). See [46] for further details on definitions and notation for force constants.

(14) The force constants in dimensionless normal coordinates are usually defined in wavenumber units by the equation  $V/hc = \sum \phi_{rst\dots} q_r q_s q_t \dots$ , where the summation over the normal coordinate indices  $r, s, t, \dots$  is unrestricted.

(15) The magnitude of  $\gamma$  is obtained from the magnitudes of the magnetic dipole moment  $\mu$  and the angular momentum (in ESR: electron spin  $\mathbf{S}$  for a  $\Sigma$ -state,  $L = 0$ ; in NMR: nuclear spin  $\mathbf{I}$ ).

(16) This gives an effective *g*-factor for a single spin ( $S = 1/2$ ) in an external static field (see Section 2.5, note 10, p. 23).

(17)  $\hat{H}_{\text{hfs}}$  is the hyperfine coupling hamiltonian.  $\hat{\mathbf{S}}$  is the electron spin operator with quantum number  $S$  (see Section 2.6.1, p. 30). The coupling constants  $a$  are usually quoted in MHz, but they are sometimes quoted in magnetic induction units (G or T) obtained by dividing by the conversion factor  $g\mu_B/h$ , which has the SI unit Hz/T;  $g_e\mu_B/h \approx 28.025 \text{ GHz T}^{-1}$  (= 2.8025 MHz G<sup>-1</sup>), where  $g_e$  is the *g*-factor for a free electron. If in liquids the hyperfine coupling is isotropic, the coupling

| <i>Name</i>                                           | <i>Symbol</i>          | <i>Definition</i>                                                        | <i>SI unit</i>                                                     | <i>Notes</i> |
|-------------------------------------------------------|------------------------|--------------------------------------------------------------------------|--------------------------------------------------------------------|--------------|
| <b>nuclear magnetic resonance (NMR):</b>              |                        |                                                                          |                                                                    |              |
| static magnetic flux density<br>of a NMR spectrometer | $B_0$                  |                                                                          | T                                                                  | 18           |
| radiofrequency magnetic<br>flux densities             | $B_1, B_2$             |                                                                          | T                                                                  | 18           |
| spin-rotation interaction<br>tensor                   | $C$                    |                                                                          | Hz                                                                 | 19           |
| spin-rotation coupling<br>constant of nucleus A       | $C_A$                  |                                                                          | Hz                                                                 | 19           |
| dipolar interaction tensor                            | $D$                    |                                                                          | Hz                                                                 | 19           |
| dipolar coupling constant<br>between nuclei A, B      | $D_{AB}$               | $D_{AB} = \frac{\mu_0 \hbar}{8\pi^2} \frac{\gamma_A \gamma_B}{r_{AB}^3}$ | Hz                                                                 | 19           |
| frequency variables of a<br>two-dimensional spectrum  | $F_1, F_2; f_1, f_2$   |                                                                          | Hz                                                                 |              |
| nuclear spin operator<br>for nucleus A                | $\hat{I}_A, \hat{S}_A$ |                                                                          | 1                                                                  | 20           |
| quantum number<br>associated with $\hat{I}_A$         | $I_A$                  |                                                                          | 1                                                                  |              |
| indirect spin coupling tensor                         | $J$                    |                                                                          | Hz                                                                 |              |
| nuclear spin-spin coupling<br>through $n$ bonds       | ${}^n J$               |                                                                          | Hz                                                                 | 21           |
| reduced nuclear spin-spin<br>coupling constant        | $K_{AB}$               | $K_{AB} = \frac{J_{AB}}{h} \frac{2\pi}{\gamma_A} \frac{2\pi}{\gamma_B}$  | T <sup>2</sup> J <sup>-1</sup> , N A <sup>-2</sup> m <sup>-3</sup> | 20           |
| eigenvalue of $\hat{I}_{Az}$                          | $m_A$                  |                                                                          | 1                                                                  | 22           |
| equilibrium macroscopic<br>magnetization per volume   | $M_0$                  |                                                                          | J T <sup>-1</sup> m <sup>-3</sup>                                  | 23           |
| nuclear quadrupole moment                             | $eQ$                   |                                                                          | C m <sup>2</sup>                                                   | 24           |

(17) (continued) constant is a scalar  $a$ . In solids the coupling is anisotropic, and the coupling constant is a 3×3 tensor  $T$ . Similar comments apply to the  $g$ -factor and to the analogous NMR parameters. A convention for the  $g$ -factor has been proposed, in which the sign of  $g$  is positive when the dipole moment is parallel to its angular momentum and negative, when it is anti parallel. Such a choice would require the  $g$ -factors for the electron orbital and spin angular momenta to be negative [25] (see Section 2.5, note 10, p. 23).

(18) The observing ( $B_1$ ) and irradiating ( $B_2$ ) radiofrequency flux densities are associated with frequencies  $\nu_1, \nu_2$  and with nutation angular frequencies  $\Omega_1, \Omega_2$  (around  $B_1, B_2$ , respectively). They are defined through  $\Omega_1 = -\gamma B_1$  and  $\Omega_2 = -\gamma B_2$  (see Section 2.3, notes 8 and 9, p. 17).

(19) The units of interaction strengths are Hz. In the context of relaxation the interaction strengths should be converted to angular frequency units (rad s<sup>-1</sup>, but commonly denoted s<sup>-1</sup>, see note 25).

(20)  $\hat{I}_A, \hat{S}_A$  have components  $\hat{I}_{Ax}, \hat{I}_{Ay}, \hat{I}_{Az}$  or  $\hat{S}_{Ax}, \hat{S}_{Ay}, \hat{S}_{Az}$ , respectively. For another way of defining angular momentum operators with unit J s (see Section 2.6.1, p. 30).

(21) Parentheses may be used to indicate the species of the nuclei coupled, e.g.  $J(^{13}\text{C}, ^1\text{H})$  or, additionally, the coupling path, e.g.  $J(\text{POCF})$ . Where no ambiguity arises, the elements involved can be, alternatively, given as subscripts, e.g.  $J_{\text{CH}}$ . The nucleus of higher mass should be given first. The same applies to the reduced coupling constant.

(22)  $M$  rather than  $m$  is frequently recommended, but most NMR practitioners use  $m$  so as to avoid confusion with magnetization.

(23) This is for a spin system in the presence of a magnetic flux density  $B_0$ .

(24) See Section 2.5, notes 14 and 15, p. 23 and 24.

| <i>Name</i>                                                  | <i>Symbol</i>         | <i>Definition</i>                                                  | <i>SI unit</i>                | <i>Notes</i> |
|--------------------------------------------------------------|-----------------------|--------------------------------------------------------------------|-------------------------------|--------------|
| electric field gradient                                      | $\mathbf{q}$          | $q_{\alpha\beta} = -\partial^2 V / \partial \alpha \partial \beta$ | $\text{V m}^{-2}$             | 25           |
| nuclear quadrupole<br>coupling constant                      | $\chi$                | $\chi = eq_{zz}Q/h$                                                | Hz                            | 19           |
| time dimensions for<br>two-dimensional NMR                   | $t_1, t_2$            |                                                                    | s                             |              |
| spin-lattice relaxation time<br>(longitudinal) for nucleus A | $T_1^A$               |                                                                    | s                             | 26, 27       |
| spin-spin relaxation time<br>(transverse) for nucleus A      | $T_2^A$               |                                                                    | s                             | 26           |
| spin wavefunctions                                           | $\alpha; \beta$       |                                                                    |                               | 28           |
| gyromagnetic ratio                                           | $\gamma$              | $\gamma = \mu/\hbar\sqrt{I(I+1)}$                                  | $\text{s}^{-1} \text{T}^{-1}$ | 15, 29       |
| chemical shift for the<br>nucleus A                          | $\delta_A$            | $\delta_A = (\nu_A - \nu_{\text{ref}})/\nu_{\text{ref}}$           | 1                             | 30           |
| nuclear Overhauser enhancement                               | $\eta$                |                                                                    | 1                             | 31           |
| nuclear magneton                                             | $\mu_N$               | $\mu_N = (m_e/m_p)\mu_B$                                           | $\text{J T}^{-1}$             | 32           |
| resonance frequency                                          | $\nu$                 |                                                                    | Hz                            | 18           |
| of a reference molecule                                      | $\nu_{\text{ref}}$    |                                                                    |                               |              |
| of the observed rf field                                     | $\nu_1$               |                                                                    |                               |              |
| of the irradiating rf field                                  | $\nu_2$               |                                                                    |                               |              |
| standardized resonance frequency<br>for nucleus A            | $\Xi_A$               |                                                                    | Hz                            | 33           |
| shielding tensor                                             | $\boldsymbol{\sigma}$ |                                                                    | 1                             | 34           |
| shielding constant                                           | $\sigma_A$            | $B_A = (1 - \sigma_A)B_0$                                          | 1                             | 34           |
| correlation time                                             | $\tau_c$              |                                                                    | s                             | 26           |

(25) The symbol  $\mathbf{q}$  is recommended by IUPAC for the field gradient tensor, with the units of  $\text{V m}^{-2}$  (see Section 2.5, p. 24). With  $\mathbf{q}$  defined in this way, the quadrupole coupling constant is  $\chi = eq_{zz}Q/h$ . It is common in NMR to denote the field gradient tensor as  $e\mathbf{q}$  and the quadrupole coupling constant as  $\chi = e^2q_{zz}Q/h$ .  $\mathbf{q}$  has principal components  $q_{XX}, q_{YY}, q_{ZZ}$ .

(26) The relaxation times and the correlation times are normally given in the units of s. Strictly speaking, this refers to  $\text{s rad}^{-1}$ .

(27) The spin-lattice relaxation time of nucleus A in the frame of reference rotating with  $\mathbf{B}_1$  is denoted  $T_{1\rho}^A$ .

(28) See Section 2.4, note 9, p. 19.

(29)  $\mu$  is the magnitude of the nuclear magnetic moment.

(30) Chemical shift (of the resonance) for the nucleus of element A (positive when the sample resonates to high frequency of the reference); for conventions, see [47]. Further information regarding solvent, references, or nucleus of interest may be given by superscripts or subscripts or in parentheses.

(31) The nuclear Overhauser effect,  $1 + \eta$ , is defined as the ratio of the signal intensity for the  $I$  nuclear spin, obtained under conditions of saturation of the  $S$  nuclear spin, and the equilibrium signal intensity for the  $I$  nuclear spin.

(32)  $m_e$  and  $m_p$  are the electron and proton mass, respectively.  $\mu_B$  is the Bohr magneton (see Section 2.5, notes 1 and 2, p. 22 and Chapter 5, p. 111).

(33) Resonance frequency for the nucleus of element A in a magnetic field such that the protons in tetramethylsilane (TMS) resonate exactly at 100 MHz.

(34) The symbols  $\sigma_A$  (and related terms of the shielding tensor and its components) should refer to shielding on an absolute scale (for theoretical work). For shielding relative to a reference, symbols such as  $\sigma_A - \sigma_{\text{ref}}$  should be used.  $B_A$  is the corresponding effective magnetic flux density (see Section 2.3, note 9, p. 17).

### 2.6.1 Symbols for angular momentum operators and quantum numbers

In the following table, all of the operator symbols denote the dimensionless ratio *angular momentum* divided by  $\hbar$ . Although this is a universal practice for the quantum numbers, some authors use the operator symbols to denote *angular momentum*, in which case the operators would have SI units: J s. The column heading “*Z-axis*” denotes the space-fixed component, and the heading “*z-axis*” denotes the molecule-fixed component along the symmetry axis (linear or symmetric top molecules), or the axis of quantization.

| <i>Angular momentum</i> <sup>1</sup> | <i>Operator symbol</i>                     | <i>Quantum number symbol</i> |               |                             | <i>Notes</i> |
|--------------------------------------|--------------------------------------------|------------------------------|---------------|-----------------------------|--------------|
|                                      |                                            | <i>Total</i>                 | <i>Z-axis</i> | <i>z-axis</i>               |              |
| electron orbital                     | $\hat{\mathbf{L}}$                         | $L$                          | $M_L$         | $\Lambda$                   | 2            |
| one electron only                    | $\hat{\mathbf{l}}$                         | $l$                          | $m_l$         | $\lambda$                   | 2            |
| electron spin                        | $\hat{\mathbf{S}}$                         | $S$                          | $M_S$         | $\Sigma$                    |              |
| one electron only                    | $\hat{\mathbf{s}}$                         | $s$                          | $m_s$         | $\sigma$                    |              |
| electron orbital plus spin           | $\hat{\mathbf{L}} + \hat{\mathbf{S}}$      |                              |               | $\Omega = \Lambda + \Sigma$ | 2            |
| nuclear orbital (rotational)         | $\hat{\mathbf{R}}$                         | $R$                          |               | $K_R, k_R$                  |              |
| nuclear spin                         | $\hat{\mathbf{I}}$                         | $I$                          | $M_I$         |                             |              |
| internal vibrational                 |                                            |                              |               |                             |              |
| spherical top                        | $\hat{\mathbf{l}}$                         | $l(l\zeta)$                  |               | $K_l$                       | 3            |
| other                                | $\hat{\mathbf{j}}, \hat{\boldsymbol{\pi}}$ |                              |               | $l(l\zeta)$                 | 2, 3         |
| sum of $R + L (+j)$                  | $\hat{\mathbf{N}}$                         | $N$                          |               | $K, k$                      | 2            |
| sum of $N + S$                       | $\hat{\mathbf{J}}$                         | $J$                          | $M_J$         | $K, k$                      | 2, 4         |
| sum of $J + I$                       | $\hat{\mathbf{F}}$                         | $F$                          | $M_F$         |                             |              |

(1) In all cases the vector operator and its components are related to the quantum numbers by eigenvalue equation analogous to:

$$\hat{\mathbf{J}}^2 \psi = J(J+1) \psi, \quad \hat{J}_Z \psi = M_J \psi, \quad \text{and} \quad \hat{J}_z \psi = K \psi,$$

where the component quantum numbers  $M_J$  and  $K$  take integral or half-odd values in the range  $-J \leq M_J \leq +J, -J \leq K \leq +J$ . (If the operator symbols are taken to represent *angular momentum*, rather than *angular momentum* divided by  $\hbar$ , the eigenvalue equations should read  $\hat{\mathbf{J}}^2 \psi = J(J+1)\hbar^2 \psi$ ,  $\hat{J}_Z \psi = M_J \hbar \psi$ , and  $\hat{J}_z \psi = K \hbar \psi$ .)  $l$  is frequently called the azimuthal quantum number and  $m_l$  the magnetic quantum number.

(2) Some authors, notably Herzberg [29–31], treat the component quantum numbers  $\Lambda, \Omega, l$  and  $K$  as taking positive or zero values only, so that each non-zero value of the quantum number labels two wavefunctions with opposite signs for the appropriate angular momentum component. When this is done, lower case  $k$  is commonly regarded as a signed quantum number, related to  $K$  by  $K = |k|$ . However, in theoretical discussions all component quantum numbers are usually treated as signed, taking both positive and negative values.

(3) There is no uniform convention for denoting the internal vibrational angular momentum;  $j, \pi, p$  and  $G$  have all been used. For symmetric top and linear molecules the component of  $j$  in the symmetry axis is always denoted by the quantum number  $l$ , where  $l$  takes values in the range  $-v \leq l \leq +v$  in steps of 2. The corresponding component of angular momentum is actually  $l\zeta\hbar$ , rather than  $l\hbar$ , where  $\zeta$  is the Coriolis  $\zeta$ -constant (see note 7, p. 26).

(4) Asymmetric top rotational states are labeled by the value of  $J$  (or  $N$  if  $S \neq 0$ ), with subscripts  $K_a, K_c$ , where the latter correlate with the  $K = |k|$  quantum number about the  $a$  and  $c$  axes in the prolate and oblate symmetric top limits respectively.

*Example*  $J_{K_a, K_c} = 5_{2,3}$  for a particular rotational level.

## 2.6.2 Symbols for symmetry operators and labels for symmetry species

### (i) Symmetry operators in space-fixed coordinates [41, 48]

|                       |                     |
|-----------------------|---------------------|
| identity              | $E$                 |
| permutation           | $P, p$              |
| space-fixed inversion | $E^*, (P)$          |
| permutation-inversion | $P^* (= PE^*), p^*$ |

The permutation operation  $P$  permutes the labels of identical nuclei.

*Example* In the  $\text{NH}_3$  molecule, if the hydrogen nuclei are labeled 1, 2 and 3, then  $P = (123)$  would symbolize the permutation where 1 is replaced by 2, 2 by 3, and 3 by 1.

The inversion operation  $E^*$  reverses the sign of all particle coordinates in the space-fixed origin, or in the molecule-fixed centre of mass if translation has been separated. It is also called the parity operator and then frequently denoted by  $P$ , although this cannot be done in parallel with  $P$  for permutation, which then should be denoted by lower case  $p$ . In field-free space and in the absence of parity violation [26], true eigenfunctions of the hamiltonian are either of positive parity  $+$  (unchanged) or of negative parity  $-$  (change sign) under  $E^*$ . The label may be used to distinguish the two nearly degenerate components formed by  $\Lambda$ -doubling (in a degenerate electronic state) or  $l$ -doubling (in a degenerate vibrational state) in linear molecules, or by  $K$ -doubling (asymmetry-doubling) in slightly asymmetric tops. For linear molecules,  $\Lambda$ - or  $l$ -doubled components may also be distinguished by the labels  $e$  or  $f$  [49]; for singlet states these correspond respectively to parity  $+$  or  $-$  for  $J$  even and vice versa for  $J$  odd (but see [49]). For linear molecules in degenerate electronic states the  $\Lambda$ -doubled levels may alternatively be labeled  $\Pi(A')$  or  $\Pi(A'')$  (or  $\Delta(A'), \Delta(A'')$  etc.) [50]. Here the labels  $A'$  or  $A''$  describe the symmetry of the electronic wavefunction at high  $J$  with respect to reflection in the plane of rotation (but see [50] for further details). The  $A'$  or  $A''$  labels are particularly useful for the correlation of states of molecules involved in reactions or photodissociation.

In relation to permutation-inversion symmetry species the superscript  $+$  or  $-$  may be used to designate parity, whereas a letter is used to designate symmetry with respect to the permutation group. One can also use the systematic notation from the theory of the symmetric group (permutation group)  $S_n$  [51], the permutation inversion group being denoted by  $S_n^*$  in this case, if one considers the full permutation group. The species is then given by the partition  $P(S_n)$  [52–54]. The examples give the species for  $S_4^*$ , where the partition is conventionally given in square brackets  $[]$ . Conventions with respect to these symbols still vary ([2.b] and [39–41, 51–54]).

*Examples*  $A_1^+$  totally symmetric species with respect to permutation, positive parity,  $[4]^+$   
 $A_1^-$  totally symmetric species with respect to permutation, negative parity,  $[4]^-$   
 $E^+$  doubly degenerate species with respect to permutation, positive parity,  $[2^2]^+$   
 $E^-$  doubly degenerate species with respect to permutation, negative parity,  $[2^2]^-$   
 $F_1^+$  triply degenerate species with respect to permutation, positive parity,  $[2,1^2]^+$   
 $F_1^-$  triply degenerate species with respect to permutation, negative parity,  $[2,1^2]^-$

The Hermann-Mauguin symbols of symmetry operations used for crystals are given in Section 2.8.1 (ii), p. 44.

### (ii) Symmetry operators in molecule-fixed coordinates (Schönflies symbols) [29–31]

|                      |                                        |
|----------------------|----------------------------------------|
| identity             | $E$                                    |
| rotation by $2\pi/n$ | $C_n$                                  |
| reflection           | $\sigma, \sigma_v, \sigma_d, \sigma_h$ |
| inversion            | $i$                                    |
| rotation-reflection  | $S_n (= C_n\sigma_h)$                  |

If  $C_n$  is the primary axis of symmetry, wavefunctions that are unchanged or change sign under the operator  $C_n$  are given species labels A or B respectively, and otherwise wavefunctions that are multiplied by  $\exp(\pm 2\pi i s/n)$  are given the species label  $E_s$ . Wavefunctions that are unchanged or change sign under  $i$  are labeled g (gerade) or u (ungerade) respectively. Wavefunctions that are unchanged or change sign under  $\sigma_h$  have species labels with a prime ' or a double prime '', respectively. For more detailed rules see [28–31].

### 2.6.3 Other symbols and conventions in optical spectroscopy

#### (i) Term symbols for atomic states

The electronic states of atoms are labeled by the value of the quantum number  $L$  for the state. The value of  $L$  is indicated by an upright capital letter: S, P, D, F, G, H, I, and K,  $\dots$ , are used for  $L = 0, 1, 2, 3, 4, 5, 6$ , and  $7, \dots$ , respectively. The corresponding lower case letters are used for the orbital angular momentum of a single electron. For a many-electron atom, the electron spin multiplicity  $(2S + 1)$  may be indicated as a left-hand superscript to the letter, and the value of the total angular momentum  $J$  as a right-hand subscript. If either  $L$  or  $S$  is zero only one value of  $J$  is possible, and the subscript is then usually suppressed. Finally, the electron configuration of an atom is indicated by giving the occupation of each one-electron orbital as in the examples below.

*Examples* B:  $(1s)^2 (2s)^2 (2p)^1$ ,  $^2P_{1/2}^\circ$   
 C:  $(1s)^2 (2s)^2 (2p)^2$ ,  $^3P_0$   
 N:  $(1s)^2 (2s)^2 (2p)^3$ ,  $^4S^\circ$

A right superscript  $^\circ$  may be used to indicate odd parity (negative parity  $-$ ). Omission of the superscript  $^\circ$  is then to be interpreted as even parity (positive parity  $+$ ). In order to avoid ambiguities it can be useful to always designate parity by  $+$  or  $-$  as a right superscript (i.e.  $^4S_{3/2}^-$  for N and  $^3P_0^+$  for C).

#### (ii) Term symbols for molecular states

The electronic states of molecules are labeled by the symmetry species label of the wavefunction in the molecular point group. These should be Latin or Greek upright capital letters. As for atoms, the spin multiplicity  $(2S + 1)$  may be indicated by a left superscript. For linear molecules the value of  $\Omega (= \Lambda + \Sigma)$  may be added as a right subscript (analogous to  $J$  for atoms). If the value of  $\Omega$  is not specified, the term symbol is taken to refer to all component states, and a right subscript r or i may be added to indicate that the components are regular (energy increases with  $\Omega$ ) or inverted (energy decreases with  $\Omega$ ) respectively.

The electronic states of molecules are also given empirical single letter labels as follows. The ground electronic state is labeled X, excited states of the same multiplicity are labeled A, B, C,  $\dots$ , in ascending order of energy, and excited states of different multiplicity are labeled with lower case letters a, b, c,  $\dots$ . In polyatomic molecules (but not diatomic molecules) it is customary to add a tilde (e.g.  $\tilde{X}$ ) to these empirical labels to prevent possible confusion with the symmetry species label.

Finally the one-electron orbitals are labeled by the corresponding lower case letters, and the electron configuration is indicated in a manner analogous to that for atoms.

*Examples* The ground state of CH is  $(1\sigma)^2 (2\sigma)^2 (3\sigma)^2 (1\pi)^1$ , X  $^2\Pi_r$ , in which the  $^2\Pi_{1/2}$  component lies below the  $^2\Pi_{3/2}$  component, as indicated by the subscript r for regular.

*Examples* (continued)

The ground state of OH is  $(1\sigma)^2(2\sigma)^2(3\sigma)^2(1\pi)^3$ ,  $X^2\Pi_i$ , in which the  $^2\Pi_{3/2}$  component lies below the  $^2\Pi_{1/2}$  component, as indicated by the subscript  $i$  for inverted.

The two lowest electronic states of CH<sub>2</sub> are  $\cdots(2a_1)^2(1b_2)^2(3a_1)^2, \tilde{a}^1A_1,$   
 $\cdots(2a_1)^2(1b_2)^2(3a_1)^1(1b_1)^1, \tilde{X}^3B_1.$

The ground state of C<sub>6</sub>H<sub>6</sub> (benzene) is  $\cdots(a_{2u})^2(e_{1g})^4, \tilde{X}^1A_{1g}.$

The vibrational states of molecules are usually indicated by giving the vibrational quantum numbers for each normal mode.

*Examples* For a bent triatomic molecule

$(0, 0, 0)$  denotes the ground state,

$(1, 0, 0)$  denotes the  $\nu_1$  state, i.e.  $\nu_1 = 1$ , and

$(1, 2, 0)$  denotes the  $\nu_1 + 2\nu_2$  state, i.e.  $\nu_1 = 1, \nu_2 = 2$ , etc.

### (iii) Notation for spectroscopic transitions

The upper and lower levels of a spectroscopic transition are indicated by a prime ' and double prime '' respectively.

*Example*  $h\nu = E' - E''$

Transitions are generally indicated by giving the excited-state label, followed by the ground-state label, separated by a dash or an arrow to indicate the direction of the transition (emission to the right, absorption to the left).

|                 |                                  |                                                                                                                        |
|-----------------|----------------------------------|------------------------------------------------------------------------------------------------------------------------|
| <i>Examples</i> | B–A                              | indicates a transition between a higher energy state B and a lower energy state A;                                     |
|                 | B→A                              | indicates emission from B to A;                                                                                        |
|                 | B←A                              | indicates absorption from A to B;                                                                                      |
|                 | A→B                              | indicates more generally a transition from the initial state A to final state B in kinetics (see Section 2.12, p. 63); |
|                 | $(0, 2, 1) \leftarrow (0, 0, 1)$ | labels the $2\nu_2 + \nu_3 - \nu_3$ hot band in a bent triatomic molecule.                                             |

A more compact notation [55] may be used to label vibronic (or vibrational) transitions in polyatomic molecules with many normal modes, in which each vibration index  $r$  is given a superscript  $v'_r$  and a subscript  $v''_r$  indicating the upper electronic and the lower electronic state values of the vibrational quantum number. When  $v'_r = v''_r = 0$  the corresponding index is suppressed.

*Examples*  $1_0^1$  denotes the transition  $(1,0,0)-(0,0,0)$ ;  
 $2_0^2 3_1^1$  denotes the transition  $(0,2,1)-(0,0,1)$ .

In order to denote transitions within the same electronic state one may use matrix notation or an arrow.

*Example*  $2_{20}$  or  $2_{2\leftarrow 0}$  denotes a vibrational transition within the electronic ground state from  $v_2 = 0$  to  $v_2 = 2$ .

For rotational transitions, the value of  $\Delta J = J' - J''$  is indicated by a letter labelling the branches of a rotational band:  $\Delta J = -2, -1, 0, 1$ , and  $2$  are labelled as the O-branch, P-branch, Q-branch, R-branch, and S-branch respectively. The changes in other quantum numbers (such as  $K$  for a symmetric top, or  $K_a$  and  $K_c$  for an asymmetric top) may be indicated by adding lower case letters as a left superscript according to the same rule.

*Example*  $^PQ$  labels a “p-type Q-branch” in a symmetric top molecule, i.e.  $\Delta K = -1, \Delta J = 0$ .

The value of  $K$  in the lower level is indicated as a right subscript, e.g.  $^PQ_{K''}$  or  $^PQ_2(5)$  indicating the transition from  $K'' = 2$  to  $K' = 1$ , the value of  $J''$  being added in parentheses.

## 2.7 ELECTROMAGNETIC RADIATION

The quantities and symbols given here have been selected on the basis of recommendations by IUPAP [4], ISO [5.f], and IUPAC [56–59], as well as by taking into account the practice in the field of laser physics. Terms used in photochemistry [60] have been considered as well, but definitions still vary. Terms used in high energy ionizing electromagnetic radiation, radiation chemistry, radiochemistry, and nuclear chemistry are not included or discussed.

| <i>Name</i>                                | <i>Symbol</i>                         | <i>Definition</i>                                       | <i>SI unit</i>                        | <i>Notes</i> |
|--------------------------------------------|---------------------------------------|---------------------------------------------------------|---------------------------------------|--------------|
| wavelength                                 | $\lambda$                             |                                                         | m                                     |              |
| speed of light                             |                                       |                                                         |                                       |              |
| in vacuum                                  | $c_0$                                 | $c_0 = 299\,792\,458\text{ m s}^{-1}$                   | $\text{m s}^{-1}$                     | 1            |
| in a medium                                | $c$                                   | $c = c_0/n$                                             | $\text{m s}^{-1}$                     | 1            |
| wavenumber in vacuum                       | $\tilde{\nu}$                         | $\tilde{\nu} = \nu/c_0 = 1/\lambda$                     | $\text{m}^{-1}$                       | 2            |
| wavenumber<br>(in a medium)                | $\sigma$                              | $\sigma = 1/\lambda$                                    | $\text{m}^{-1}$                       |              |
| frequency                                  | $\nu$                                 | $\nu = c/\lambda$                                       | Hz                                    |              |
| angular frequency,<br>pulsatance           | $\omega$                              | $\omega = 2\pi\nu$                                      | $\text{s}^{-1}$ , $\text{rad s}^{-1}$ |              |
| refractive index                           | $n$                                   | $n = c_0/c$                                             | 1                                     |              |
| Planck constant                            | $h$                                   |                                                         | J s                                   |              |
| Planck constant divided by $2\pi$          | $\hbar$                               | $\hbar = h/2\pi$                                        | J s                                   |              |
| radiant energy                             | $Q, W$                                |                                                         | J                                     | 3            |
| radiant energy density                     | $\rho, w$                             | $\rho = \text{d}Q/\text{d}V$                            | $\text{J m}^{-3}$                     | 3            |
| spectral radiant energy density            |                                       |                                                         |                                       | 3            |
| in terms of frequency                      | $\rho_\nu, w_\nu$                     | $\rho_\nu = \text{d}\rho/\text{d}\nu$                   | $\text{J m}^{-3} \text{ Hz}^{-1}$     |              |
| in terms of wavenumber                     | $\rho_{\tilde{\nu}}, w_{\tilde{\nu}}$ | $\rho_{\tilde{\nu}} = \text{d}\rho/\text{d}\tilde{\nu}$ | $\text{J m}^{-2}$                     |              |
| in terms of wavelength                     | $\rho_\lambda, w_\lambda$             | $\rho_\lambda = \text{d}\rho/\text{d}\lambda$           | $\text{J m}^{-4}$                     |              |
| radiant power (radiant energy<br>per time) | $P, \Phi$                             | $P = \text{d}Q/\text{d}t$                               | W                                     | 3<br>3       |
| radiant intensity                          | $I_e$                                 | $I_e = \text{d}P/\text{d}\Omega$                        | $\text{W sr}^{-1}$                    | 3, 4         |
| radiant excitance                          | $M$                                   | $M = \text{d}P/\text{d}A_{\text{source}}$               | $\text{W m}^{-2}$                     | 3, 4         |

(1) When there is no risk of ambiguity the subscript denoting vacuum is often omitted.  $n$  denotes the refraction index of the medium.

(2) The unit  $\text{cm}^{-1}$  is widely used for the quantity wavenumber in vacuum.

(3) The symbols for the quantities such as *radiant energy* and *intensity* are also used for the corresponding quantities concerning visible radiation, i.e. luminous quantities and photon quantities. Subscripts e for energetic, v for visible, and p for photon may be added whenever confusion between these quantities might otherwise occur. The units used for luminous quantities are derived from the base unit candela (cd) (see Section 3.3, p. 87).

|                 |                    |                                        |                                          |
|-----------------|--------------------|----------------------------------------|------------------------------------------|
| <i>Examples</i> | radiant intensity  | $I_e, I_e = \text{d}P/\text{d}\Omega,$ | SI unit: $\text{W sr}^{-1}$              |
|                 | luminous intensity | $I_v,$                                 | SI unit: cd                              |
|                 | photon intensity   | $I_p,$                                 | SI unit: $\text{s}^{-1} \text{ sr}^{-1}$ |

The radiant intensity  $I_e$  should be distinguished from the plain intensity or irradiance  $I$  (see note 5). Additional subscripts to distinguish absorbed (abs), transmitted (tr) or reflected (refl) quantities may be added, if necessary.

(4) The radiant intensity is the radiant power per solid angle in the direction of the point from which the source is being observed. The radiant excitance is the total emitted radiant power per area  $A_{\text{source}}$  of the radiation source, for all wavelengths. The radiance is the radiant intensity per area of

| <i>Name</i>                                    | <i>Symbol</i>                      | <i>Definition</i>                                                   | <i>SI unit</i>                    | <i>Notes</i> |
|------------------------------------------------|------------------------------------|---------------------------------------------------------------------|-----------------------------------|--------------|
| radiance                                       | $L$                                | $I_e = \int L \cos \Theta \, dA_{\text{source}}$                    | $\text{W sr}^{-1} \text{ m}^{-2}$ | 3, 4         |
| intensity, irradiance                          | $I, E$                             | $I = dP/dA$                                                         | $\text{W m}^{-2}$                 | 3, 5         |
| spectral intensity,<br>spectral irradiance     | $I_{\tilde{\nu}}, E_{\tilde{\nu}}$ | $I_{\tilde{\nu}} = dI/d\tilde{\nu}$                                 | $\text{W m}^{-1}$                 | 6            |
| fluence                                        | $F, (H)$                           | $F = \int I dt = \int (dP/dA) dt$                                   | $\text{J m}^{-2}$                 | 7            |
| Einstein coefficient,<br>spontaneous emission  | $A_{ij}$                           | $dN_j/dt = -\sum_i A_{ij} N_j$                                      | $\text{s}^{-1}$                   | 8, 9         |
| stimulated or induced<br>emission              | $B_{ij}$                           | $dN_j/dt = -\sum_i \rho_{\tilde{\nu}}(\tilde{\nu}_{ij}) B_{ij} N_j$ | $\text{s kg}^{-1}$                |              |
| absorption                                     | $B_{ji}$                           | $dN_i/dt = -\sum_j \rho_{\tilde{\nu}}(\tilde{\nu}_{ij}) B_{ji} N_i$ | $\text{s kg}^{-1}$                |              |
| emissivity, emittance                          | $\varepsilon$                      | $\varepsilon = M/M_{\text{bb}}$                                     | 1                                 | 10           |
| Stefan-Boltzmann constant                      | $\sigma$                           | $M_{\text{bb}} = \sigma T^4$                                        | $\text{W m}^{-2} \text{ K}^{-4}$  | 10           |
| étendue (throughput,<br>light gathering power) | $E, (e)$                           | $E = A\Omega = P/L$                                                 | $\text{m}^2 \text{ sr}$           | 11           |

(4) (continued) radiation source;  $\Theta$  is the angle between the normal to the area element and the direction of observation as seen from the source.

(5) The intensity or irradiance is the radiation power per area that is received at a surface. Intensity, symbol  $I$ , is usually used in discussions involving collimated beams of light, as in applications of the Beer-Lambert law for spectrometric analysis. Intensity of electromagnetic radiation can also be defined as the modulus of the Poynting vector (see Section 2.3, p. 17 and Section 7.4, p. 148). In photochemistry the term intensity is sometimes used as an alias for radiant intensity and must not be understood as an irradiance, for which the symbol  $E$  is preferred [60].

(6) Spectral quantities may be defined with respect to frequency  $\nu$ , wavelength  $\lambda$ , or wavenumber  $\tilde{\nu}$ ; see the entry for spectral radiant energy density in this table.

(7) Fluence is used in photochemistry to specify the energy per area delivered in a given time interval (for instance by a laser pulse); fluence is the time integral of the fluence rate. Sometimes distinction must be made between irradiance and fluence rate [60]; fluence rate reduces to irradiance for a light beam incident from a single direction perpendicularly to the surface. The time integral of irradiance is called radiant exposure.

(8) The indices  $i$  and  $j$  refer to individual states;  $E_j > E_i$ ,  $E_j - E_i = hc\tilde{\nu}_{ij}$ , and  $B_{ji} = B_{ij}$  in the defining equations. The coefficients  $B$  are defined here using energy density  $\rho_{\tilde{\nu}}$  in terms of wavenumber; they may alternatively be defined using energy density in terms of frequency  $\rho_{\nu}$ , in which case  $B$  has SI units  $\text{m kg}^{-1}$ , and  $B_{\nu} = c_0 B_{\tilde{\nu}}$  where  $B_{\nu}$  is defined using frequency and  $B_{\tilde{\nu}}$  using wavenumber. The defining equations refer to the *partial* contributions to the rate of change.

(9) The relation between the Einstein coefficients  $A$  and  $B_{\tilde{\nu}}$  is  $A = 8\pi hc_0 \tilde{\nu}^3 B_{\tilde{\nu}}$ . The Einstein stimulated absorption or emission coefficient  $B$  may also be related to the transition moment between the states  $i$  and  $j$ ; for an electric dipole transition the relation is

$$B_{\tilde{\nu},ij} = \frac{8\pi^3}{3h^2 c_0 (4\pi\epsilon_0)} \sum_{\rho} |\langle i | \mu_{\rho} | j \rangle|^2$$

in which the sum over  $\rho$  runs over the three space-fixed cartesian axes, and  $\mu_{\rho}$  is a space-fixed component of the dipole moment operator. Again, these equations are based on a wavenumber definition of the Einstein coefficient  $B$  (i.e.  $B_{\tilde{\nu}}$  rather than  $B_{\nu}$ ).

(10) The emittance of a sample is the ratio of the radiant excitance emitted by the sample to the radiant excitance emitted by a black body at the same temperature;  $M_{\text{bb}}$  is the latter quantity. See Chapter 5, p. 112 for the value of the Stefan-Boltzmann constant.

| <i>Name</i>                              | <i>Symbol</i>       | <i>Definition</i>                         | <i>SI unit</i>               | <i>Notes</i> |
|------------------------------------------|---------------------|-------------------------------------------|------------------------------|--------------|
| resolution                               | $\delta\tilde{\nu}$ |                                           | $\text{m}^{-1}$              | 2, 12, 13    |
| resolving power                          | $R$                 | $R = \tilde{\nu}/\delta\tilde{\nu}$       | 1                            | 13           |
| free spectral range                      | $\Delta\tilde{\nu}$ | $\Delta\tilde{\nu} = 1/2l$                | $\text{m}^{-1}$              | 2, 14        |
| finesse                                  | $f$                 | $f = \Delta\tilde{\nu}/\delta\tilde{\nu}$ | 1                            | 14           |
| quality factor                           | $Q$                 | $Q = 2\pi\nu \frac{W}{-dW/dt}$            | 1                            | 14, 15       |
| first radiation constant                 | $c_1$               | $c_1 = 2\pi hc_0^2$                       | $\text{W m}^2$               |              |
| second radiation constant                | $c_2$               | $c_2 = hc_0/k_B$                          | $\text{K m}$                 | 16           |
| transmittance, transmission factor       | $\tau, T$           | $\tau = P_{\text{tr}}/P_0$                | 1                            | 17, 18       |
| absorptance, absorption factor           | $\alpha$            | $\alpha = P_{\text{abs}}/P_0$             | 1                            | 17, 18       |
| reflectance, reflection factor           | $\rho, R$           | $\rho = P_{\text{refl}}/P_0$              | 1                            | 17, 18       |
| (decadic) absorbance                     | $A_{10}, A$         | $A_{10} = -\lg(1 - \alpha_i)$             | 1                            | 17–20        |
| napierian absorbance                     | $A_e, B$            | $A_e = -\ln(1 - \alpha_i)$                | 1                            | 17–20        |
| absorption coefficient, (linear) decadic | $a, K$              | $a = A_{10}/l$                            | $\text{m}^{-1}$              | 18, 21       |
| (linear) napierian                       | $\alpha$            | $\alpha = A_e/l$                          | $\text{m}^{-1}$              | 18, 21       |
| molar (decadic)                          | $\varepsilon$       | $\varepsilon = a/c = A_{10}/cl$           | $\text{m}^2 \text{mol}^{-1}$ | 18, 21, 22   |
| molar napierian                          | $\kappa$            | $\kappa = \alpha/c = A_e/cl$              | $\text{m}^2 \text{mol}^{-1}$ | 18, 21, 22   |

(11) Étendue is a characteristic of an optical instrument. It is a measure of the light gathering power, i.e. the power transmitted per radiance of the source.  $A$  is the area of the source or image stop;  $\Omega$  is the solid angle accepted from each point of the source by the aperture stop.

(12) The precise definition of resolution depends on the lineshape, but usually resolution is taken as the full line width at half maximum intensity (FWHM) on a wavenumber,  $\delta\tilde{\nu}$ , or frequency,  $\delta\nu$ , scale. Frequently the use of resolving power, of dimension 1, is preferable.

(13) This quantity characterizes the performance of a spectrometer, or the degree to which a spectral line (or a laser beam) is monochromatic. It may also be defined using frequency  $\nu$ , or wavelength  $\lambda$ .

(14) These quantities characterize a Fabry-Perot cavity, or a laser cavity.  $l$  is the cavity spacing, and  $2l$  is the round-trip path length. The free spectral range is the wavenumber interval between successive longitudinal cavity modes.

(15)  $W$  is the energy stored in the cavity, and  $-dW/dt$  is the rate of decay of stored energy.  $Q$  is also related to the linewidth of a single cavity mode:  $Q = \nu/\delta\nu = \tilde{\nu}/\delta\tilde{\nu}$ . Thus high  $Q$  cavities give narrow linewidths.

(16)  $k_B$  is the Boltzmann constant (see Section 2.9, p. 45 and Chapter 5, p. 111).

(17) If scattering and luminescence can be neglected,  $\tau + \alpha + \rho = 1$ . In optical spectroscopy internal properties (denoted by subscript i) are defined to exclude surface effects and effects of the cuvette such as reflection losses, so that  $\tau_i + \alpha_i = 1$ , if scattering and luminescence can be neglected. This leads to the customary form of the Beer-Lambert law,  $P_{\text{tr}}/P_0 = I_{\text{tr}}/I_0 = \tau_i = 1 - \alpha_i = \exp(-\kappa cl)$ . Hence  $A_e = -\ln(\tau_i)$ ,  $A_{10} = -\lg(\tau_i)$ .

(18) In spectroscopy all of these quantities are commonly taken to be defined in terms of the spectral intensity,  $I_{\tilde{\nu}}(\tilde{\nu})$ , hence they are all regarded as functions of wavenumber  $\tilde{\nu}$  (or frequency  $\nu$ ) across the spectrum. Thus, for example, the absorption coefficient  $\alpha(\tilde{\nu})$  as a function of wavenumber  $\tilde{\nu}$  defines the absorption spectrum of the sample; similarly  $T(\tilde{\nu})$  defines the transmittance spectrum. Spectroscopists use  $I(\tilde{\nu})$  instead of  $I_{\tilde{\nu}}(\tilde{\nu})$ .

(19) The definitions given here relate the absorbance  $A_{10}$  or  $A_e$  to the *internal* absorptance  $\alpha_i$  (see

| <i>Name</i>                       | <i>Symbol</i>                 | <i>Definition</i>                                                                                | <i>SI unit</i>                    | <i>Notes</i> |
|-----------------------------------|-------------------------------|--------------------------------------------------------------------------------------------------|-----------------------------------|--------------|
| net absorption cross section      | $\sigma_{\text{net}}$         | $\sigma_{\text{net}} = \kappa/N_{\text{A}}$                                                      | $\text{m}^2$                      | 23           |
| absorption coefficient            |                               |                                                                                                  |                                   |              |
| integrated over $\tilde{\nu}$     | $A, \bar{A}$                  | $A = \int \kappa(\tilde{\nu}) \, \text{d}\tilde{\nu}$                                            | $\text{m mol}^{-1}$               | 23, 24       |
|                                   | $S$                           | $S = A/N_{\text{A}}$                                                                             | $\text{m}$                        | 23, 24       |
|                                   | $\bar{S}$                     | $\bar{S} = (1/pl) \int \ln(I_0/I) \, \text{d}\tilde{\nu}$                                        | $\text{Pa}^{-1} \text{ m}^{-2}$   | 23–25        |
| integrated over $\ln \tilde{\nu}$ | $\Gamma$                      | $\Gamma = \int \kappa(\tilde{\nu}) \tilde{\nu}^{-1} \, \text{d}\tilde{\nu}$                      | $\text{m}^2 \text{ mol}^{-1}$     | 23, 24       |
| integrated net absorption         |                               |                                                                                                  |                                   |              |
| cross section                     | $G_{\text{net}}$              | $G_{\text{net}} = \int \sigma_{\text{net}}(\tilde{\nu}) \tilde{\nu}^{-1} \, \text{d}\tilde{\nu}$ | $\text{m}^2$                      | 23, 24       |
| absorption index,                 |                               |                                                                                                  |                                   |              |
| imaginary refractive index        | $k$                           | $k = \alpha/4\pi\tilde{\nu}$                                                                     | 1                                 | 26           |
| complex refractive index          | $\hat{n}$                     | $\hat{n} = n + ik$                                                                               | 1                                 |              |
| molar refraction,                 |                               |                                                                                                  |                                   |              |
| molar refractivity                | $R$                           | $R = \left( \frac{n^2 - 1}{n^2 + 2} \right) V_{\text{m}}$                                        | $\text{m}^3 \text{ mol}^{-1}$     |              |
| angle of optical rotation         | $\alpha$                      |                                                                                                  | 1, rad                            | 27           |
| specific optical rotatory power   | $[\alpha]_{\lambda}^{\theta}$ | $[\alpha]_{\lambda}^{\theta} = \alpha/\gamma l$                                                  | $\text{rad m}^2 \text{ kg}^{-1}$  | 27           |
| molar optical rotatory power      | $\alpha_{\text{m}}$           | $\alpha_{\text{m}} = \alpha/cl$                                                                  | $\text{rad m}^2 \text{ mol}^{-1}$ | 27           |

(19) (continued) note 17). However the subscript i on the absorbance  $\alpha$  is often omitted. Experimental data must include corrections for reflections, scattering and luminescence, if the absorbance is to have an absolute meaning. In practice the absorbance is measured as the logarithm of the ratio of the light transmitted through a reference cell (with solvent only) to that transmitted through a sample cell.

(20) In reference [57] the symbol  $A$  is used for decadic absorbance, and  $B$  for napierian absorbance.

(21)  $l$  is the absorbing path length, and  $c$  is the amount (of substance) concentration.

(22) The molar decadic absorption coefficient  $\varepsilon$  is sometimes called the “extinction coefficient” in the published literature. Unfortunately numerical values of the “extinction coefficient” are often quoted without specifying units; the absence of units usually means that the units are  $\text{mol}^{-1} \text{ dm}^3 \text{ cm}^{-1}$  (see also [61]). The word “extinction” should properly be reserved for the sum of the effects of absorption, scattering, and luminescence.

(23) Note that these quantities give the net absorption coefficient  $\kappa$ , the net absorption cross section  $\sigma_{\text{net}}$ , and the net values of  $A, S, \bar{S}, \Gamma$ , and  $G_{\text{net}}$ , in the sense that they are the sums of effects due to absorption and induced emission (see the discussion in Section 2.7.1, p. 38).

(24) The definite integral defining these quantities may be specified by the limits of integration in parentheses, e.g.  $G(\tilde{\nu}_1, \tilde{\nu}_2)$ . In general the integration is understood to be taken over an absorption line or an absorption band.  $A, \bar{S}$ , and  $\Gamma$  are measures of the strength of the band in terms of amount concentration;  $G_{\text{net}} = \Gamma/N_{\text{A}}$  and  $S = A/N_{\text{A}}$  are corresponding molecular quantities. For a single spectral line the relation of these quantities to the Einstein transition probabilities is discussed in Section 2.7.1, p. 38. The symbol  $\bar{A}$  may be used for the integrated absorption coefficient  $A$  when there is a possibility of confusion with the Einstein spontaneous emission coefficient  $A_{ij}$ .

The integrated absorption coefficient of an electronic transition is often expressed in terms of the oscillator strength or “ $f$ -value”, which is dimensionless, or in terms of the Einstein transition probability  $A_{ij}$  between the states involved, with SI unit  $\text{s}^{-1}$ . Whereas  $A_{ij}$  has a simple and universally accepted meaning (see note 9, p. 35), there are differing uses of  $f$ . A common practical conversion is given by the equation

$$f_{ij} = [(4\pi\varepsilon_0) m_e c_0 / 8\pi^2 e^2] \lambda^2 A_{ij} \quad \text{or} \quad f_{ij} \approx (1.4992 \times 10^{-14}) (A_{ij}/\text{s}^{-1}) (\lambda/\text{nm})^2$$

in which  $\lambda$  is the transition wavelength, and  $i$  and  $j$  refer to individual states. For strongly allowed electronic transitions  $f$  is of order unity.

### 2.7.1 Quantities and symbols concerned with the measurement of absorption intensity

In most experiments designed to measure the intensity of spectral absorption, the measurement gives the net absorption due to the effects of absorption from the lower energy level  $m$  to the upper energy level  $n$ , minus induced emission from  $n$  to  $m$ . Since the populations depend on the temperature, so does the measured net absorption. This comment applies to all the quantities defined in the table to measure absorption intensity, although for transitions where  $hc_0\tilde{\nu} \gg kT$  the temperature dependence is small and for  $\tilde{\nu} > 1000 \text{ cm}^{-1}$  at ordinary temperatures induced emission can generally be neglected.

In a more fundamental approach one defines the cross section  $\sigma_{ji}(\tilde{\nu})$  for an induced radiative transition from the state  $i$  to the state  $j$  (in either absorption or emission). For an ideal absorption experiment with only the lower state  $i$  populated the integrated absorption cross section for the transition  $j \leftarrow i$  is given by

$$G_{ji} = \int \sigma_{ji}(\tilde{\nu}) \tilde{\nu}^{-1} d\tilde{\nu} = \int \sigma_{ji}(\nu) \nu^{-1} d\nu$$

If the upper and lower energy levels are degenerate, the observed line strength is given by summing over transitions between all states  $i$  in the lower energy level  $m$  and all states  $j$  in the upper energy level  $n$ , multiplying each term by the fractional population  $p_i$  in the appropriate initial state. Neglecting induced emission this gives

$$G_{\text{net}}(n \leftarrow m) = \sum_{i,j} p_i G_{ji}$$

---

(Notes continued)

(25) The quantity  $\bar{S}$  is only used for gases; it is defined in a manner similar to  $A$ , except that the partial pressure  $p$  of the gas replaces the concentration  $c$ . At low pressures  $p_i \approx c_i RT$ , so that  $\bar{S}$  and  $A$  are related by the equation  $\bar{S} \approx A/RT$ . Thus if  $\bar{S}$  is used to report line or band intensities, the temperature should be specified.  $I_0$  is the incident,  $I$  the transmitted intensity, thus  $\ln(I_0/I) = -\ln(I/I_0) = -\ln(1 - P_{\text{abs}}/P_0) = A_e$  (see also notes 17 and 19, p. 36).

(26)  $\alpha$  in the definition is the napierian absorption coefficient.

(27) The sign convention for the angle of optical rotation is as follows:  $\alpha$  is positive if the plane of polarization is rotated clockwise as viewed looking towards the light source. If the rotation is anti clockwise, then  $\alpha$  is negative. The optical rotation due to a solute in solution may be specified by a statement of the type

$$\alpha(589.3 \text{ nm}, 20^\circ\text{C}, \text{ sucrose}, 10 \text{ g dm}^{-3} \text{ in H}_2\text{O}, 10 \text{ cm path}) = +0.6647^\circ$$

The same information may be conveyed by quoting either the specific optical rotatory power  $\alpha/\gamma l$ , or the molar optical rotatory power  $\alpha/cl$ , where  $\gamma$  is the mass concentration,  $c$  is the amount (of substance) concentration, and  $l$  is the path length. Most tabulations give the specific optical rotatory power, denoted  $[\alpha]_\lambda^\theta$ . The wavelength of light used  $\lambda$  (frequently the sodium D line) and the Celsius temperature  $\theta$  are conventionally written as a subscript and superscript to the specific rotatory power  $[\alpha]$ . For pure liquids and solids  $[\alpha]_\lambda^\theta$  is similarly defined as  $[\alpha]_\lambda^\theta = \alpha/\rho l$ , where  $\rho$  is the mass density.

Specific optical rotatory powers are customarily called *specific rotations*, and are unfortunately usually quoted without units. The absence of units may usually be taken to mean that the units are  $^\circ \text{ cm}^3 \text{ g}^{-1} \text{ dm}^{-1}$  for pure liquids and solutions, or  $^\circ \text{ cm}^3 \text{ g}^{-1} \text{ mm}^{-1}$  for solids, where  $^\circ$  is used as a symbol for degrees of plane angle.

If induced emission is significant then the net integrated cross section becomes

$$G_{\text{net}}(n \leftarrow m) = \sum_{i,j} (p_i - p_j) G_{ji} = \left( \frac{p_m}{d_m} - \frac{p_n}{d_n} \right) \sum_{i,j} G_{ji}$$

Here  $p_i$  and  $p_j$  denote the fractional populations of states  $i$  and  $j$  ( $p_i = \exp\{-E_i/kT\}/q$  in thermal equilibrium, where  $q$  is the partition function);  $p_m$  and  $p_n$  denote the corresponding fractional populations of the energy levels, and  $d_m$  and  $d_n$  the degeneracies ( $p_i = p_m/d_m$ , etc.). The absorption intensity  $G_{ji}$ , and the Einstein coefficients  $A_{ij}$  and  $B_{ji}$ , are fundamental measures of the line strength between the individual states  $i$  and  $j$ ; they are related to each other by the general equations

$$G_{ji} = hB_{\tilde{\nu},ji} = (h/c_0) B_{\nu,ji} = A_{ij}/(8\pi c_0 \tilde{\nu}^3)$$

Finally, for an electric dipole transition these quantities are related to the square of the transition moment by the equation

$$G_{ji} = hB_{\tilde{\nu},ji} = A_{ij}/(8\pi c_0 \tilde{\nu}^3) = \frac{8\pi^3}{3hc_0(4\pi\epsilon_0)} |M_{ji}|^2$$

where the transition moment  $M_{ji}$  is given by

$$|M_{ji}|^2 = \sum_{\rho} |\langle i | \mu_{\rho} | j \rangle|^2$$

Here the sum is over the three space-fixed cartesian axes and  $\mu_{\rho}$  is a space-fixed component of the electric dipole moment. Inserting numerical values for the constants the relation between  $G_{ji}$  and  $M_{ji}$  may be expressed in a practical representation as

$$(G_{ji}/\text{pm}^2) \approx 41.6238 |M_{ji}/\text{D}|^2$$

where  $1 \text{ D} \approx 3.335641 \times 10^{-30} \text{ C m}$  (D is the symbol of the debye) [62].

Net integrated absorption band intensities are usually characterized by one of the quantities  $A, S, \bar{S}, \Gamma$ , or  $G_{\text{net}}$  as defined in the table. The relation between these quantities is given by the (approximate) equations

$$G_{\text{net}} = \Gamma/N_A \approx A/(\tilde{\nu}_0 N_A) \approx S/\tilde{\nu}_0 \approx \bar{S} (kT/\tilde{\nu}_0)$$

However, only the first equality is exact. The relation to  $A$ ,  $\bar{S}$  and  $S$  involves dividing by the band centre wavenumber  $\tilde{\nu}_0$  for a band, to correct for the fact that  $A$ ,  $\bar{S}$  and  $S$  are obtained by integrating over wavenumber rather than integrating over the logarithm of wavenumber used for  $G_{\text{net}}$  and  $\Gamma$ . This correction is only approximate for a band (although negligible error is involved for single-line intensities in gases). The relation to  $\bar{S}$  involves the assumption that the gas is ideal (which is approximately true at low pressures), and also involves the temperature. Thus the quantities  $\Gamma$  and  $G_{\text{net}}$  are most simply related to more fundamental quantities such as the Einstein transition probabilities and the transition moment, and are the preferred quantities for reporting integrated line or band intensities.

The situation is further complicated by some authors using the symbol  $S$  for any of the above quantities, particularly for any of the quantities here denoted  $A, S$  and  $\bar{S}$ . It is therefore particularly important to define quantities and symbols used in reporting integrated intensities.

For transitions between individual states any of the more fundamental quantities  $G_{ji}$ ,  $B_{\tilde{\nu},ji}$ ,  $A_{ij}$ , or  $|M_{ji}|$  may be used; the relations are as given above, and are exact. The integrated absorption coefficient  $A$  should not be confused with the Einstein coefficient  $A_{ij}$  (nor with absorbance, for which the symbol  $A$  is also used). Where such confusion might arise, we recommend writing  $\bar{A}$  for the band intensity expressed as an integrated absorption coefficient over wavenumber.

The SI unit and commonly used units of  $A$ ,  $S$ ,  $\bar{S}$ ,  $\Gamma$  and  $G$  are as in the table below. Also given in the table are numerical transformation coefficients in commonly used units, from  $A$ ,  $S$ ,  $\bar{S}$ , and  $\Gamma$  to  $G_{\text{net}}$ .

| <i>Quantity</i> | <i>SI unit</i>                  | <i>Common unit</i>                | <i>Transformation coefficient</i>                                                                                                          |
|-----------------|---------------------------------|-----------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------|
| $A, \bar{A}$    | $\text{m mol}^{-1}$             | $\text{km mol}^{-1}$              | $(G/\text{pm}^2) = 16.605\ 40 \frac{A/(\text{km mol}^{-1})}{\tilde{\nu}_0/\text{cm}^{-1}}$                                                 |
| $\bar{S}$       | $\text{Pa}^{-1} \text{ m}^{-2}$ | $\text{atm}^{-1} \text{ cm}^{-2}$ | $(G/\text{pm}^2) = 1.362\ 603 \times 10^{-2} \frac{(\bar{S}/\text{atm}^{-1} \text{ cm}^{-2})(T/\text{K})}{(\tilde{\nu}_0/\text{cm}^{-1})}$ |
| $S$             | $\text{m}$                      | $\text{cm}$                       | $(G/\text{pm}^2) = 10^{20} \frac{(S/\text{cm})}{\tilde{\nu}_0/\text{cm}^{-1}}$                                                             |
| $\Gamma$        | $\text{m}^2 \text{ mol}^{-1}$   | $\text{cm}^2 \text{ mol}^{-1}$    | $(G/\text{pm}^2) = 1.660\ 540 \times 10^{-4} (\Gamma/\text{cm}^2 \text{ mol}^{-1})$                                                        |
| $G$             | $\text{m}^2$                    | $\text{pm}^2$                     |                                                                                                                                            |

Quantities concerned with spectral absorption intensity and relations among these quantities are discussed in references [62–65], and a list of published measurements of line intensities and band intensities for gas phase infrared spectra may be found in references [63–65]. For relations between spectroscopic absorption intensities and kinetic radiative transition rates see also [66].

## 2.7.2 Conventions for absorption intensities in condensed phases

Provided transmission measurements are accurately corrected for reflection and other losses, the absorbance, absorption coefficient, and integrated absorption coefficient,  $A$ , that are described above may be used for condensed phases. The corrections typically used are adequate for weak and medium bands in neat liquids and solids and for bands of solutes in dilute solution. In order to make the corrections accurately for strong bands in neat liquids and solids, it is necessary to determine the real and imaginary refractive indices  $n$  and  $k$ , of the sample throughout the measured spectral range. Then, the resulting  $n$  and  $k$  themselves provide a complete description of the absorption intensity in the condensed phase. For liquids, this procedure requires knowledge of  $n$  and  $k$  of the cell windows. Reflection spectra are also processed to yield  $n$  and  $k$ . For non-isotropic solids all intensity properties must be defined with respect to specific crystal axes. If the  $n$  and  $k$  spectra are known, spectra of any optical property or measured quantity of the sample may be obtained from them. Physicists prefer to use the complex relative permittivity (see Section 2.3, p. 16),  $\hat{\epsilon}_{\text{r}} = \epsilon_{\text{r}}' + i\epsilon_{\text{r}}'' = \text{Re } \hat{\epsilon}_{\text{r}} + i\text{Im } \hat{\epsilon}_{\text{r}}$  instead of the complex refractive index  $\hat{n}$ .  $\epsilon_{\text{r}}' = \text{Re } \hat{\epsilon}_{\text{r}}$  denotes the real part and  $\epsilon_{\text{r}}'' = \text{Im } \hat{\epsilon}_{\text{r}}$  the imaginary part of  $\hat{\epsilon}_{\text{r}}$ , respectively. They are related through  $\hat{\epsilon}_{\text{r}} = \hat{n}^2$ , so that  $\epsilon_{\text{r}}' = n^2 - k^2$  and  $\epsilon_{\text{r}}'' = 2nk$ .

The refractive indices and relative permittivities are properties of the bulk phase. In order to obtain information about the molecules in the liquid free from dielectric effects of the bulk, the local field that acts on the molecules,  $\mathbf{E}_{\text{loc}}$ , must be determined as a function of the applied field  $\mathbf{E}$ . A simple relation is the Lorentz local field,  $\mathbf{E}_{\text{loc}} = \mathbf{E} + \mathbf{P}/3\epsilon_0$ , where  $\mathbf{P}$  is the dielectric polarization. This local field is based on the assumption that long-range interactions are isotropic, so it is realistic only for liquids and isotropic solids.

Use of this local field gives the Lorentz-Lorenz formula (this relation is usually called the Clausius-Mossotti formula when applied to static fields) generalized to treat absorbing materials at

any wavenumber

$$\frac{\hat{\varepsilon}_r(\tilde{\nu}) - 1}{\hat{\varepsilon}_r(\tilde{\nu}) + 2} = \frac{1}{3\varepsilon_0 V_m} \hat{\alpha}_m(\tilde{\nu})$$

Here  $V_m$  is the molar volume, and  $\hat{\alpha}_m$  is the complex molar polarizability (see Section 2.3, note 7, p. 16). The imaginary part  $\alpha_m''$  of the complex molar polarizability describes the absorption by molecules in the liquid, corrected for the long-range isotropic dielectric effects but influenced by the anisotropic environment formed by the first few nearest neighbor molecules. The real part  $\alpha_m'$  is the molar polarizability (see sections 2.3 and 2.5) in the limit of infinite frequency.

The integrated absorption coefficient of a molecular absorption band in the condensed phase is described by [67, 68] (see note 1, below)

$$C_j = \frac{1}{4\pi\varepsilon_0} \int_{\text{band } j} \tilde{\nu} \alpha_m''(\tilde{\nu}) d\tilde{\nu}$$

Theoretical analysis usually assumes that the measured band includes the transition  $j \leftarrow i$  with band centre wavenumber  $\tilde{\nu}_0$  and all of its hot-band transitions. Then

$$C_j = \frac{N_A \pi}{3hc_0(4\pi\varepsilon_0)} \tilde{\nu}_0 g(|M_{ji}|^2)$$

where  $\tilde{\nu}_0 g(|M_{ji}|^2)$  is the population-weighted sum over all contributing transitions of the wavenumber times the square of the electric dipole moment of the transition ( $g$  is in general a temperature dependent function which includes effects from the inhomogeneous band structure). The traditional relation between the gas and liquid phase values of the absorption coefficient of a given band  $j$  is the Polo-Wilson equation [69]

$$A_{\text{liq}} = \frac{(\bar{n}^2 + 2)^2}{9\bar{n}} A_{\text{gas}}$$

where  $\bar{n}$  is the estimated average value of  $n$  through the absorption band. This relation is valid under the following conditions:

- (i) bands are sufficiently weak, i.e.  $2nk + k^2 \ll n^2$ ;
- (ii) the double harmonic approximation yields a satisfactory theoretical approximation of the band structure; in this case the function  $g$  is nearly temperature independent and the gas and liquid phase values of  $g$  are equal;
- (iii)  $\tilde{\nu}_0 |M_{ji}|^2$  are identical in the gas and liquid phases.

A more recent and more general relation that requires only the second and the third of these conditions is  $A_{\text{gas}} = 8\pi^2 C_j$ . It follows that for bands that meet all three of the conditions

$$A_{\text{liq}} = 8\pi^2 C_j \frac{(\bar{n}^2 + 2)^2}{9\bar{n}}$$

This relation shows that, while the refractive index  $n$  is automatically incorporated in the integrated absorption coefficient  $C_j$  for a liquid, the traditional absorption coefficient  $A$  needs to be further corrected for permittivity or refraction.

---

(1) The SI unit of this quantity is  $\text{m mol}^{-1}$ . In the Gaussian system the definition of  $C_j$  as  $C_j = \int_{\text{band } j} \tilde{\nu} \alpha_m''(\tilde{\nu}) d\tilde{\nu}$  is often adopted, with unit  $\text{cm mol}^{-1}$ .

## 2.8 SOLID STATE

The quantities and their symbols given here have been selected from more extensive lists of IUPAP [4] and ISO [5.n]. See also the *International Tables for Crystallography*, Volume A [70].

| <i>Name</i>                                                          | <i>Symbol</i>                                                                        | <i>Definition</i>                                                                                                                                                                              | <i>SI unit</i>  | <i>Notes</i> |
|----------------------------------------------------------------------|--------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------|--------------|
| lattice vector,                                                      |                                                                                      |                                                                                                                                                                                                |                 |              |
| Bravais lattice vector                                               | $\mathbf{R}, \mathbf{R}_0$                                                           |                                                                                                                                                                                                | m               |              |
| fundamental translation vectors for the crystal lattice              | $\mathbf{a}_1; \mathbf{a}_2; \mathbf{a}_3, \mathbf{a}; \mathbf{b}; \mathbf{c}$       | $\mathbf{R} = n_1 \mathbf{a}_1 + n_2 \mathbf{a}_2 + n_3 \mathbf{a}_3$                                                                                                                          | m               | 1            |
| (angular) fundamental translation vectors for the reciprocal lattice | $\mathbf{b}_1; \mathbf{b}_2; \mathbf{b}_3, \mathbf{a}^*; \mathbf{b}^*; \mathbf{c}^*$ | $\mathbf{a}_i \cdot \mathbf{b}_k = 2\pi\delta_{ik}$                                                                                                                                            | m <sup>-1</sup> | 2            |
| (angular) reciprocal lattice vector                                  | $\mathbf{G}$                                                                         | $\mathbf{G} = h_1 \mathbf{b}_1 + h_2 \mathbf{b}_2 + h_3 \mathbf{b}_3 = h_1 \mathbf{a}^* + h_2 \mathbf{b}^* + h_3 \mathbf{c}^*$<br>$\mathbf{G} \cdot \mathbf{R} = 2\pi m = 2\pi \sum_i n_i h_i$ | m <sup>-1</sup> | 1, 3         |
| unit cell lengths                                                    | $a; b; c$                                                                            |                                                                                                                                                                                                | m               |              |
| unit cell angles                                                     | $\alpha; \beta; \gamma$                                                              |                                                                                                                                                                                                | rad, 1          |              |
| reciprocal unit cell lengths                                         | $a^*; b^*; c^*$                                                                      |                                                                                                                                                                                                | m <sup>-1</sup> |              |
| reciprocal unit cell angles                                          | $\alpha^*; \beta^*; \gamma^*$                                                        |                                                                                                                                                                                                | rad, 1          |              |
| fractional coordinates                                               | $x; y; z$                                                                            | $x = X/a$                                                                                                                                                                                      | 1               | 4            |
| atomic scattering factor                                             | $f$                                                                                  | $f = E_a/E_e$                                                                                                                                                                                  | 1               | 5            |
| structure factor                                                     | $F(h, k, l)$                                                                         | $F = \sum_{n=1}^N f_n e^{2\pi i(hx_n + ky_n + lz_n)}$                                                                                                                                          | 1               | 6            |
| with indices $h, k, l$                                               |                                                                                      |                                                                                                                                                                                                |                 |              |
| lattice plane spacing                                                | $d$                                                                                  |                                                                                                                                                                                                | m               |              |
| Bragg angle                                                          | $\theta$                                                                             | $n\lambda = 2d \sin \theta$                                                                                                                                                                    | rad, 1          | 7            |
| order of reflection                                                  | $n$                                                                                  |                                                                                                                                                                                                | 1               |              |
| order parameters,                                                    |                                                                                      |                                                                                                                                                                                                |                 |              |
| short range                                                          | $\sigma$                                                                             |                                                                                                                                                                                                | 1               |              |
| long range                                                           | $s$                                                                                  |                                                                                                                                                                                                | 1               |              |
| Burgers vector                                                       | $\mathbf{b}$                                                                         |                                                                                                                                                                                                | m               |              |
| particle position vector                                             | $\mathbf{r}_j, \mathbf{R}_j$                                                         |                                                                                                                                                                                                | m               | 8            |
| equilibrium position                                                 | $\mathbf{R}_0$                                                                       |                                                                                                                                                                                                | m               |              |
| vector of an ion                                                     |                                                                                      |                                                                                                                                                                                                |                 |              |
| displacement vector of an ion                                        | $\mathbf{u}$                                                                         | $\mathbf{u} = \mathbf{R} - \mathbf{R}_0$                                                                                                                                                       | m               |              |
| Debye-Waller factor                                                  | $B, D$                                                                               | $D = e^{-2\langle(\mathbf{q}\cdot\mathbf{u})^2\rangle}$                                                                                                                                        | 1               | 9            |

- (1)  $n_1, n_2$  and  $n_3$  are integers.  $a, b$  and  $c$  are also called the lattice constants.
- (2) Reciprocal lattice vectors are sometimes defined by  $\mathbf{a}_i \cdot \mathbf{b}_k = \delta_{ik}$ .
- (3)  $m$  is an integer with  $m = n_1 h + n_2 k + n_3 l$ .
- (4)  $X$  denotes the coordinate of dimension length.
- (5)  $E_a$  and  $E_e$  denote the scattering amplitudes for the atom and the isolated electron, respectively.
- (6)  $N$  is the number of atoms in the unit cell.
- (7)  $\lambda$  is the wavelength of the incident radiation.
- (8) To distinguish between electron and ion position vectors, lower case and capital letters are used respectively. The subscript  $j$  relates to particle  $j$ .
- (9)  $\hbar\mathbf{q}$  is the momentum transfer in the scattering of neutrons,  $\langle \rangle$  denotes thermal averaging.

| <i>Name</i>                                | <i>Symbol</i>                        | <i>Definition</i>                                                       | <i>SI unit</i>                           | <i>Notes</i> |
|--------------------------------------------|--------------------------------------|-------------------------------------------------------------------------|------------------------------------------|--------------|
| Debye angular wavenumber                   | $q_D, k_D$                           | $k_D = (C_i 6\pi^2)^{1/3}$                                              | $\text{m}^{-1}$                          | 10           |
| Debye angular frequency                    | $\omega_D$                           | $\omega_D = k_D c_0$                                                    | $\text{s}^{-1}$                          | 10           |
| Debye frequency                            | $\nu_D$                              | $\nu_D = \omega_D/2\pi$                                                 | $\text{s}^{-1}$                          |              |
| Debye wavenumber                           | $\tilde{\nu}_D$                      | $\tilde{\nu}_D = \nu_D/c_0$                                             | $\text{m}^{-1}$                          | 10           |
| Debye temperature                          | $\Theta_D$                           | $\Theta_D = \hbar\nu_D/k_B$                                             | K                                        |              |
| Grüneisen parameter                        | $\gamma, \Gamma$                     | $\gamma = \alpha V/\kappa C_V$                                          | 1                                        | 11           |
| Madelung constant                          | $\alpha, \mathcal{M}$                | $E_{\text{coul}} = \frac{\alpha N_A z_+ z_- e^2}{4\pi\epsilon_0 R_0}$   | 1                                        | 12           |
| density of states                          | $N_E$                                | $N_E = dN(E)/dE$                                                        | $\text{J}^{-1} \text{m}^{-3}$            | 13           |
| (spectral) density of<br>vibrational modes | $N_\omega, g$                        | $N_\omega = dN(\omega)/d\omega$                                         | $\text{s m}^{-3}$                        | 14           |
| resistivity tensor                         | $\boldsymbol{\rho}, (\rho_{ik})$     | $\mathbf{E} = \boldsymbol{\rho} \cdot \mathbf{j}$                       | $\Omega \text{ m}$                       | 15           |
| conductivity tensor                        | $\boldsymbol{\sigma}, (\sigma_{ik})$ | $\mathbf{j} = \boldsymbol{\sigma} \cdot \mathbf{E}$                     | $\text{S m}^{-1}$                        | 15           |
| residual resistivity                       | $\rho_R$                             |                                                                         | $\Omega \text{ m}$                       |              |
| thermal conductivity tensor                | $\lambda_{ik}$                       | $\mathbf{J}_q = -\boldsymbol{\lambda} \cdot \nabla T$                   | $\text{W m}^{-1} \text{K}^{-1}$          | 15           |
| relaxation time                            | $\tau$                               | $\tau = l/v_F$                                                          | s                                        | 16           |
| Lorenz coefficient                         | $L$                                  | $L = \lambda/\sigma T$                                                  | $\text{V}^2 \text{K}^{-2}$               | 17           |
| Hall coefficient                           | $A_H, R_H$                           | $\mathbf{E} = \rho \mathbf{j} + R_H(\mathbf{B} \times \mathbf{j})$      | $\text{m}^3 \text{C}^{-1}$               |              |
| thermoelectric force                       | $E$                                  |                                                                         | V                                        | 18           |
| Peltier coefficient                        | $\Pi$                                | $E = \Pi \frac{\Delta T}{T}$                                            | V                                        | 18           |
| Thomson coefficient                        | $\mu, (\tau)$                        | $\mu = \Pi/T$                                                           | $\text{V K}^{-1}$                        |              |
| number density,<br>number concentration    | $C, n, p$                            |                                                                         | $\text{m}^{-3}$                          | 19           |
| band gap energy                            | $E_g$                                |                                                                         | J                                        | 20           |
| donor ionization energy                    | $E_d$                                |                                                                         | J                                        | 20           |
| acceptor ionization energy                 | $E_a$                                |                                                                         | J                                        | 20           |
| Fermi energy                               | $E_F, \epsilon_F$                    | $\epsilon_F = \lim_{T \rightarrow 0} \mu$                               | J                                        | 20           |
| work function,<br>electron work function   | $\Phi$                               | $\Phi = E_\infty - E_F$                                                 | J                                        | 21           |
| angular wave vector,<br>propagation vector | $\mathbf{k}, \mathbf{q}$             | $k = 2\pi/\lambda$                                                      | $\text{m}^{-1}$                          | 22           |
| Bloch function                             | $u_k(\mathbf{r})$                    | $\psi(\mathbf{r}) = u_k(\mathbf{r}) \exp(i\mathbf{k} \cdot \mathbf{r})$ | $\text{m}^{-3/2}$                        | 23           |
| charge density of electrons                | $\rho$                               | $\rho(\mathbf{r}) = -e\psi^*(\mathbf{r})\psi(\mathbf{r})$               | $\text{C m}^{-3}$                        | 23, 24       |
| effective mass                             | $m^*$                                |                                                                         | kg                                       | 25           |
| mobility                                   | $\mu$                                | $\mathbf{v}_{\text{drift}} = \mu \mathbf{E}$                            | $\text{m}^2 \text{V}^{-1} \text{s}^{-1}$ | 25           |
| mobility ratio                             | $b$                                  | $b = \mu_n/\mu_p$                                                       | 1                                        |              |
| diffusion coefficient                      | $D$                                  | $\mathbf{j} = -D\nabla C$                                               | $\text{m}^2 \text{s}^{-1}$               | 25, 26       |
| diffusion length                           | $L$                                  | $L = \sqrt{D\tau}$                                                      | m                                        | 25, 26       |
| characteristic (Weiss)<br>temperature      | $\theta, \theta_W$                   |                                                                         | K                                        |              |
| Curie temperature                          | $T_C$                                |                                                                         | K                                        |              |
| Néel temperature                           | $T_N$                                |                                                                         | K                                        |              |

(10)  $C_i$  is the ion density,  $c_0$  is the speed of light in vacuum.  $q_d$  is equal to  $2\pi$  times the inverse of the Debye cut-off wavelength of the elastic lattice wave.

(11)  $\alpha$  is the cubic expansion coefficient,  $V$  the volume,  $\kappa$  the isothermal compressibility, and  $C_V$  the heat capacity at constant volume.

(12)  $E_{\text{coul}}$  is the electrostatic interaction energy per mole of ion pairs with charges  $z_+e$  and  $-z_-e$ .

### 2.8.1 Symbols for planes and directions in crystals

|                                                                                  |                              |
|----------------------------------------------------------------------------------|------------------------------|
| Miller indices of a crystal face, or of a single net plane                       | $(hkl)$ or $(h_1h_2h_3)$     |
| indices of the Bragg reflection from the set of parallel net planes              | $hkl$ or $h_1h_2h_3$         |
| indices of a set of all symmetrically equivalent crystal faces,<br>or net planes | $\{hkl\}$ or $\{h_1h_2h_3\}$ |
| indices of a lattice direction (zone axis)                                       | $[uvw]$                      |
| indices of a set of symmetrically equivalent lattice directions                  | $\langle uvw \rangle$        |

In each of these cases, when the letter symbol is replaced by numbers it is customary to omit the commas. For a single plane or crystal face, or a specific direction, a negative number is indicated by a bar over the number.

*Example*  $(\bar{1}10)$  denotes the parallel planes  $h = -1$ ,  $k = 1$ ,  $l = 0$ .

#### (i) Crystal lattice symbols

|              |       |
|--------------|-------|
| primitive    | P     |
| face-centred | F     |
| body-centred | I     |
| base-centred | A;B;C |
| rhombohedral | R     |

#### (ii) Hermann-Mauguin symbols of symmetry operations

| Operation           | Symbol          | Examples                                                  |
|---------------------|-----------------|-----------------------------------------------------------|
| $n$ -fold rotation  | $n$             | 1; 2; 3; 4; 6                                             |
| $n$ -fold inversion | $\bar{n}$       | $\bar{1}$ ; $\bar{2}$ ; $\bar{3}$ ; $\bar{4}$ ; $\bar{6}$ |
| $n$ -fold screw     | $n_k$           | $2_1$ ; $3_1$ ; $3_2$ ;...                                |
| reflection          | $m$             |                                                           |
| glide               | $a; b; c; n; d$ |                                                           |

---

(Notes continued)

(13)  $N(E)$  is the total number of states of electronic energy less than  $E$ , divided by the volume.

(14)  $N(\omega)$  is the total number of vibrational modes with circular frequency less than  $\omega$ , divided by the volume.

(15) Tensors may be replaced by their corresponding scalar quantities in isotropic media.  $\mathbf{J}_q$  is the heat flux vector or thermal current density.

(16) The definition applies to electrons in metals;  $l$  is the mean free path, and  $v_F$  is the electron velocity on the Fermi sphere.

(17)  $\lambda$  and  $\sigma$  are the thermal and electrical conductivities in isotropic media.

(18) The substances to which the symbol applies are denoted by subscripts. The thermoelectric force is an electric potential difference induced by the gradient of the chemical potential.

(19) Specific number densities are denoted by subscripts: for electrons  $n_n$ ,  $n_-(n)$ ; for holes  $n_p$ ,  $n_+(p)$ ; for donors  $n_d$ ; for acceptors  $n_a$ ; for the intrinsic number density  $n_i$  ( $n_i^2 = n_+n_-$ ).

(20) The commonly used unit for this quantity is eV.  $\mu$  is the chemical potential per entity.

(21)  $E_\infty$  is the electron energy at rest at infinite distance [71].

(22)  $\mathbf{k}$  is used for particles,  $\mathbf{q}$  for phonons. Here,  $\lambda$  is the wavelength.

(23)  $\psi(\mathbf{r})$  is a one-electron wavefunction.

(24) The total charge density is obtained by summing over all electrons.

(25) Subscripts n and p or  $-$  and  $+$  may be added to denote electrons and holes, respectively.

(26)  $\mathbf{j}$  is the particle flux density.  $D$  is the diffusion coefficient and  $\tau$  the lifetime.

## 2.9 STATISTICAL THERMODYNAMICS

The names and symbols given here are in agreement with those recommended by IUPAP [4] and by ISO [5.h].

| <i>Name</i>                                                | <i>Symbol</i>                                                                               | <i>Definition</i>                                                                      | <i>SI unit</i>                    | <i>Notes</i> |
|------------------------------------------------------------|---------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|-----------------------------------|--------------|
| number of entities                                         | $N$                                                                                         |                                                                                        | 1                                 |              |
| number density of entities,<br>number concentration        | $C$                                                                                         | $C = N/V$                                                                              | $\text{m}^{-3}$                   |              |
| Avogadro constant                                          | $L, N_{\text{A}}$                                                                           | $L = N/n$                                                                              | $\text{mol}^{-1}$                 | 1            |
| Boltzmann constant                                         | $k, k_{\text{B}}$                                                                           |                                                                                        | $\text{J K}^{-1}$                 |              |
| (molar) gas constant                                       | $R$                                                                                         | $R = Lk$                                                                               | $\text{J K}^{-1} \text{mol}^{-1}$ |              |
| molecular position vector                                  | $\mathbf{r}(x, y, z)$                                                                       |                                                                                        | $\text{m}$                        |              |
| molecular velocity<br>vector                               | $\mathbf{c}(c_x, c_y, c_z),$<br>$\mathbf{u}(u_x, u_y, u_z),$<br>$\mathbf{v}(v_x, v_y, v_z)$ | $\mathbf{c} = \text{d}\mathbf{r}/\text{d}t$                                            | $\text{m s}^{-1}$                 |              |
| molecular momentum<br>vector                               | $\mathbf{p}(p_x, p_y, p_z)$                                                                 | $\mathbf{p} = m\mathbf{c}$                                                             | $\text{kg m s}^{-1}$              | 2            |
| velocity distribution<br>function                          | $f(c_x)$                                                                                    | $f = \left(\frac{m}{2\pi kT}\right)^{1/2} \exp\left(-\frac{mc_x^2}{2kT}\right)$        | $\text{m}^{-1} \text{s}$          | 2            |
| speed distribution<br>function                             | $F(c)$                                                                                      | $F = 4\pi c^2 \left(\frac{m}{2\pi kT}\right)^{3/2} \exp\left(-\frac{mc^2}{2kT}\right)$ | $\text{m}^{-1} \text{s}$          | 2            |
| average speed                                              | $\bar{c}, \bar{u}, \bar{v},$<br>$\langle c \rangle, \langle u \rangle, \langle v \rangle$   | $\bar{c} = \int cF(c) \text{d}c$                                                       | $\text{ms}^{-1}$                  |              |
| generalized coordinate                                     | $q$                                                                                         |                                                                                        | (varies)                          | 3            |
| generalized momentum                                       | $p$                                                                                         | $p = \partial L / \partial \dot{q}$                                                    | (varies)                          | 3            |
| volume in phase space                                      | $\Omega$                                                                                    | $\Omega = (1/h) \int p \text{d}q$                                                      | 1                                 |              |
| probability                                                | $P, p$                                                                                      |                                                                                        | 1                                 |              |
| statistical weight,<br>degeneracy                          | $g, d, W, \omega, \beta$                                                                    |                                                                                        | 1                                 | 4            |
| (cumulative) number<br>of states                           | $W, N$                                                                                      | $W(E) = \sum_i \text{H}(E - E_i)$                                                      | 1                                 | 5, 6         |
| density of states                                          | $\rho(E)$                                                                                   | $\rho(E) = \text{d}W(E)/\text{d}E$                                                     | $\text{J}^{-1}$                   |              |
| partition function,<br>sum over states,<br>single molecule | $q, z$                                                                                      | $q = \sum_i g_i \exp(-E_i/kT)$                                                         | 1                                 | 6            |
| canonical ensemble,<br>(system, or assembly)               | $Q, Z$                                                                                      | $Q = \sum_i g_i \exp(-E_i/kT)$                                                         | 1                                 | 6            |

(1)  $n$  is the amount of substance (or the chemical amount, enplethy). While the symbol  $N_{\text{A}}$  is used to honour Amedeo Avogadro, the symbol  $L$  is used in honour of Josef Loschmidt.

(2)  $m$  is the mass of the particle.

(3) If  $q$  is a length then  $p$  is a momentum. In the definition of  $p$ ,  $L$  denotes the Lagrangian.

(4)  $\beta$  is sometimes used for a spin statistical weight and the degeneracy is also called polytropy. It is the number of linearly independent energy eigenfunctions for the same energy.

(5)  $\text{H}(x)$  is the Heaviside function (see Section 4.2, p. 107),  $W$  or  $W(E)$  is the total number of quantum states with energy less than  $E$ .

(6)  $E_i$  denotes the energy of the  $i$ th level of a molecule or quantum system under consideration and  $g_i$  denotes its degeneracy.

| <i>Name</i>                                                   | <i>Symbol</i>                   | <i>Definition</i>                                      | <i>SI units</i>   | <i>Notes</i> |
|---------------------------------------------------------------|---------------------------------|--------------------------------------------------------|-------------------|--------------|
| microcanonical ensemble                                       | $\Omega, z, Z$                  | $Z = \sum_{i=1}^Z 1$                                   | 1                 |              |
| partition function, sum over states, grand canonical ensemble | $\Xi$                           |                                                        | 1                 |              |
| symmetry number                                               | $\sigma, s$                     |                                                        | 1                 |              |
| reciprocal energy parameter to replace temperature            | $\beta$                         | $\beta = 1/kT$                                         | J <sup>-1</sup>   |              |
| characteristic temperature                                    | $\Theta, \theta$                |                                                        | K                 | 7            |
| absolute activity                                             | $\lambda$                       | $\lambda_B = \exp(\mu_B/RT)$                           | 1                 | 8            |
| density operator                                              | $\hat{\rho}, \hat{\sigma}$      | $\hat{\rho} = \sum_k p_k  \Psi_k\rangle\langle\Psi_k $ | 1                 | 9            |
| density matrix                                                | $\mathbf{P}, \boldsymbol{\rho}$ | $\mathbf{P} = \{P_{mn}\}$                              | 1                 | 10           |
| element                                                       | $P_{mn}, \rho_{mn}$             | $P_{mn} = \langle\phi_m \hat{\rho} \phi_n\rangle$      | 1                 | 10           |
| statistical entropy                                           | $S$                             | $S = -k \sum_i p_i \ln p_i$                            | J K <sup>-1</sup> | 11           |

(7) Particular characteristic temperatures are denoted with subscripts, e.g. rotational  $\Theta_r = hc\tilde{B}/k$ , vibrational  $\Theta_v = hc\tilde{\nu}/k$ , Debye  $\Theta_D = hc\tilde{\nu}_D/k$ , Einstein  $\Theta_E = hc\tilde{\nu}_E/k$ .  $\Theta$  is to be preferred over  $\theta$  to avoid confusion with Celsius temperature.

(8) The definition applies to entities B.  $\mu_B$  is the chemical potential (see Section 2.11, p. 57).

(9)  $|\Psi_k\rangle$  refers to the quantum state  $k$  of the system and  $p_k$  to the probability of this state in an ensemble. If  $p_k = 1$  for a given state  $k$  one speaks of a pure state, otherwise of a mixture.

(10) The density matrix  $\mathbf{P}$  is defined by its matrix elements  $P_{mn}$  in a set of basis states  $\phi_m$ . Alternatively, one can write  $P_{mn} = \sum_k p_k c_m^{(k)} c_n^{(k)*}$ , where  $c_m^{(k)}$  is the (complex) coefficient of  $\phi_m$  in the expansion of  $|\Psi_k\rangle$  in the basis states  $\{\phi_i\}$ .

(11) In the expression for the statistical entropy,  $k$  is the Boltzmann constant and  $p_i$  the average population or probability for a quantum level. The equilibrium (maximum) entropy for a microcanonical ensemble results then as  $S = k \ln Z$ . A variety of other ensembles beyond the microcanonical, canonical, or grand canonical can be defined.

## 2.10 GENERAL CHEMISTRY

The symbols given by IUPAP [4] and by ISO [5.d,5.h] are in agreement with the recommendations given here.

| <i>Name</i>                                                            | <i>Symbol</i> | <i>Definition</i>             | <i>SI unit</i>               | <i>Notes</i> |
|------------------------------------------------------------------------|---------------|-------------------------------|------------------------------|--------------|
| number of entities<br>(e.g. molecules, atoms,<br>ions, formula units)  | $N$           |                               | 1                            |              |
| amount of substance, amount<br>(chemical amount)                       | $n$           | $n_B = N_B/L$                 | mol                          | 1, 2         |
| Avogadro constant                                                      | $L, N_A$      |                               | $\text{mol}^{-1}$            | 3            |
| mass of atom,<br>atomic mass                                           | $m_a, m$      |                               | kg                           |              |
| mass of entity<br>(molecule, formula unit)                             | $m, m_f$      |                               | kg                           | 4            |
| atomic mass constant                                                   | $m_u$         | $m_u = m_a(^{12}\text{C})/12$ | kg                           | 5            |
| molar mass                                                             | $M$           | $M_B = m/n_B$                 | $\text{kg mol}^{-1}$         | 2, 6, 7      |
| molar mass constant                                                    | $M_u$         | $M_u = m_u N_A$               | $\text{g mol}^{-1}$          | 7, 8         |
| relative molecular mass,<br>(relative molar mass,<br>molecular weight) | $M_r$         | $M_r = m_f/m_u$               | 1                            | 8            |
| relative atomic mass,<br>(atomic weight)                               | $A_r$         | $A_r = m_a/m_u$               | 1                            | 8            |
| molar volume                                                           | $V_m$         | $V_{m,B} = V/n_B$             | $\text{m}^3 \text{mol}^{-1}$ | 2, 6, 7      |
| mass fraction                                                          | $w$           | $w_B = m_B / \sum_i m_i$      | 1                            | 2            |

(1) The words “of substance” may be replaced by the specification of the entity, e.g. “amount of oxygen atoms” or “amount of oxygen (or dioxygen,  $\text{O}_2$ ) molecules”. Note that “amount of oxygen” is ambiguous and should be used only if the meaning is clear from the context (see also the discussion in Section 2.10.1 (v), p. 53).

*Example* When the amount of  $\text{O}_2$  is equal to 3 mol,  $n(\text{O}_2) = 3 \text{ mol}$ , the amount of  $(1/2) \text{O}_2$  is equal to 6 mol, and  $n((1/2) \text{O}_2) = 6 \text{ mol}$ . Thus  $n((1/2) \text{O}_2) = 2n(\text{O}_2)$ .

(2) The definition applies to entities B which should always be indicated by a subscript or in parentheses, e.g.  $n_B$  or  $n(\text{B})$ . When the chemical composition is written out, parentheses should be used,  $n(\text{O}_2)$ .

(3) The symbol  $N_A$  is used to honour Amedeo Avogadro, the symbol  $L$  is used in honour of Josef Loschmidt.

(4) Note that “formula unit” is not a unit (see examples in Section 2.10.1 (iii), p. 50).

(5)  $m_u$  is equal to the unified atomic mass unit, with symbol u, i.e.  $m_u = 1 \text{ u}$  (see Section 3.7, p. 92). The dalton, symbol Da, is used as an alternative name for the unified atomic mass unit.

(6) The definition applies to pure substance, where  $m$  is the total mass and  $V$  is the total volume. However, corresponding quantities may also be defined for a mixture as  $m/n$  and  $V/n$ , where  $n = \sum_i n_i$ . These quantities are called the mean molar mass and the mean molar volume respectively.

(7) These names, which include the word “molar”, unfortunately use the name of a *unit* in the description of a *quantity*, which in principle is to be avoided.

(8) For historical reasons the terms “molecular weight” and “atomic weight” are still used. For molecules  $M_r$  is the relative molecular mass or “molecular weight”. For atoms  $M_r$  is the relative atomic mass or “atomic weight”, and the symbol  $A_r$  may be used.  $M_r$  may also be called the relative molar mass,  $M_{r,B} = M_B/M_u$ , where  $M_u = 1 \text{ g mol}^{-1}$ . The standard atomic weights are listed in Section 6.2, p. 117.

| <i>Name</i>                                                  | <i>Symbol</i>  | <i>Definition</i>                | <i>SI unit</i>       | <i>Notes</i> |
|--------------------------------------------------------------|----------------|----------------------------------|----------------------|--------------|
| volume fraction                                              | $\phi$         | $\phi_B = V_B / \sum_i V_i$      | 1                    | 2, 9         |
| mole fraction, amount-of-substance fraction, amount fraction | $x, y$         | $x_B = n_B / \sum_i n_i$         | 1                    | 2, 10        |
| (total) pressure                                             | $p, (P)$       |                                  | Pa                   | 2, 11        |
| partial pressure                                             | $p_B$          | $p_B = y_B p$                    | Pa                   | 12           |
| mass concentration, (mass density)                           | $\gamma, \rho$ | $\gamma_B = m_B / V$             | kg m <sup>-3</sup>   | 2, 13, 14    |
| number concentration, number density of entities             | $C, n$         | $C_B = N_B / V$                  | m <sup>-3</sup>      | 2, 13, 15    |
| amount concentration, concentration                          | $c, [B]$       | $c_B = n_B / V$                  | mol m <sup>-3</sup>  | 2, 13, 16    |
| solubility                                                   | $s$            | $s_B = c_B$ (saturated solution) | mol m <sup>-3</sup>  | 2, 17        |
| molality                                                     | $m, b$         | $m_B = n_B / m_A$                | mol kg <sup>-1</sup> | 2, 7, 14     |
| surface concentration                                        | $\Gamma$       | $\Gamma_B = n_B / A$             | mol m <sup>-2</sup>  | 2, 18        |
| stoichiometric number                                        | $\nu$          |                                  | 1                    | 19           |
| extent of reaction, advancement                              | $\xi$          | $\xi = (n_B - n_{B,0}) / \nu_B$  | mol                  | 19           |
| degree of reaction                                           | $\alpha$       | $\alpha = \xi / \xi_{\max}$      | 1                    | 20           |

(9) Here,  $V_B$  and  $V_i$  are the volumes of appropriate components prior to mixing. As other definitions are possible, e.g. ISO 31 [5], the term should not be used in accurate work without spelling out the definition.

(10) For condensed phases  $x$  is used, and for gaseous mixtures  $y$  may be used [72].

(11) Pressures are often expressed in the non-SI unit bar, where 1 bar = 10<sup>5</sup> Pa. The standard pressure  $p^\ominus = 1 \text{ bar} = 10^5 \text{ Pa}$  (see Section 2.11.1 (v), p. 62, Section 7.2, p. 138, and the conversion table on p. 233). Pressures are often expressed in millibar or hectopascal, where 1 mbar = 10<sup>-3</sup> bar = 100 Pa = 1 hPa.

(12) The symbol and the definition apply to molecules B, which should be specified. In real (non-ideal) gases, care is required in defining the partial pressure.

(13)  $V$  is the volume of the mixture. Quantities that describe compositions of mixtures can be found in [72].

(14) In this definition the symbol  $m$  is used with two different meanings:  $m_B$  denotes the *molality* of solute B,  $m_A$  denotes the *mass* of solvent A (thus the unit mol kg<sup>-1</sup>). This confusion of notation is avoided by using the symbol  $b$  for molality. A solution of molality 1 mol kg<sup>-1</sup> is occasionally called a 1 molal solution, denoted 1 m solution; however, the symbol m must not be treated as a symbol for the unit mol kg<sup>-1</sup> in combination with other units.

(15) The term number concentration and symbol  $C$  is preferred for mixtures. Care must be taken not to use the symbol  $n$  where it may be mistakenly interpreted to denote amount of substance.

(16) “Amount concentration” is an abbreviation of “amount-of-substance concentration”. (The Clinical Chemistry Division of IUPAC recommends that amount-of-substance concentration be abbreviated to “substance concentration” [14, 73].) The word “concentration” is normally used alone where there is no risk of confusion, as in conjunction with the name (or symbol) of a chemical substance, or as contrast to molality (see Section 2.13, p. 70). In polymer science the word “concentration” and the symbol  $c$  is normally used for mass concentration. In the older literature this quantity was often called *molarity*, a usage that should be avoided due to the risk of confusion with the quantity *molality*. Units commonly used for amount concentration are mol L<sup>-1</sup> (or mol dm<sup>-3</sup>), mmol L<sup>-1</sup>,  $\mu\text{mol L}^{-1}$  etc., often denoted M, mM,  $\mu\text{M}$  etc. (pronounced molar, millimolar, micromolar).

## 2.10.1 Other symbols and conventions in chemistry

### (i) The symbols for the chemical elements

The symbols for the chemical elements are (in most cases) derived from their Latin names and consist of one or two letters which should always be printed in Roman (upright) type. A complete list is given in Section 6.2, p. 117. The symbol is not followed by a full stop except at the end of a sentence.

*Examples* I, U, Pa, C

The symbols have two different meanings (which also reflects on their use in chemical formulae and equations):

(a) On a *microscopic* level they can denote an atom of the element. For example, Cl denotes a chlorine atom having 17 protons and 18 or 20 neutrons (giving a mass number of 35 or 37), the difference being ignored. Its mass is on average 35.4527 u in terrestrial samples.

(b) On a *macroscopic* level they denote a sample of the element. For example, Fe denotes a sample of iron, and He a sample of helium gas. They may also be used as a shorthand to denote the element: “Fe is one of the most common elements in the Earth’s crust.”

The term *nuclide* implies an atom of specified atomic number (proton number) and mass number (nucleon number). A nuclide may be specified by attaching the mass number as a left superscript to the *symbol* for the element, as in  $^{14}\text{C}$ , or added after the *name* of the element, as in carbon-14. Nuclides having the same atomic number but different mass numbers are called isotopic nuclides or *isotopes*, as in  $^{12}\text{C}$ ,  $^{14}\text{C}$ . If no left superscript is attached, the symbol is read as including all isotopes in natural abundance:  $n(\text{Cl}) = n(^{35}\text{Cl}) + n(^{37}\text{Cl})$ . Nuclides having the same mass number but different atomic numbers are called isobaric nuclides or *isobars*:  $^{14}\text{C}$ ,  $^{14}\text{N}$ . The atomic number may be attached as a left subscript:  $^{14}_6\text{C}$ ,  $^{14}_7\text{N}$ .

The ionic charge number is denoted by a right superscript, by the sign alone when the charge number is equal to plus one or minus one.

|                 |                                        |                                                  |
|-----------------|----------------------------------------|--------------------------------------------------|
| <i>Examples</i> | $\text{Na}^+$                          | a sodium positive ion (cation)                   |
|                 | $^{79}\text{Br}^-$                     | a bromine-79 negative ion (anion, bromide ion)   |
|                 | $\text{Al}^{3+}$ or $\text{Al}^{+3}$   | aluminium triply positive ion                    |
|                 | $3 \text{S}^{2-}$ or $3 \text{S}^{-2}$ | three sulfur doubly negative ions (sulfide ions) |

$\text{Al}^{3+}$  is commonly used in chemistry and recommended by [74]. The forms  $\text{Al}^{+3}$  and  $\text{S}^{-2}$ , although widely used, are obsolete [74], as well as the old notation  $\text{Al}^{+++}$ ,  $\text{S}^=$ , and  $\text{S}^{--}$ .

---

(Notes continued)

(16) (continued) Thus M is often treated as a symbol for  $\text{mol L}^{-1}$ .

(17) Solubility may also be expressed in any units corresponding to quantities that denote relative composition, such as mass fraction, amount fraction, molality, volume fraction, etc.

(18) A denotes the surface area.

(19) The stoichiometric number is defined through the stoichiometric equation. It is negative for reactants and positive for products. The values of the stoichiometric numbers depend on how the reaction equation is written (see Section 2.10.1 (iv), p. 52).  $n_{\text{B},0}$  denotes the value of  $n_{\text{B}}$  at “zero time”, when  $\xi = 0$  mol.

(20)  $\xi_{\text{max}}$  is the value of  $\xi$  when at least one of the reactants is exhausted. For specific reactions, terms such as “degree of dissociation” and “degree of ionization” etc. are commonly used.

The right superscript position is also used to convey other information. Excited electronic states may be denoted by an asterisk.

*Examples*  $\text{H}^*$ ,  $\text{Cl}^*$

Oxidation numbers are denoted by positive or negative Roman numerals or by zero (see also Section 2.10.1 (iv), p. 52).

*Examples*  $\text{Mn}^{\text{VII}}$ , manganese(VII),  $\text{O}^{-\text{II}}$ ,  $\text{Ni}^0$

The positions and meanings of indices around the symbol of the element are summarized as follows:

|                   |                                                             |
|-------------------|-------------------------------------------------------------|
| left superscript  | mass number                                                 |
| left subscript    | atomic number                                               |
| right superscript | charge number, oxidation number, excitation                 |
| right subscript   | number of atoms per entity (see Section 2.10.1 (iii) below) |

## (ii) Symbols for particles and nuclear reactions

|             |                                       |               |                          |                       |                                   |
|-------------|---------------------------------------|---------------|--------------------------|-----------------------|-----------------------------------|
| proton      | $\text{p}$ , $\text{p}^+$             | positron      | $\text{e}^+$ , $\beta^+$ | triton                | $\text{t}$                        |
| antiproton  | $\bar{\text{p}}$                      | positive muon | $\mu^+$                  | helion                | $\text{h}$ ( $^3\text{He}^{2+}$ ) |
| neutron     | $\text{n}$                            | negative muon | $\mu^-$                  | $\alpha$ -particle    | $\alpha$ ( $^4\text{He}^{2+}$ )   |
| antineutron | $\bar{\text{n}}$                      | photon        | $\gamma$                 | (electron) neutrino   | $\nu_{\text{e}}$                  |
| electron    | $\text{e}$ , $\text{e}^-$ , $\beta^-$ | deuteron      | $\text{d}$               | electron antineutrino | $\bar{\nu}_{\text{e}}$            |

Particle symbols are printed in Roman (upright) type (but see Chapter 6, p. 113). The electric charge of particles may be indicated by adding the superscript  $+$ ,  $-$ , or  $0$ ; e.g.  $\text{p}^+$ ,  $\text{n}^0$ ,  $\text{e}^-$ , etc. If the symbols  $\text{p}$  and  $\text{e}$  are used without a charge, they refer to the positive proton and negative electron respectively. A summary of recommended names for muonium and hydrogen atoms and their ions can be found in [75].

The meaning of the symbolic expression indicating a nuclear reaction should be as follows:

$$\begin{array}{ccc} \text{initial} & \left( \begin{array}{cc} \text{incoming particles} & , \quad \text{outgoing particles} \\ \text{nuclide} & \text{or quanta} \end{array} \right. & \left. \begin{array}{cc} & \\ & \text{or quanta} \end{array} \right) & \text{final} \\ & & & \text{nuclide} \end{array}$$

*Examples*  $^{14}\text{N}(\alpha, \text{p})^{17}\text{O}$ ,  $^{59}\text{Co}(\text{n}, \gamma)^{60}\text{Co}$   
 $^{23}\text{Na}(\gamma, 3\text{n})^{20}\text{Na}$ ,  $^{31}\text{P}(\gamma, \text{pn})^{29}\text{Si}$

One can also use the standard notation from kinetics (see Section 2.12, p. 63).

*Examples*  $^{14}_7\text{N} + \alpha \rightarrow ^{17}_8\text{O} + \text{p}$   
 $\text{n} \rightarrow \text{p} + \text{e} + \bar{\nu}_{\text{e}}$

## (iii) Chemical formulae

As in the case of chemical symbols of elements, chemical formulae have two different meanings:

(a) On a *microscopic* level they denote one atom, one molecule, one ion, one radical, etc. The number of atoms in an entity (always an integer) is indicated by a right subscript, the numeral 1 being omitted. Groups of atoms may be enclosed in parentheses. Charge numbers of ions and excitation symbols are added as right superscripts to the formula. The radical nature of an entity may be expressed by adding a dot to the symbol. The nomenclature for radicals, ions, radical ions, and related species is described in [76, 77].

*Examples* Xe, N<sub>2</sub>, C<sub>6</sub>H<sub>6</sub>, Na<sup>+</sup>, SO<sub>4</sub><sup>2-</sup>  
 (CH<sub>3</sub>)<sub>3</sub>COH (a 2-methyl-2-propanol molecule)  
 NO<sub>2</sub><sup>\*</sup> (an excited nitrogen dioxide molecule)  
 NO (a nitrogen oxide molecule)  
 NO<sup>•</sup> (a nitrogen oxide molecule, stressing its free radical character)

In writing the formula for a complex ion, spacing for charge number can be added (staggered arrangement), as well as parentheses: SO<sub>4</sub><sup>2-</sup>, (SO<sub>4</sub>)<sup>2-</sup>. The staggered arrangement is now recommended [74].

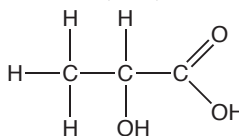
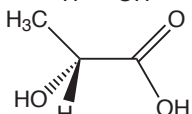
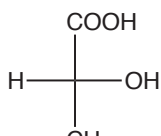
Specific electronic states of entities (atoms, molecules, ions) can be denoted by giving the electronic term symbol (see Section 2.6.3, p. 32) in parentheses. Vibrational and rotational states can be specified by giving the corresponding quantum numbers (see Section 2.6, p. 25 and 33).

*Examples* Hg(<sup>3</sup>P<sub>1</sub>) a mercury atom in the triplet-P-one state  
 HF(*v* = 2, *J* = 6) a hydrogen fluoride molecule in the vibrational state  
*v* = 2 and the rotational state *J* = 6  
 H<sub>2</sub>O<sup>+</sup>(<sup>2</sup>A<sub>1</sub>) a water molecule ion in the doublet-A-one state

(b) On a *macroscopic* level a formula denotes a sample of a chemical substance (not necessarily stable, or capable of existing in isolated form). The chemical composition is denoted by right subscripts (not necessarily integers; the numeral 1 being omitted). A “formula unit” (which is *not* a unit of a quantity!) is an entity specified as a group of atoms (see (iv) and (v) below).

*Examples* Na, Na<sup>+</sup>, NaCl, Fe<sub>0.91</sub>S, XePtF<sub>6</sub>, NaCl

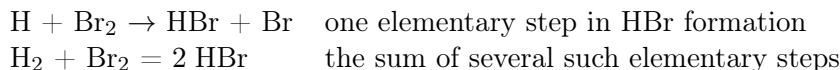
The formula can be used in expressions like ρ(H<sub>2</sub>SO<sub>4</sub>), mass density of sulfuric acid. When specifying amount of substance the formula is often multiplied with a factor, normally a small integer or a fraction, see examples in (iv) and (v). Less formally, the formula is often used as a shorthand (“reacting with H<sub>2</sub>SO<sub>4</sub>”). Chemical formulae may be written in different ways according to the information they convey [15, 74, 78, 79]:

| Formula             | Information conveyed            | Example                                                                               | Notes |
|---------------------|---------------------------------|---------------------------------------------------------------------------------------|-------|
| empirical           | stoichiometric proportion only  | CH <sub>2</sub> O                                                                     | 1     |
| molecular           | in accord with molecular mass   | C <sub>3</sub> H <sub>6</sub> O <sub>3</sub>                                          |       |
| structural          | structural arrangement of atoms | CH <sub>3</sub> CH(OH)COOH                                                            | 1     |
| connectivity        | connectivity                    |  | 2     |
| stereochemical      | stereochemical configuration    |  | 2     |
|                     | Fischer projection              |  | 3     |
| resonance structure | electronic arrangement          |  | 2, 4  |

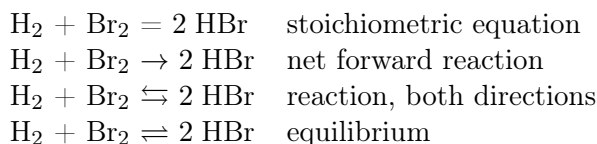
(1) Molecules differing only in isotopic composition are called isotopomers or isotopologues. For example, CH<sub>2</sub>O, CHDO, CD<sub>2</sub>O and CH<sub>2</sub><sup>17</sup>O are all isotopomers or isotopologues of the formaldehyde molecule. It has been suggested [16] to reserve isotopomer for molecules of the same isotopic

#### (iv) Equations for chemical reactions

(a) On a *microscopic* level the reaction equation represents an elementary reaction (an event involving single atoms, molecules, and radicals), or the sum of a set of such reactions. Stoichiometric numbers are  $\pm 1$  (sometimes  $\pm 2$ ). A single arrow is used to connect reactants and products in an elementary reaction. An equal sign is used for the “net” reaction, the result of a set of elementary reactions (see Section 2.12.1, p. 68).



(b) On a *macroscopic* level, different symbols are used connecting the reactants and products in the reaction equation, with the following meanings:

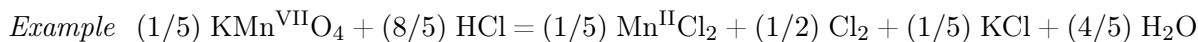


The two-sided arrow  $\leftrightarrow$  should not be used for reactions to avoid confusion with resonance structures (see Section 2.10.1 (iii), p. 50).

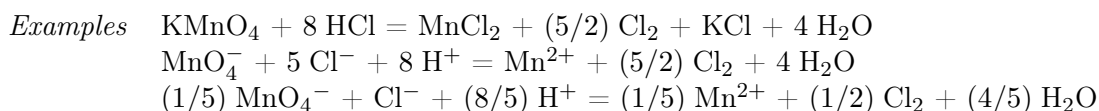
Stoichiometric numbers are not unique. One and the same reaction can be expressed in different ways (without any change in meaning).

*Examples* The formation of hydrogen bromide from the elements can equally well be written in either of these two ways  
 $(1/2) \text{H}_2 + (1/2) \text{Br}_2 = \text{HBr}$   
 $\text{H}_2 + \text{Br}_2 = 2 \text{HBr}$

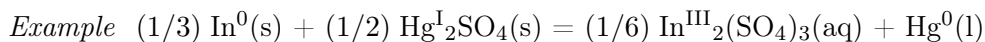
Ions not taking part in a reaction (“spectator ions”) are often removed from an equation. Redox equations are often written so that the value of the stoichiometric number for the electrons transferred (which are normally omitted from the overall equation) is equal to  $\pm 1$ .



The oxidation of chloride by permanganate in acid solution can thus be represented in several (equally correct) ways.



Similarly a reaction in an electrochemical cell may be written so that the electron number of an electrochemical cell reaction,  $z$ , (see Section 2.13, p. 71) is equal to one:



where the symbols in parentheses denote the state of aggregation (see Section 2.10.1 (vi), p. 54).

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(Notes continued)

(1) (continued) composition but different structure, such as  $\text{CD}_3\text{CH}(\text{OH})\text{COOH}$  and  $\text{CH}_3\text{CD}(\text{OD})\text{COOD}$  for which one also uses isotope-isomer.

(2) The lines in the connectivity, stereochemical, or resonance structure formula represent single, double, or triple bonds. They are also used as representing lone-pair electrons.

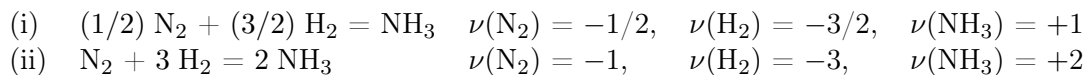
(3) In the Fischer projection the horizontal substituents are interpreted as above the plane of the paper whereas the vertical substituents are interpreted as behind the plane of the paper.

(4) The two-sided arrow represents the arrangement of valence electrons in resonance structures, such as benzene  $\text{C}_6\text{H}_6$  and does not signify a reaction equation (see Section 2.10.1 (iv), this page).

**Symbolic Notation:** A general chemical equation can be written as

$$0 = \sum_j \nu_j B_j$$

where  $B_j$  denotes a species in the reaction and  $\nu_j$  the corresponding stoichiometric number (negative for reactants and positive for products). The ammonia synthesis is equally well expressed in these two possible ways:



The changes  $\Delta n_j = n_j - n_{j,0}$  in the amounts of any reactant and product  $j$  during the course of the reaction is governed by one parameter, the extent of reaction  $\xi$ , through the equation

$$n_j = n_{j,0} + \nu_j \xi$$

The extent of reaction depends on how the reaction is written, but it is independent of which entity in the reaction is used in the definition. Thus, for reaction (i), when  $\xi = 2$  mol, then  $\Delta n(\text{N}_2) = -1$  mol,  $\Delta n(\text{H}_2) = -3$  mol and  $\Delta n(\text{NH}_3) = +2$  mol. For reaction (ii), when  $\Delta n(\text{N}_2) = -1$  mol then  $\xi = 1$  mol.

**Matrix Notation:** For multi-reaction systems it is convenient to write the chemical equations in matrix form

$$\mathbf{A} \boldsymbol{\nu} = \mathbf{0}$$

where  $\mathbf{A}$  is the conservation (or formula) matrix with elements  $A_{ij}$  representing the number of atoms of the  $i$ th element in the  $j$ th reaction species (reactant or product entity) and  $\boldsymbol{\nu}$  is the stoichiometric number matrix with elements  $\nu_{jk}$  being the stoichiometric numbers of the  $j$ th reaction species in the  $k$ th reaction. When there are  $N_s$  reacting species involved in the system consisting of  $N_e$  elements  $\mathbf{A}$  becomes an  $N_e \times N_s$  matrix. Its nullity,  $N(\mathbf{A}) = N_s - \text{rank}(\mathbf{A})$ , gives the number of independent chemical reactions,  $N_r$ , and the  $N_s \times N_r$  stoichiometric number matrix,  $\boldsymbol{\nu}$ , can be determined as the null space of  $\mathbf{A}$ .  $\mathbf{0}$  is an  $N_e \times N_r$  zero matrix [80].

## (v) Amount of substance and the specification of entities

The quantity “amount of substance” or “chemical amount” (“Stoffmenge” in German, “quantité de matière” in French) has been used by chemists for a long time without a proper name. It was simply referred to as the “number of moles”. This practice should be abandoned: the name of a physical quantity should not contain the name of a unit (few would use “number of metres” as a synonym for “length”).

The amount of substance is proportional to the number of specified elementary entities of that substance; the proportionality constant is the same for all substances and is the reciprocal of the Avogadro constant. The elementary entities may be chosen as convenient, not necessarily as physically real individual particles. In the examples below,  $(1/2) \text{ Cl}_2$ ,  $(1/5) \text{ KMnO}_4$ , etc. are artificial in the sense that no such elementary entities exists. Since the amount of substance and all physical quantities derived from it depend on this choice it is essential to specify the entities to avoid ambiguities.

|                 |                                                  |                                                            |
|-----------------|--------------------------------------------------|------------------------------------------------------------|
| <i>Examples</i> | $n_{\text{Cl}}, n(\text{Cl})$                    | amount of Cl, amount of chlorine atoms                     |
|                 | $n(\text{Cl}_2)$                                 | amount of $\text{Cl}_2$ , amount of chlorine molecules     |
|                 | $n(\text{H}_2\text{SO}_4)$                       | amount of (entities) $\text{H}_2\text{SO}_4$               |
|                 | $n((1/5) \text{KMnO}_4)$                         | amount of (entities) $(1/5) \text{KMnO}_4$                 |
|                 | $M(\text{P}_4)$                                  | molar mass of tetraphosphorus $\text{P}_4$                 |
|                 | $c_{\text{Cl}^-}, c(\text{Cl}^-), [\text{Cl}^-]$ | amount concentration of $\text{Cl}^-$                      |
|                 | $\rho(\text{H}_2\text{SO}_4)$                    | mass density of sulfuric acid                              |
|                 | $\Lambda(\text{MgSO}_4)$                         | molar conductivity of (entities of) $\text{MgSO}_4$        |
|                 | $\Lambda((1/2) \text{MgSO}_4)$                   | molar conductivity of (entities of) $(1/2) \text{MgSO}_4$  |
|                 | $\lambda(\text{Mg}^{2+})$                        | ionic conductivity of (entities of) $\text{Mg}^{2+}$       |
|                 | $\lambda((1/2) \text{Mg}^{2+})$                  | ionic conductivity of (entities of) $(1/2) \text{Mg}^{2+}$ |

Using definitions of various quantities we can derive equations like

$$\begin{aligned}
 n((1/5) \text{KMnO}_4) &= 5n(\text{KMnO}_4) \\
 \lambda((1/2) \text{Mg}^{2+}) &= (1/2)\lambda(\text{Mg}^{2+}) \\
 [(1/2) \text{H}_2\text{SO}_4] &= 2 [\text{H}_2\text{SO}_4]
 \end{aligned}$$

(See also examples in Section 3.3, p. 88.)

Note that “amount of sulfur” is an ambiguous statement, because it might imply  $n(\text{S})$ ,  $n(\text{S}_8)$ , or  $n(\text{S}_2)$ , etc. In most cases analogous statements are less ambiguous. Thus for compounds the implied entity is usually the molecule or the common formula entity, and for solid metals it is the atom.

|                 |                                                                         |
|-----------------|-------------------------------------------------------------------------|
| <i>Examples</i> | “2 mol of water” implies $n(\text{H}_2\text{O}) = 2 \text{ mol}$        |
|                 | “0.5 mol of sodium chloride” implies $n(\text{NaCl}) = 0.5 \text{ mol}$ |
|                 | “3 mmol of iron” implies $n(\text{Fe}) = 3 \text{ mmol}$                |

Such statements should be avoided whenever there might be ambiguity.

In the equation  $pV = nRT$  and in equations involving colligative properties, the entity implied in the definition of  $n$  should be an independently translating particle (a whole molecule for a gas), whose nature is unimportant. Quantities that describe compositions of mixtures can be found in [72].

## (vi) States of aggregation

The following one-, two- or three-letter symbols are used to represent the states of aggregation of chemical species [1.j]. The letters are appended to the formula symbol in parentheses, and should be printed in Roman (upright) type without a full stop (period).

|              |                                            |     |                    |
|--------------|--------------------------------------------|-----|--------------------|
| a, ads       | species adsorbed on a surface              | g   | gas or vapor       |
| am           | amorphous solid                            | l   | liquid             |
| aq           | aqueous solution                           | lc  | liquid crystal     |
| aq, $\infty$ | aqueous solution at infinite dilution      | mon | monomeric form     |
| cd           | condensed phase<br>(i.e., solid or liquid) | n   | nematic phase      |
| cr           | crystalline                                | pol | polymeric form     |
| f            | fluid phase<br>(i.e., gas or liquid)       | s   | solid              |
|              |                                            | sln | solution           |
|              |                                            | vit | vitreous substance |

|                 |                                           |                                                           |
|-----------------|-------------------------------------------|-----------------------------------------------------------|
| <i>Examples</i> | $\text{HCl(g)}$                           | hydrogen chloride in the gaseous state                    |
|                 | $C_V(\text{f})$                           | heat capacity of a fluid at constant volume               |
|                 | $V_m(\text{lc})$                          | molar volume of a liquid crystal                          |
|                 | $U(\text{cr})$                            | internal energy of a crystalline solid                    |
|                 | $\text{MnO}_2(\text{am})$                 | manganese dioxide as an amorphous solid                   |
|                 | $\text{MnO}_2(\text{cr, I})$              | manganese dioxide as crystal form I                       |
|                 | $\text{NaOH(aq)}$                         | aqueous solution of sodium hydroxide                      |
|                 | $\text{NaOH(aq, } \infty)$                | aqueous solution of sodium hydroxide at infinite dilution |
|                 | $\Delta_f H^\circ(\text{H}_2\text{O, l})$ | standard enthalpy of formation of liquid water            |

The symbols g, l, etc. to denote gas phase, liquid phase, etc., are also sometimes used as a right superscript, and the Greek letter symbols  $\alpha$ ,  $\beta$ , etc. may be similarly used to denote phase  $\alpha$ , phase  $\beta$ , etc. in a general notation.

|                 |                                  |                                                          |
|-----------------|----------------------------------|----------------------------------------------------------|
| <i>Examples</i> | $V_m^{\text{l}}, V_m^{\text{s}}$ | molar volume of the liquid phase, ... of the solid phase |
|                 | $S_m^{\alpha}, S_m^{\beta}$      | molar entropy of phase $\alpha$ , ... of phase $\beta$   |

## 2.11 CHEMICAL THERMODYNAMICS

The names and symbols of the more generally used quantities given here are also recommended by IUPAP [4] and by ISO [5.d,5.h]. Additional information can be found in [1.d,1.j] and [81].

| <i>Name</i>                            | <i>Symbol</i>              | <i>Definition</i>                                | <i>SI unit</i>                     | <i>Notes</i> |
|----------------------------------------|----------------------------|--------------------------------------------------|------------------------------------|--------------|
| heat                                   | $Q, q$                     |                                                  | J                                  | 1            |
| work                                   | $W, w$                     |                                                  | J                                  | 1            |
| internal energy                        | $U$                        | $dU = dQ + dW$                                   | J                                  | 1            |
| enthalpy                               | $H$                        | $H = U + pV$                                     | J                                  |              |
| thermodynamic temperature              | $T, (\Theta)$              |                                                  | K                                  |              |
| International temperature              | $T_{90}$                   |                                                  | K                                  | 2            |
| Celsius temperature                    | $\theta, t$                | $\theta/^{\circ}\text{C} = T/\text{K} - 273.15$  | $^{\circ}\text{C}$                 | 3            |
| entropy                                | $S$                        | $dS = dQ_{\text{rev}}/T$                         | $\text{J K}^{-1}$                  |              |
| Helmholtz energy, (Helmholtz function) | $A, F$                     | $A = U - TS$                                     | J                                  | 4            |
| Gibbs energy, (Gibbs function)         | $G$                        | $G = H - TS$                                     | J                                  |              |
| Massieu function                       | $J$                        | $J = -A/T$                                       | $\text{J K}^{-1}$                  |              |
| Planck function                        | $Y$                        | $Y = -G/T$                                       | $\text{J K}^{-1}$                  |              |
| surface tension                        | $\gamma, \sigma$           | $\gamma = (\partial G / \partial A_s)_{T,p,n_i}$ | $\text{J m}^{-2}, \text{N m}^{-1}$ |              |
| molar quantity $X$                     | $X_m, (\bar{X})$           | $X_m = X/n$                                      | $[X]/\text{mol}$                   | 5, 6         |
| specific quantity $X$                  | $x$                        | $x = X/m$                                        | $[X]/\text{kg}$                    | 5, 6         |
| pressure coefficient                   | $\beta$                    | $\beta = (\partial p / \partial T)_V$            | $\text{Pa K}^{-1}$                 |              |
| relative pressure coefficient          | $\alpha_p$                 | $\alpha_p = (1/p)(\partial p / \partial T)_V$    | $\text{K}^{-1}$                    |              |
| compressibility, isothermal            | $\kappa_T$                 | $\kappa_T = -(1/V)(\partial V / \partial p)_T$   | $\text{Pa}^{-1}$                   |              |
| isentropic                             | $\kappa_S$                 | $\kappa_S = -(1/V)(\partial V / \partial p)_S$   | $\text{Pa}^{-1}$                   |              |
| linear expansion coefficient           | $\alpha_l$                 | $\alpha_l = (1/l)(\partial l / \partial T)$      | $\text{K}^{-1}$                    |              |
| cubic expansion coefficient            | $\alpha, \alpha_V, \gamma$ | $\alpha = (1/V)(\partial V / \partial T)_p$      | $\text{K}^{-1}$                    | 7            |
| heat capacity, at constant pressure    | $C_p$                      | $C_p = (\partial H / \partial T)_p$              | $\text{J K}^{-1}$                  |              |
| at constant volume                     | $C_V$                      | $C_V = (\partial U / \partial T)_V$              | $\text{J K}^{-1}$                  |              |

(1) In the differential form,  $d$  denotes an inexact differential. The given equation in integrated form is  $\Delta U = Q + W$ .  $Q > 0$  and  $W > 0$  indicate an increase in the energy of the system.

(2) This temperature is defined by the “International Temperature Scale of 1990 (ITS-90)” for which specific definitions are prescribed [82–84]. However, the CIPM, in its 94<sup>th</sup> Meeting of October 2005, has approved a Recommendation T3 (2005) of the Comité Consultatif de Thermométrie, where the ITS-90 becomes one of several “mises en pratique” of the kelvin definition. The Technical Annex (2005) is available as Doc. CCT\_05\_33 [85]. It concerns the definition of a reference isotopic composition of hydrogen and water, when used for the realization of fixed points of the “mises en pratique”.

(3) This quantity is sometimes misnamed “centigrade temperature”.

(4) The symbol  $F$  is sometimes used as an alternate symbol for the Helmholtz energy.

(5) The definition applies to pure substance. However, the concept of molar and specific quantities (see Section 1.4, p. 6) may also be applied to mixtures.  $n$  is the amount of substance (see Section

| <i>Name</i>                                            | <i>Symbol</i>            | <i>Definition</i>                                         | <i>SI unit</i>                      | <i>Notes</i> |
|--------------------------------------------------------|--------------------------|-----------------------------------------------------------|-------------------------------------|--------------|
| ratio of heat capacities                               | $\gamma, (\kappa)$       | $\gamma = C_p/C_V$                                        | 1                                   |              |
| Joule-Thomson coefficient                              | $\mu, \mu_{JT}$          | $\mu = (\partial T/\partial p)_H$                         | K Pa <sup>-1</sup>                  |              |
| thermal power                                          | $\Phi, P$                | $\Phi = dQ/dt$                                            | W                                   |              |
| virial coefficient,                                    |                          |                                                           |                                     |              |
| second                                                 | $B$                      | $pV_m = RT(1 + B/V_m +$                                   | m <sup>3</sup> mol <sup>-1</sup>    | 8            |
| third                                                  | $C$                      | $C/V_m^2 + \dots)$                                        | m <sup>6</sup> mol <sup>-2</sup>    | 8            |
| van der Waals                                          | $a$                      | $(p + a/V_m^2)(V_m - b) = RT$                             | J m <sup>3</sup> mol <sup>-2</sup>  | 9            |
| coefficients                                           | $b$                      |                                                           | m <sup>3</sup> mol <sup>-1</sup>    | 9            |
| compression factor,<br>(compressibility factor)        | $Z$                      | $Z = pV_m/RT$                                             | 1                                   |              |
| partial molar<br>quantity $X$                          | $X_B, (\bar{X}_B)$       | $X_B = (\partial X/\partial n_B)_{T,p,n_{j \neq B}}$      | [ $X$ ]/mol                         | 10           |
| chemical potential,<br>(partial molar Gibbs<br>energy) | $\mu$                    | $\mu_B = (\partial G/\partial n_B)_{T,p,n_{j \neq B}}$    | J mol <sup>-1</sup>                 | 11           |
| standard chemical<br>potential                         | $\mu^\ominus, \mu^\circ$ |                                                           | J mol <sup>-1</sup>                 | 12           |
| absolute activity                                      | $\lambda$                | $\lambda_B = \exp(\mu_B/RT)$                              | 1                                   | 11           |
| (relative) activity                                    | $a$                      | $a_B = \exp\left(\frac{\mu_B - \mu_B^\ominus}{RT}\right)$ | 1                                   | 11, 13       |
| standard partial<br>molar enthalpy                     | $H_B^\ominus$            | $H_B^\ominus = \mu_B^\ominus + TS_B^\ominus$              | J mol <sup>-1</sup>                 | 11, 12       |
| standard partial<br>molar entropy                      | $S_B^\ominus$            | $S_B^\ominus = -(\partial \mu_B^\ominus/\partial T)_p$    | J mol <sup>-1</sup> K <sup>-1</sup> | 11, 12       |

(Notes continued)

(5) (continued) 2.10, notes 1 and 2, p. 47).

(6)  $X$  is an extensive quantity, whose SI unit is [ $X$ ]. In the case of molar quantities the entities should be specified.

*Example*  $V_{m,B} = V_m(B) = V/n_B$  denotes the molar volume of B

(7) This quantity is also called the coefficient of thermal expansion, or the expansivity coefficient.

(8) Another set of “pressure virial coefficients” may be defined by

$$pV_m = RT(1 + B_p p + C_p p^2 + \dots)$$

(9) For a gas satisfying the van der Waals equation of state, given in the definition, the second virial coefficient is related to the parameters  $a$  and  $b$  in the van der Waals equation by

$$B = b - a/RT$$

(10) The symbol applies to entities B which should be specified. The bar may be used to distinguish partial molar  $X$  from  $X$  when necessary.

*Example* The partial molar volume of Na<sub>2</sub>SO<sub>4</sub> in aqueous solution may be denoted  $\bar{V}(\text{Na}_2\text{SO}_4, \text{aq})$ , in order to distinguish it from the volume of the solution  $V(\text{Na}_2\text{SO}_4, \text{aq})$ .

(11) The definition applies to entities B which should be specified. The chemical potential can be defined equivalently by the corresponding partial derivatives of other thermodynamic functions ( $U$ ,  $H$ ,  $A$ ).

(12) The symbol  $^\ominus$  or  $^\circ$  is used to indicate standard. They are equally acceptable. Definitions of standard states are discussed in Section 2.11.1 (iv), p. 61. Whenever a standard chemical potential

| <i>Name</i>                               | <i>Symbol</i>        | <i>Definition</i>                                                                             | <i>SI unit</i>                                             | <i>Notes</i>     |
|-------------------------------------------|----------------------|-----------------------------------------------------------------------------------------------|------------------------------------------------------------|------------------|
| standard reaction Gibbs energy (function) | $\Delta_r G^\ominus$ | $\Delta_r G^\ominus = \sum_B \nu_B \mu_B^\ominus$                                             | J mol <sup>-1</sup>                                        | 12, 14<br>15, 16 |
| affinity of reaction                      | $A, \mathcal{A}$     | $A = -(\partial G / \partial \xi)_{p,T}$<br>$= -\sum_B \nu_B \mu_B$                           | J mol <sup>-1</sup>                                        | 15               |
| standard reaction enthalpy                | $\Delta_r H^\ominus$ | $\Delta_r H^\ominus = \sum_B \nu_B H_B^\ominus$                                               | J mol <sup>-1</sup>                                        | 12, 14<br>15, 16 |
| standard reaction entropy                 | $\Delta_r S^\ominus$ | $\Delta_r S^\ominus = \sum_B \nu_B S_B^\ominus$                                               | J mol <sup>-1</sup> K <sup>-1</sup>                        | 12, 14, 15       |
| reaction quotient                         | $Q$                  | $Q = \prod_B a_B^{\nu_B}$                                                                     | 1                                                          | 17               |
| equilibrium constant                      | $K^\ominus, K$       | $K^\ominus = \exp(-\Delta_r G^\ominus / RT)$                                                  | 1                                                          | 12, 15, 18       |
| equilibrium constant, pressure basis      | $K_p$                | $K_p = \prod_B p_B^{\nu_B}$                                                                   | Pa <sup><math>\sum \nu_B</math></sup>                      | 15, 19           |
| concentration basis                       | $K_c$                | $K_c = \prod_B c_B^{\nu_B}$                                                                   | (mol m <sup>-3</sup> ) <sup><math>\sum \nu_B</math></sup>  | 15, 19           |
| molality basis                            | $K_m$                | $K_m = \prod_B m_B^{\nu_B}$                                                                   | (mol kg <sup>-1</sup> ) <sup><math>\sum \nu_B</math></sup> | 15, 19           |
| fugacity                                  | $f, \tilde{p}$       | $f_B = \lambda_B \lim_{p \rightarrow 0} (p_B / \lambda_B)_T$                                  | Pa                                                         | 11               |
| fugacity coefficient                      | $\phi$               | $\phi_B = f_B / p_B$                                                                          | 1                                                          |                  |
| Henry's law constant                      | $k_H$                | $k_{H,B} = \lim_{x_B \rightarrow 0} (f_B / x_B)$<br>$= (\partial f_B / \partial x_B)_{x_B=0}$ | Pa                                                         | 11, 20           |

(Notes continued)

(12) (continued)  $\mu^\ominus$  or a standard equilibrium constant  $K^\ominus$  or other standard quantity is used, the standard state must be specified.

(13) In the defining equation given here, the pressure dependence of the activity has been neglected as is often done for condensed phases at atmospheric pressure.

An equivalent definition is  $a_B = \lambda_B / \lambda_B^\ominus$ , where  $\lambda_B^\ominus = \exp(\mu_B^\ominus / RT)$ . The definition of  $\mu^\ominus$  depends on the choice of the standard state (see Section 2.11.1 (iv), p. 61).

(14) The symbol r indicates reaction in general. In particular cases r can be replaced by another appropriate subscript, e.g.  $\Delta_f H^\ominus$  denotes the standard molar enthalpy of formation; see Section 2.11.1 (i), p. 59 below for a list of subscripts.  $\Delta_r$  can be interpreted as operator symbol  $\Delta_r \stackrel{\text{def}}{=} \partial / \partial \xi$ .

(15) The reaction must be specified for which this quantity applies.

(16) Reaction enthalpies (and reaction energies in general) are usually quoted in kJ mol<sup>-1</sup>. In the older literature kcal mol<sup>-1</sup> is also common, however, various calories exist. For the thermochemical calorie, 1 kcal = 4.184 kJ (see Section 7.2, p. 137).

(17) This quantity applies in general to a system which is not in equilibrium.

(18) This quantity is equal to  $Q$  in equilibrium, when the affinity is zero. It is dimensionless and its value depends on the choice of standard state, which must be specified. ISO [5.h] and the IUPAC Thermodynamics Commission [81] recommend the symbol  $K^\ominus$  and the name “standard equilibrium constant”. Many chemists prefer the symbol  $K$  and the name “thermodynamic equilibrium constant”.

(19) These quantities are not in general dimensionless. One can define in an analogous way an equilibrium constant in terms of fugacity  $K_f$ , etc. At low pressures  $K_p$  is approximately related to  $K^\ominus$  by the equation  $K^\ominus \approx K_p / (p^\ominus)^{\sum \nu_B}$ , and similarly in dilute solutions  $K_c$  is approximately related to  $K^\ominus$  by  $K^\ominus \approx K_c / (c^\ominus)^{\sum \nu_B}$ ; however, the exact relations involve fugacity coefficients or activity coefficients [81].

| <i>Name</i>                | <i>Symbol</i> | <i>Definition</i>                                    | <i>SI unit</i>       | <i>Notes</i> |
|----------------------------|---------------|------------------------------------------------------|----------------------|--------------|
| activity coefficient       |               |                                                      |                      |              |
| referenced to Raoult's law | $f$           | $f_B = a_B/x_B$                                      | 1                    | 11, 21       |
| referenced to Henry's law  |               |                                                      |                      |              |
| molality basis             | $\gamma_m$    | $a_{m,B} = \gamma_{m,B} m_B / m^\ominus$             | 1                    | 11, 22       |
| concentration basis        | $\gamma_c$    | $a_{c,B} = \gamma_{c,B} c_B / c^\ominus$             | 1                    | 11, 22       |
| mole fraction basis        | $\gamma_x$    | $a_{x,B} = \gamma_{x,B} x_B$                         | 1                    | 10, 21       |
| ionic strength,            |               |                                                      |                      |              |
| molality basis             | $I_m, I$      | $I_m = (1/2) \sum_i m_i z_i^2$                       | mol kg <sup>-1</sup> |              |
| concentration basis        | $I_c, I$      | $I_c = (1/2) \sum_i c_i z_i^2$                       | mol m <sup>-3</sup>  |              |
| osmotic coefficient,       |               |                                                      |                      |              |
| molality basis             | $\phi_m$      | $\phi_m = \frac{\mu_A^* - \mu_A}{RT M_A \sum_B m_B}$ | 1                    | 23, 24       |
| mole fraction basis        | $\phi_x$      | $\phi_x = -\frac{\mu_B^* - \mu_B}{RT \ln x_B}$       | 1                    | 22, 23       |
| osmotic pressure           | $\Pi$         | $\Pi = -(RT/V_A) \ln a_A$                            | Pa                   | 23, 25       |

(Notes continued)

(19) (continued) The equilibrium constant of dissolution of an electrolyte (describing the equilibrium between excess solid phase and solvated ions) is often called a solubility product, denoted  $K_{\text{sol}}$  or  $K_s$  (or  $K_{\text{sol}}^\ominus$  or  $K_s^\ominus$  as appropriate). In a similar way the equilibrium constant for an acid dissociation is often written  $K_a$ , for base hydrolysis  $K_b$ , and for water dissociation  $K_w$ .

(20) Henry's law is sometimes expressed in terms of molalities or concentration and then the corresponding units of Henry's law constant are Pa kg mol<sup>-1</sup> or Pa m<sup>3</sup> mol<sup>-1</sup>, respectively.

(21) This quantity applies to pure phases, substances in mixtures, or solvents.

(22) This quantity applies to solutes.

(23) A is the solvent, B is one or more solutes.

(24) The entities B are independent solute molecules, ions, etc. regardless of their nature. Their amount is sometimes expressed in osmoles (meaning a mole of osmotically active entities), but this use is discouraged.

(25) This definition of osmotic pressure applies to an incompressible fluid.

### 2.11.1 Other symbols and conventions in chemical thermodynamics

A more extensive description of this subject can be found in [81].

#### (i) Symbols used as subscripts to denote a physical chemical process or reaction

These symbols should be printed in Roman (upright) type, without a full stop (period).

|                          |     |
|--------------------------|-----|
| adsorption               | ads |
| atomization              | at  |
| combustion reaction      | c   |
| dilution (of a solution) | dil |
| displacement             | dpl |

|                                                      |     |
|------------------------------------------------------|-----|
| formation reaction                                   | f   |
| immersion                                            | imm |
| melting, fusion (solid $\rightarrow$ liquid)         | fus |
| mixing of fluids                                     | mix |
| reaction in general                                  | r   |
| solution (of solute in solvent)                      | sol |
| sublimation (solid $\rightarrow$ gas)                | sub |
| transition (between two phases)                      | trs |
| triple point                                         | tp  |
| vaporization, evaporation (liquid $\rightarrow$ gas) | vap |

### (ii) Recommended superscripts

|                                     |                     |
|-------------------------------------|---------------------|
| activated complex, transition state | ‡, ≠                |
| apparent                            | app                 |
| excess quantity                     | E                   |
| ideal                               | id                  |
| infinite dilution                   | $\infty$            |
| pure substance                      | *                   |
| standard                            | $\ominus$ , $\circ$ |

### (iii) Examples of the use of the symbol $\Delta$

The symbol  $\Delta$  denotes a change in an extensive thermodynamic quantity for a process. The addition of a subscript to the  $\Delta$  denotes a change in the property.

*Examples*     $\Delta_{\text{vap}}H = \Delta_{\text{l}}^{\text{g}}H = H(\text{g}) - H(\text{l})$  for the molar enthalpy of vaporization.  
 $\Delta_{\text{vap}}H = 40.7 \text{ kJ mol}^{-1}$  for water at 100 °C under its own vapor pressure.  
This can also be written  $\Delta H_{\text{vap}}$ , but this usage is not recommended.

The subscript r is used to denote changes associated with a *chemical reaction*. Symbols such as  $\Delta_{\text{r}}H$  are defined by the equation

$$\Delta_{\text{r}}H = \sum_{\text{B}} \nu_{\text{B}}H_{\text{B}} = (\partial H / \partial \xi)_{T,p}$$

It is thus essential to specify the stoichiometric reaction equation when giving numerical values for such quantities in order to define the extent of reaction  $\xi$  and the value of the stoichiometric numbers  $\nu_{\text{B}}$ .

*Example*     $\text{N}_2(\text{g}) + 3 \text{H}_2(\text{g}) = 2 \text{NH}_3(\text{g}), \quad \Delta_{\text{r}}H^{\ominus}(298.15 \text{ K}) = -92.4 \text{ kJ mol}^{-1}$   
 $\Delta_{\text{r}}S^{\ominus}(298.15 \text{ K}) = -199 \text{ J mol}^{-1} \text{ K}^{-1}$

The  $\text{mol}^{-1}$  in the units identifies the quantities in this example as the change per extent of reaction. They may be called the molar enthalpy and entropy of reaction, and a subscript m may be added to the symbol to emphasize the difference from the integral quantities if desired.

The *standard reaction quantities* are particularly important. They are defined by the equations

$$\begin{aligned} \Delta_{\text{r}}H^{\ominus} &= \sum_{\text{B}} \nu_{\text{B}}H_{\text{B}}^{\ominus} \\ \Delta_{\text{r}}S^{\ominus} &= \sum_{\text{B}} \nu_{\text{B}}S_{\text{B}}^{\ominus} \\ \Delta_{\text{r}}G^{\ominus} &= \sum_{\text{B}} \nu_{\text{B}}\mu_{\text{B}}^{\ominus} \end{aligned}$$

It is important to specify notation with care for these symbols. The relation to the affinity of the reaction is

$$-A = \Delta_r G = \Delta_r G^\circ + RT \ln \left( \prod_B a_B^{\nu_B} \right)$$

and the relation to the standard equilibrium constant is  $\Delta_r G^\circ = -RT \ln K^\circ$ . The product of the activities is the reaction quotient  $Q$ , see p. 58.

The term *combustion* and symbol c denote the complete oxidation of a substance. For the definition of complete oxidation of substances containing elements other than C, H and O see [86]. The corresponding reaction equation is written so that the stoichiometric number  $\nu$  of the substance is  $-1$ .

*Example* The standard enthalpy of combustion of gaseous methane is  
 $\Delta_c H^\circ(\text{CH}_4, \text{g}, 298.15 \text{ K}) = -890.3 \text{ kJ mol}^{-1}$ , implying the reaction  
 $\text{CH}_4(\text{g}) + 2 \text{O}_2(\text{g}) \rightarrow \text{CO}_2(\text{g}) + 2 \text{H}_2\text{O}(\text{l})$ .

The term *formation* and symbol f denote the formation of the substance from elements in their reference state (usually the most stable state of each element at the chosen temperature and standard pressure). The corresponding reaction equation is written so that the stoichiometric number  $\nu$  of the substance is  $+1$ .

*Example* The standard entropy of formation of crystalline mercury(II) chloride is  
 $\Delta_f S^\circ(\text{HgCl}_2, \text{cr}, 298.15 \text{ K}) = -154.3 \text{ J mol}^{-1} \text{K}^{-1}$ , implying the reaction  
 $\text{Hg}(\text{l}) + \text{Cl}_2(\text{g}) \rightarrow \text{HgCl}_2(\text{cr})$ .

The term *atomization*, symbol at, denotes a process in which a substance is separated into its constituent atoms in the ground state in the gas phase. The corresponding reaction equation is written so that the stoichiometric number  $\nu$  of the substance is  $-1$ .

*Example* The standard (internal) energy of atomization of liquid water is  
 $\Delta_{\text{at}} U^\circ(\text{H}_2\text{O}, \text{l}) = 625 \text{ kJ mol}^{-1}$ , implying the reaction  
 $\text{H}_2\text{O}(\text{l}) \rightarrow 2 \text{H}(\text{g}) + \text{O}(\text{g})$ .

#### (iv) Standard states [1.j] and [81]

The standard chemical potential of substance B at temperature  $T$ ,  $\mu_B^\circ(T)$ , is the value of the chemical potential under standard conditions, specified as follows. Three differently defined standard states are recognized.

*For a gas phase.* The standard state for a gaseous substance, whether pure or in a gaseous mixture, is the (hypothetical) state of the pure substance B in the gaseous phase at the standard pressure  $p = p^\circ$  and exhibiting ideal gas behavior. The standard chemical potential is defined as

$$\mu_B^\circ(T) = \lim_{p \rightarrow 0} [\mu_B(T, p, y_B, \dots) - RT \ln(y_B p / p^\circ)]$$

*For a pure phase, or a mixture, or a solvent, in the liquid or solid state.* The standard state for a liquid or solid substance, whether pure or in a mixture, or for a solvent, is the state of the pure substance B in the liquid or solid phase at the standard pressure  $p = p^\circ$ . The standard chemical potential is defined as

$$\mu_B(T) = \mu_B^*(T, p^\circ)$$

*For a solute in solution.* For a solute in a liquid or solid solution the standard state is referenced to the ideal dilute behavior of the solute. It is the (hypothetical) state of solute B at the standard molality  $m^\circ$ , standard pressure  $p^\circ$ , and behaving like the infinitely dilute solution. The standard chemical potential is defined as

$$\mu_{\text{B}}^{\ominus}(T) = [\mu_{\text{B}}(T, p^{\ominus}, m_{\text{B}}, \dots) - RT \ln(m_{\text{B}}/m^{\ominus})]^{\infty}$$

The chemical potential of the solute B as a function of the molality  $m_{\text{B}}$  at constant pressure  $p = p^{\ominus}$  is then given by the expression

$$\mu_{\text{B}}(m_{\text{B}}) = \mu_{\text{B}}^{\ominus} + RT \ln(m_{\text{B}}\gamma_{m,\text{B}}/m^{\ominus})$$

Sometimes (amount) concentration  $c$  is used as a variable in place of molality  $m$ ; both of the above equations then have  $c$  in place of  $m$  throughout. Occasionally mole fraction  $x$  is used in place of  $m$ ; both of the above equations then have  $x$  in place of  $m$  throughout, and  $x^{\ominus} = 1$ . Although the standard state of a solute is always referenced to ideal dilute behavior, the definition of the standard state and the value of the standard chemical potential  $\mu^{\ominus}$  are different depending on whether molality  $m$ , concentration  $c$ , or mole fraction  $x$  is used as a variable.

### (v) Standard pressures, molality, and concentration

In principle one may choose any values for the standard pressure  $p^{\ominus}$ , the standard molality  $m^{\ominus}$ , and the standard concentration  $c^{\ominus}$ , although the choice must be specified. For example, in tabulating data appropriate to high pressure chemistry it may be convenient to choose a value of  $p^{\ominus} = 100 \text{ MPa}$  ( $= 1 \text{ kbar}$ ).

In practice, however, the most common choice is

$$\begin{aligned} p^{\ominus} &= 0.1 \text{ MPa} = 100 \text{ kPa} \quad (= 1 \text{ bar}) \\ m^{\ominus} &= 1 \text{ mol kg}^{-1} \\ c^{\ominus} &= 1 \text{ mol dm}^{-3} \end{aligned}$$

These values for  $m^{\ominus}$  and  $c^{\ominus}$  are universally accepted. The value for  $p^{\ominus} = 100 \text{ kPa}$ , is the IUPAC recommendation since 1982 [1.j], and is recommended for tabulating thermodynamic data. Prior to 1982 the standard pressure was usually taken to be  $p^{\ominus} = 101\,325 \text{ Pa}$  ( $= 1 \text{ atm}$ , called the *standard atmosphere*). In any case, the value for  $p^{\ominus}$  should be specified.

The conversion of values corresponding to different  $p^{\ominus}$  is described in [87–89]. The newer value of  $p^{\ominus}$ , 100 kPa is sometimes called the *standard state pressure*.

### (vi) Biochemical standard states

Special standard states that are close to physiological conditions are often chosen. The biochemical standard state is often chosen at  $[\text{H}^+] = 10^{-7} \text{ mol dm}^{-3}$ . The concentrations of the solutes may be grouped together as for example, the total phosphate concentration rather than the concentration of each component, ( $\text{H}_3\text{PO}_4$ ,  $\text{H}_2\text{PO}_4^-$ ,  $\text{HPO}_4^{2-}$ ,  $\text{PO}_4^{3-}$ ), separately. Standard and other reference states must be specified with care [90,91].

### (vii) Thermodynamic properties

Values of many thermodynamic quantities represent basic chemical properties of substances and serve for further calculations. Extensive tabulations exist, e.g. [92–96]. Special care has to be taken in reporting the data and their uncertainties [97,98].

### (viii) Reference state (of an element)

The state in which the element is stable at the chosen standard state pressure and for a given temperature [16].

## 2.12 CHEMICAL KINETICS AND PHOTOCHEMISTRY

The recommendations given here are based on previous IUPAC recommendations [1.c,1.k] and [17], which are not in complete agreement. Recommendations regarding photochemistry are given in [60] and for recommendations on reporting of chemical kinetics data see also [99]. A glossary of terms used in chemical kinetics has been given in [100].

| <i>Name</i>                                                             | <i>Symbol</i> | <i>Definition</i>                           | <i>SI unit</i>                                     | <i>Notes</i> |
|-------------------------------------------------------------------------|---------------|---------------------------------------------|----------------------------------------------------|--------------|
| rate of change of quantity $X$                                          | $\dot{X}$     | $\dot{X} = dX/dt$                           | (varies)                                           | 1            |
| rate of conversion                                                      | $\dot{\xi}$   | $\dot{\xi} = d\xi/dt$                       | $\text{mol s}^{-1}$                                | 2            |
| rate of concentration change<br>(due to chemical reaction)              | $r_B, v_B$    | $r_B = dc_B/dt$                             | $\text{mol m}^{-3} \text{s}^{-1}$                  | 3, 4         |
| rate of reaction (based on<br>amount concentration)                     | $v, v_c$      | $v = \nu_B^{-1} dc_B/dt$<br>$= \dot{\xi}/V$ | $\text{mol m}^{-3} \text{s}^{-1}$                  | 2, 4         |
| rate of reaction (based on<br>number concentration),<br>(reaction rate) | $v, v_C$      | $v_C = \nu_B^{-1} dC_B/dt$                  | $\text{m}^{-3} \text{s}^{-1}$                      |              |
| partial order of reaction                                               | $m_B, n_B$    | $v = k \prod_B c_B^{m_B}$                   | 1                                                  | 5            |
| overall order of reaction                                               | $m, n$        | $m = \sum_B m_B$                            | 1                                                  |              |
| rate constant,<br>rate coefficient                                      | $k, k(T)$     | $v = k \prod_B c_B^{m_B}$                   | $(\text{m}^3 \text{mol}^{-1})^{m-1} \text{s}^{-1}$ | 6            |

(1) E.g. rate of pressure change  $\dot{p} = dp/dt$ , for which the SI unit is  $\text{Pa s}^{-1}$ , rate of entropy change  $dS/dt$  with SI unit  $\text{J K}^{-1} \text{s}^{-1}$ .

(2) The reaction must be specified, for which this quantity applies, by giving the stoichiometric equation.

(3) The symbol and the definition apply to entities B.

(4) Note that  $r_B$  and  $v$  can also be defined on the basis of partial pressure, number concentration, surface concentration, etc., with analogous definitions. If necessary differently defined rates of reaction can be distinguished by a subscript, e.g.  $v_p = \nu_B^{-1} dp_B/dt$ , etc. Note that the rate of reaction can only be defined for a reaction of known and time-independent stoichiometry, in terms of a specified reaction equation; also the second equation for the rate of reaction follows from the first only if the volume  $V$  is constant and  $c_B$  is uniform throughout (more generally, the definition applies to local concentrations). The derivatives must be those due to the chemical reaction considered; in open systems, such as flow systems, effects due to input and output processes must be taken into account separately, as well as transport processes in general by the equation

$$(dc_B/dt)_{\text{total}} = (dc_B/dt)_{\text{reaction}} + (dc_B/dt)_{\text{transport}}$$

(5) The symbol applies to reactant B. The symbol  $m$  is used to avoid confusion with  $n$  for amount of substance. The order of reaction is only defined if the particular rate law applies.  $m$  is a real number.

(6) Rate constants  $k$  and pre-exponential factors  $A$  are usually quoted in either  $(\text{dm}^3 \text{mol}^{-1})^{m-1} \text{s}^{-1}$  or on a molecular scale in  $(\text{cm}^3)^{m-1} \text{s}^{-1}$  or  $(\text{cm}^3 \text{molecule}^{-1})^{m-1} \text{s}^{-1}$ . Note that “molecule” is not a unit, but is often included for clarity, although this does not conform to accepted usage.  $k(T)$  is written to stress temperature dependence. Rate constants are frequently quoted as decadic logarithms.

|                |                       |                                                             |    |                                                            |
|----------------|-----------------------|-------------------------------------------------------------|----|------------------------------------------------------------|
| <i>Example</i> | second order reaction | $k = 10^{8.2} \text{ cm}^3 \text{ mol}^{-1} \text{ s}^{-1}$ | or | $\lg(k/\text{cm}^3 \text{ mol}^{-1} \text{ s}^{-1}) = 8.2$ |
|                | or alternatively      | $k = 10^{-12.6} \text{ cm}^3 \text{ s}^{-1}$                | or | $\lg(k/\text{cm}^3 \text{ s}^{-1}) = -12.6$                |

| <i>Name</i>                              | <i>Symbol</i>                                     | <i>Definition</i>                                                  | <i>SI unit</i>                                     | <i>Notes</i> |
|------------------------------------------|---------------------------------------------------|--------------------------------------------------------------------|----------------------------------------------------|--------------|
| rate constant of unimolecular reaction   | $k_{\text{uni}}, k_{\text{uni}}(T, c_{\text{M}})$ | $v = k_{\text{uni}}c_{\text{B}}$                                   | $\text{s}^{-1}$                                    | 7            |
| at high pressure                         | $k_{\infty}$                                      | $k_{\text{uni}}(c_{\text{M}} \rightarrow \infty)$                  | $\text{s}^{-1}$                                    | 7            |
| at low pressure                          | $k_0$                                             | $v = k_0 c_{\text{M}} c_{\text{B}}$                                | $\text{m}^3 \text{mol}^{-1} \text{s}^{-1}$         | 7            |
| Boltzmann constant                       | $k, k_{\text{B}}$                                 |                                                                    | $\text{J K}^{-1}$                                  |              |
| half life                                | $t_{1/2}$                                         | $c(t_{1/2}) = c(0)/2$                                              | $\text{s}$                                         |              |
| relaxation time, lifetime, mean life     | $\tau$                                            | $\Delta c(\tau) = \Delta c(0)/e$                                   | $\text{s}$                                         | 8            |
| (Arrhenius) activation energy            | $E_{\text{A}}, E_{\text{a}}$                      | $E_{\text{A}} = RT^2 d(\ln k)/dT$                                  | $\text{J mol}^{-1}$                                | 9            |
| pre-exponential factor, frequency factor | $A$                                               | $A = k \exp(E_{\text{A}}/RT)$                                      | $(\text{m}^3 \text{mol}^{-1})^{m-1} \text{s}^{-1}$ | 9, 10        |
| hard sphere radius                       | $r$                                               |                                                                    | $\text{m}$                                         |              |
| collision diameter                       | $d_{\text{AB}}$                                   | $d_{\text{AB}} = r_{\text{A}} + r_{\text{B}}$                      | $\text{m}$                                         |              |
| collision cross section                  | $\sigma$                                          | $\sigma = \pi d_{\text{AB}}^2$                                     | $\text{m}^2$                                       | 11           |
| mean relative speed between A and B      | $\bar{c}_{\text{AB}}$                             | $\bar{c}_{\text{AB}} = (8kT/\pi\mu)^{1/2}$                         | $\text{m s}^{-1}$                                  | 12           |
| collision frequency of A with A          | $z_{\text{A}}(\text{A})$                          | $z_{\text{A}}(\text{A}) = \sqrt{2} C_{\text{A}} \sigma \bar{c}$    | $\text{s}^{-1}$                                    | 11           |
| of A with B                              | $z_{\text{A}}(\text{B})$                          | $z_{\text{A}}(\text{B}) = C_{\text{B}} \sigma \bar{c}_{\text{AB}}$ | $\text{s}^{-1}$                                    | 11           |

(7) The rates of unimolecular reactions show a dependence upon the concentration  $c_{\text{M}}$  of a collision partner M. One writes  $k_{\text{uni}}(T, c_{\text{M}})$  to emphasize the temperature and “pressure” dependence. At high  $c_{\text{M}} (\rightarrow \infty)$  the dependence vanishes. At low  $c_{\text{M}} (\rightarrow 0)$  it gives a partial order 1 of reaction with respect to  $c_{\text{M}}$ . In this case one defines the second order rate constant  $k_0$ .

(8)  $\tau$  is defined as the time interval in which a concentration perturbation  $\Delta c$  falls to  $1/e$  of its initial value  $\Delta c(0)$ . If some initial concentration of a substance decays to zero as  $t \rightarrow \infty$ , as in radioactive decay, then the relaxation time is the average lifetime of that substance (radioisotope). This lifetime or decay time must be distinguished from the half life.

(9) One may use as defining equation

$$E_{\text{A}} = -Rd(\ln k)/d(1/T)$$

The term Arrhenius activation energy is to be used only for the empirical quantity defined in the table. Other empirical equations with different “activation energies”, such as

$$k(T) = A' T^n \exp(-E_{\text{a}}'/RT)$$

are also being used. In such expressions  $A'$ ,  $n$ , and  $E_{\text{a}}'$  are taken to be temperature independent parameters.

The term activation energy is also used for an energy threshold appearing in the electronic potential (the height of the electronic energy barrier). For this “activation energy” the symbol  $E_0$  and the term threshold energy is preferred, but  $E_{\text{a}}$  is also commonly used. Furthermore,  $E_0$  may or may not include a correction for zero point energies of reactants and the transition state.

It is thus recommended to specify in any given context exactly which activation energy is meant and to reserve (Arrhenius) activation energy only and exactly for the quantity defined in the table.  $E_{\text{A}}$  depends on temperature and may be written  $E_{\text{A}}(T)$ .

(10)  $A$  is dependent on temperature and may be written  $A(T)$ .

(11) The collision cross section  $\sigma$  is a constant in the hard sphere collision model, but generally it is energy dependent. One may furthermore define a temperature dependent average collision cross section (see note 16).  $C$  denotes the number concentration.

(12)  $\mu$  is the reduced mass,  $\mu = m_{\text{A}}m_{\text{B}}/(m_{\text{A}} + m_{\text{B}})$ .

| <i>Name</i>                                           | <i>Symbol</i>                                  | <i>Definition</i>                                                                                                                 | <i>SI unit</i>                             | <i>Notes</i> |
|-------------------------------------------------------|------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------|--------------|
| collision density,<br>collision number<br>of A with A | $Z_{AA}$                                       | $Z_{AA} = C_A z_A(A)$                                                                                                             | $\text{s}^{-1} \text{m}^{-3}$              | 13           |
| of A with B                                           | $Z_{AB}$                                       | $Z_{AB} = C_A z_A(B)$                                                                                                             | $\text{s}^{-1} \text{m}^{-3}$              | 13           |
| collision frequency<br>factor                         | $z_{AB}$                                       | $z_{AB} = Z_{AB}/L c_A c_B$                                                                                                       | $\text{m}^3 \text{mol}^{-1} \text{s}^{-1}$ | 13           |
| mean free path                                        | $\lambda$                                      | $\lambda = \bar{c}/z_A$                                                                                                           | m                                          |              |
| impact parameter                                      | $b$                                            |                                                                                                                                   | m                                          | 14           |
| scattering angle                                      | $\theta$                                       |                                                                                                                                   | 1, rad                                     | 15           |
| differential cross<br>section                         | $I_{ji}$                                       | $I_{ji} = d\sigma_{ji}/d\Omega$                                                                                                   | $\text{m}^2 \text{sr}^{-1}$                | 16           |
| total cross section                                   | $\sigma_{ji}$                                  | $\sigma_{ji} = \int I_{ji} d\Omega$                                                                                               | $\text{m}^2$                               | 16           |
| scattering matrix                                     | $\mathbf{S}$                                   |                                                                                                                                   | 1                                          | 17           |
| transition probability                                | $P_{ji}$                                       | $P_{ji} =  S_{ji} ^2$                                                                                                             | 1                                          | 16, 17       |
| standard enthalpy<br>of activation                    | $\Delta^\ddagger H^\ominus, \Delta H^\ddagger$ | $k(T) = \frac{kT}{h} \exp\left(\frac{\Delta^\ddagger S^\ominus}{R}\right) \exp\left(\frac{-\Delta^\ddagger H^\ominus}{RT}\right)$ | $\text{J mol}^{-1}$                        | 18           |
| volume of activation                                  | $\Delta^\ddagger V^\ominus, \Delta V^\ddagger$ | $\Delta^\ddagger V^\ominus = -RT(\partial(\ln k)/\partial p)_T$                                                                   | $\text{m}^3 \text{mol}^{-1}$               |              |
| standard internal<br>energy of activation             | $\Delta^\ddagger U^\ominus, \Delta U^\ddagger$ |                                                                                                                                   | $\text{J mol}^{-1}$                        | 18           |
| standard entropy<br>of activation                     | $\Delta^\ddagger S^\ominus, \Delta S^\ddagger$ |                                                                                                                                   | $\text{J mol}^{-1} \text{K}^{-1}$          | 18           |

(13)  $Z_{AA}$  and  $Z_{AB}$  are the total number of AA or AB collisions per time and volume in a system containing only A molecules, or containing molecules of two types, A and B. Three-body collisions can be treated in a similar way.

(14) The impact parameter  $b$  characterizes an individual collision between two particles; it is defined as the distance of closest approach that would result if the particle trajectories were undeflected by the collision.

(15)  $\theta = 0$  implies no deflection.

(16) In all these matrix quantities the first index refers to the final and the second to the initial channel.  $i$  and  $j$  denote reactant and product channels, respectively, and  $\Omega$  denotes solid angle;  $d\sigma_{ji}/d\Omega$  is equal to (scattered particle current per solid angle) divided by (incident particle current per area). Elastic scattering implies  $i = j$ . Both  $I_{ji}$  and  $\sigma_{ji}$  depend on the total energy of relative motion, and may be written  $I_{ji}(E)$  and  $\sigma_{ji}(E)$ . General collision theory leads to an expression of a rate coefficient in terms of the energy dependent total reaction cross section

$$k_{ji}(T) = \left(\frac{8k_B T}{\pi\mu}\right)^{1/2} \int_0^\infty \left(\frac{E}{k_B T}\right) \sigma_{ji}(E) \exp(-E/k_B T) \left(\frac{dE}{k_B T}\right)$$

where  $E$  is the translational collision energy and  $\mu$  the reduced mass (see note 12). The integral can be interpreted as the thermally averaged collision cross section  $\langle\sigma_{ji}\rangle$ , which is to be used in the calculation of the collision frequency under thermal conditions (see note 11).

(17) The scattering matrix  $\mathbf{S}$  is used in quantum scattering theory [101].  $\mathbf{S}$  is a unitary matrix  $\mathbf{S}\mathbf{S}^\dagger = \mathbf{1}$ .  $P_{ji} = |S_{ji}|^2$  is the probability that collision partners incident in channel  $i$  will emerge in channel  $j$ .

(18) The quantities  $\Delta^\ddagger H^\ominus$ ,  $\Delta^\ddagger U^\ominus$ ,  $\Delta^\ddagger S^\ominus$  and  $\Delta^\ddagger G^\ominus$  are used in the transition state theory of chemical reactions. They are appropriately used only in connection with elementary reactions. The relation between the rate constant  $k$  and these quantities is more generally

$$k = \kappa(k_B T/h) \exp(-\Delta^\ddagger G^\ominus/RT)$$

| <i>Name</i>                                                              | <i>Symbol</i>                                  | <i>Definition</i>                                                                     | <i>SI unit</i>                 | <i>Notes</i> |
|--------------------------------------------------------------------------|------------------------------------------------|---------------------------------------------------------------------------------------|--------------------------------|--------------|
| standard Gibbs energy of activation                                      | $\Delta^\ddagger G^\ominus, \Delta G^\ddagger$ | $\Delta^\ddagger G^\ominus = \Delta^\ddagger H^\ominus - T \Delta^\ddagger S^\ominus$ | J mol <sup>-1</sup>            | 18           |
| molecular partition function for transition state                        | $q^\ddagger, q^\ddagger(T)$                    | $k_\infty = \kappa \frac{k_B T}{h} \frac{q^\ddagger}{q_A} \exp(-E_0/k_B T)$           | 1                              | 18           |
| transmission coefficient                                                 | $\kappa, \gamma$                               | $\kappa = \frac{k_\infty h}{k_B T} \frac{q_A}{q^\ddagger} \exp(E_0/k_B T)$            | 1                              | 18           |
| molecular partition function per volume                                  | $\tilde{q}$                                    | $\tilde{q} = q/V$                                                                     | m <sup>-3</sup>                | 18           |
| specific rate constant of unimolecular reaction at $E, J$                | $k_{\text{uni}}(E, J)$                         | $k_{\text{uni}}(E, J) = -\frac{d(\ln c_A(E, J))}{dt}$                                 | s <sup>-1</sup>                | 19           |
| specific rate constant of bimolecular reaction at collision energy $E_t$ | $k_{\text{bi}}(E_t)$                           | $k_{\text{bi}}(E_t) = \sigma(E_t) \sqrt{2E_t/\mu}$                                    | m <sup>3</sup> s <sup>-1</sup> | 11           |
| density of states                                                        | $\rho(E, J, \dots)$                            | $\rho(E, J, \dots) = dN(E, J, \dots)/dE$                                              | J <sup>-1</sup>                | 19           |
| number (sum) of states                                                   | $N(E), W(E)$                                   | $N(E) = \int_0^E \rho(E') dE'$                                                        | 1                              | 19           |
| density of states of transition state                                    | $\rho^\ddagger(E)$                             | $\rho^\ddagger(E) = dN^\ddagger(E)/dE$                                                | J <sup>-1</sup>                | 19           |
| number of states of transition state                                     | $N^\ddagger(E), W^\ddagger(E)$                 | $N^\ddagger(E) = \int_{E_0}^E \rho^\ddagger(E') dE'$                                  | 1                              | 19           |
| number of open adiabatic reaction channels                               | $W(E)$                                         | $W(E) = \sum_a H(E - V_a^{\text{max}})$                                               | 1                              | 19           |
| Michaelis constant                                                       | $K_M$                                          | $v = \frac{V [S]}{K_M + [S]}$                                                         | mol m <sup>-3</sup>            | 20           |
| rate (coefficient) matrix                                                | $\mathbf{K}, K_{fi}$                           | $-\frac{d\mathbf{c}}{dt} = \mathbf{K} \mathbf{c}$                                     | s <sup>-1</sup>                | 21           |
| quantum yield, photochemical yield                                       | $\Phi, \phi$                                   |                                                                                       | 1                              | 22           |
| fluorescence rate constant $k_f$                                         |                                                | $\frac{d[h\nu]_f}{dt} = k_f c^*$                                                      | s <sup>-1</sup>                | 23, 24       |
| natural lifetime                                                         | $\tau_0$                                       | $\tau_0 = 1/k_f$                                                                      | s                              | 23           |
| natural linewidth                                                        | $\Gamma, \Gamma_f$                             | $\Gamma = \hbar k_f$                                                                  | J                              | 23           |
| predissociation linewidth                                                | $\Gamma_p, \Gamma_{\text{diss}}$               | $\Gamma_p = \hbar k_p$                                                                | J                              | 23           |

(18) (continued) where  $k$  has the dimensions of a first-order rate constant. A rate constant for a reaction of order  $n$  is obtained by multiplication with  $(c^\ominus)^{1-n}$ . For a bimolecular reaction of an ideal gas multiply by  $V^\ominus = (c^\ominus)^{-1} = (kT/p^\ominus)$ .  $\kappa$  is a transmission coefficient, and  $\Delta^\ddagger G^\ominus = \Delta^\ddagger H^\ominus - T \Delta^\ddagger S^\ominus$ . The standard symbol  $^\ominus$  is sometimes omitted, and these quantities are frequently written  $\Delta^\ddagger H$ ,  $\Delta^\ddagger U$ ,  $\Delta^\ddagger S$  and  $\Delta^\ddagger G$ . However, the omission of the specification of a standard state leads to ambiguity in the values of these quantities. The choice of  $p^\ominus$  and  $c^\ominus$  in general affects the values of  $\Delta^\ddagger H^\ominus$ ,  $\Delta^\ddagger S^\ominus$  and  $\Delta^\ddagger G^\ominus$ .

The statistical mechanical formulation of transition state theory results in the equation for  $k_\infty$  as given in the table for a unimolecular reaction in the high pressure limit, and for a bimolecular reaction one has (often with  $\kappa = 1$  assumed)

(Notes continued)

(18) (continued)

$$k_{\text{bi}} = \kappa \frac{k_{\text{B}}T}{h} \frac{\tilde{q}^\ddagger}{\tilde{q}_{\text{A}}\tilde{q}_{\text{B}}} \exp(-E_0/k_{\text{B}}T)$$

where  $q^\ddagger$  is the partition function of the transition state and  $\tilde{q}^\ddagger, \tilde{q}_{\text{A}}, \tilde{q}_{\text{B}}$  are partition functions per volume for the transition state and the reaction partners A and B.  $E_0$  is the threshold energy for reaction, below which no reaction is assumed to occur. In transition state theory, it is the difference of the zero-point level of the transition state and the zero-point level of reactants.

(19) In the theory of unimolecular reactions, it is important to consider rate constants for an ensemble of molecules at well defined energy  $E$  (and possibly other good quantum numbers such as angular momentum  $J$  etc.) with concentration  $c_{\text{A}}(E, J, \dots)$ . Expressions equivalent to transition state theory arise for these in the framework of quasi-equilibrium theory, Rice-Ramsperger-Kassel-Marcus (RRKM) theory of the statistical adiabatic channel model (SACM). The unimolecular rate constant is then given by the RRKM expression

$$k(E) = \frac{N^\ddagger(E)}{h\rho(E)}$$

$\rho(E)$  is the density of states of the molecule,  $N(E)$  is the total number of states of the molecule with energy less than  $E$ ,

$$N(E) = \sum_i H(E - E_i) = \int_0^E \rho(E') dE'$$

and  $N^\ddagger(E)$  is the corresponding number of states of the transition state. In the framework of the statistical adiabatic channel model one has

$$k(E, J) = \kappa \frac{W(E, J)}{h\rho(E, J)}$$

$V_a^{\text{max}}$  is the maximum of the potential of adiabatic channel  $a$ ,  $H(x)$  is the Heaviside function (see Section 4.2, p. 107) and  $J$  is the angular momentum quantum number [102]. There may be further constants of the motion beyond  $J$  appearing here.

(20) The Michaelis constant arises in the treatment of a mechanism of enzyme catalysis



where E is the enzyme, S the substrate, ES the enzyme-substrate complex and P the product.  $V$  is called limiting rate [100].

(21) In generalized first-order kinetics, the rate equation can be written as a matrix equation, with the concentration vector  $\mathbf{c} = (c_1, c_2, \dots, c_n)^T$  and the first-order rate coefficients  $K_{fi}$  as matrix elements.

(22) The quantum yield  $\phi$  is defined in general by the *number of defined events* divided by *number of photons absorbed* [60]. For a photochemical reaction it can be defined as

$$\phi = \left| \frac{d\xi/dt}{dn_\gamma/dt} \right|$$

which is the *rate of conversion* divided by the *rate of photon absorption*.

(23) For exponential decay by spontaneous emission (fluorescence) one has for the decay of the excited state concentration  $c^*$

$$\frac{d[h\nu]}{dt} = -\frac{dc^*}{dt} = k_{\text{f}}c^*$$

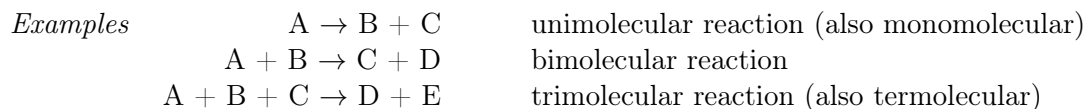
The full width at half maximum  $\Gamma$  of a lorentzian absorption line is related to the rate constant of fluorescence. For a predissociation  $\Gamma_{\text{p}}$  is related to the predissociation rate constant  $k_{\text{p}}$ . However, linewidths may also have other contributions in practice.

### 2.12.1 Other symbols, terms, and conventions used in chemical kinetics

Additional descriptions can be found in [100].

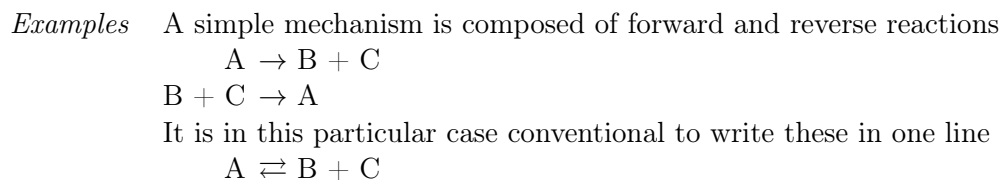
#### (i) Elementary reactions

Reactions that occur at the molecular level in one step are called *elementary reactions*. It is conventional to define them as unidirectional, written with a simple arrow always from left to right. The number of relevant reactant particles on the left hand side is called the *molecularity* of the elementary reaction.

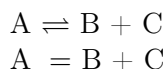


#### (ii) Composite mechanisms

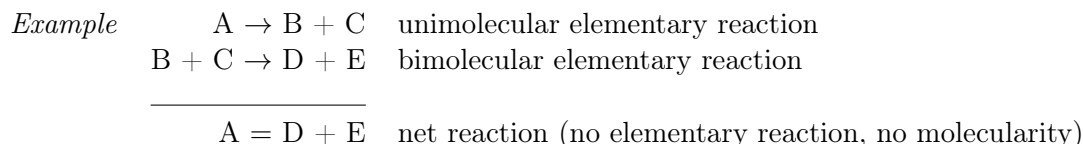
A reaction that involves more than one elementary reaction is said to occur by a composite mechanism. The terms complex mechanism, indirect mechanism, and stepwise mechanism are also commonly used. Special types of mechanisms include chain-reaction mechanisms, catalytic reaction mechanisms, etc.



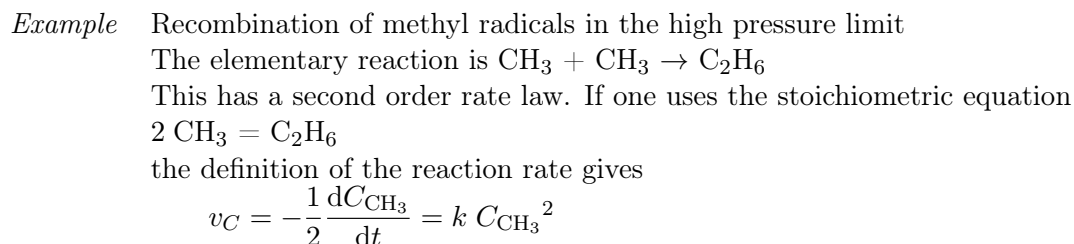
However, it is useful in kinetics to distinguish this from a net reaction, which is written either with two one-sided arrows or an “equal” sign



When one combines a composite mechanism to obtain a net reaction, one should not use the simple arrow in the resulting equation.



It is furthermore useful to distinguish the stoichiometric equation defining the reaction rate and rate constant from the equation defining the elementary reaction and rate law.

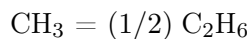


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(Notes continued)

(24) The einstein is sometimes either used as the unit of amount of substance of photons, where one einstein corresponds to 1 mol of photons, or as the unit of energy where one einstein corresponds to the energy  $Lh\nu$  of 1 mol of monochromatic photons with frequency  $\nu$ .

(continued) If one uses the stoichiometric equation



the definition of the reaction rate gives

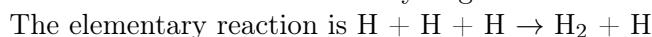
$$v_C = -\frac{dC_{\text{CH}_3}}{dt} = k' C_{\text{CH}_3}^2$$

Hence  $k' = 2k$ .

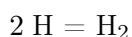
Similar to reaction enthalpies, rate coefficients are only defined with a given stoichiometric equation. It is recommended always to state explicitly the stoichiometric equation and the differential equation for the reaction rate to avoid ambiguities. If no other statement is made, the stoichiometric coefficients should be as in the reaction equation (for the elementary reaction). However, it is not always possible to use a reaction equation as stoichiometric equation.

*Example*

The trimolecular reaction for hydrogen atoms



This has a third order rate law. The stoichiometric equation is



and the definition of the reaction rate gives

$$v_c = -\frac{1}{2} \frac{d[\text{H}]}{dt} = k [\text{H}]^3$$

The bimolecular reaction  $\text{H} + \text{H} \rightarrow \text{H}_2$  with the same stoichiometric equation is a *different* elementary reaction (a very unlikely one), not the same as the trimolecular reaction. Other examples include catalyzed reactions such as  $\text{X} + \text{A} \rightarrow \text{X} + \text{B}$  with the stoichiometric equation  $\text{A} = \text{B}$  and the rate  $v_c = -d[\text{A}]/dt = k[\text{A}][\text{X}]$  ( $\neq d[\text{X}]/dt = 0$ ). Again, the unimolecular reaction  $\text{A} \rightarrow \text{B}$  exists, but would be a *different* elementary reaction with the same stoichiometry.

## 2.13 ELECTROCHEMISTRY

Electrochemical concepts, terminology and symbols are more extensively described in [1.i] and [103–107]. For the field of electrochemical engineering see [108], for semiconductor electrochemistry and photo-electrochemical energy conversion see [109], for corrosion nomenclature see [110], for impedances in electrochemical systems see [111], for electrokinetic phenomena see [112], for electrochemical biosensors see [113], and for chemically modified electrodes (CME) see [114].

| <i>Name</i>                        | <i>Symbol</i>            | <i>Definition</i>                                                    | <i>SI unit</i>       | <i>Notes</i> |
|------------------------------------|--------------------------|----------------------------------------------------------------------|----------------------|--------------|
| elementary charge                  | $e$                      | proton charge                                                        | C                    |              |
| Faraday constant                   | $F$                      | $F = eN_A$                                                           | C mol <sup>-1</sup>  |              |
| charge number of an ion            | $z$                      | $z_B = Q_B/e$                                                        | 1                    | 1            |
| ionic strength,<br>molality basis  | $I_m, I$                 | $I_m = (1/2) \sum_i m_i z_i^2$                                       | mol kg <sup>-1</sup> |              |
| concentration basis                | $I_c, I$                 | $I_c = (1/2) \sum_i c_i z_i^2$                                       | mol m <sup>-3</sup>  | 2            |
| mean ionic molality                | $m_{\pm}$                | $m_{\pm}^{(\nu_+ + \nu_-)} = m_+^{\nu_+} m_-^{\nu_-}$                | mol kg <sup>-1</sup> | 3            |
| mean ionic activity<br>coefficient | $\gamma_{\pm}$           | $\gamma_{\pm}^{(\nu_+ + \nu_-)} = \gamma_+^{\nu_+} \gamma_-^{\nu_-}$ | 1                    | 3            |
| mean ionic activity                | $a_{\pm}$                | $a_{\pm} = m_{\pm} \gamma_{\pm} / m^{\ominus}$                       | 1                    | 3, 4, 5      |
| activity of an electrolyte         | $a(A_{\nu_+} B_{\nu_-})$ | $a(A_{\nu_+} B_{\nu_-}) = a_{\pm}^{(\nu_+ + \nu_-)}$                 | 1                    | 3            |
| pH                                 | pH                       | $\text{pH} = -\lg(a_{\text{H}^+})$                                   | 1                    | 5, 6         |
| outer electric potential           | $\psi$                   |                                                                      | V                    | 7            |
| surface electric potential         | $\chi$                   |                                                                      | V                    | 8            |
| inner electric potential           | $\phi$                   | $\phi = \chi + \psi$                                                 | V                    |              |
| Volta potential difference         | $\Delta\psi$             | $\Delta_{\alpha}^{\beta} \psi = \psi^{\beta} - \psi^{\alpha}$        | V                    | 9            |

(1) The definition applies to entities B.

(2) To avoid confusion with the cathodic current, symbol  $I_c$  (note roman subscript), the symbol  $I$  or sometimes  $\mu$  (when the current is denoted by  $I$ ) is used for ionic strength based on concentration.

(3)  $\nu_+$  and  $\nu_-$  are the numbers of cations and anions per formula unit of an electrolyte  $A_{\nu_+} B_{\nu_-}$ .

*Example* For  $\text{Al}_2(\text{SO}_4)_3$ ,  $\nu_+ = 2$  and  $\nu_- = 3$ .

$m_+$  and  $m_-$ , and  $\gamma_+$  and  $\gamma_-$ , are the molalities and activity coefficients of cations and anions. If the molality of  $A_{\nu_+} B_{\nu_-}$  is  $m$ , then  $m_+ = \nu_+ m$  and  $m_- = \nu_- m$ . A similar definition is used on a concentration scale for the mean ionic concentration  $c_{\pm}$ .

(4)  $m^{\ominus} = 1 \text{ mol kg}^{-1}$ .

(5) For an individual ion, neither the activity  $a_+, a_-$  nor the activity coefficient  $\gamma_+, \gamma_-$  is experimentally measurable.

(6) The definition of pH is discussed in Section 2.13.1 (viii), p. 75. The symbol pH is an exception to the general rules for the symbols of physical quantities (Section 1.3.1, p. 5) in that it is a two-letter symbol and it is always printed in Roman (upright) type.

(7)  $\psi^{\beta}$  is the electrostatic potential of phase  $\beta$  due to the electric charge  $Q$  of the phase. It may be calculated from classical electrostatics. For example, for a conducting sphere with excess charge  $Q$  and radius  $r$ , placed in vacuo,  $\psi = Q/4\pi\epsilon_0 r$ .

(8) The surface potential is defined as the work (divided by the charge) necessary to transfer an ideal (i.e. massless) positive charge through a dipole layer at the interphase between phase  $\alpha$  and  $\beta$ . The absolute value of  $\chi$  cannot be determined, only differences are measurable.

(9)  $\Delta\psi$  is the outer potential difference due to the charge on phases  $\alpha$  and  $\beta$ . It is a measurable quantity.

| <i>Name</i>                                                     | <i>Symbol</i>                      | <i>Definition</i>                                                                                                                                                                             | <i>SI unit</i>      | <i>Notes</i> |
|-----------------------------------------------------------------|------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------|--------------|
| Galvani potential difference                                    | $\Delta\phi$                       | $\Delta_{\alpha}^{\beta}\phi = \phi^{\beta} - \phi^{\alpha}$                                                                                                                                  | V                   | 10           |
| electrochemical potential                                       | $\tilde{\mu}_{\text{B}}^{\alpha}$  | $\tilde{\mu}_{\text{B}}^{\alpha} = (\partial G/\partial n_{\text{B}}^{\alpha})_{T,p,n_{j \neq \text{B}}} = \mu_{\text{B}}^{\alpha\ominus} + RT \ln a_{\text{B}} + z_{\text{B}}F\phi^{\alpha}$ | J mol <sup>-1</sup> | 1, 11        |
| surface charge density                                          | $\sigma$                           | $\sigma = Q_{\text{s}}/A$                                                                                                                                                                     | C m <sup>-2</sup>   | 12           |
| electrode potential                                             | $E, U$                             |                                                                                                                                                                                               | V                   | 13           |
| potential difference of an electrochemical cell, cell potential | $E_{\text{cell}}, U_{\text{cell}}$ | $E_{\text{cell}} = E_{\text{R}} - E_{\text{L}}$                                                                                                                                               | V                   | 14           |
| electron number of an electrochemical reaction (charge number)  | $z, n$                             | $z =  \nu_{\text{e}} $                                                                                                                                                                        | 1                   | 15           |
| standard electrode potential                                    | $E^{\ominus}$                      | $E^{\ominus} = -\Delta_{\text{r}}G^{\ominus}/zF$                                                                                                                                              | V                   | 16, 17       |

(10)  $\Delta\phi$  is the inner potential difference between points within the bulk phases  $\alpha$  and  $\beta$ ; it is measurable only if the phases are of identical composition.

(11) The electrochemical potential of ion B in a phase is the partial molar Gibbs energy of the ion. The special name “electrochemical potential”, and the tilde on the symbol  $\tilde{\mu}_{\text{B}}$ , are used as a reminder that for an ion the partial molar Gibbs energy depends on the inner potential  $\phi$  as well as on the chemical composition. The difference of electrochemical potential of ion B between two phases  $\alpha$  and  $\beta$  of different chemical composition and different inner potential is given by

$$\tilde{\mu}_{\text{B}}^{\alpha} - \tilde{\mu}_{\text{B}}^{\beta} = \mu_{\text{B}}^{\alpha\ominus} - \mu_{\text{B}}^{\beta\ominus} + RT \ln (a_{\text{B}}^{\alpha}/a_{\text{B}}^{\beta}) + z_{\text{B}}F(\phi^{\alpha} - \phi^{\beta})$$

It is not generally possible to separate the term depending on the composition from the term depending on the inner potential, because it is not possible to measure either the relative activity  $a_{\text{B}}$  of an ion, or the inner potential difference between two phases of different chemical composition. The electrochemical potential, however, is always well defined, and in a multiphase system with phases at different inner potentials, equilibrium with regard to the transfer of ion B between phases is achieved when its electrochemical potential is the same in all phases.

(12)  $Q_{\text{s}}$  is the charge on one side of the interface,  $A$  is the surface area. For the case of metal and semiconductor electrodes, by convention the charge refers to the electrode side.

(13) The absolute value of the electrode potential cannot be measured, so  $E$  is always reported relative to the potential of some reference electrode, e.g. that of a standard hydrogen electrode (SHE) (see Section 2.13.1 (vi), p. 74). The concept of an *absolute* electrode potential is discussed in [106].

(14) The two electrodes of an electrochemical cell can be distinguished by the subscripts L (left) and R (right), or 1 and 2. This facilitates the representation of the cell in a so-called cell diagram (see Section 2.13.1 (iii), p. 73).  $E_{\text{L}}$  and  $E_{\text{R}}$  are the electrode potentials of these two electrodes. In electrochemical engineering, the cell voltage is exclusively denoted by  $U$ . The limiting value of  $E_{\text{cell}}$  for zero current flowing through the cell, all local charge transfer and chemical equilibria being established, was formerly called emf (electromotive force). The name electromotive force and the symbol emf are no longer recommended, since a potential difference is not a force.

(15)  $z$  (or  $n$ ) is the number of electrons in the balanced electrode reaction as written. It is a positive integer, equal to  $|\nu_{\text{e}}|$ , when  $\nu_{\text{e}}$  is the stoichiometric number of the electron in the electrode reaction.  $n$  is commonly used where there is no risk of confusion with amount of substance.

(16) The symbols  $^{\ominus}$  and  $^{\circ}$  are both used to indicate standard state; they are equally acceptable.

(17) The standard potential is the value of the *equilibrium potential* of an electrode under standard conditions.  $\Delta_{\text{r}}G^{\ominus}$  is the standard Gibbs energy of this electrode reaction, written as a reduction, with respect to that of the standard hydrogen electrode (see also Section 2.13.1 (vi), p. 74). The electrode potential attains its *equilibrium value* if no current flows through the cell, and all local charge

| Name                                                             | Symbol             | Definition                                                           | SI unit           | Notes     |
|------------------------------------------------------------------|--------------------|----------------------------------------------------------------------|-------------------|-----------|
| equilibrium electrode potential (of an electrochemical reaction) | $E_{\text{eq}}$    | $E_{\text{eq}} = E^\circ - (RT/zF) \sum_i \nu_i \ln a_i$             | V                 | 15–18     |
| formal potential                                                 | $E^{\circ'}$       | $E_{\text{eq}} = E^{\circ'} - (RT/zF) \sum_i \nu_i \ln(c_i/c^\circ)$ | V                 | 15, 19    |
| liquid junction potential                                        | $E_j$              |                                                                      |                   | 20        |
| electric current                                                 | $I$                | $I = dQ/dt$                                                          | A                 | 21        |
| electric current density                                         | $j, \mathbf{j}$    | $j = I/A$                                                            | A m <sup>-2</sup> | 22        |
| faradaic current                                                 | $I_F$              | $I_F = I_c + I_a$                                                    | A                 | 23        |
| reduction rate constant                                          | $k_c$              | $I_c = -nFAk_c \prod_B (c_B')^{n_B}$                                 | (varies)          | 1, 15, 24 |
| oxidation rate constant                                          | $k_a$              | $I_a = nFAk_a \prod_B (c_B')^{n_B}$                                  | (varies)          | 1, 15, 24 |
| transfer coefficient                                             | $\alpha, \alpha_c$ | $\alpha_c = -(RT/nF)d(\ln k_c)/dE$                                   | 1                 | 15, 25    |
| overpotential                                                    | $\eta, E_\eta$     | $\eta = E - E_{\text{eq}}$                                           | V                 |           |
| Tafel slope                                                      | $b$                | $b = (\partial E / \partial \ln  I_F )_{c_i, T, p}$                  | V                 | 26        |
| mass transfer coefficient                                        | $k_d$              | $k_{d,B} =  \nu_B  I_{\text{lim},B} / nFcA$                          | m s <sup>-1</sup> | 1, 15, 27 |

(17) (continued) transfer equilibria across phase boundaries that are represented in the cell diagram (except at possible electrolyte-electrolyte junctions) and local chemical equilibria within the phases are established.

(18) This is the Nernst equation.  $\sum \nu_i \ln a_i$  refers to the electrode reaction, where  $a_i$  are the activities of the species taking part in it;  $\nu_i$  are the stoichiometric numbers of these species in the equation for the electrode reaction, written as a reduction,  $\nu_i$  is positive for products and negative for reactants. The equilibrium potential is also called *Nernst potential*, or *reversible potential*.

(19) It is  $E^{\circ'}$  which is calculated in electrochemical experiments when the concentrations of the various species are known, but not their activities. Its value depends on the composition of the electrolyte solution. The argument of  $\ln$  is dimensionless, the concentration  $c_i$  is normalized through division by the standard concentration  $c^\circ$ , usually  $c^\circ = 1 \text{ mol dm}^{-3}$  for soluble species.

(20)  $E_j$  is the Galvani potential difference between two electrolyte solutions in contact.

(21)  $Q$  is the charge transferred through the external circuit of the cell.

(22) Formally, the current density is a vector,  $dI = \mathbf{j} \cdot \mathbf{e}_n dA$  (see Section 2.3, note 2, p. 16).

(23)  $I_F$  is the current through the electrode|solution interface, resulting from the charge transfer due to an electrode reaction proceeding as reactants +  $ne^- \rightarrow$  products.  $I_c$  is the *cathodic* partial current due to the reduction reaction,  $I_a$  is the *anodic* partial current due to the oxidation reaction. By definition  $I_c$  is negative and  $I_a$  positive. At the equilibrium potential,  $I_a = -I_c = I_0$  (the exchange current) and  $j_a = -j_c = j_0$  (the exchange current density).  $I$  or  $j$  may achieve limiting values, indicated by the subscript  $\text{lim}$ , in addition to the subscripts F, c, or a.

(24) For a first-order reaction the SI unit is m s<sup>-1</sup>. Here  $n$  (or  $z$ ) is the number of electrons transferred in the electrochemical reaction,  $c_B'$  is the concentration at the interphase,  $n_B$  is the order of the reaction with respect to entity B. The formerly used symbols  $k_{\text{red}}, k_f$  and  $\overrightarrow{k}$  (for  $k_c$ ) and  $k_{\text{ox}}, k_b$  and  $\overleftarrow{k}$  (for  $k_a$ ) are not recommended.

(25)  $\alpha$  or  $\alpha_c$  is also named *cathodic transfer coefficient*. Analogously the *anodic transfer coefficient* is defined as  $\alpha_a = (RT/nF)d(\ln k_a)/dE$ . At the same potential  $\alpha_a = 1 - \alpha_c$ . The *symmetry factor*,  $\beta$ , is the cathodic transfer coefficient,  $\alpha_i$  of an elementary reaction step  $i$ , in which only one electron is transferred.

(26) The Tafel slope  $b$  is an experimental quantity from which kinetic information can be derived.

(27) The mass transfer coefficient is the flux divided by the concentration. For steady-state mass transfer,  $k_{d,B} = D_B/\delta_B$  where  $\delta_B$  is the diffusion layer thickness (which is model dependent) and  $D_B$  is the diffusion coefficient of entity B. For more information see [17].

| <i>Name</i>                                            | <i>Symbol</i>      | <i>Definition</i>                                                                                   | <i>SI unit</i>                           | <i>Notes</i> |
|--------------------------------------------------------|--------------------|-----------------------------------------------------------------------------------------------------|------------------------------------------|--------------|
| electrokinetic potential<br>( $\zeta$ -potential)      | $\zeta$            |                                                                                                     | V                                        |              |
| conductivity                                           | $\kappa, (\sigma)$ | $\mathbf{j} = \kappa \mathbf{E}$                                                                    | $\text{S m}^{-1}$                        | 22, 28       |
| electric mobility                                      | $u, (m)$           | $u_{\text{B}} =  v_{\text{B}} / \mathbf{E} $                                                        | $\text{m}^2 \text{V}^{-1} \text{s}^{-1}$ | 1, 29        |
| ionic conductivity,<br>molar conductivity<br>of an ion | $\lambda$          | $\lambda_{\text{B}} =  z_{\text{B}}  F u_{\text{B}}$                                                | $\text{S m}^2 \text{mol}^{-1}$           | 1, 30, 31    |
| molar conductivity                                     | $\Lambda$          | $\Lambda(\text{A}_{\nu+}\text{B}_{\nu-}) = \nu_+ \lambda_+ + \nu_- \lambda_-$                       | $\text{S m}^2 \text{mol}^{-1}$           | 1, 30, 31    |
| transport number                                       | $t$                | $t_{\text{B}} = \lambda_{\text{B}} c_{\text{B}} / \sum_i \lambda_i c_i = j_{\text{B}} / \sum_i j_i$ | 1                                        | 1            |

(28) Conductivity was formerly called specific conductance.  $\mathbf{E}$  is the electric field strength vector.

(29)  $v_{\text{B}}$  is the migration velocity of entities B and  $|\mathbf{E}|$  is the electric field strength within the phase concerned.

(30) The unit  $\text{S cm}^2 \text{mol}^{-1}$  is often used for molar conductivity. The conductivity is equal to  $\kappa = \sum \lambda_i c_i$ .

(31) It is important to specify the entity to which molar conductivity refers. It is standard practice to choose the entity to be  $1/z_{\text{B}}$  of an ion of charge number  $z_{\text{B}}$ , in order to normalize ion charge, so that for example molar conductivities for potassium, barium and lanthanum ions would be quoted as  $\lambda(\text{K}^+)$ ,  $\lambda((1/2) \text{Ba}^{2+})$ , or  $\lambda((1/3) \text{La}^{3+})$  the so-called molecular conductivity of *equivalent entities* (formerly called *equivalent conductivity*) [115].

### 2.13.1 Sign and notation conventions in electrochemistry <sup>1</sup>

#### (i) Electrochemical cells

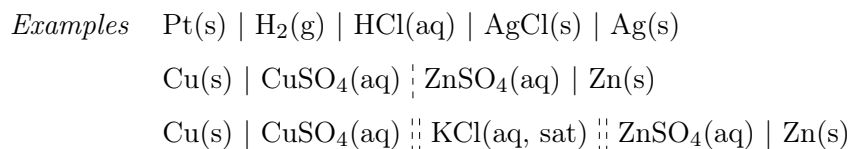
Electrochemical cells consist of at least two (usually metallic) electron conductors in contact with ionic conductors (electrolytes). The current flow through electrochemical cells may be zero or non-zero. Electrochemical cells with current flow can operate either as *galvanic cells*, in which chemical reactions occur spontaneously and chemical energy is converted into electrical energy, or as *electrolytic cells*, in which electrical energy is converted into chemical energy. In both cases part of the energy becomes converted into (positive or negative) heat.

#### (ii) Electrode

There are at present two usages for the term electrode, namely either (i) the electron conductor (usually a metal) connected to the external circuits or (ii) the half cell consisting of one electron conductor and at least one ionic conductor. The latter version has usually been favored in electrochemistry.

#### (iii) Representation of electrochemical cells

Electrochemical cells are represented by diagrams such as those in the following examples:



A single vertical bar ( $\mid$ ) should be used to represent a phase boundary, a dashed vertical bar ( $\parallel$ ) to represent a junction between miscible liquids, and double dashed vertical bars ( $\parallel \parallel$ ) to represent a liquid junction, in which the liquid junction potential is assumed to be eliminated.

<sup>1</sup> These are in accordance with the “Stockholm Convention” of 1953 [103].

#### (iv) Potential difference of an electrochemical cell

The potential difference of an electrochemical cell is measured between a metallic conductor attached to the right-hand electrode of the cell diagram and an identical metallic conductor attached to the left-hand electrode. Electric potential differences can be measured only between two pieces of material of the same composition. In practice, these are almost always two pieces of copper, attached to the electrodes of the cell.

At a boundary copper | electrode material there is a contact potential difference; its value is incorporated in the constant to which the electrode potential is referred.

Due to the different rates of diffusion of anions and cations from one solution to the other, liquid junction potentials,  $E_j$ , appear whenever two solutions of different composition are immiscible or are separated by a diaphragm or some other means to avoid mixing. If equilibrium is not established at these junctions, the cells may include unknown junction potentials. Salt bridges (see, for example, the third cell in the previous Section 2.13.1 (iii)) are commonly employed to minimize or to stabilize the contributions of (liquid) junction potentials at the interface of two miscible electrolyte solutions to the measured potential difference. At the interface of two immiscible electrolyte solutions, a thermodynamic distribution potential can be established due to equilibrium charge partition.

#### (v) Standard potential of an electrochemical cell reaction

If no current flows through the cell, and all local charge transfer and local chemical equilibria of each electrode reaction are established, the potential difference of the cell is related to the Gibbs energy of the overall cell reaction by the equation

$$\Delta_r G = -zFE_{\text{cell,eq}}$$

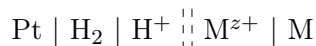
assuming that junction potentials are negligible. If the reaction of the electrochemical cell as written takes place spontaneously,  $\Delta_r G$  is negative and  $E_{\text{cell}}$  positive. If the cell is written with the electrodes in the inverse order  $\Delta_r G$  is positive and  $E_{\text{cell}}$  is negative. The equilibrium potential of the cell, i.e. when no current flows, is given by

$$E_{\text{cell,eq}} = E^\circ_{\text{cell,eq}} - \frac{RT}{zF} \sum_i \nu_i \ln a_i$$

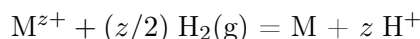
$a_i$  are the activities of the species taking part in the cell reaction and  $\nu_i$  are the stoichiometric numbers of these species in the equation written for the cell reaction (see also notes 15 and 16, p. 71).

#### (vi) Standard electrode potential (of an electrochemical reaction)

The *standard potential* of an electrochemical reaction, abbreviated as standard potential, is defined as the standard potential of a hypothetical cell, in which the electrode (half-cell) at the left of the cell diagram is the *standard hydrogen electrode* (SHE) and the electrode at the right is the electrode in question. This implies that the cell reaction always involves the oxidation of molecular hydrogen. The standard hydrogen electrode consists of a platinum electrode in contact with a solution of  $\text{H}^+$  at unit activity and saturated with  $\text{H}_2$  gas with a fugacity referred to the standard pressure of  $10^5$  Pa (see Section 2.11.1 (v), p. 62). For a metallic electrode in equilibrium with solvated ions the cell diagram is



and relates to the reaction



This diagram may be abbreviated  $E(\text{M}^{z+}/\text{M})$ , but the order of these symbols should not be reversed. Note that the standard hydrogen electrode as defined is limited to aqueous solutions. For more information on measuring electrode potentials in aqueous and non-aqueous systems, see [103, 107].

### (vii) Anode, cathode

The terms anode and cathode may only be applied to electrochemical cells through which a net current flows. In a cell at equilibrium the terms plus pole and minus pole are used. An *anode* is an electrode at which the predominating electrochemical reaction is an oxidation; electrons are produced (in a galvanic cell) or extracted (in an electrolytic cell). A *cathode* is an electrode at which the predominating electrochemical reaction is a reduction which consumes the electrons from the anode, and that reach the cathode through the external circuit. Since electrons flow from lower to higher (more positive) electric potentials, in a galvanic cell they flow from the anode (the negative electrode) to the cathode (the positive electrode), while in an electrolytic cell the electrons extracted from the anode (the positive electrode) by the external source flow to the cathode (the negative electrode).

Note that in rechargeable batteries such as the lead-acid battery (lead accumulator), the positive electrode is a cathode during discharge and an anode during charge, and the negative electrode is the anode during discharge and the cathode during charge. In order to avoid confusion, it is recommended that in rechargeable batteries only the terms positive and negative electrodes (or plates) be used. For more detailed information on electrode potentials, see [109].

### (viii) Definition of pH [116]

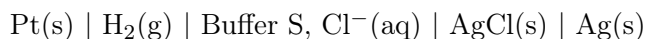
The quantity pH is defined in terms of the activity of hydrogen(1+) ions (hydrogen ions) in solution:

$$\text{pH} = \text{p}a_{\text{H}^+} = -\lg(a_{\text{H}^+}) = -\lg(m_{\text{H}^+}\gamma_{\text{m}, \text{H}^+}/m^\ominus)$$

where  $a_{\text{H}^+}$  is the activity of the hydrogen(1+) (hydrogen ion) in solution,  $\text{H}^+(\text{aq})$ , and  $\gamma_{\text{m}, \text{H}^+}$  is the activity coefficient of  $\text{H}^+(\text{aq})$  on the molality basis at molality  $m_{\text{H}^+}$ . The symbol  $\text{p}$  is interpreted as an operator ( $\text{p}x = -\lg x$ ) [116, 117] (see Section 4.1, p. 103) with the unique exception of the symbol pH [117]. The symbol pH is also an exception to the rules for symbols and quantities (see Section 1.3.1, p. 5). The standard molality  $m^\ominus$  is chosen to be equal to 1 mol kg<sup>-1</sup>. Since pH is defined in terms of a quantity that cannot be measured independently, the above equation can be only regarded as a *notional* definition.

The establishment of primary pH standards requires the application of the concept of the “primary method of measurement” [116], assuring full traceability of the results of all measurements and their uncertainties. Any limitation in the theory or determination of experimental variables must be included in the estimated uncertainty of the method.

The primary method for the measurement of pH involves the use of a cell without transference, known as the *Harned cell* [118]:



Application of the Nernst equation to the above leads to the relationship

$$E = E^\ominus - \frac{RT \ln 10}{F} \lg[(m_{\text{H}^+}\gamma_{\text{H}^+}/m^\ominus)(m_{\text{Cl}^-}\gamma_{\text{Cl}^-}/m^\ominus)]$$

where  $E$  is the potential difference of the cell and  $E^\ominus$  the standard potential of the AgCl|Ag electrode. This equation can be rearranged to yield

$$-\lg(a_{\text{H}^+}\gamma_{\text{Cl}^-}) = \frac{(E - E^\ominus)}{(RT \ln 10)/F} + \lg(m_{\text{Cl}^-}/m^\ominus)$$

Measurements of  $E$  are made and the quantity  $-\lg(a_{\text{H}^+}\gamma_{\text{Cl}^-})$  is obtained by extrapolation to  $m_{\text{Cl}^-}/m^\ominus = 0$ . The value of  $\gamma_{\text{Cl}^-}$  is calculated using the Bates-Guggenheim convention [117] based on Debye-Hückel theory. Then  $-\lg(a_{\text{H}^+})$  is calculated and identified as pH(PS), where PS signifies primary standard. The uncertainties in the two estimates are typically  $\pm 0.001$  in  $-\lg(a_{\text{H}^+}\gamma_{\text{Cl}^-})^\ominus$ , and  $\pm 0.003$  in pH, respectively.

Materials for primary standard pH buffers must also meet the appropriate requirements for reference materials including chemical purity and stability, and applicability of the Bates-Guggenheim convention to the estimation of  $\lg(\gamma_{\text{Cl}^-})$ . This convention requires that the ionic strength must be  $\leq 0.1 \text{ mol kg}^{-1}$ . Primary standard buffers should also lead to small liquid junction potentials when used in cells with liquid junctions. Secondary standards, pH(SS), are also available, but carry a greater uncertainty in measured values.

Practical pH measurements generally use cells involving liquid junctions in which, consequently, liquid junction potentials,  $E_j$ , are present [116]. Measurements of pH are not normally performed using the Pt|H<sub>2</sub> electrode, but rather the glass (or other H<sup>+</sup>-selective) electrode, whose response factor ( $dE/d\text{pH}$ ) usually deviates from the Nernst slope. The associated uncertainties are significantly larger than those associated with the fundamental measurements using the Harned cell. Nonetheless, incorporation of the uncertainties for the primary method, and for all subsequent measurements, permits the uncertainties for all procedures to be linked to the primary standards by an unbroken chain of comparisons.

Reference values for standards in D<sub>2</sub>O and aqueous-organic solvent mixtures exist [119].

## 2.14 COLLOID AND SURFACE CHEMISTRY

The recommendations given here are based on more extensive IUPAC recommendations [1.e–1.h] and [120–123]. Catalyst characterization is described in [124] and quantities related to macromolecules in [125].

| <i>Name</i>                                    | <i>Symbol</i>                 | <i>Definition</i>                                  | <i>SI unit</i>                     | <i>Notes</i> |
|------------------------------------------------|-------------------------------|----------------------------------------------------|------------------------------------|--------------|
| specific surface area                          | $a_s, a, s$                   | $a_s = A/m$                                        | $\text{m}^2 \text{kg}^{-1}$        | 1            |
| surface amount of B                            | $n_B^s$                       |                                                    | mol                                | 2            |
| adsorbed amount of B                           | $n_B^a$                       |                                                    | mol                                | 2            |
| surface excess amount of B                     | $n_B^\sigma$                  |                                                    | mol                                | 3            |
| surface excess<br>concentration of B           | $\Gamma_B, (\Gamma_B^\sigma)$ | $\Gamma_B = n_B^\sigma / A$                        | $\text{mol m}^{-2}$                | 3            |
| total surface excess<br>concentration          | $\Gamma, (\Gamma^\sigma)$     | $\Gamma = \sum_i \Gamma_i$                         | $\text{mol m}^{-2}$                |              |
| area per molecule                              | $a, \sigma$                   | $a_B = A/N_B^a$                                    | $\text{m}^2$                       | 4            |
| area per molecule in a<br>filled monolayer     | $a_m, \sigma_m$               | $a_{m,B} = A/N_{m,B}$                              | $\text{m}^2$                       | 4            |
| surface coverage                               | $\theta$                      | $\theta = N_B^a / N_{m,B}$                         | 1                                  | 4            |
| contact angle                                  | $\theta$                      |                                                    | rad, 1                             |              |
| film thickness                                 | $t, h, \delta$                |                                                    | m                                  |              |
| thickness of (surface or<br>interfacial) layer | $\tau, \delta, t$             |                                                    | m                                  |              |
| surface tension,<br>interfacial tension        | $\gamma, \sigma$              | $\gamma = (\partial G / \partial A_s)_{T,p,n_i}$   | $\text{N m}^{-1}, \text{J m}^{-2}$ |              |
| film tension                                   | $\Sigma_f$                    | $\Sigma_f = (\partial G / \partial A_f)_{T,p,n_i}$ | $\text{N m}^{-1}$                  | 5            |
| Debye length of the<br>diffuse layer           | $L_D$                         | $L_D = \kappa^{-1}$                                | m                                  | 6            |
| average molar masses                           |                               |                                                    |                                    |              |
| number-average                                 | $M_n$                         | $M_n = \sum_i n_i M_i / \sum_i n_i$                | $\text{kg mol}^{-1}$               |              |
| mass-average                                   | $M_m, M_w$                    | $M_m = \sum_i n_i M_i^2 / \sum_i n_i M_i$          | $\text{kg mol}^{-1}$               |              |
| z-average                                      | $M_z$                         | $M_z = \sum_i n_i M_i^3 / \sum_i n_i M_i^2$        | $\text{kg mol}^{-1}$               |              |
| sedimentation coefficient                      | $s$                           | $s = v/a$                                          | s                                  | 7            |
| van der Waals constant                         | $\lambda$                     |                                                    | J                                  |              |
| retarded van der Waals<br>constant             | $\beta, B$                    |                                                    | J                                  |              |
| van der Waals-Hamaker<br>constant              | $A_H$                         |                                                    | J                                  |              |
| surface pressure                               | $\pi$                         | $\pi = \gamma^0 - \gamma$                          | $\text{N m}^{-1}$                  | 8            |

(1) The subscript s designates any surface area where absorption or deposition of species may occur.  $m$  designates the mass of a solid absorbent.

(2) The value of  $n_B^s$  depends on the thickness assigned to the surface layer (see also note 1, p. 47).

(3) The values of  $n_B^\sigma$  and  $\Gamma_B$  depend on the convention used to define the position of the Gibbs surface. They are given by the excess amount of B or surface concentration of B over values that would apply if each of the two bulk phases were homogeneous right up to the Gibbs dividing surface.

## Additional recommendations

The superscript  $s$  denotes the properties of a surface or interfacial layer. In the presence of adsorption it may be replaced by the superscript  $a$ .

|                 |                                       |                           |
|-----------------|---------------------------------------|---------------------------|
| <i>Examples</i> | Helmholtz energy of interfacial layer | $A^s$                     |
|                 | amount of adsorbed substance          | $n^a$                     |
|                 | amount of adsorbed $O_2$              | $n^a(O_2)$ or $n(O_2, a)$ |
|                 | area per molecule B in a monolayer    | $a(B), a_B$               |

The superscript  $\sigma$  is used to denote a surface excess property relative to the Gibbs surface.

|                |                                |              |
|----------------|--------------------------------|--------------|
| <i>Example</i> | surface excess amount          | $n_B^\sigma$ |
|                | (or Gibbs surface amount) of B |              |

In general the values of  $\Gamma_A$  and  $\Gamma_B$  depend on the position chosen for the Gibbs dividing surface. However, two quantities,  $\Gamma_B^{(A)}$  and  $\Gamma_B^{(n)}$  (and correspondingly  $n_B^{\sigma(A)}$  and  $n_B^{\sigma(n)}$ ), may be defined in a way that is invariant to this choice (see [1.e]).  $\Gamma_B^{(A)}$  is called the *relative* surface excess concentration of B with respect to A, or more simply the relative adsorption of B; it is the value of  $\Gamma_B$  when the surface is chosen to make  $\Gamma_A = 0$ .  $\Gamma_B^{(n)}$  is called the *reduced* surface excess concentration of B, or more simply the reduced adsorption of B; it is the value of  $\Gamma_B$  when the surface is chosen to make the total excess  $\Gamma = \sum_i \Gamma_i = 0$ .

Properties of phases ( $\alpha, \beta, \gamma$ ) may be denoted by corresponding superscript indices.

|                 |                                                         |                        |
|-----------------|---------------------------------------------------------|------------------------|
| <i>Examples</i> | surface tension of phase $\alpha$                       | $\gamma^\alpha$        |
|                 | interfacial tension between phases $\alpha$ and $\beta$ | $\gamma^{\alpha\beta}$ |

Symbols of thermodynamic quantities divided by surface area are usually the corresponding lower case letters; an alternative is to use a circumflex.

|                |                              |                            |
|----------------|------------------------------|----------------------------|
| <i>Example</i> | interfacial entropy per area | $s^s(= \hat{s}^s) = S^s/A$ |
|----------------|------------------------------|----------------------------|

The following abbreviations are used in colloid chemistry:

|     |                                      |
|-----|--------------------------------------|
| ccc | critical coagulation concentration   |
| cmc | critical micellisation concentration |
| iep | isoelectric point                    |
| pzc | point of zero charge                 |

---

(Notes continued)

(3) (continued) See [1.e], and also additional recommendations on this page.

(4)  $N_B^a$  is the number of adsorbed molecules ( $N_B^a = L n_B^a$ ), and  $N_{m,B}^a$  is the number of adsorbed molecules in a filled monolayer. The definition applies to entities B.

(5) In the equation,  $A_f$  is the area of the film.

(6) The characteristic Debye length [1.e] and [123] or Debye screening length [123]  $L_D$  appears in Gouy-Chapman theory and in the theory of semiconductor space-charge.

(7) In the definition,  $v$  is the velocity of sedimentation and  $a$  is the acceleration of free fall or centrifugation. The symbol for a limiting sedimentation coefficient is  $[s]$ , for a reduced sedimentation coefficient  $s^\circ$ , and for a reduced limiting sedimentation coefficient  $[s^\circ]$ ; see [1.e] for further details.

(8) In the definition,  $\gamma^0$  is the surface tension of the clean surface and  $\gamma$  that of the covered surface.

### 2.14.1 Surface structure [126]

#### (i) Single-crystal surface and vector designations

3D Miller indices  $(hkl)$  are used to specify a surface termination with a  $(hkl)$  bulk plane. Families of symmetry-equivalent surfaces are designated  $\{hkl\}$ . Vector directions are designated  $[hkl]$  and families of symmetry-equivalent vector directions  $\langle hkl \rangle$ .

#### (ii) Notations for stepped surfaces (for ideally terminated bulk lattices)

Stepped surfaces (often consisting of mono-atomic height steps and terraces of arbitrary width) with general Miller indices  $(hkl)$  are viewed as composed of alternating low-Miller-index facets  $(h_s k_s l_s)$  for the step faces and  $(h_t k_t l_t)$  for the terrace planes. The step notation designates such a surface as

$$(hkl) = n(h_t k_t l_t) \times (h_s k_s l_s)$$

where  $n$  measures the terrace width in units of atom rows.

*Example* The (755) surface of an fcc solid, with (111) oriented terraces and (100) oriented step faces is designated in the step notation as  $6(111) \times (100)$ .

The microfacet notation allows more complex surface terminations, using decomposition into three independent low-index facet orientations  $(h_1 k_1 l_1)$ ,  $(h_2 k_2 l_2)$  and  $(h_3 k_3 l_3)$ :

$$(hkl) = a_\lambda^1(h_1 k_1 l_1) + a_\mu^2(h_2 k_2 l_2) + a_\nu^3(h_3 k_3 l_3).$$

The factors  $a^\beta$  are simple vectorial decomposition coefficients of  $(hkl)$  onto the three microfacet Miller-index vectors, while the subscripts  $\lambda, \mu$  and  $\nu$  indicate the relative facet sizes in terms of 2D unit cells on each facet.

*Example* The (10,8,7) surface of an fcc solid, with (111) oriented terraces and steps zigzagging with alternating orientations  $(11\bar{1})$  and  $(100)$ , is designated in the microfacet notation as  $(10,8,7) = [(15/2)_{15}(111) + (1/2)_1(11\bar{1}) + 2_2(100)]$ .

#### (iii) Notations for superlattices at surfaces

Matrix notation: a 2D superlattice with basis vectors  $\mathbf{b}_1, \mathbf{b}_2$  on a bulk-like substrate with 2D surface basis vectors  $\mathbf{a}_1, \mathbf{a}_2$  satisfies the relation  $\mathbf{b}_1 = m_{11}\mathbf{a}_1 + m_{12}\mathbf{a}_2$  and  $\mathbf{b}_2 = m_{21}\mathbf{a}_1 + m_{22}\mathbf{a}_2$ , thus defining a matrix  $M = \begin{pmatrix} m_{11} & m_{12} \\ m_{21} & m_{22} \end{pmatrix}$  that uniquely specifies the superlattice. If the matrix elements are all integers, the superlattice is termed commensurate with the substrate lattice.

Wood notation: in many cases the superlattice can be labeled as  $i \left( \frac{b_1}{a_1} \times \frac{b_2}{a_2} \right) R\alpha^\circ$ , where  $i$  is c for centered superlattices or  $i$  is p for primitive superlattices (p is often dropped) and  $\alpha$  is a rotation angle relative to the substrate basis vectors (this part is dropped when  $\alpha = 0$ ).

*Example* The black and white chess board is a superlattice relative to the underlying set of all squares. This superlattice can be designated with a matrix  $\begin{pmatrix} 1 & 1 \\ -1 & 1 \end{pmatrix}$ , which is equivalent to the Wood label  $(\sqrt{2} \times \sqrt{2}) R45^\circ$  or with the alternative description  $c(2 \times 2)$ .

#### (iv) Symbols

|                                  |                                          |
|----------------------------------|------------------------------------------|
| $\mathbf{a}_1, \mathbf{a}_2$     | 2D substrate basis vector                |
| $\mathbf{a}_1^*, \mathbf{a}_2^*$ | 2D reciprocal substrate basis vector     |
| $\mathbf{b}_1, \mathbf{b}_2$     | 2D superlattice basis vectors            |
| $\mathbf{b}_1^*, \mathbf{b}_2^*$ | 2D reciprocal superlattice basis vectors |
| $\mathbf{g}$                     | 2D reciprocal lattice vector             |
| $hk$                             | 2D Miller indices                        |
| $hkl$                            | 3D Miller indices                        |
| $M$                              | matrix for superlattice notation         |
| $z$                              | coordinate perpendicular to surface      |
| $\theta$                         | surface coverage                         |
| $\Theta_D$                       | Debye temperature                        |
| $\Delta\phi_W$                   | work function change                     |
| $\lambda_e$                      | electron mean free path                  |
| $\phi_W$                         | work function                            |

## 2.15 TRANSPORT PROPERTIES

The names and symbols recommended here are in agreement with those recommended by IUPAP [4] and ISO [5.m]. Further information on transport phenomena in electrochemical systems can also be found in [104].

The following symbols are used in the definitions: mass ( $m$ ), time ( $t$ ), volume ( $V$ ), area ( $A$ ), density ( $\rho$ ), speed ( $v$ ), length ( $l$ ), viscosity ( $\eta$ ), pressure ( $p$ ), acceleration of free fall ( $g$ ), cubic expansion coefficient ( $\alpha$ ), thermodynamic temperature ( $T$ ), surface tension ( $\gamma$ ), speed of sound ( $c$ ), mean free path ( $\lambda$ ), frequency ( $f$ ), thermal diffusivity ( $a$ ), coefficient of heat transfer ( $h$ ), thermal conductivity ( $k$ ), specific heat capacity at constant pressure ( $c_p$ ), diffusion coefficient ( $D$ ), mole fraction ( $x$ ), mass transfer coefficient ( $k_d$ ), permeability ( $\mu$ ), electric conductivity ( $\kappa$ ), and magnetic flux density ( $B$ ).

| <i>Name</i>                   | <i>Symbol</i>       | <i>Definition</i>                                   | <i>SI unit</i>                           | <i>Notes</i> |
|-------------------------------|---------------------|-----------------------------------------------------|------------------------------------------|--------------|
| flux of mass $m$              | $q_m$               | $q_m = dm/dt$                                       | $\text{kg s}^{-1}$                       | 1            |
| flux density of mass $m$      | $J_m$               | $J_m = q_m/A$                                       | $\text{kg m}^{-2} \text{s}^{-1}$         | 2, 3         |
| heat flux,<br>thermal power   | $\Phi, P$           | $\Phi = dQ/dt$                                      | W                                        | 1            |
| heat flux density             | $J_q$               | $J_q = \Phi/A$                                      | $\text{W m}^{-2}$                        | 3            |
| thermal conductance           | $G$                 | $G = \Phi/\Delta T$                                 | $\text{W K}^{-1}$                        |              |
| thermal resistance            | $R$                 | $R = 1/G$                                           | $\text{K W}^{-1}$                        |              |
| thermal conductivity          | $\lambda, k$        | $\lambda = -J_q/(dT/dl)$                            | $\text{W m}^{-1} \text{K}^{-1}$          |              |
| coefficient of heat transfer  | $h, (k, K, \alpha)$ | $h = J_q/\Delta T$                                  | $\text{W m}^{-2} \text{K}^{-1}$          |              |
| thermal diffusivity           | $a$                 | $a = \lambda/\rho c_p$                              | $\text{m}^2 \text{s}^{-1}$               |              |
| diffusion coefficient         | $D$                 | $D = -J_n/(dc/dl)$                                  | $\text{m}^2 \text{s}^{-1}$               | 4            |
| viscosity,<br>shear viscosity | $\eta$              | $\eta = -\tau_{xz}(\partial V_x/\partial z)^{-1}$   | $\text{Pa s}$                            | 5            |
| bulk viscosity                | $\kappa$            | $\kappa = -\tau_{xx}(\nabla \cdot \mathbf{V})^{-1}$ | $\text{Pa s}$                            | 5            |
| thermal diffusion coefficient | $D^T$               | $D^T = J_x c^{-1}(dT/dx)^{-1}$                      | $\text{m}^2 \text{K}^{-1} \text{s}^{-1}$ |              |

(1) A vector quantity,  $\mathbf{q}_m = (dm/dt) \mathbf{e}$ , where  $\mathbf{e}$  is the unit vector in the direction of the flow, can be defined, sometimes called the “flow rate” of  $m$ . Similar definitions apply to any extensive quantity.

(2) The flux density of molecules,  $J_N$ , determines either the rate at which the surface would be covered if each molecule stuck, or the rate of effusion through a hole in the surface. In studying the exposure,  $\int J_N dt$ , of a surface to a gas, surface scientists find it useful to use the product of pressure and time as a measure of the exposure, since this product is equal to the number flux density,  $J_N$ , times the time  $J_N t = C(\bar{u}/4)pt$ , where  $C$  is the number density of molecules and  $\bar{u}$  their average speed. The unit langmuir (symbol L) corresponds to the exposure of a surface to a gas at  $10^{-6}$  Torr for 1 s.

(3) In previous editions, the term “flux” was used for what is now called “flux density”, in agreement with IUPAP [4]. The term “density” in the name of an intensive physical quantity usually implies “extensive quantity divided by volume” for scalar quantities but “extensive quantity divided by area” for vector quantities denoting flow or flux.

(4)  $c$  is the amount concentration.

(5) See also Section 2.2, p. 15;  $\tau$  is the shear stress tensor.

### 2.15.1 Transport characteristic numbers: Quantities of dimension one

| <i>Name</i>                         | <i>Symbol</i> | <i>Definition</i>                                                                                                | <i>Notes</i> |
|-------------------------------------|---------------|------------------------------------------------------------------------------------------------------------------|--------------|
| Reynolds number                     | $Re$          | $Re = \rho v l / \eta$                                                                                           |              |
| Euler number                        | $Eu$          | $Eu = \Delta p / \rho v^2$                                                                                       |              |
| Froude number                       | $Fr$          | $Fr = v / (lg)^{1/2}$                                                                                            |              |
| Grashof number                      | $Gr$          | $Gr = l^3 g \alpha \Delta T \rho^2 / \eta^2$                                                                     |              |
| Weber number                        | $We$          | $We = \rho v^2 l / \gamma$                                                                                       |              |
| Mach number                         | $Ma$          | $Ma = v / c$                                                                                                     |              |
| Knudsen number                      | $Kn$          | $Kn = \lambda / l$                                                                                               |              |
| Strouhal number                     | $Sr$          | $Sr = l f / v$                                                                                                   |              |
| Fourier number                      | $Fo$          | $Fo = at / l^2$                                                                                                  |              |
| Péclet number                       | $Pe$          | $Pe = vl / a$                                                                                                    |              |
| Rayleigh number                     | $Ra$          | $Ra = l^3 g \alpha \Delta T \rho / \eta a$                                                                       |              |
| Nusselt number                      | $Nu$          | $Nu = hl / k$                                                                                                    |              |
| Stanton number                      | $St$          | $St = h / \rho v c_p$                                                                                            |              |
| Fourier number for<br>mass transfer | $Fo^*$        | $Fo^* = Dt / l^2$                                                                                                | 1            |
| Péclet number for<br>mass transfer  | $Pe^*$        | $Pe^* = vl / D$                                                                                                  | 1            |
| Grashof number for<br>mass transfer | $Gr^*$        | $Gr^* = l^3 g \left( \frac{\partial \rho}{\partial x} \right)_{T,p} \left( \frac{\Delta x \rho}{\eta^2} \right)$ | 1            |
| Nusselt number for<br>mass transfer | $Nu^*$        | $Nu^* = k_d l / D$                                                                                               | 1, 2         |
| Stanton number for<br>mass transfer | $St^*$        | $St^* = k_d / v$                                                                                                 | 1            |
| Prandtl number                      | $Pr$          | $Pr = \eta / \rho a$                                                                                             |              |
| Schmidt number                      | $Sc$          | $Sc = \eta / \rho D$                                                                                             |              |
| Lewis number                        | $Le$          | $Le = a / D$                                                                                                     |              |
| magnetic Reynolds number            | $Rm, Re_m$    | $Rm = v \mu \kappa l$                                                                                            |              |
| Alfvén number                       | $Al$          | $Al = v(\rho \mu)^{1/2} / B$                                                                                     |              |
| Hartmann number                     | $Ha$          | $Ha = Bl(\kappa / \eta)^{1/2}$                                                                                   |              |
| Cowling number                      | $Co$          | $Co = B^2 / \mu \rho v^2$                                                                                        |              |

(1) This quantity applies to the transport of matter in binary mixtures [72].

(2) The name Sherwood number and symbol  $Sh$  have been widely used for this quantity.

### **3 DEFINITIONS AND SYMBOLS FOR UNITS**



### 3.1 THE INTERNATIONAL SYSTEM OF UNITS (SI)

The International System of Units (SI) was adopted by the 11th General Conference of Weights and Measures (CGPM) in 1960 [3]. It is a coherent system of units built from seven *SI base units*, one for each of the seven dimensionally independent base quantities (see Section 1.2, p. 4); they are: metre, kilogram, second, ampere, kelvin, mole, and candela for the base quantities length, mass, time, electric current, thermodynamic temperature, amount of substance, and luminous intensity, respectively. The definitions of the SI base units are given in Section 3.3, p. 87. The *SI derived units* are expressed as products of powers of the base units, analogous to the corresponding relations between physical quantities. If no numerical factors other than 1 are used in the defining equations for derived units, then the derived units defined in this way are called *coherent derived units*. The SI base units and the derived units without any multiple or sub-multiple prefixes form a coherent set of units, and are called the *coherent SI units*.

In the International System of Units there is only one coherent SI unit for each physical quantity. This is either the appropriate SI base unit itself (see Section 3.2, p. 86) or the appropriate SI derived unit (see sections 3.4 and 3.5, p. 89 and 90). However, any of the approved decimal prefixes, called *SI prefixes*, may be used to construct decimal multiples or submultiples of SI units (see Section 3.6, p. 91). The SI units and the decimal multiples and submultiples constructed with the SI prefixes are called the complete set of SI units, or simply the SI units, or the units of the SI.

It is recommended that only units of the SI be used in science and technology (with SI prefixes where appropriate). Where there are special reasons for making an exception to this rule, it is recommended always to define the units in terms of SI units.

### 3.2 NAMES AND SYMBOLS FOR THE SI BASE UNITS

The symbols listed here are internationally agreed and shall not be changed in other languages or scripts. See sections 1.3 and 1.4, p. 5 and p. 6 on the printing of symbols for units.

| <i>Base quantity</i>      | <i>SI base unit</i> |               |
|---------------------------|---------------------|---------------|
|                           | <i>Name</i>         | <i>Symbol</i> |
| length                    | metre               | m             |
| mass                      | kilogram            | kg            |
| time                      | second              | s             |
| electric current          | ampere              | A             |
| thermodynamic temperature | kelvin              | K             |
| amount of substance       | mole                | mol           |
| luminous intensity        | candela             | cd            |

### 3.3 DEFINITIONS OF THE SI BASE UNITS

The following definitions of the seven SI base units are adopted by the General Conference on Weights and Measures (CGPM) [3].

**metre** (symbol: m)

**The metre is the length of path traveled by light in vacuum during a time interval of  $1/299\,792\,458$  of a second.**

(17<sup>th</sup> CGPM, 1983)

**kilogram** (symbol: kg)

**The kilogram is the unit of mass; it is equal to the mass of the international prototype of the kilogram.**

(3<sup>rd</sup> CGPM, 1901) <sup>1</sup>

**second** (symbol: s)

**The second is the duration of 9 192 631 770 periods of the radiation corresponding to the transition between the two hyperfine levels of the ground state of the caesium 133 atom.**

(13<sup>th</sup> CGPM, 1967)

This definition refers to a caesium atom at rest at a temperature of 0 K.

(CIPM, 1997)

In this definition it is understood that the Cs atom at a temperature of  $T = 0$  K is unperturbed by black-body radiation. The frequency of primary frequency standards should therefore be corrected for the frequency shift due to the ambient radiation, as stated by the Consultative Committee for Time and Frequency (CCTF, 1999).

**ampere** (symbol: A)

**The ampere is that constant current which, if maintained in two straight parallel conductors of infinite length, of negligible circular cross-section, and placed 1 metre apart in vacuum, would produce between these conductors a force equal to  $2 \times 10^{-7}$  newton per metre of length.**

(9<sup>th</sup> CGPM, 1948)

**kelvin** (symbol: K)

**The kelvin, unit of thermodynamic temperature, is the fraction  $1/273.16$  of the thermodynamic temperature of the triple point of water.**

(13<sup>th</sup> CGPM, 1967)

This definition refers to water having the isotopic composition defined exactly by the following amount-of-substance ratios: 0.000 155 76 mole of  $^2\text{H}$  per mole of  $^1\text{H}$ , 0.000 379 9 mole of  $^{17}\text{O}$  per mole of  $^{16}\text{O}$ , and 0.002 005 2 mole of  $^{18}\text{O}$  per mole of  $^{16}\text{O}$ .

(CIPM, 2005) <sup>2</sup>

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<sup>1</sup> The kilogram is the only base unit which is not defined by a measurement on a system defined by natural microscopic constants or an experimental setup derived from such a system. Rather it is defined by a human artefact (the international prototype of the kilogram). Therefore, alternative definitions of the kilogram are under current discussion [127–129].

<sup>2</sup> See also Section 2.11, note 2, p. 56.

**mole** (symbol: mol)

**1. The mole is the amount of substance of a system which contains as many elementary entities as there are atoms in 0.012 kilogram of carbon 12; its symbol is “mol”.**

**2. When the mole is used, the elementary entities must be specified and may be atoms, molecules, ions, electrons, other particles, or specified groups of such particles.**

(14<sup>th</sup> CGPM, 1971)

In this definition, it is understood that unbound atoms of carbon 12, at rest and in their ground state, are referred to.

(CIPM, 1980)

*Examples of the use of the mole*

1 mol of  $^1\text{H}_2$  contains about  $6.022 \times 10^{23}$   $^1\text{H}_2$  molecules, or  $12.044 \times 10^{23}$   $^1\text{H}$  atoms

1 mol of HgCl has a mass of 236.04 g

1 mol of  $\text{Hg}_2\text{Cl}_2$  has a mass of 472.09 g

1 mol of  $\text{Hg}_2^{2+}$  has a mass of 401.18 g and a charge of 192.97 kC

1 mol of  $\text{Fe}_{0.91}\text{S}$  has a mass of 82.88 g

1 mol of  $\text{e}^-$  has a mass of 548.58  $\mu\text{g}$  and a charge of  $-96.49$  kC

1 mol of photons whose frequency is  $5 \times 10^{14}$  Hz

has an energy of about 199.5 kJ

Specification of the entity does not imply that the entities are identical: one may have 1 mol of an isotope mixture or gas mixture.

**candela** (symbol: cd)

**The candela is the luminous intensity, in a given direction, of a source that emits monochromatic radiation of frequency  $540 \times 10^{12}$  hertz and that has a radiant intensity in that direction of 1/683 watt per steradian.**

(16<sup>th</sup> CGPM, 1979)

### 3.4 SI DERIVED UNITS WITH SPECIAL NAMES AND SYMBOLS

| Derived quantity                                                | Name           | Symbol             | SI derived unit                      |                                                        | Notes |
|-----------------------------------------------------------------|----------------|--------------------|--------------------------------------|--------------------------------------------------------|-------|
|                                                                 |                |                    | Expressed in terms of other SI units |                                                        |       |
| plane angle                                                     | radian         | rad                | $\text{m m}^{-1}$                    | = 1                                                    | 1     |
| solid angle                                                     | steradian      | sr                 | $\text{m}^2 \text{m}^{-2}$           | = 1                                                    | 1     |
| frequency                                                       | hertz          | Hz                 | $\text{s}^{-1}$                      |                                                        | 2     |
| force                                                           | newton         | N                  | $\text{m kg s}^{-2}$                 |                                                        |       |
| pressure, stress                                                | pascal         | Pa                 | $\text{N m}^{-2}$                    | = $\text{m}^{-1} \text{kg s}^{-2}$                     |       |
| energy, work, heat                                              | joule          | J                  | $\text{N m}$                         | = $\text{m}^2 \text{kg s}^{-2}$                        |       |
| power, radiant flux                                             | watt           | W                  | $\text{J s}^{-1}$                    | = $\text{m}^2 \text{kg s}^{-3}$                        |       |
| electric charge                                                 | coulomb        | C                  | $\text{A s}$                         |                                                        |       |
| electric potential,<br>electromotive force,<br>electric tension | volt           | V                  | $\text{J C}^{-1}$                    | = $\text{m}^2 \text{kg s}^{-3} \text{A}^{-1}$          |       |
| electric resistance                                             | ohm            | $\Omega$           | $\text{V A}^{-1}$                    | = $\text{m}^2 \text{kg s}^{-3} \text{A}^{-2}$          |       |
| electric conductance                                            | siemens        | S                  | $\Omega^{-1}$                        | = $\text{m}^{-2} \text{kg}^{-1} \text{s}^3 \text{A}^2$ |       |
| electric capacitance                                            | farad          | F                  | $\text{C V}^{-1}$                    | = $\text{m}^{-2} \text{kg}^{-1} \text{s}^4 \text{A}^2$ |       |
| magnetic flux                                                   | weber          | Wb                 | $\text{V s}$                         | = $\text{m}^2 \text{kg s}^{-2} \text{A}^{-1}$          |       |
| magnetic flux density                                           | tesla          | T                  | $\text{Wb m}^{-2}$                   | = $\text{kg s}^{-2} \text{A}^{-1}$                     |       |
| inductance                                                      | henry          | H                  | $\text{V A}^{-1} \text{s}$           | = $\text{m}^2 \text{kg s}^{-2} \text{A}^{-2}$          |       |
| Celsius temperature                                             | degree Celsius | $^{\circ}\text{C}$ |                                      |                                                        | 3     |
| luminous flux                                                   | lumen          | lm                 | $\text{cd sr}$                       | = cd                                                   |       |
| illuminance                                                     | lux            | lx                 | $\text{lm m}^{-2}$                   | = $\text{cd m}^{-2}$                                   |       |
| activity, (radioactivity)<br>referred to a radio-<br>nuclide    | becquerel      | Bq                 | $\text{s}^{-1}$                      |                                                        | 4     |
| absorbed dose,<br>kerma                                         | gray           | Gy                 | $\text{J kg}^{-1}$                   | = $\text{m}^2 \text{s}^{-2}$                           | 4     |
| dose equivalent<br>(dose equivalent index)                      | sievert        | Sv                 | $\text{J kg}^{-1}$                   | = $\text{m}^2 \text{s}^{-2}$                           | 4     |
| catalytic activity                                              | katal          | kat                | $\text{mol s}^{-1}$                  |                                                        | 4, 5  |

(1) Radian and steradian are derived units. Since they are then of dimension 1, this leaves open the possibility of including them or omitting them in expressions of SI derived units. In practice this means that rad and sr may be used when appropriate and may be omitted if clarity is not lost.

(2) For angular frequency and for angular velocity the unit  $\text{rad s}^{-1}$ , or simply  $\text{s}^{-1}$ , should be used, and this may *not* be replaced with Hz. The unit Hz shall be used *only* for frequency in the sense of cycles per second.

(3) The Celsius temperature  $t$  is defined by the equation

$$t/^{\circ}\text{C} = T/\text{K} - 273.15$$

The SI unit of Celsius temperature is the degree Celsius,  $^{\circ}\text{C}$ , which is equal to the kelvin, K.  $^{\circ}\text{C}$  shall be treated as a single symbol, with no space between the  $^{\circ}$  sign and the C. The symbol  $^{\circ}\text{K}$ , and the symbol  $^{\circ}$ , shall no longer be used for the unit of thermodynamic temperature.

(4) Becquerel is the basic unit to be used in nuclear and radiochemistry; becquerel, gray, and sievert are admitted for reasons of safeguarding human health [3].

(5) When the amount of a catalyst cannot be expressed as a number of elementary entities, an amount of substance, or a mass, a “catalytic activity” can still be defined as a property of the catalyst measured by a catalyzed rate of conversion under specified optimized conditions. The katal,  $1 \text{ kat} = 1 \text{ mol s}^{-1}$ , should replace the “(enzyme) unit”,  $1 \text{ U} = \mu\text{mol min}^{-1} \approx 16.67 \text{ nkat}$  [130].

### 3.5 SI DERIVED UNITS FOR OTHER QUANTITIES

This table gives examples of other SI derived units; the list is merely illustrative.

| SI derived unit          |                                     |                                                                       |       |
|--------------------------|-------------------------------------|-----------------------------------------------------------------------|-------|
| Derived quantity         | Symbol                              | Expressed in terms of SI base units                                   | Notes |
| efficiency               | W W <sup>-1</sup>                   | = 1                                                                   |       |
| area                     | m <sup>2</sup>                      |                                                                       |       |
| volume                   | m <sup>3</sup>                      |                                                                       |       |
| speed, velocity          | m s <sup>-1</sup>                   |                                                                       |       |
| angular velocity         | rad s <sup>-1</sup>                 | = s <sup>-1</sup>                                                     |       |
| acceleration             | m s <sup>-2</sup>                   |                                                                       |       |
| moment of force          | N m                                 | = m <sup>2</sup> kg s <sup>-2</sup>                                   |       |
| repetency, wavenumber    | m <sup>-1</sup>                     |                                                                       | 1     |
| density, mass density    | kg m <sup>-3</sup>                  |                                                                       |       |
| specific volume          | m <sup>3</sup> kg <sup>-1</sup>     |                                                                       |       |
| amount concentration     | mol m <sup>-3</sup>                 |                                                                       | 2     |
| molar volume             | m <sup>3</sup> mol <sup>-1</sup>    |                                                                       |       |
| heat capacity, entropy   | J K <sup>-1</sup>                   | = m <sup>2</sup> kg s <sup>-2</sup> K <sup>-1</sup>                   |       |
| molar heat capacity,     | J K <sup>-1</sup> mol <sup>-1</sup> | = m <sup>2</sup> kg s <sup>-2</sup> K <sup>-1</sup> mol <sup>-1</sup> |       |
| molar entropy            |                                     |                                                                       |       |
| specific heat capacity,  | J K <sup>-1</sup> kg <sup>-1</sup>  | = m <sup>2</sup> s <sup>-2</sup> K <sup>-1</sup>                      |       |
| specific entropy         |                                     |                                                                       |       |
| molar energy             | J mol <sup>-1</sup>                 | = m <sup>2</sup> kg s <sup>-2</sup> mol <sup>-1</sup>                 |       |
| specific energy          | J kg <sup>-1</sup>                  | = m <sup>2</sup> s <sup>-2</sup>                                      |       |
| energy density           | J m <sup>-3</sup>                   | = m <sup>-1</sup> kg s <sup>-2</sup>                                  |       |
| surface tension          | N m <sup>-1</sup>                   | = kg s <sup>-2</sup>                                                  |       |
| heat flux density,       | W m <sup>-2</sup>                   | = kg s <sup>-3</sup>                                                  |       |
| irradiance               |                                     |                                                                       |       |
| thermal conductivity     | W m <sup>-1</sup> K <sup>-1</sup>   | = m kg s <sup>-3</sup> K <sup>-1</sup>                                |       |
| kinematic viscosity,     | m <sup>2</sup> s <sup>-1</sup>      |                                                                       |       |
| diffusion coefficient    |                                     |                                                                       |       |
| dynamic viscosity,       | Pa s                                | = m <sup>-1</sup> kg s <sup>-1</sup>                                  |       |
| shear viscosity          |                                     |                                                                       |       |
| electric charge density  | C m <sup>-3</sup>                   | = m <sup>-3</sup> s A                                                 |       |
| electric current density | A m <sup>-2</sup>                   |                                                                       |       |
| conductivity             | S m <sup>-1</sup>                   | = m <sup>-3</sup> kg <sup>-1</sup> s <sup>3</sup> A <sup>2</sup>      |       |
| molar conductivity       | S m <sup>2</sup> mol <sup>-1</sup>  | = kg <sup>-1</sup> s <sup>3</sup> A <sup>2</sup> mol <sup>-1</sup>    |       |
| permittivity             | F m <sup>-1</sup>                   | = m <sup>-3</sup> kg <sup>-1</sup> s <sup>4</sup> A <sup>2</sup>      |       |
| permeability             | H m <sup>-1</sup>                   | = m kg s <sup>-2</sup> A <sup>-2</sup>                                |       |
| electric field strength  | V m <sup>-1</sup>                   | = m kg s <sup>-3</sup> A <sup>-1</sup>                                |       |
| magnetic field strength  | A m <sup>-1</sup>                   |                                                                       |       |
| exposure (X and γ rays)  | C kg <sup>-1</sup>                  | = kg <sup>-1</sup> s A                                                |       |
| absorbed dose rate       | Gy s <sup>-1</sup>                  | = m <sup>2</sup> s <sup>-3</sup>                                      |       |

(1) The word “wavenumber” denotes the quantity “reciprocal wavelength”. Its widespread use to denote the unit cm<sup>-1</sup> should be discouraged.

(2) The words “amount concentration” are an abbreviation for “amount-of-substance concentration”. When there is not likely to be any ambiguity this quantity may be called simply “concentration”.

### 3.6 SI PREFIXES AND PREFIXES FOR BINARY MULTIPLES

The following prefixes [3] are used to denote decimal multiples and submultiples of SI units.

| <i>Submultiple</i> | <i>Prefix</i> |               | <i>Multiple</i> | <i>Prefix</i> |               |
|--------------------|---------------|---------------|-----------------|---------------|---------------|
|                    | <i>Name</i>   | <i>Symbol</i> |                 | <i>Name</i>   | <i>Symbol</i> |
| $10^{-1}$          | deci          | d             | $10^1$          | deca          | da            |
| $10^{-2}$          | centi         | c             | $10^2$          | hecto         | h             |
| $10^{-3}$          | milli         | m             | $10^3$          | kilo          | k             |
| $10^{-6}$          | micro         | $\mu$         | $10^6$          | mega          | M             |
| $10^{-9}$          | nano          | n             | $10^9$          | giga          | G             |
| $10^{-12}$         | pico          | p             | $10^{12}$       | tera          | T             |
| $10^{-15}$         | femto         | f             | $10^{15}$       | peta          | P             |
| $10^{-18}$         | atto          | a             | $10^{18}$       | exa           | E             |
| $10^{-21}$         | zepto         | z             | $10^{21}$       | zetta         | Z             |
| $10^{-24}$         | yocto         | y             | $10^{24}$       | yotta         | Y             |

Prefix symbols shall be printed in Roman (upright) type with no space between the prefix and the unit symbol.

*Example* kilometre, km

When a prefix is used with a unit symbol, the combination is taken as a new symbol that can be raised to any power without the use of parentheses.

*Examples*  $1 \text{ cm}^3 = (10^{-2} \text{ m})^3 = 10^{-6} \text{ m}^3$   
 $1 \mu\text{s}^{-1} = (10^{-6} \text{ s})^{-1} = 10^6 \text{ s}^{-1}$   
 $1 \text{ V/cm} = 1 \text{ V}/(10^{-2} \text{ m}) = 10^2 \text{ V/m}$   
 $1 \text{ mmol/dm}^3 = 10^{-3} \text{ mol}/(10^{-3} \text{ m}^3) = 1 \text{ mol m}^{-3}$

A prefix shall never be used on its own, and prefixes are not to be combined into compound prefixes.

*Example* pm, not  $\mu\mu\text{m}$

The names and symbols of decimal multiples and submultiples of the SI base unit of mass, the kilogram, symbol kg, which already contains a prefix, are constructed by adding the appropriate prefix to the name gram and symbol g.

*Examples* mg, not  $\mu\text{kg}$ ; Mg, not kkg

The International Electrotechnical Commission (IEC) has standardized the following prefixes for binary multiples, mainly used in information technology, to be distinguished from the SI prefixes for decimal multiples [7].

| <i>Multiple</i>         | <i>Prefix</i> |               | <i>Origin</i> |
|-------------------------|---------------|---------------|---------------|
|                         | <i>Name</i>   | <i>Symbol</i> |               |
| $(2^{10})^1 = (1024)^1$ | kibi          | Ki            | kilobinary    |
| $(2^{10})^2 = (1024)^2$ | mebi          | Mi            | megabinary    |
| $(2^{10})^3 = (1024)^3$ | gibi          | Gi            | gigabinary    |
| $(2^{10})^4 = (1024)^4$ | tebi          | Ti            | terabinary    |
| $(2^{10})^5 = (1024)^5$ | pebi          | Pi            | petabinary    |
| $(2^{10})^6 = (1024)^6$ | exbi          | Ei            | exabinary     |
| $(2^{10})^7 = (1024)^7$ | zebi          | Zi            | zettabinary   |
| $(2^{10})^8 = (1024)^8$ | yobi          | Yi            | yottabinary   |

### 3.7 NON-SI UNITS ACCEPTED FOR USE WITH THE SI

The following units are not part of the SI, but it is recognized by the CGPM [3] that they will continue to be used in appropriate contexts. SI prefixes may be attached to some of these units, such as millilitre, mL; megaelectronvolt, MeV; kilotonne, kt. A more extensive list of non-SI units, with conversion factors to the corresponding SI units, is given in Chapter 7, p. 129.

| <i>Physical quantity</i>                                | <i>Unit accepted for use with the SI</i> |                                               |                                                       | <i>Notes</i> |
|---------------------------------------------------------|------------------------------------------|-----------------------------------------------|-------------------------------------------------------|--------------|
|                                                         | <i>Name</i>                              | <i>Symbol</i>                                 | <i>Value in SI units</i>                              |              |
| time                                                    | minute                                   | min                                           | = 60 s                                                |              |
| time                                                    | hour                                     | h                                             | = 3600 s                                              |              |
| time                                                    | day                                      | d                                             | = 86 400 s                                            |              |
| plane angle                                             | degree                                   | °, deg                                        | = ( $\pi/180$ ) rad                                   |              |
| plane angle                                             | minute                                   | '                                             | = ( $\pi/10\,800$ ) rad                               |              |
| plane angle                                             | second                                   | "                                             | = ( $\pi/648\,000$ ) rad                              |              |
| volume                                                  | litre                                    | l, L                                          | = 1 dm <sup>3</sup> = 10 <sup>-3</sup> m <sup>3</sup> | 1            |
| mass                                                    | tonne                                    | t                                             | = 1 Mg = 10 <sup>3</sup> kg                           |              |
| level of a field quantity,<br>level of a power quantity | neper                                    | Np                                            | = ln e = (1/2) ln e <sup>2</sup> = 1                  | 2            |
| level of a field quantity,<br>level of a power quantity | bel                                      | B                                             |                                                       | 2            |
| energy                                                  | electronvolt                             | eV (= $e \cdot 1\text{ V}$ )                  | = 1.602 176 487(40) × 10 <sup>-19</sup> J             | 3            |
| mass                                                    | dalton,<br>unified atomic<br>mass unit   | Da<br>u (= $m_{\text{a}}(^{12}\text{C})/12$ ) | = 1.660 538 782(83) × 10 <sup>-27</sup> kg<br>= 1 Da  | 3, 4         |
| length                                                  | nautical mile                            | M                                             | = 1852 m                                              | 5            |
|                                                         | astronomical<br>unit                     | ua                                            | = 1.495 978 706 91(6) × 10 <sup>11</sup> m            | 6            |

(1) The alternative symbol L is the only exception of the general rule that symbols for units shall be printed in lower case letters unless they are derived from a personal name. In order to avoid the risk of confusion between the letter l and the number 1, the use of L is accepted. However, only the lower case l is used by ISO and IEC.

(2) For logarithmic ratio quantities and their units, see [131].

(3) The values of these units in terms of the corresponding SI units are not exact, since they depend on the values of the physical constants  $e$  (for electronvolt) and  $m_{\text{a}}(^{12}\text{C})$  or  $N_{\text{A}}$  (for the unified atomic mass unit), which are determined by experiment, see Chapter 5, p. 111.

(4) The dalton, with symbol Da, and the unified atomic mass unit, with symbol u, are alternative names for the same unit. The dalton may be combined with SI prefixes to express the masses of large or small entities.

(5) There is no agreed symbol for the nautical mile. The SI Brochure uses the symbol M.

(6) The astronomical unit is a unit of length approximately equal to the mean Earth-Sun distance. Its value is such that, when used to describe the motion of bodies in the Solar System, the heliocentric gravitational constant is (0.017 202 098 95)<sup>2</sup> ua<sup>3</sup> d<sup>-2</sup> (see also [3]).

### 3.8 COHERENT UNITS AND CHECKING DIMENSIONS

If equations between numerical values have the same form as equations between physical quantities, then the system of units defined in terms of base units avoids numerical factors between units, and is said to be a *coherent system*. For example, the kinetic energy  $T$  of a particle of mass  $m$  moving with a speed  $v$  is defined by the equation

$$T = (1/2) mv^2$$

but the SI unit of kinetic energy is the joule, defined by the equation

$$J = \text{kg (m/s)}^2 = \text{kg m}^2 \text{ s}^{-2}$$

where it is to be noted that the factor  $(1/2)$  is omitted. In fact the joule, symbol J, is simply a special name and symbol for the product of units  $\text{kg m}^2 \text{ s}^{-2}$ .

The International System (SI) is a coherent system of units. The advantage of a coherent system of units is that if the value of each quantity is substituted for the quantity symbol in any quantity equation, then the units may be canceled, leaving an equation between numerical values which is exactly similar (including all numerical factors) to the original equation between the quantities. Checking that the units cancel in this way is sometimes described as checking the dimensions of the equation.

The use of a coherent system of units is not essential. In particular the use of multiple or submultiple prefixes destroys the coherence of the SI, but is nonetheless often convenient.

### 3.9 FUNDAMENTAL PHYSICAL CONSTANTS USED AS UNITS

Sometimes fundamental physical constants, or other well defined physical quantities, are used as though they were units in certain specialized fields of science. For example, in astronomy it may be more convenient to express the mass of a star in terms of the mass of the sun, and to express the period of the planets in their orbits in terms of the period of the earth's orbit, rather than to use SI units. In atomic and molecular physics it is similarly more convenient to express masses in terms of the electron mass,  $m_e$ , or in terms of the unified atomic mass unit,  $1 \text{ u} = m_{\text{u}} = m(^{12}\text{C}/12)$ , and to express charges in terms of the elementary charge  $e$ , rather than to use SI units. One reason for using such physical quantities as though they were units is that the nature of the experimental measurements or calculations in the specialized field may be such that results are naturally obtained in such terms, and can only be converted to SI units at a later stage. When physical quantities are used as units in this way their relation to the SI must be determined by experiment, which is subject to uncertainty, and the conversion factor may change as new and more precise experiments are developed. Another reason for using such units is that uncertainty in the conversion factor to the SI may be greater than the uncertainty in the ratio of the measurements expressed in terms of the physical constant as a unit. Both reasons make it preferable to present experimental results without converting to SI units.

Three such physical quantities that have been recognized as units by the CIPM are the electronvolt (eV), the dalton (Da) or the unified atomic mass unit (u), and the astronomical unit (ua), listed below [3]. The electronvolt is the product of a fundamental constant (the elementary charge,  $e$ ) and the SI unit of potential difference (the volt, V). The dalton is related to the mass of the carbon-12 nuclide, and is thus a fundamental constant. The astronomical unit is a more arbitrarily defined constant that is convenient to astronomers. However, there are many other physical quantities or fundamental constants that are sometimes used in this way as though they were units, so that it is hardly possible to list them all.

| <i>Physical quantity</i> | <i>Name of unit</i>                 | <i>Symbol for unit</i> | <i>Value in SI units</i>                                         | <i>Notes</i> |
|--------------------------|-------------------------------------|------------------------|------------------------------------------------------------------|--------------|
| energy                   | electronvolt                        | eV                     | $1 \text{ eV} = 1.602\,176\,487(40) \times 10^{-19} \text{ J}$   | 1            |
| mass                     | dalton,<br>unified atomic mass unit | Da, u                  | $1 \text{ Da} = 1.660\,538\,782(83) \times 10^{-27} \text{ kg}$  | 2            |
| length                   | astronomical unit                   | ua                     | $1 \text{ ua} = 1.495\,978\,706\,91(6) \times 10^{11} \text{ m}$ | 3            |

(1) The electronvolt is the kinetic energy acquired by an electron in passing through a potential barrier of 1 V in vacuum.

(2) The dalton and the unified atomic mass unit are alternative names for the same unit. The dalton may be combined with the SI prefixes to express the masses of large molecules in kilodalton (kDa) or megadalton (MDa).

(3) The value of the astronomical unit in SI units is defined such that, when used to describe the motion of bodies in the solar system, the heliocentric gravitational constant is  $(0.017\,202\,098\,95)^2 \text{ ua}^3 \text{ d}^{-2}$ . The value must be obtained from experiment, and is therefore not known exactly (see also [3]).

#### 3.9.1 Atomic units [22] (see also Section 7.3, p. 143)

One particular group of physical constants that are used as though they were units deserve special mention. These are the so-called *atomic units* and arise in calculations of electronic wavefunctions for atoms and molecules, i.e. in quantum chemistry. The first five atomic units in the table below have special names and symbols. Only four of these are independent; all others may be derived by multiplication and division in the usual way, and the table includes a number of examples. The

relation between the five named atomic units may be expressed by any one of the equations

$$E_h = \hbar^2/m_e a_0^2 = e^2/4\pi\epsilon_0 a_0 = m_e e^4/(4\pi\epsilon_0)^2 \hbar^2$$

The relation of atomic units to the corresponding SI units involves the values of the fundamental physical constants, and is therefore not exact. The numerical values in the table are from the CODATA compilation [23] and based on the fundamental constants given in Chapter 5, p. 111. The numerical results of calculations in theoretical chemistry are frequently quoted in atomic units, or as numerical values in the form *physical quantity* divided by *atomic unit*, so that the reader may make the conversion using the current best estimates of the physical constants.

| <i>Physical quantity</i>      | <i>Name of unit</i>                  | <i>Symbol for unit</i> | <i>Value in SI units</i>                                                          | <i>Notes</i> |
|-------------------------------|--------------------------------------|------------------------|-----------------------------------------------------------------------------------|--------------|
| mass                          | electron mass                        | $m_e$                  | $= 9.109\,382\,15(45) \times 10^{-31} \text{ kg}$                                 |              |
| charge                        | elementary charge                    | $e$                    | $= 1.602\,176\,487(40) \times 10^{-19} \text{ C}$                                 |              |
| action,<br>(angular momentum) | Planck constant<br>divided by $2\pi$ | $\hbar$                | $= 1.054\,571\,628(53) \times 10^{-34} \text{ J s}$                               | 1            |
| length                        | Bohr radius                          | $a_0$                  | $= 5.291\,772\,085\,9(36) \times 10^{-11} \text{ m}$                              | 1            |
| energy                        | Hartree energy                       | $E_h$                  | $= 4.359\,743\,94(22) \times 10^{-18} \text{ J}$                                  | 1            |
| time                          |                                      | $\hbar/E_h$            | $= 2.418\,884\,326\,505(16) \times 10^{-17} \text{ s}$                            |              |
| speed                         |                                      | $a_0 E_h/\hbar$        | $= 2.187\,691\,254\,1(15) \times 10^6 \text{ m s}^{-1}$                           | 2            |
| force                         |                                      | $E_h/a_0$              | $= 8.238\,722\,06(41) \times 10^{-8} \text{ N}$                                   |              |
| linear momentum               |                                      | $\hbar/a_0$            | $= 1.992\,851\,565(99) \times 10^{-24} \text{ N s}$                               |              |
| electric current              |                                      | $e E_h/\hbar$          | $= 6.623\,617\,63(17) \times 10^{-3} \text{ A}$                                   |              |
| electric field                |                                      | $E_h/ea_0$             | $= 5.142\,206\,32(13) \times 10^{11} \text{ V m}^{-1}$                            |              |
| electric dipole<br>moment     |                                      | $ea_0$                 | $= 8.478\,352\,81(21) \times 10^{-30} \text{ C m}$                                |              |
| electric quadrupole<br>moment |                                      | $ea_0^2$               | $= 4.486\,551\,07(11) \times 10^{-40} \text{ C m}^2$                              |              |
| electric polarizability       |                                      | $e^2 a_0^2/E_h$        | $= 1.648\,777\,253\,6(34) \times 10^{-41} \text{ C}^2 \text{ m}^2 \text{ J}^{-1}$ |              |
| 1st hyper-polarizability      |                                      | $e^3 a_0^3/E_h^2$      | $= 3.206\,361\,533(81) \times 10^{-53} \text{ C}^3 \text{ m}^3 \text{ J}^{-2}$    |              |
| 2nd hyper-polarizability      |                                      | $e^4 a_0^4/E_h^3$      | $= 6.235\,380\,95(31) \times 10^{-65} \text{ C}^4 \text{ m}^4 \text{ J}^{-3}$     |              |
| magnetic flux<br>density      |                                      | $\hbar/ea_0^2$         | $= 2.350\,517\,382(59) \times 10^5 \text{ T}$                                     |              |
| magnetic dipole<br>moment     |                                      | $e\hbar/m_e$           | $= 1.854\,801\,830(46) \times 10^{-23} \text{ J T}^{-1}$                          | 3            |
| magnetizability               |                                      | $e^2 a_0^2/m_e$        | $= 7.891\,036\,433(27) \times 10^{-29} \text{ J T}^{-2}$                          |              |

(1)  $\hbar = h/2\pi$ ;  $a_0 = 4\pi\epsilon_0 \hbar^2/m_e e^2$ ;  $E_h = \hbar^2/m_e a_0^2$ .

(2) The numerical value of the speed of light, when expressed in atomic units, is equal to the reciprocal of the fine-structure constant  $\alpha$ ;

$c/(\text{au of speed}) = c\hbar/a_0 E_h = \alpha^{-1} = 137.035\,999\,679(94)$ .

(3) The atomic unit of magnetic dipole moment is twice the Bohr magneton,  $\mu_B$ .

### 3.9.2 The equations of quantum chemistry expressed in terms of reduced quantities using atomic units

It is customary to write the equations of quantum chemistry in terms of reduced quantities. Thus energies are expressed as reduced energies  $E^*$ , distances as reduced distances  $r^*$ , masses as reduced masses  $m^*$ , charges as reduced charges  $Q^*$ , and angular momenta as reduced angular momenta  $J^*$ , where the reduced quantities are given by the equations

$$E^* = E/E_h, \quad r^* = r/a_0, \quad m^* = m/m_e, \quad Q^* = Q/e, \quad \text{and} \quad J^* = J/\hbar \quad (1)$$

The reduced quantity in each case is the dimensionless ratio of the actual quantity to the corresponding atomic unit. The advantage of expressing all the equations in terms of reduced quantities is that the equations are simplified since all the physical constants disappear from the equations (although this simplification is achieved at the expense of losing the advantage of dimensional checking, since all reduced quantities are dimensionless). For example the Schrödinger equation for the hydrogen atom, expressed in the usual physical quantities, has the form

$$-(\hbar^2/2m_e)\nabla_r^2 \psi(r, \theta, \phi) + V(r)\psi(r, \theta, \phi) = E\psi(r, \theta, \phi) \quad (2)$$

Here  $r$ ,  $\theta$ , and  $\phi$  are the coordinates of the electron, and the operator  $\nabla_r$  involves derivatives  $\partial/\partial r$ ,  $\partial/\partial\theta$ , and  $\partial/\partial\phi$ . However in terms of reduced quantities the corresponding equation has the form

$$-(1/2)\nabla_{r^*}^2 \psi(r^*, \theta, \phi) + V^*(r^*)\psi(r^*, \theta, \phi) = E^*\psi(r^*, \theta, \phi) \quad (3)$$

where  $\nabla_{r^*}$  involves derivatives  $\partial/\partial r^*$ ,  $\partial/\partial\theta$ , and  $\partial/\partial\phi$ . This may be shown by substituting the reduced (starred) quantities for the actual (unstarred) quantities in Equation (2) which leads to Equation (3).

In the field of quantum chemistry it is customary to write all equations in terms of reduced (starred) quantities, so that all quantities become dimensionless, and all fundamental constants such as  $e$ ,  $m_e$ ,  $\hbar$ ,  $E_h$ , and  $a_0$  disappear from the equations. As observed above this simplification is achieved at the expense of losing the possibility of dimensional checking. To compare the results of a numerical calculation with experiment it is of course necessary to transform the calculated values of the reduced quantities back to the values of the actual quantities using Equation (1).

Unfortunately it is also customary not to use the star that has been used here, but instead to use exactly the same symbol for the dimensionless reduced quantities and the actual quantities. This makes it impossible to write equations such as (1). (It is analogous to the situation that would arise if we were to use exactly the same symbol for  $h$  and  $\hbar$ , where  $\hbar = h/2\pi$ , thus making it impossible to write the relation between  $h$  and  $\hbar$ .) This may perhaps be excused on the grounds that it becomes tedious to include a star on the symbol for every physical quantity when writing the equations in quantum chemistry, but it is important that readers unfamiliar with the field should realize what has been done. It is also important to realize how the values of quantities “expressed in atomic units”, i.e. the values of reduced quantities, may be converted back to the values of the original quantities in SI units.

It is also customary to make statements such as “in atomic units  $e$ ,  $m_e$ ,  $\hbar$ ,  $E_h$ , and  $a_0$  are all equal to 1”, which is not a correct statement. The correct statement would be that in atomic units the elementary charge is equal to 1  $e$ , the mass of an electron is equal to 1  $m_e$ , etc. The difference between equations such as (3), which contain no fundamental constants, and (2) which do contain fundamental constants, concerns the quantities rather than the units. In (3) all the quantities are dimensionless reduced quantities, defined by (1), whereas in (2) the quantities are the usual (dimensioned) physical quantities with which we are familiar in other circumstances.

Finally, many authors make no use of the symbols for the atomic units listed in the tables above, but instead use the symbol “a.u.” or “au” for all atomic units. This custom should not be followed. It leads to confusion, just as it would if we were to write “SI” as a symbol for every SI unit, or “CGS” as a symbol for every CGS unit.

*Examples* For the hydrogen molecule the equilibrium bond length  $r_e$ , and the dissociation energy  $D_e$ , are given by

|                  |            |                           |
|------------------|------------|---------------------------|
| $r_e = 2.1 a_0$  | <i>not</i> | $r_e = 2.1 \text{ a.u.}$  |
| $D_e = 0.16 E_h$ | <i>not</i> | $D_e = 0.16 \text{ a.u.}$ |

### 3.10 DIMENSIONLESS QUANTITIES

Values of dimensionless physical quantities, more properly called “quantities of dimension one”, are often expressed in terms of mathematically exactly defined values denoted by special symbols or abbreviations, such as % (percent). These symbols are then treated as units, and are used as such in calculations.

#### 3.10.1 Fractions (relative values, yields, and efficiencies)

Fractions such as relative uncertainty, amount-of-substance fraction  $x$  (also called amount fraction), mass fraction  $w$ , and volume fraction  $\varphi$  (see Section 2.10, p. 47 for all these quantities), are sometimes expressed in terms of the symbols in the table below.

| <i>Name</i> | <i>Symbol</i> | <i>Value</i> | <i>Example</i>                                                                                   |
|-------------|---------------|--------------|--------------------------------------------------------------------------------------------------|
| percent     | %             | $10^{-2}$    | The isotopic abundance of carbon-13 expressed as an amount-of-substance fraction is $x = 1.1$ %. |
| permille    | ‰             | $10^{-3}$    | The mass fraction of water in a sample is $w = 2.3$ ‰.                                           |

These multiples of the unit one are not part of the SI and ISO recommends that these symbols should never be used. They are also frequently used as units of “concentration” without a clear indication of the type of fraction implied, e.g. amount-of-substance fraction, mass fraction or volume fraction. To avoid ambiguity they should be used only in a context where the meaning of the quantity is carefully defined. Even then, the use of an appropriate SI unit ratio may be preferred.

*Examples*      The mass fraction  $w = 1.5 \times 10^{-6} = 1.5$  mg/kg.  
The amount-of-substance fraction  $x = 3.7 \times 10^{-2} = 3.7$  % or  $x = 37$  mmol/mol.  
Atomic absorption spectroscopy shows the aqueous solution to contain a mass concentration of nickel  $\rho(\text{Ni}) = 2.6$  mg dm $^{-3}$ , which is approximately equivalent to a mass fraction  $w(\text{Ni}) = 2.6 \times 10^{-6}$ .

Note the importance of using the recommended name and symbol for the quantity in each of the above examples. Statements such as “the concentration of nickel was  $2.6 \times 10^{-6}$ ” are ambiguous and should be avoided.

The last example illustrates the approximate equivalence of  $\rho/\text{mg dm}^{-3}$  and  $w/10^{-6}$  in aqueous solution, which follows from the fact that the mass density of a dilute aqueous solution is always approximately  $1.0$  g cm $^{-3}$ . Dilute solutions are often measured or calibrated to a known mass concentration in mg dm $^{-3}$ , and this unit is then to be preferred to using ppm (or other corresponding abbreviations, which are language dependent) to specify a mass fraction.

#### 3.10.2 Deprecated usage

Adding extra labels to % and similar symbols, such as % (V/V) (meaning % by volume) should be avoided. Qualifying labels may be added to symbols for physical quantities, but never to units.

*Example*      A mass fraction  $w = 0.5$  %, but *not*  $0.5$  % ( $m/m$ ).

The symbol % should not be used in combination with other units. In table headings and in labeling the axes of graphs the use of % in the denominator is to be avoided. Although one would write  $x(^{13}\text{C}) = 1.1$  %, the notation  $100\ x$  is to be preferred to  $x/\%$  in tables and graphs (see for example Section 6.3, column 5, p. 122).

The further symbols listed in the table below are also found in the literature, but their use is not recommended. Note that the names and symbols for  $10^{-9}$  and  $10^{-12}$  in this table are here based on the American system of names. In other parts of the world, a billion often stands for  $10^{12}$  and a trillion for  $10^{18}$ . Note also that the symbol ppt is sometimes used for part per thousand, and sometimes for part per trillion. In 1948 the word billion had been proposed for  $10^{12}$  and trillion for  $10^{18}$  [132]. Although ppm, ppb, ppt and alike are widely used in various applications of analytical and environmental chemistry, it is suggested to abandon completely their use because of the ambiguities involved. These units are unnecessary and can be easily replaced by SI-compatible quantities such as pmol/mol (picomole per mole), which are unambiguous. The last column contains suggested replacements (similar replacements can be formulated as mg/g,  $\mu\text{g/g}$ , pg/g etc.).

| <i>Name</i>                                 | <i>Symbol</i> | <i>Value</i> | <i>Examples</i>                                                                                                                                                                                                    | <i>Replacement</i>  |
|---------------------------------------------|---------------|--------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------|
| part per hundred                            | pph, %        | $10^{-2}$    | The degree of dissociation is 1.5 %.                                                                                                                                                                               |                     |
| part per thousand,<br>permille <sup>1</sup> | ppt, ‰        | $10^{-3}$    | An approximate preindustrial value of the CO <sub>2</sub> content of the Earth's atmosphere was 0.275 ‰ (0.275 ppt).<br>The element Ti has a mass fraction 5.65 ‰ (5.65×10 <sup>3</sup> ppm) in the Earth's crust. | mmol/mol<br>mg/g    |
| part per million                            | ppm           | $10^{-6}$    | The volume fraction of helium is 20 ppm.                                                                                                                                                                           | $\mu\text{mol/mol}$ |
| part per hundred million                    | pphm          | $10^{-8}$    | The mass fraction of impurity in the metal was less than 5 pphm.                                                                                                                                                   |                     |
| part per billion                            | ppb           | $10^{-9}$    | The air quality standard for ozone is a volume fraction of $\varphi = 120$ ppb.                                                                                                                                    | nmol/mol            |
| part per trillion                           | ppt           | $10^{-12}$   | The natural background volume fraction of NO in air was found to be $\varphi = 140$ ppt.                                                                                                                           | pmol/mol            |
| part per quadrillion                        | ppq           | $10^{-15}$   |                                                                                                                                                                                                                    | fmol/mol            |

<sup>1</sup> The permille is also spelled per mill, permill, per mil, permil, per mille, or promille.

### 3.10.3 Units for logarithmic quantities: neper, bel, and decibel

In some fields, especially in acoustics and telecommunications, special names are given to the number 1 when expressing physical quantities defined in terms of the logarithm of a ratio [131]. For a damped linear oscillation the amplitude of a quantity as a function of time is given by

$$F(t) = A e^{-\delta t} \cos \omega t = A \operatorname{Re}[e^{(-\delta + i\omega)t}]$$

From this relation it is clear that the coherent SI unit for the decay coefficient  $\delta$  and the angular frequency  $\omega$  is the second to the power of minus one,  $\text{s}^{-1}$ . However, the special names neper, Np, and radian, rad (see Section 2.1, p. 13, Section 3.4, p. 89, and Section 3.7, p. 92), are used for the units of the dimensionless products  $\delta t$  and  $\omega t$ , respectively. Thus the quantities  $\delta$  and  $\omega$  may be expressed in the units Np/s and rad/s, respectively. Used in this way the neper, Np, and the radian, rad, may both be thought of as special names for the number 1.

In the fields of acoustics and signal transmission, signal power levels and signal amplitude levels (or field level) are usually expressed as the decadic or the napierian logarithm of the ratio of the power  $P$  to a reference power  $P_0$ , or of the field  $F$  to a reference field  $F_0$ . Since power is often proportional to the square of the field or amplitude (when the field acts on equal impedances in linear systems) it is convenient to define the power level and the field level to be equal in such a case. This is done by defining the field level and the power level according to the relations

$$L_F = \ln(F/F_0), \quad \text{and} \quad L_P = (1/2) \ln(P/P_0)$$

so that if  $(P/P_0) = (F/F_0)^2$  then  $L_P = L_F$ . The above equations may be written in the form

$$L_F = \ln(F/F_0) \text{ Np}, \quad \text{and} \quad L_P = (1/2) \ln(P/P_0) \text{ Np}$$

The bel, B, and its more frequently used submultiple the decibel, dB, are used when the field and power levels are calculated using decadic logarithms according to the relations

$$L_P = \lg(P/P_0) \text{ B} = 10 \lg(P/P_0) \text{ dB}$$

and

$$L_F = 2 \lg(F/F_0) \text{ B} = 20 \lg(F/F_0) \text{ dB}$$

The relation between the bel and the neper follows from comparing these equations with the preceeding equations. We obtain

$$L_F = \ln(F/F_0) \text{ Np} = 2 \lg(F/F_0) \text{ B} = \ln(10) \lg(F/F_0) \text{ Np}$$

giving

$$1 \text{ B} = 10 \text{ dB} = (1/2) \ln(10) \text{ Np} \approx 1.151 \text{ 293 Np}$$

In practice the bel is hardly ever used. Only the decibel is used, to represent the decadic logarithm, particularly in the context of acoustics, and in labeling the controls of power amplifiers. Thus the statement  $L_P = n \text{ dB}$  implies that  $10 \lg(P/P_0) = n$ .

The general use of special units for logarithmic quantities is discussed in [131]. The quantities power level and field level, and the units bel, decibel and neper, are given in the table and notes that follow.

| <i>Name</i> | <i>Quantity</i>          | <i>Numerical value multiplied by unit</i>                                         | <i>Notes</i> |
|-------------|--------------------------|-----------------------------------------------------------------------------------|--------------|
| field level | $L_F = \ln(F/F_0)$       | $= \ln(F/F_0) \text{ Np} = 2 \lg(F/F_0) \text{ B} = 20 \lg(F/F_0) \text{ dB}$     | 1–3          |
| power level | $L_P = (1/2) \ln(P/P_0)$ | $= (1/2) \ln(P/P_0) \text{ Np} = \lg(P/P_0) \text{ B} = 10 \lg(P/P_0) \text{ dB}$ | 4–6          |

(1)  $F_0$  is a reference field quantity, which should be specified.

(2) In the context of acoustics the field level is called the sound pressure level and given the symbol  $L_p$ , and the reference pressure  $p_0 = 20 \mu\text{Pa}$ .

(3) For example, when  $L_F = 1 \text{ Np}$ ,  $F/F_0 = e \approx 2.718 \text{ 281 8}$ .

(4)  $P_0$  is a reference power, which should be specified. The factor  $1/2$  is included in the definition to make  $L_P \hat{=} L_F$ .

(5) In the context of acoustics the power level is called the sound power level and given the symbol  $L_W$ , and the reference power  $P_0 = 1 \text{ pW}$ .

(6) For example, when  $L_P = 1 \text{ B} = 10 \text{ dB}$ ,  $P/P_0 = 10$ ; and when  $L_P = 2 \text{ B} = 20 \text{ dB}$ ,  $P/P_0 = 100$ ; etc.



## 4 RECOMMENDED MATHEMATICAL SYMBOLS



## 4.1 PRINTING OF NUMBERS AND MATHEMATICAL SYMBOLS [5.a]

1. Numbers in general shall be printed in Roman (upright) type. The decimal sign between digits in a number should be a point (e.g. 2.3) or a comma (e.g. 2,3). When the decimal sign is placed before the first significant digit of a number, a zero shall always precede the decimal sign. To facilitate the reading of long numbers the digits may be separated into groups of three about the decimal sign, using only a thin space (but never a point or a comma, nor any other symbol). However, when there are only four digits before or after the decimal marker we recommend that no space is required and no space should be used.

*Examples*    2573.421 736    or    2573,421 736    or    0.257 342 173 6 $\times 10^4$     or  
                  0,257 342 173 6 $\times 10^4$   
                  32 573.4215    or    32 573,4215

2. Numerical values of physical quantities which have been experimentally determined are usually subject to some uncertainty. The experimental uncertainty should always be specified. The magnitude of the uncertainty may be represented as follows

*Examples*     $l = [5.3478 - 0.0064, 5.3478 + 0.0064] \text{ cm}$   
                   $l = 5.3478(32) \text{ cm}$

In the first example the range of uncertainty is indicated directly as  $[a - b, a + b]$ . It is recommended that this notation should be used only with the meaning that the interval  $[a - b, a + b]$  contains the true value with a high degree of certainty, such that  $b \geq 2\sigma$ , where  $\sigma$  denotes the standard uncertainty or standard deviation (see Chapter 8, p. 149).

In the second example,  $a(c)$ , the range of uncertainty  $c$  indicated in parentheses is assumed to apply to the least significant digits of  $a$ . It is recommended that this notation be reserved for the meaning that  $b$  represents  $1\sigma$  in the final digits of  $a$ .

3. Letter symbols for mathematical constants (e.g.  $e$ ,  $\pi$ ,  $i = \sqrt{-1}$ ) shall be printed in Roman (upright) type, but letter symbols for numbers other than constants (e.g. quantum numbers) should be printed in italic (sloping) type, similar to physical quantities.
4. Symbols for specific mathematical functions and operators (e.g.  $\ln$ ,  $\lg$ ,  $\exp$ ,  $\sin$ ,  $\cos$ ,  $d$ ,  $\delta$ ,  $\Delta$ ,  $\nabla$ , ...) shall be printed in Roman type, but symbols for a general function (e.g.  $f(x)$ ,  $F(x, y)$ , ...) shall be printed in italic type.
5. The operator  $p$  (as in  $pa_{\text{H}^+}$ ,  $pK = -\lg K$  etc., see Section 2.13.1 (viii), p. 75) shall be printed in Roman type.
6. Symbols for symmetry species in group theory (e.g.  $S$ ,  $P$ ,  $D$ , ...,  $s$ ,  $p$ ,  $d$ , ...,  $\Sigma$ ,  $\Pi$ ,  $\Delta$ , ...,  $A_{1g}$ ,  $B_2''$ , ...) shall be printed in Roman (upright) type when they represent the state symbol for an atom or a molecule, although they are often printed in italic type when they represent the symmetry species of a point group.

7. Vectors and matrices shall be printed in bold italic type.

*Examples* force ***F***, electric field ***E***, position vector ***r***

Ordinary italic type is used to denote the magnitude of the corresponding vector.

*Example*  $r = |\mathbf{r}|$

Tensor quantities may be printed in bold face italic sans-serif type.

*Examples* ***S***, ***T***

Vectors may alternatively be characterized by an arrow,  $\vec{A}$ ,  $\vec{a}$  and second-rank tensors by a double arrow,  $\vec{\vec{S}}$ ,  $\vec{\vec{T}}$ .

## 4.2 SYMBOLS, OPERATORS, AND FUNCTIONS [5.k]

| <i>Description</i>                                           | <i>Symbol</i>                                                         | <i>Notes</i> |
|--------------------------------------------------------------|-----------------------------------------------------------------------|--------------|
| <b>signs and symbols</b>                                     |                                                                       |              |
| equal to                                                     | $=$                                                                   |              |
| not equal to                                                 | $\neq$                                                                |              |
| identically equal to                                         | $\equiv$                                                              |              |
| equal by definition to                                       | $\stackrel{\text{def}}{=}, :=$                                        |              |
| approximately equal to                                       | $\approx$                                                             |              |
| asymptotically equal to                                      | $\simeq$                                                              |              |
| corresponds to                                               | $\hat{=}$                                                             |              |
| proportional to                                              | $\sim, \propto$                                                       |              |
| tends to, approaches                                         | $\rightarrow$                                                         |              |
| infinity                                                     | $\infty$                                                              |              |
| less than                                                    | $<$                                                                   |              |
| greater than                                                 | $>$                                                                   |              |
| less than or equal to                                        | $\leq$                                                                |              |
| greater than or equal to                                     | $\geq$                                                                |              |
| much less than                                               | $\ll$                                                                 |              |
| much greater than                                            | $\gg$                                                                 |              |
| <b>operations</b>                                            |                                                                       |              |
| plus                                                         | $+$                                                                   |              |
| minus                                                        | $-$                                                                   |              |
| plus or minus                                                | $\pm$                                                                 |              |
| minus or plus                                                | $\mp$                                                                 |              |
| $a$ multiplied by $b$                                        | $a\ b, ab, a \cdot b, a \times b$                                     | 1            |
| $a$ divided by $b$                                           | $a/b, ab^{-1}, \frac{a}{b}$                                           | 2            |
| magnitude of $a$                                             | $ a $                                                                 |              |
| $a$ to the power $n$                                         | $a^n$                                                                 |              |
| square root of $a$ , and of $a^2 + b^2$                      | $\sqrt{a}, a^{1/2}, \text{ and } \sqrt{a^2 + b^2}, (a^2 + b^2)^{1/2}$ |              |
| $n$ th root of $a$                                           | $a^{1/n}, \sqrt[n]{a}$                                                |              |
| mean value of $a$                                            | $\langle a \rangle, \bar{a}$                                          |              |
| sign of $a$ (equal to $a/ a $ if $a \neq 0$ , 0 if $a = 0$ ) | $\text{sgn } a$                                                       |              |
| $n$ factorial                                                | $n!$                                                                  |              |
| binominal coefficient, $n!/p!(n-p)!$                         | $C_p^n, \binom{n}{p}$                                                 |              |
| sum of $a_i$                                                 | $\sum a_i, \sum_i a_i, \sum_{i=1}^n a_i$                              |              |
| product of $a_i$                                             | $\prod a_i, \prod_i a_i, \prod_{i=1}^n a_i$                           |              |
| <b>functions</b>                                             |                                                                       |              |
| sine of $x$                                                  | $\sin x$                                                              |              |
| cosine of $x$                                                | $\cos x$                                                              |              |
| tangent of $x$                                               | $\tan x$                                                              |              |
| cotangent of $x$                                             | $\cot x$                                                              |              |

(1) When multiplication is indicated by a dot, the dot shall be half high:  $a \cdot b$ .

(2)  $a : b$  is as well used for “divided by”. However, this symbol is mainly used to express ratios such as length scales in maps.

| <i>Description</i>                           | <i>Symbol</i>                                                                                                                                                                                                                                                                                                                             | <i>Notes</i> |
|----------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------|
| arc sine of $x$                              | $\arcsin x$                                                                                                                                                                                                                                                                                                                               | 3            |
| arc cosine of $x$                            | $\arccos x$                                                                                                                                                                                                                                                                                                                               | 3            |
| arc tangent of $x$                           | $\arctan x$                                                                                                                                                                                                                                                                                                                               | 3            |
| arc cotangent of $x$                         | $\operatorname{arccot} x$                                                                                                                                                                                                                                                                                                                 | 3            |
| hyperbolic sine of $x$                       | $\sinh x$                                                                                                                                                                                                                                                                                                                                 |              |
| hyperbolic cosine of $x$                     | $\cosh x$                                                                                                                                                                                                                                                                                                                                 |              |
| hyperbolic tangent of $x$                    | $\tanh x$                                                                                                                                                                                                                                                                                                                                 |              |
| hyperbolic cotangent of $x$                  | $\coth x$                                                                                                                                                                                                                                                                                                                                 |              |
| area hyperbolic sine of $x$                  | $\operatorname{arsinh} x$                                                                                                                                                                                                                                                                                                                 | 3            |
| area hyperbolic cosine of $x$                | $\operatorname{arcosh} x$                                                                                                                                                                                                                                                                                                                 | 3            |
| area hyperbolic tangent of $x$               | $\operatorname{artanh} x$                                                                                                                                                                                                                                                                                                                 | 3            |
| area hyperbolic cotangent of $x$             | $\operatorname{arcoth} x$                                                                                                                                                                                                                                                                                                                 | 3            |
| base of natural logarithms                   | $e$                                                                                                                                                                                                                                                                                                                                       |              |
| exponential of $x$                           | $\exp x, e^x$                                                                                                                                                                                                                                                                                                                             |              |
| logarithm to the base $a$ of $x$             | $\log_a x$                                                                                                                                                                                                                                                                                                                                | 4            |
| natural logarithm of $x$                     | $\ln x, \log_e x$                                                                                                                                                                                                                                                                                                                         | 4            |
| logarithm to the base 10 of $x$              | $\lg x, \log_{10} x$                                                                                                                                                                                                                                                                                                                      | 4            |
| logarithm to the base 2 of $x$               | $\operatorname{lb} x, \log_2 x$                                                                                                                                                                                                                                                                                                           | 4            |
| greatest integer $\leq x$                    | $\operatorname{ent} x$                                                                                                                                                                                                                                                                                                                    |              |
| integer part of $x$                          | $\operatorname{int} x$                                                                                                                                                                                                                                                                                                                    |              |
| integer division                             | $\operatorname{int}(n/m)$                                                                                                                                                                                                                                                                                                                 |              |
| remainder after integer division             | $n/m - \operatorname{int}(n/m)$                                                                                                                                                                                                                                                                                                           |              |
| change in $x$                                | $\Delta x = x(\text{final}) - x(\text{initial})$                                                                                                                                                                                                                                                                                          |              |
| infinitesimal variation of $f$               | $\delta f$                                                                                                                                                                                                                                                                                                                                |              |
| limit of $f(x)$ as $x$ tends to $a$          | $\lim_{x \rightarrow a} f(x)$                                                                                                                                                                                                                                                                                                             |              |
| 1st derivative of $f$                        | $df/dx, f', (d/dx)f$                                                                                                                                                                                                                                                                                                                      |              |
| 2nd derivative of $f$                        | $d^2 f/dx^2, f''$                                                                                                                                                                                                                                                                                                                         |              |
| $n$ th derivative of $f$                     | $d^n f/dx^n, f^{(n)}$                                                                                                                                                                                                                                                                                                                     |              |
| partial derivative of $f$                    | $\partial f/\partial x, \partial_x f, D_x f$                                                                                                                                                                                                                                                                                              |              |
| total differential of $f$                    | $df$                                                                                                                                                                                                                                                                                                                                      |              |
| inexact differential of $f$                  | $\bar{d}f$                                                                                                                                                                                                                                                                                                                                | 5            |
| first derivative of $x$ with respect to time | $\dot{x}, dx/dt$                                                                                                                                                                                                                                                                                                                          |              |
| integral of $f(x)$                           | $\int f(x) dx, \int dx f(x)$                                                                                                                                                                                                                                                                                                              |              |
| Kronecker delta                              | $\delta_{ij} = \begin{cases} 1 & \text{if } i = j \\ 0 & \text{if } i \neq j \end{cases}$                                                                                                                                                                                                                                                 |              |
| Levi-Civita symbol                           | $\varepsilon_{ijk} = \begin{cases} 1 & \text{if } ijk \text{ is a cyclic permutation of } 123 \\ \varepsilon_{123} = \varepsilon_{231} = \varepsilon_{312} = 1 \\ -1 & \text{if } ijk \text{ is an anticyclic permutation of } 123 \\ \varepsilon_{132} = \varepsilon_{321} = \varepsilon_{213} = -1 \\ 0 & \text{otherwise} \end{cases}$ |              |
| Dirac delta function (distribution)          | $\delta(x), \int f(x)\delta(x) dx = f(0)$                                                                                                                                                                                                                                                                                                 |              |

(3) These are the inverse of the parent function, i.e.  $\arcsin x$  is the operator inverse of  $\sin x$ .

(4) For positive  $x$ .

(5) Notation used in thermodynamics, see Section 2.11, note 1, p. 56.

| <i>Description</i>                                                                      | <i>Symbol</i>                                                                                                     | <i>Notes</i> |
|-----------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------|--------------|
| unit step function, Heaviside function                                                  | $\varepsilon(x)$ , $H(x)$ , $h(x)$ ,<br>$\varepsilon(x) = 1$ for $x > 0$ , $\varepsilon(x) = 0$ for $x < 0$ .     |              |
| gamma function                                                                          | $\Gamma(x) = \int_0^{\infty} t^{x-1} e^{-t} dt$<br>$\Gamma(n+1) = (n)!$ for positive integers $n$                 |              |
| convolution of functions $f$ and $g$                                                    | $f * g = \int_{-\infty}^{+\infty} f(x-x')g(x') dx'$                                                               |              |
| <b>complex numbers</b>                                                                  |                                                                                                                   |              |
| square root of $-1$ , $\sqrt{-1}$                                                       | $i$                                                                                                               |              |
| real part of $z = a + i b$                                                              | $\text{Re } z = a$                                                                                                |              |
| imaginary part of $z = a + i b$                                                         | $\text{Im } z = b$                                                                                                |              |
| modulus of $z = a + i b$ ,<br>absolute value of $z = a + i b$                           | $ z  = (a^2 + b^2)^{1/2}$                                                                                         |              |
| argument of $z = a + i b$                                                               | $\arg z$ ; $\tan(\arg z) = b/a$                                                                                   |              |
| complex conjugate of $z = a + i b$                                                      | $z^* = a - i b$                                                                                                   |              |
| <b>vectors</b>                                                                          |                                                                                                                   |              |
| vector $\mathbf{a}$                                                                     | $\mathbf{a}$ , $\vec{a}$                                                                                          |              |
| cartesian components of $\mathbf{a}$                                                    | $a_x, a_y, a_z$                                                                                                   |              |
| unit vectors in cartesian coordinate system                                             | $\mathbf{e}_x, \mathbf{e}_y, \mathbf{e}_z$ or $\mathbf{i}, \mathbf{j}, \mathbf{k}$                                |              |
| scalar product                                                                          | $\mathbf{a} \cdot \mathbf{b}$                                                                                     |              |
| vector or cross product                                                                 | $\mathbf{a} \times \mathbf{b}$ , $(\mathbf{a} \wedge \mathbf{b})$                                                 |              |
| nabla operator, del operator                                                            | $\nabla = \mathbf{e}_x \partial/\partial x + \mathbf{e}_y \partial/\partial y + \mathbf{e}_z \partial/\partial z$ |              |
| Laplacian operator                                                                      | $\nabla^2, \Delta = \partial^2/\partial x^2 + \partial^2/\partial y^2 + \partial^2/\partial z^2$                  |              |
| gradient of a scalar field $V$                                                          | $\text{grad } V, \nabla V$                                                                                        |              |
| divergence of a vector field $\mathbf{A}$                                               | $\text{div } \mathbf{A}, \nabla \cdot \mathbf{A}$                                                                 |              |
| rotation of a vector field $\mathbf{A}$                                                 | $\text{rot } \mathbf{A}, \nabla \times \mathbf{A}, (\text{curl } \mathbf{A})$                                     |              |
| <b>matrices</b>                                                                         |                                                                                                                   |              |
| matrix of element $A_{ij}$                                                              | $\mathbf{A}$                                                                                                      |              |
| product of matrices $\mathbf{A}$ and $\mathbf{B}$                                       | $\mathbf{AB}$ , where $(\mathbf{AB})_{ik} = \sum_j A_{ij} B_{jk}$                                                 |              |
| unit matrix                                                                             | $\mathbf{E}, \mathbf{I}$                                                                                          |              |
| inverse of a square matrix $\mathbf{A}$                                                 | $\mathbf{A}^{-1}$                                                                                                 |              |
| transpose of matrix $\mathbf{A}$                                                        | $\mathbf{A}^T, \tilde{\mathbf{A}}$                                                                                |              |
| complex conjugate of matrix $\mathbf{A}$                                                | $\mathbf{A}^*$                                                                                                    |              |
| conjugate transpose (adjoint) of $\mathbf{A}$<br>(hermitian conjugate of $\mathbf{A}$ ) | $\mathbf{A}^H, \mathbf{A}^\dagger$ , where $(\mathbf{A}^\dagger)_{ij} = A_{ji}^*$                                 |              |
| trace of a square matrix $\mathbf{A}$                                                   | $\sum_i A_{ii}, \text{tr } \mathbf{A}$                                                                            |              |
| determinant of a square matrix $\mathbf{A}$                                             | $\det \mathbf{A},  \mathbf{A} $                                                                                   |              |
| <b>sets and logical operators</b>                                                       |                                                                                                                   |              |
| $p$ and $q$ (conjunction sign)                                                          | $p \wedge q$                                                                                                      |              |
| $p$ or $q$ or both (disjunction sign)                                                   | $p \vee q$                                                                                                        |              |
| negation of $p$ , not $p$                                                               | $\neg p$                                                                                                          |              |
| $p$ implies $q$                                                                         | $p \Rightarrow q$                                                                                                 |              |
| $p$ is equivalent to $q$                                                                | $p \Leftrightarrow q$                                                                                             |              |

| <i>Description</i>          | <i>Symbol</i>   | <i>Notes</i> |
|-----------------------------|-----------------|--------------|
| $A$ is contained in $B$     | $A \subset B$   |              |
| union of $A$ and $B$        | $A \cup B$      |              |
| intersection of $A$ and $B$ | $A \cap B$      |              |
| $x$ belongs to $A$          | $x \in A$       |              |
| $x$ does not belong to $A$  | $x \notin A$    |              |
| the set $A$ contains $x$    | $A \ni x$       |              |
| $A$ but not $B$             | $A \setminus B$ |              |

## 5 FUNDAMENTAL PHYSICAL CONSTANTS



The data given in this table are from the CODATA recommended values of the fundamental physical constants 2006 [23] (online at <http://physics.nist.gov/constants>) and from the 2006 compilation of the Particle Data Group [133] (online at <http://pdg.lbl.gov>), see notes for details. The standard deviation uncertainty in the least significant digits is given in parentheses.

| <i>Quantity</i>                                                               | <i>Symbol</i>                               | <i>Value</i>                                             | <i>Notes</i> |
|-------------------------------------------------------------------------------|---------------------------------------------|----------------------------------------------------------|--------------|
| magnetic constant                                                             | $\mu_0$                                     | $4\pi \times 10^{-7} \text{ H m}^{-1}$ (defined)         | 1            |
| speed of light in vacuum                                                      | $c_0, c$                                    | $299\,792\,458 \text{ m s}^{-1}$ (defined)               |              |
| electric constant                                                             | $\varepsilon_0 = 1/\mu_0 c_0^2$             | $8.854\,187\,817\dots \times 10^{-12} \text{ F m}^{-1}$  | 1, 2         |
| characteristic impedance<br>of vacuum                                         | $Z_0 = \mu_0 c_0$                           | $376.730\,313\,461\dots \Omega$                          | 2            |
| Planck constant                                                               | $h$                                         | $6.626\,068\,96(33) \times 10^{-34} \text{ J s}$         |              |
|                                                                               | $\hbar = h/2\pi$                            | $1.054\,571\,628(53) \times 10^{-34} \text{ J s}$        |              |
|                                                                               | $\hbar c_0$                                 | $1.986\,445\,501(99) \times 10^{-25} \text{ J m}$        |              |
| Fermi coupling constant                                                       | $G_F/(\hbar c_0)^3$                         | $1.166\,37(1) \times 10^{-5} \text{ GeV}^{-2}$           | 3            |
| weak mixing angle $\theta_W$                                                  | $\sin^2 \theta_W$                           | $0.222\,55(56)$                                          | 4, 5         |
| elementary charge                                                             | $e$                                         | $1.602\,176\,487(40) \times 10^{-19} \text{ C}$          |              |
| electron mass                                                                 | $m_e$                                       | $9.109\,382\,15(45) \times 10^{-31} \text{ kg}$          |              |
| proton mass                                                                   | $m_p$                                       | $1.672\,621\,637(83) \times 10^{-27} \text{ kg}$         |              |
| neutron mass                                                                  | $m_n$                                       | $1.674\,927\,211(84) \times 10^{-27} \text{ kg}$         |              |
| atomic mass constant                                                          | $m_u = 1 \text{ u}$                         | $1.660\,538\,782(83) \times 10^{-27} \text{ kg}$         | 6            |
| Avogadro constant                                                             | $L, N_A$                                    | $6.022\,141\,79(30) \times 10^{23} \text{ mol}^{-1}$     | 7            |
| Boltzmann constant                                                            | $k, k_B$                                    | $1.380\,650\,4(24) \times 10^{-23} \text{ J K}^{-1}$     |              |
| Faraday constant                                                              | $F = Le$                                    | $9.648\,533\,99(24) \times 10^4 \text{ C mol}^{-1}$      |              |
| molar gas constant                                                            | $R$                                         | $8.314\,472(15) \text{ J K}^{-1} \text{ mol}^{-1}$       |              |
| zero of the Celsius scale                                                     |                                             | $273.15 \text{ K}$ (defined)                             |              |
| molar volume of ideal gas,<br>$p = 100 \text{ kPa}, t = 0^\circ \text{C}$     | $V_m$                                       | $22.710\,981(40) \text{ dm}^3 \text{ mol}^{-1}$          |              |
| molar volume of ideal gas,<br>$p = 101.325 \text{ kPa}, t = 0^\circ \text{C}$ |                                             | $22.413\,996(39) \text{ dm}^3 \text{ mol}^{-1}$          |              |
| standard atmosphere                                                           |                                             | $101\,325 \text{ Pa}$ (defined)                          |              |
| fine-structure constant                                                       | $\alpha = \mu_0 e^2 c_0 / 2\hbar$           | $7.297\,352\,537\,6(50) \times 10^{-3}$                  |              |
|                                                                               | $\alpha^{-1}$                               | $137.035\,999\,676(94)$                                  |              |
| Bohr radius                                                                   | $a_0 = 4\pi\varepsilon_0 \hbar^2 / m_e e^2$ | $5.291\,772\,085\,9(36) \times 10^{-11} \text{ m}$       |              |
| Hartree energy                                                                | $E_h = \hbar^2 / m_e a_0^2$                 | $4.359\,743\,94(22) \times 10^{-18} \text{ J}$           |              |
| Rydberg constant                                                              | $R_\infty = E_h / 2\hbar c_0$               | $1.097\,373\,156\,852\,7(73) \times 10^7 \text{ m}^{-1}$ |              |
| Bohr magneton                                                                 | $\mu_B = e\hbar / 2m_e$                     | $9.274\,009\,15(23) \times 10^{-24} \text{ J T}^{-1}$    |              |
| electron magnetic moment                                                      | $\mu_e$                                     | $-9.284\,763\,77(23) \times 10^{-24} \text{ J T}^{-1}$   |              |
| Landé $g$ -factor for<br>free electron                                        | $g_e = 2\mu_e / \mu_B$                      | $-2.002\,319\,304\,362\,2(15)$                           |              |
| nuclear magneton                                                              | $\mu_N = e\hbar / 2m_p$                     | $5.050\,783\,24(13) \times 10^{-27} \text{ J T}^{-1}$    |              |

(1)  $\text{H m}^{-1} = \text{N A}^{-2} = \text{N s}^2 \text{ C}^{-2}$ ;  $\text{F m}^{-1} = \text{C}^2 \text{ J}^{-1} \text{ m}^{-1}$ .

(2)  $\varepsilon_0$  and  $Z_0$  may be calculated exactly from the defined values of  $\mu_0$  and  $c_0$ .

(3) The value of the Fermi coupling constant is recommended by the Particle Data Group [133].

(4) With the weak mixing angle  $\theta_W$ ,  $\sin^2 \theta_W$  is sometimes called Weinberg parameter. There are a number of schemes differing in the masses used to determine  $\sin^2 \theta_W$  (see Chapter 10 in [133]). The value given here for  $\sin^2 \theta_W$  [23] is based on the on-shell scheme which uses  $\sin^2 \theta_W = 1 - (m_W/m_Z)^2$ , where the quantities  $m_W$  and  $m_Z$  are the masses of the  $W^\pm$ - and  $Z^0$ -bosons, respectively.

(5) The Particle Data Group [133] gives  $m_W = 80.403(29) \text{ GeV}/c_0^2$ ,  $m_Z = 91.1876(21) \text{ GeV}/c_0^2$  and recommends  $\sin^2 \theta_W = 0.231\,22(15)$ , based on the  $\overline{\text{MS}}$  scheme. The corresponding value in

| <i>Quantity</i>                                                      | <i>Symbol</i>                     | <i>Value</i>                                                               | <i>Notes</i> |
|----------------------------------------------------------------------|-----------------------------------|----------------------------------------------------------------------------|--------------|
| proton magnetic moment                                               | $\mu_p$                           | $1.410\,606\,662(37) \times 10^{-26} \text{ J T}^{-1}$                     |              |
| proton gyromagnetic ratio                                            | $\gamma_p = 4\pi\mu_p/h$          | $2.675\,222\,099(70) \times 10^8 \text{ s}^{-1} \text{ T}^{-1}$            |              |
| shielded proton magnetic moment (H <sub>2</sub> O, sphere, 25 °C)    | $\mu_p'/\mu_B$                    | $1.520\,993\,128(17) \times 10^{-3}$                                       |              |
| shielded proton gyromagnetic ratio (H <sub>2</sub> O, sphere, 25 °C) | $\gamma_p'/2\pi$                  | $42.576\,388\,1(12) \text{ MHz T}^{-1}$                                    |              |
| Stefan-Boltzmann constant                                            | $\sigma = 2\pi^5 k^4/15h^3 c_0^2$ | $5.670\,400(40) \times 10^{-8} \text{ W m}^{-2} \text{ K}^{-4}$            |              |
| first radiation constant                                             | $c_1 = 2\pi h c_0^2$              | $3.741\,771\,18(19) \times 10^{-16} \text{ W m}^2$                         |              |
| second radiation constant                                            | $c_2 = hc_0/k$                    | $1.438\,775\,2(25) \times 10^{-2} \text{ m K}$                             |              |
| Newtonian constant of gravitation                                    | $G$                               | $6.674\,28(67) \times 10^{-11} \text{ m}^3 \text{ kg}^{-1} \text{ s}^{-2}$ |              |
| standard acceleration of gravity                                     | $g_n$                             | $9.806\,65 \text{ m s}^{-2} \text{ (defined)}$                             |              |

(5) (continued) the on-shell scheme is  $\sin^2 \theta_W = 0.223\,06(33)$ . The effective parameter also depends on the energy range or momentum transfer considered.

(6) u is the (unified) atomic mass unit (see Section 3.9, p. 94).

(7) See [134] and other papers in the same special issue of *Metrologia* on the precise measurement of the Avogadro constant.

## Values of common mathematical constants

| <i>Mathematical constant</i>                   | <i>Symbol</i> | <i>Value</i>                | <i>Notes</i> |
|------------------------------------------------|---------------|-----------------------------|--------------|
| ratio of circumference to diameter of a circle | $\pi$         | $3.141\,592\,653\,59 \dots$ | 1            |
| base of natural logarithms                     | e             | $2.718\,281\,828\,46 \dots$ |              |
| natural logarithm of 10                        | $\ln 10$      | $2.302\,585\,092\,99 \dots$ |              |

(1) A mnemonic for  $\pi$ , based on the number of letters in words of the English language, is:

*“How I like a drink, alcoholic of course, after the heavy lectures involving quantum mechanics!”*

There are similar mnemonics in poem form in French:

*“Que j’aime à faire apprendre ce nombre utile aux sages!  
Immortel Archimède, artiste ingénieur,  
Qui de ton jugement peut priser la valeur?  
Pour moi, ton problème eut de pareils avantages.”*

and German:

*“Wie? O! Dies  $\pi$   
Macht ernstlich so vielen viele Müh’!  
Lernt immerhin, Jünglinge, leichte Verselein,  
Wie so zum Beispiel dies dürfte zu merken sein!”*

See the Japanese [2.e] and Russian [2.c] editions for further mnemonics.

## 6 PROPERTIES OF PARTICLES, ELEMENTS, AND NUCLIDES

The symbols for particles, chemical elements, and nuclides have been discussed in Section 2.10.1 (ii), p. 50. The Particle Data Group [133] recommends the use of italic symbols for particles and this has been adopted by many physicists (see also Section 1.6, p. 7).



## 6.1 PROPERTIES OF SELECTED PARTICLES

The data given in this table are from the CODATA recommended values of the fundamental physical constants 2006 [23] (online at <http://physics.nist.gov/constants>) and from the 2006 compilation of the Particle Data Group (PDG) [133] (online at <http://pdg.lbl.gov>), see notes for details. The standard deviation uncertainty in the least significant digits is given in parentheses.

| Name               | Symbol    | Spin<br>$I$ | Charge<br>number<br>$z$ | Mass                                    |                           | Notes |
|--------------------|-----------|-------------|-------------------------|-----------------------------------------|---------------------------|-------|
|                    |           |             |                         | $m/u$                                   | $mc_0^2/\text{MeV}$       |       |
| photon             | $\gamma$  | 1           | 0                       | 0                                       | 0                         |       |
| neutrino           | $\nu_e$   | 1/2         | 0                       | $\approx 0$                             | $\approx 0$               | 1, 2  |
| electron           | $e^-$     | 1/2         | -1                      | $5.485\,799\,094\,3(23) \times 10^{-4}$ | $0.510\,998\,910(13)$     | 3     |
| muon               | $\mu^\pm$ | 1/2         | $\pm 1$                 | $0.113\,428\,926\,4(30)$                | $105.658\,369\,2(94)$     | 2     |
| pion               | $\pi^\pm$ | 0           | $\pm 1$                 | $0.149\,834\,76(37)$                    | $139.570\,18(35)$         | 2     |
| pion               | $\pi^0$   | 0           | 0                       | $0.144\,903\,35(64)$                    | $134.9766(6)$             | 2     |
| proton             | $p$       | 1/2         | 1                       | $1.007\,276\,466\,77(10)$               | $938.272\,013(23)$        |       |
| neutron            | $n$       | 1/2         | 0                       | $1.008\,664\,915\,97(43)$               | $939.565\,346(23)$        |       |
| deuteron           | $d$       | 1           | 1                       | $2.013\,553\,212\,724(78)$              | $1875.612\,793(47)$       |       |
| triton             | $t$       | 1/2         | 1                       | $3.015\,500\,713\,4(25)$                | $2808.920\,906(70)$       | 4     |
| helion             | $h$       | 1/2         | 2                       | $3.014\,932\,247\,3(26)$                | $2808.391\,383(70)$       | 4     |
| $\alpha$ -particle | $\alpha$  | 0           | 2                       | $4.001\,506\,179\,127(62)$              | $3727.379\,109(93)$       |       |
| Z-boson            | $Z^0$     | 1           | 0                       |                                         | $91.1876(21) \times 10^3$ | 2, 5  |
| W-boson            | $W^\pm$   | 1           | $\pm 1$                 |                                         | $80.403(29) \times 10^3$  | 2, 5  |

(1) The neutrino and antineutrino may perhaps have a small mass,  $m_{\bar{\nu}_e} < 2 \text{ eV}/c_0^2$  [133]. In addition to the electron neutrino  $\nu_e$  one finds also a tau neutrino,  $\nu_\tau$ , and a myon neutrino,  $\nu_\mu$  (and their antiparticles  $\bar{\nu}$ ).

(2) These data are from the Particle Data Group [133].

(3) The electron is sometimes denoted by  $e$  or as a  $\beta$ -particle by  $\beta^-$ . Its anti particle  $e^+$  (positron, also  $\beta^+$ ) has the same mass as the electron  $e^-$  but opposite charge and opposite magnetic moment.

(4) Triton is the  $^3\text{H}^+$ , and helion the  $^3\text{He}^{2+}$  particle.

(5)  $Z^0$  and  $W^\pm$  are gauge bosons [133].

|                    | <i>Symbol</i>    | <i>Magnetic moment</i>          | <i>Mean life</i> <sup>1</sup> |              |
|--------------------|------------------|---------------------------------|-------------------------------|--------------|
| <i>Name</i>        |                  | $\mu/\mu_{\text{N}}$            | $\tau/\text{s}$               | <i>Notes</i> |
| photon             | $\gamma$         | 0                               |                               |              |
| neutrino           | $\nu_{\text{e}}$ | $\approx 0$                     |                               | 2, 6         |
| electron           | $\text{e}^{-}$   | $-1.001\ 159\ 652\ 181\ 11(74)$ |                               | 7, 8         |
| muon               | $\mu^{+}$        | $8.890\ 596\ 98(23)$            | $2.197\ 03(4)\times 10^{-6}$  | 2, 8, 9      |
| pion               | $\pi^{\pm}$      | 0                               | $2.6033(5)\times 10^{-8}$     | 2            |
| pion               | $\pi^0$          | 0                               | $8.4(6)\times 10^{-17}$       | 2            |
| proton             | p                | $2.792\ 847\ 356(23)$           |                               | 8, 10        |
| neutron            | n                | $-1.913\ 042\ 73(45)$           | $885.7(8)$                    | 8            |
| deuteron           | d                | $0.857\ 438\ 230\ 8(72)$        |                               | 8            |
| triton             | t                | $2.978\ 962\ 448(38)$           |                               | 8, 11        |
| helion             | h                | $-2.127\ 497\ 718(25)$          |                               | 8, 12        |
| $\alpha$ -particle | $\alpha$         | 0                               |                               |              |

<sup>1</sup> The PDG [133] gives the mean life ( $\tau$ ) values, see also Section 2.12, note 8, p. 64.

(6) The Particle Data Group [133] gives  $\mu/\mu_B < 0.9 \times 10^{-10}$ .

(7) The value of the magnetic moment is given in Bohr magnetons  $\mu/\mu_B$ ,  $\mu_B = e\hbar/2m_e$ .

In nuclear physics and chemistry the masses of particles are often quoted as their energy equivalents (usually in megaelectronvolts). The unified atomic mass unit corresponds to 931.494 028(23) MeV [23].

Atom-like pairs of a positive particle and an electron are sometimes sufficiently stable to be treated as individual entities with special names.

*Examples*    positronium ( $e^+e^-$ ; Ps)     $m(\text{Ps}) = 1.097\,152\,515\,21(46) \times 10^{-3} \text{ u}$   
                   muonium ( $\mu^+e^-$ ; Mu)     $m(\text{Mu}) = 0.113\,977\,490\,9(29) \text{ u}$

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(Notes continued)

(8) The sign of the magnetic moment is defined with respect to the direction of the spin angular momentum.

(9)  $\mu^-$  and  $\mu^+$  have the same mass but opposite charge and opposite magnetic moment.

(10) The shielded proton magnetic moment,  $\mu_p'$ , is given by  $\mu_p'/\mu_N = 2.792\,775\,598(30)$  ( $\text{H}_2\text{O}$ , sphere, 25 °C).

(11) The half life,  $t_{1/2}$ , of the triton is about 12.3 a (see Section 2.12, p. 64) with a corresponding mean life,  $\tau$ , of 17.7 a.

(12) This is the shielded helion magnetic moment,  $\mu_h'$ , given as  $\mu_h'/\mu_N$  (gas, sphere, 25 °C).

## 6.2 STANDARD ATOMIC WEIGHTS OF THE ELEMENTS 2005

As agreed by the IUPAC Commission on Atomic Weights and Isotopic Abundances (CAWIA) in 1979 [135] the relative atomic mass (generally called atomic weight [136]) of an element, E, can be defined for any specified sample. It is the average mass of the atoms in the sample divided by the unified atomic mass unit<sup>1</sup> or alternatively the molar mass of its atoms divided by the molar mass constant  $M_{\text{u}} = N_{\text{A}}m_{\text{u}} = 1 \text{ g mol}^{-1}$ :

$$A_{\text{r}}(\text{E}) = \overline{m}_{\text{a}}(\text{E})/\text{u} = M(\text{E})/M_{\text{u}}$$

The variations in isotopic composition of many elements in samples of different origin limit the precision to which an atomic weight can be given. The standard atomic weights revised biennially by the CAWIA are meant to be applicable for normal materials. This means that to a high level of confidence the atomic weight of an element in any normal sample will be within the uncertainty limits of the tabulated value. By “normal” it is meant here that the material is a reasonably possible source of the element or its compounds in commerce for industry and science and that it has not been subject to significant modification of isotopic composition within a geologically brief period [137]. This, of course, excludes materials studied themselves for very anomalous isotopic composition. New statistical guidelines have been formulated and used to provide uncertainties on isotopic abundances in the isotopic composition of the elements 1997 [138].

Table 6.2 below lists the atomic weights of the elements 2005 [139] and the term symbol  $^{2S+1}L_J$  for the atomic ground state [140] in the order of the atomic number. The atomic weights have been recommended by the IUPAC Commission on Isotopic Abundances and Atomic Weights (CIAAW) in 2005 [139] and apply to elements as they exist naturally on earth. An electronic version of the CIAAW Table of Standard Atomic Weights 2005 can be found on the CIAAW web page at [http://www.ciaaw.org/atomic\\_weights4.htm](http://www.ciaaw.org/atomic_weights4.htm). The list includes the approved names of elements 110 and 111 (Ds and Rg) [141,142]. The symbol Rg has also been used for “rare gas”. For a history of recommended atomic weight values from 1882 to 1997, see [143].

The atomic weights of many elements depend on the origin and treatment of the materials [138]. The notes to this table explain the types of variation to be expected for individual elements. When used with due regard to the notes the values are considered reliable to  $\pm$  the figure given in parentheses being applicable to the last digit. For elements without a characteristic terrestrial isotopic composition no standard atomic weight is recommended. The atomic mass of its most stable isotope can be found in Section 6.3 below.

| <i>Symbol</i> | <i>Atomic number</i> | <i>Name</i> | <i>Atomic weight<br/>(Relative atomic mass)</i> | <i>Ground state<br/>term symbol</i> | <i>Note</i> |
|---------------|----------------------|-------------|-------------------------------------------------|-------------------------------------|-------------|
| H             | 1                    | hydrogen    | 1.007 94(7)                                     | $^2\text{S}_{1/2}$                  | g, m, r     |
| He            | 2                    | helium      | 4.002 602(2)                                    | $^1\text{S}_0$                      | g, r        |
| Li            | 3                    | lithium     | [6.941(2)] <sup>†</sup>                         | $^2\text{S}_{1/2}$                  | g, m, r     |
| Be            | 4                    | beryllium   | 9.012 182(3)                                    | $^1\text{S}_0$                      |             |
| B             | 5                    | boron       | 10.811(7)                                       | $^2\text{P}_{1/2}^{\circ}$          | g, m, r     |
| C             | 6                    | carbon      | 12.0107(8)                                      | $^3\text{P}_0$                      | g, r        |
| N             | 7                    | nitrogen    | 14.0067(2)                                      | $^4\text{S}_{3/2}^{\circ}$          | g, r        |
| O             | 8                    | oxygen      | 15.9994(3)                                      | $^3\text{P}_2$                      | g, r        |
| F             | 9                    | fluorine    | 18.998 403 2(5)                                 | $^2\text{P}_{3/2}^{\circ}$          |             |
| Ne            | 10                   | neon        | 20.1797(6)                                      | $^1\text{S}_0$                      | g, m        |
| Na            | 11                   | sodium      | 22.989 769 28(2)                                | $^2\text{S}_{1/2}$                  |             |
| Mg            | 12                   | magnesium   | 24.3050(6)                                      | $^1\text{S}_0$                      |             |

<sup>1</sup> Note that the atomic mass constant  $m_{\text{u}}$  is equal to the Dalton, Da, or the unified atomic mass unit, u, and is defined in terms of the mass of the carbon-12 atom:  $m_{\text{u}} = 1 \text{ u} = 1 \text{ Da} = m_{\text{a}}(^{12}\text{C})/12$ .

| <i>Symbol</i> | <i>Atomic<br/>number</i> | <i>Name</i>          | <i>Atomic weight<br/>(Relative atomic mass)</i> | <i>Ground state<br/>term symbol</i> | <i>Note</i> |
|---------------|--------------------------|----------------------|-------------------------------------------------|-------------------------------------|-------------|
| Al            | 13                       | aluminium (aluminum) | 26.981 538 6(8)                                 | $2P_{1/2}^o$                        |             |
| Si            | 14                       | silicon              | 28.0855(3)                                      | $3P_0$                              | r           |
| P             | 15                       | phosphorus           | 30.973 762(2)                                   | $4S_{3/2}^o$                        |             |
| S             | 16                       | sulfur               | 32.065(5)                                       | $3P_2$                              | g, r        |
| Cl            | 17                       | chlorine             | 35.453(2)                                       | $2P_{3/2}^o$                        | g, m, r     |
| Ar            | 18                       | argon                | 39.948(1)                                       | $1S_0$                              | g, r        |
| K             | 19                       | potassium            | 39.0983(1)                                      | $2S_{1/2}$                          |             |
| Ca            | 20                       | calcium              | 40.078(4)                                       | $1S_0$                              | g           |
| Sc            | 21                       | scandium             | 44.955 912(6)                                   | $2D_{3/2}$                          |             |
| Ti            | 22                       | titanium             | 47.867(1)                                       | $3F_2$                              |             |
| V             | 23                       | vanadium             | 50.9415(1)                                      | $4F_{3/2}$                          |             |
| Cr            | 24                       | chromium             | 51.9961(6)                                      | $7S_3$                              |             |
| Mn            | 25                       | manganese            | 54.938 045(5)                                   | $6S_{5/2}$                          |             |
| Fe            | 26                       | iron                 | 55.845(2)                                       | $5D_4$                              |             |
| Co            | 27                       | cobalt               | 58.933 195(5)                                   | $4F_{9/2}$                          |             |
| Ni            | 28                       | nickel               | 58.6934(2)                                      | $3F_4$                              |             |
| Cu            | 29                       | copper               | 63.546(3)                                       | $2S_{1/2}$                          | r           |
| Zn            | 30                       | zinc                 | 65.409(4)                                       | $1S_0$                              |             |
| Ga            | 31                       | gallium              | 69.723(1)                                       | $2P_{1/2}^o$                        |             |
| Ge            | 32                       | germanium            | 72.64(1)                                        | $3P_0$                              |             |
| As            | 33                       | arsenic              | 74.921 60(2)                                    | $4S_{3/2}^o$                        |             |
| Se            | 34                       | selenium             | 78.96(3)                                        | $3P_2$                              | r           |
| Br            | 35                       | bromine              | 79.904(1)                                       | $2P_{3/2}^o$                        |             |
| Kr            | 36                       | krypton              | 83.798(2)                                       | $1S_0$                              | g, m        |
| Rb            | 37                       | rubidium             | 85.4678(3)                                      | $2S_{1/2}$                          | g           |
| Sr            | 38                       | strontium            | 87.62(1)                                        | $1S_0$                              | g, r        |
| Y             | 39                       | yttrium              | 88.905 85(2)                                    | $2D_{3/2}$                          |             |
| Zr            | 40                       | zirconium            | 91.224(2)                                       | $3F_2$                              | g           |
| Nb            | 41                       | niobium              | 92.906 38(2)                                    | $6D_{1/2}$                          |             |
| Mo            | 42                       | molybdenum           | 95.94(2)                                        | $7S_3$                              | g           |
| Tc            | 43                       | technetium           |                                                 | $6S_{5/2}$                          | A           |
| Ru            | 44                       | ruthenium            | 101.07(2)                                       | $5F_5$                              | g           |
| Rh            | 45                       | rhodium              | 102.905 50(2)                                   | $4F_{9/2}$                          |             |
| Pd            | 46                       | palladium            | 106.42(1)                                       | $1S_0$                              | g           |
| Ag            | 47                       | silver               | 107.8682(2)                                     | $2S_{1/2}$                          | g           |
| Cd            | 48                       | cadmium              | 112.411(8)                                      | $1S_0$                              | g           |
| In            | 49                       | indium               | 114.818(3)                                      | $2P_{1/2}^o$                        |             |
| Sn            | 50                       | tin                  | 118.710(7)                                      | $3P_0$                              | g           |
| Sb            | 51                       | antimony             | 121.760(1)                                      | $4S_{3/2}^o$                        | g           |
| Te            | 52                       | tellurium            | 127.60(3)                                       | $3P_2$                              | g           |
| I             | 53                       | iodine               | 126.904 47(3)                                   | $2P_{3/2}^o$                        |             |
| Xe            | 54                       | xenon                | 131.293(6)                                      | $1S_0$                              | g, m        |
| Cs            | 55                       | caesium (cesium)     | 132.905 451 9(2)                                | $2S_{1/2}$                          |             |
| Ba            | 56                       | barium               | 137.327(7)                                      | $1S_0$                              |             |
| La            | 57                       | lanthanum            | 138.905 47(7)                                   | $2D_{3/2}$                          | g           |
| Ce            | 58                       | cerium               | 140.116(1)                                      | $1G_4^o$                            | g           |

| <i>Symbol</i> | <i>Atomic<br/>number</i> | <i>Name</i>  | <i>Atomic weight<br/>(Relative atomic mass)</i> | <i>Ground state<br/>term symbol</i> | <i>Note</i> |
|---------------|--------------------------|--------------|-------------------------------------------------|-------------------------------------|-------------|
| Pr            | 59                       | praseodymium | 140.907 65(2)                                   | $4I_{9/2}^o$                        |             |
| Nd            | 60                       | neodymium    | 144.242(3)                                      | $5I_4$                              | g           |
| Pm            | 61                       | promethium   |                                                 | $6H_{5/2}^o$                        | A           |
| Sm            | 62                       | samarium     | 150.36(2)                                       | $7F_0$                              | g           |
| Eu            | 63                       | europium     | 151.964(1)                                      | $8S_{7/2}^o$                        | g           |
| Gd            | 64                       | gadolinium   | 157.25(3)                                       | $9D_2^o$                            | g           |
| Tb            | 65                       | terbium      | 158.925 35(2)                                   | $6H_{15/2}^o$                       |             |
| Dy            | 66                       | dysprosium   | 162.500(1)                                      | $5I_8$                              | g           |
| Ho            | 67                       | holmium      | 164.930 32(2)                                   | $4I_{15/2}^o$                       |             |
| Er            | 68                       | erbium       | 167.259(3)                                      | $3H_6$                              | g           |
| Tm            | 69                       | thulium      | 168.934 21(2)                                   | $2F_{7/2}^o$                        |             |
| Yb            | 70                       | ytterbium    | 173.04(3)                                       | $1S_0$                              | g           |
| Lu            | 71                       | lutetium     | 174.967(1)                                      | $2D_{3/2}$                          | g           |
| Hf            | 72                       | hafnium      | 178.49(2)                                       | $3F_2$                              |             |
| Ta            | 73                       | tantalum     | 180.947 88(2)                                   | $4F_{3/2}$                          |             |
| W             | 74                       | tungsten     | 183.84(1)                                       | $5D_0$                              |             |
| Re            | 75                       | rhenium      | 186.207(1)                                      | $6S_{5/2}$                          |             |
| Os            | 76                       | osmium       | 190.23(3)                                       | $5D_4$                              | g           |
| Ir            | 77                       | iridium      | 192.217(3)                                      | $4F_{9/2}$                          |             |
| Pt            | 78                       | platinum     | 195.084(9)                                      | $3D_3$                              |             |
| Au            | 79                       | gold         | 196.966 569(4)                                  | $2S_{1/2}$                          |             |
| Hg            | 80                       | mercury      | 200.59(2)                                       | $1S_0$                              |             |
| Tl            | 81                       | thallium     | 204.3833(2)                                     | $2P_{1/2}^o$                        |             |
| Pb            | 82                       | lead         | 207.2(1)                                        | $3P_0$                              | g, r        |
| Bi            | 83                       | bismuth      | 208.980 40(1)                                   | $4S_{3/2}^o$                        |             |
| Po            | 84                       | polonium     |                                                 | $3P_2$                              | A           |
| At            | 85                       | astatine     |                                                 | $2P_{3/2}^o$                        | A           |
| Rn            | 86                       | radon        |                                                 | $1S_0$                              | A           |
| Fr            | 87                       | francium     |                                                 | $2S_{1/2}$                          | A           |
| Ra            | 88                       | radium       |                                                 | $1S_0$                              | A           |
| Ac            | 89                       | actinium     |                                                 | $2D_{3/2}$                          | A           |
| Th            | 90                       | thorium      | 232.038 06(2)                                   | $3F_2$                              | g, Z        |
| Pa            | 91                       | protactinium | 231.035 88(2)                                   | $4K_{11/2}$                         | Z           |
| U             | 92                       | uranium      | 238.028 91(3)                                   | $5L_6$                              | g, m, Z     |
| Np            | 93                       | neptunium    |                                                 | $6L_{11/2}$                         | A           |
| Pu            | 94                       | plutonium    |                                                 | $7F_0$                              | A           |
| Am            | 95                       | americium    |                                                 | $8S_{7/2}^o$                        | A           |
| Cm            | 96                       | curium       |                                                 | $9D_2^o$                            | A           |
| Bk            | 97                       | berkelium    |                                                 | $6H_{15/2}^o$                       | A           |
| Cf            | 98                       | californium  |                                                 | $5I_8$                              | A           |
| Es            | 99                       | einsteinium  |                                                 | $4I_{15/2}^o$                       | A           |
| Fm            | 100                      | fermium      |                                                 | $3H_6$                              | A           |
| Md            | 101                      | mendelevium  |                                                 | $2F_{7/2}^o$                        | A           |
| No            | 102                      | nobelium     |                                                 | $1S_0$                              | A           |

| <i>Symbol</i> | <i>Atomic<br/>number</i> | <i>Name</i>   | <i>Atomic weight<br/>(Relative atomic mass)</i> | <i>Ground state<br/>term symbol</i> | <i>Note</i> |
|---------------|--------------------------|---------------|-------------------------------------------------|-------------------------------------|-------------|
| Lr            | 103                      | lawrencium    |                                                 |                                     | A           |
| Rf            | 104                      | rutherfordium |                                                 |                                     | A           |
| Db            | 105                      | dubnium       |                                                 |                                     | A           |
| Sg            | 106                      | seaborgium    |                                                 |                                     | A           |
| Bh            | 107                      | bohrium       |                                                 |                                     | A           |
| Hs            | 108                      | hassium       |                                                 |                                     | A           |
| Mt            | 109                      | meitnerium    |                                                 |                                     | A           |
| Ds            | 110                      | darmstadtium  |                                                 |                                     | A           |
| Rg            | 111                      | roentgenium   |                                                 |                                     | A           |

<sup>†</sup> Commercially available Li materials have atomic weights that range between 6.939 and 6.996; if a more accurate value is required, it must be determined for the specific material.

(g) **G**eological specimens are known in which the element has an isotopic composition outside the limits for normal material. The difference between the atomic weight of the element in such specimens and that given in the table may exceed the stated uncertainty.

(m) **M**odified isotopic compositions may be found in commercially available material because it has been subjected to an undisclosed or inadvertent isotopic fractionation. Substantial deviations in atomic weight of the element from that given in the table can occur.

(r) **R**ange in isotopic composition of normal terrestrial material prevents a more precise  $A_r(E)$  being given; the tabulated  $A_r(E)$  value and uncertainty should be applicable to normal material.

(A) Radioactive element without stable nuclide that lacks a characteristic terrestrial isotopic composition. In the IUPAC Periodic Table of the Elements on the inside back cover, a value in brackets indicates the mass number of the longest-lived isotope of the element (see also the table of nuclide masses, Section 6.3).

(Z) An element without stable nuclide(s), exhibiting a range of characteristic terrestrial compositions of long-lived radionuclide(s) such that a meaningful atomic weight can be given.

## 6.3 PROPERTIES OF NUCLIDES

The table contains the following properties of naturally occurring and some unstable nuclides:

Column

- 1  $Z$  is the atomic number (number of protons) of the nuclide.
- 2 Symbol of the element.
- 3  $A$  is the mass number of the nuclide. The asterisk \* denotes an unstable nuclide (for elements without naturally occurring isotopes it is the most stable nuclide) and the # sign a nuclide of sufficiently long lifetime (greater than  $10^5$  years) [144] to enable the determination of its isotopic abundance.
- 4 The atomic mass is given in unified atomic mass units,  $1 \text{ u} = m_{\text{a}}(^{12}\text{C})/12$ , together with the standard errors in parentheses and applicable to the last digits quoted. The data were extracted from a more extensive list of the AME2003 atomic mass evaluation [145, 146].
- 5 Representative isotopic compositions are given as amount-of-substance fractions (mole fractions),  $x$ , of the corresponding atoms in percents. According to the opinion of CAWIA, they represent the isotopic composition of chemicals or materials most commonly encountered in the laboratory. They may not, therefore, correspond to the most abundant natural material [138]. It must be stressed that those values should be used to determine the average properties of chemicals or materials of unspecified natural terrestrial origin, though no actual sample having the exact composition listed may be available. The values listed here are from the 2001 CAWIA review as given in column 9 of ref. [147] as representative isotopic composition. This reference uses the *Atomic Mass Evaluation 1993* [148, 149]. There is an inconsistency in this Table because column 4 uses the most recent masses AME2003 [145, 146], whereas column 5 is based on previous atomic masses [148, 149]. When precise work is to be undertaken, such as assessment of individual properties, samples with more precisely known isotopic abundances (such as listed in column 8 of ref. [147]) should be obtained or suitable measurements should be made. The uncertainties given in parentheses are applicable to the last digits quoted and cover the range of probable variations in the materials as well as experimental errors. For additional data and background information on ranges of isotope-abundance variations in natural and anthropogenic material, see [149, 150].
- 6  $I$  is the nuclear spin quantum number. A plus sign indicates positive parity and a minus sign indicates negative parity. Parentheses denotes uncertain values; all values have been taken from the NUBASE evaluation [144].
- 7 Under magnetic moment the maximum  $z$ -component expectation value of the magnetic dipole moment,  $m$ , in nuclear magnetons is given. The positive or negative sign implies that the orientation of the magnetic dipole with respect to the angular momentum corresponds to the rotation of a positive or negative charge, respectively. The data were extracted from the compilation by N. J. Stone [151]. An asterisk \* indicates that more than one value is given in the original compilation; \*\* indicates that an older value exists with a higher stated precision. The absence of a plus or minus sign means that the sign has not been determined by the experimenter.
- 8 Under quadrupole moment, the electric quadrupole moment area (see Section 2.5, notes 14 and 15 on p. 23 and 24) is given in units of square femtometres,  $1 \text{ fm}^2 = 10^{-30} \text{ m}^2$ , although most of the tables quote them in barn (b),  $1 \text{ b} = 10^{-28} \text{ m}^2 = 100 \text{ fm}^2$ . The positive sign implies a prolate nucleus, the negative sign an oblate nucleus. The data are taken from N. J. Stone [151]. An asterisk \* indicates that more than one value is given in the original compilation; \*\* indicates that an older value exists with a higher stated precision. The absence of a plus or minus sign means that the sign has not been determined by the experimenter.

| <i>Z</i> | <i>Symbol</i>   | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br><i>100 x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|-----------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 1        | H<br>(D)<br>(T) | 1        | 1.007 825 032 07(10)                          | 99.9885(70)                                            | 1/2+                                       | +2.792 847 34(3)                                            | +0.286(2)*                                                     |
|          |                 | 2        | 2.014 101 777 8(4)                            | 0.0115(70)                                             | 1+                                         | +0.857 438 228(9)                                           |                                                                |
|          |                 | 3*       | 3.016 049 277 7(25)                           |                                                        | 1/2+                                       | +2.978 962 44(4)                                            |                                                                |
| 2        | He              | 3        | 3.016 029 319 1(26)                           | 0.000 134(3)                                           | 1/2+                                       | −2.127 497 72(3)                                            |                                                                |
|          |                 | 4        | 4.002 603 254 15(6)                           | 99.999 866(3)                                          | 0+                                         | 0                                                           |                                                                |
| 3        | Li              | 6        | 6.015 122 795(16)                             | 7.59(4)                                                | 1+                                         | +0.822 047 3(6)*                                            | −0.082(2)*                                                     |
|          |                 | 7        | 7.016 004 55(8)                               | 92.41(4)                                               | 3/2−                                       | +3.256 427(2)*                                              | −4.06(8)*                                                      |
| 4        | Be              | 9        | 9.012 182 2(4)                                | 100                                                    | 3/2−                                       | −1.177 432(3)*                                              | +5.29(4)*                                                      |
| 5        | B               | 10       | 10.012 937 0(4)                               | 19.9(7)                                                | 3+                                         | +1.800 644 78(6)                                            | +8.47(6)                                                       |
|          |                 | 11       | 11.009 305 4(4)                               | 80.1(7)                                                | 3/2−                                       | +2.688 648 9(10)                                            | +4.07(3)                                                       |
| 6        | C               | 12       | 12 (by definition)                            | 98.93(8)                                               | 0+                                         | 0                                                           |                                                                |
|          |                 | 13       | 13.003 354 837 8(10)                          | 1.07(8)                                                | 1/2−                                       | +0.702 411 8(14)                                            |                                                                |
|          |                 | 14*      | 14.003 241 989(4)                             |                                                        | 0+                                         | 0                                                           |                                                                |
| 7        | N               | 14       | 14.003 074 004 8(6)                           | 99.636(20)                                             | 1+                                         | +0.403 761 00(6)                                            | +2.001(10)*                                                    |
|          |                 | 15       | 15.000 108 898 2(7)                           | 0.364(20)                                              | 1/2−                                       | −0.283 188 84(5)                                            |                                                                |
| 8        | O               | 16       | 15.994 914 619 56(16)                         | 99.757(16)                                             | 0+                                         | 0                                                           | −2.578*                                                        |
|          |                 | 17       | 16.999 131 70(12)                             | 0.038(1)                                               | 5/2+                                       | −1.893 79(9)                                                |                                                                |
|          |                 | 18       | 17.999 161 0(7)                               | 0.205(14)                                              | 0+                                         | 0                                                           |                                                                |
| 9        | F               | 19       | 18.998 403 22(7)                              | 100                                                    | 1/2+                                       | +2.628 868(8)                                               |                                                                |
| 10       | Ne              | 20       | 19.992 440 175 4(19)                          | 90.48(3)                                               | 0+                                         | 0                                                           | +10.3(8)                                                       |
|          |                 | 21       | 20.993 846 68(4)                              | 0.27(1)                                                | 3/2+                                       | −0.661 797(5)                                               |                                                                |
|          |                 | 22       | 21.991 385 114(19)                            | 9.25(3)                                                | 0+                                         | 0                                                           |                                                                |
| 11       | Na              | 23       | 22.989 769 280 9(29)                          | 100                                                    | 3/2+                                       | +2.217 655 6(6)*                                            | +10.45(10)*                                                    |
| 12       | Mg              | 24       | 23.985 041 700(14)                            | 78.99(4)                                               | 0+                                         | 0                                                           | +19.9(2)*                                                      |
|          |                 | 25       | 24.985 836 92(3)                              | 10.00(1)                                               | 5/2+                                       | −0.855 45(8)                                                |                                                                |
|          |                 | 26       | 25.982 592 929(30)                            | 11.01(3)                                               | 0+                                         | 0                                                           |                                                                |
| 13       | Al              | 27       | 26.981 538 63(12)                             | 100                                                    | 5/2+                                       | +3.641 506 9(7)                                             | +14.66(10)*                                                    |
| 14       | Si              | 28       | 27.976 926 532 5(19)                          | 92.223(19)                                             | 0+                                         | 0                                                           |                                                                |
|          |                 | 29       | 28.976 494 700(22)                            | 4.685(8)                                               | 1/2+                                       | −0.555 29(3)                                                |                                                                |
|          |                 | 30       | 29.973 770 17(3)                              | 3.092(11)                                              | 0+                                         | 0                                                           |                                                                |
| 15       | P               | 31       | 30.973 761 63(20)                             | 100                                                    | 1/2+                                       | +1.131 60(3)                                                |                                                                |
| 16       | S               | 32       | 31.972 071 00(15)                             | 94.99(26)                                              | 0+                                         | 0                                                           | −6.4(10)*                                                      |
|          |                 | 33       | 32.971 458 76(15)                             | 0.75(2)                                                | 3/2+                                       | +0.643 821 2(14)                                            |                                                                |
|          |                 | 34       | 33.967 866 90(12)                             | 4.25(24)                                               | 0+                                         | 0                                                           |                                                                |
|          |                 | 36       | 35.967 080 76(20)                             | 0.01(1)                                                | 0+                                         | 0                                                           |                                                                |
| 17       | Cl              | 35       | 34.968 852 68(4)                              | 75.76(10)                                              | 3/2+                                       | +0.821 874 3(4)                                             | 8.50(11)*                                                      |
|          |                 | 37       | 36.965 902 59(5)                              | 24.24(10)                                              | 3/2+                                       | +0.684 123 6(4)                                             | −6.44(7)*                                                      |
| 18       | Ar              | 36       | 35.967 545 106(29)                            | 0.3365(30)                                             | 0+                                         | 0                                                           |                                                                |
|          |                 | 38       | 37.962 732 4(4)                               | 0.0632(5)                                              | 0+                                         | 0                                                           |                                                                |
|          |                 | 40       | 39.962 383 122 5(29)                          | 99.6003(30)                                            | 0+                                         | 0                                                           |                                                                |
| 19       | K               | 39       | 38.963 706 68(20)                             | 93.2581(44)                                            | 3/2+                                       | +0.391 47(3)**                                              | +5.85                                                          |
|          |                 | 40#      | 39.963 998 48(21)                             | 0.0117(1)                                              | 4−                                         | −1.298 100(3)                                               | −7.3(1)*                                                       |
|          |                 | 41       | 40.961 825 76(21)                             | 6.7302(44)                                             | 3/2+                                       | +0.214 870 1(2)**                                           | +7.11(7)*                                                      |

| <i>Z</i> | <i>Symbol</i> | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br>100 <i>x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 20       | Ca            | 40       | 39.962 590 98(22)                             | 96.941(156)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 42       | 41.958 618 01(27)                             | 0.647(23)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 43       | 42.958 766 6(3)                               | 0.135(10)                                              | 7/2−                                       | −1.317 643(7)*                                              | −5.5(1)**                                                      |
|          |               | 44       | 43.955 481 8(4)                               | 2.086(110)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 46       | 45.953 692 6(24)                              | 0.004(3)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 48#      | 47.952 534(4)                                 | 0.187(21)                                              | 0+                                         | 0                                                           |                                                                |
| 21       | Sc            | 45       | 44.955 911 9(9)                               | 100                                                    | 7/2−                                       | +4.756 487(2)                                               | −15.6(3)*                                                      |
| 22       | Ti            | 46       | 45.952 631 6(9)                               | 8.25(3)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 47       | 46.951 763 1(9)                               | 7.44(2)                                                | 5/2−                                       | −0.788 48(1)                                                | +30.0(20)*                                                     |
|          |               | 48       | 47.947 946 3(9)                               | 73.72(3)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 49       | 48.947 870 0(9)                               | 5.41(2)                                                | 7/2−                                       | −1.104 17(1)                                                | 24.7(11)*                                                      |
|          |               | 50       | 49.944 791 2(9)                               | 5.18(2)                                                | 0+                                         | 0                                                           |                                                                |
| 23       | V             | 50#      | 49.947 158 5(11)                              | 0.250(4)                                               | 6+                                         | +3.345 688 9(14)                                            | 21.0(40)*                                                      |
|          |               | 51       | 50.943 959 5(11)                              | 99.750(4)                                              | 7/2−                                       | +5.148 705 7(2)                                             | −4.3(5)*                                                       |
| 24       | Cr            | 50       | 49.946 044 2(11)                              | 4.345(13)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 52       | 51.940 507 5(8)                               | 83.789(18)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 53       | 52.940 649 4(8)                               | 9.501(17)                                              | 3/2−                                       | −0.474 54(3)                                                | −15.0(50)*                                                     |
|          |               | 54       | 53.938 880 4(8)                               | 2.365(7)                                               | 0+                                         | 0                                                           |                                                                |
| 25       | Mn            | 55       | 54.938 045 1(7)                               | 100                                                    | 5/2−                                       | +3.468 717 90(9)                                            | +33.0(10)*                                                     |
| 26       | Fe            | 54       | 53.939 610 5(7)                               | 5.845(35)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 56       | 55.934 937 5(7)                               | 91.754(36)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 57       | 56.935 394 0(7)                               | 2.119(10)                                              | 1/2−                                       | +0.090 623 00(9)*                                           |                                                                |
|          |               | 58       | 57.933 275 6(8)                               | 0.282(4)                                               | 0+                                         | 0                                                           |                                                                |
| 27       | Co            | 59       | 58.933 195 0(7)                               | 100                                                    | 7/2−                                       | +4.627(9)                                                   | +41.0(10)*                                                     |
| 28       | Ni            | 58       | 57.935 342 9(7)                               | 68.0769(89)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 60       | 59.930 786 4(7)                               | 26.2231(77)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 61       | 60.931 056 0(7)                               | 1.1399(6)                                              | 3/2−                                       | −0.750 02(4)                                                | +16.2(15)                                                      |
|          |               | 62       | 61.928 345 1(6)                               | 3.6345(17)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 64       | 63.927 966 0(7)                               | 0.9256(9)                                              | 0+                                         | 0                                                           |                                                                |
| 29       | Cu            | 63       | 62.929 597 5(6)                               | 69.15(15)                                              | 3/2−                                       | 2.227 345 6(14)*                                            | −21.1(4)*                                                      |
|          |               | 65       | 64.927 789 5(7)                               | 30.85(15)                                              | 3/2−                                       | 2.381 61(19)*                                               | −19.5(4)                                                       |
| 30       | Zn            | 64       | 63.929 142 2(7)                               | 48.268(321)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 66       | 65.926 033 4(10)                              | 27.975(77)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 67       | 66.927 127 3(10)                              | 4.102(21)                                              | 5/2−                                       | +0.875 204 9(11)*                                           | +15.0(15)                                                      |
|          |               | 68       | 67.924 844 2(10)                              | 19.024(123)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 70       | 69.925 319 3(21)                              | 0.631(9)                                               | 0+                                         | 0                                                           |                                                                |
| 31       | Ga            | 69       | 68.925 573 6(13)                              | 60.108(9)                                              | 3/2−                                       | +2.016 59(5)                                                | +16.50(8)*                                                     |
|          |               | 71       | 70.924 701 3(11)                              | 39.892(9)                                              | 3/2−                                       | +2.562 27(2)                                                | +10.40(8)*                                                     |
| 32       | Ge            | 70       | 69.924 247 4(11)                              | 20.38(18)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 72       | 71.922 075 8(18)                              | 27.31(26)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 73       | 72.923 458 9(18)                              | 7.76(8)                                                | 9/2+                                       | −0.879 467 7(2)                                             | −17.0(30)                                                      |
|          |               | 74       | 73.921 177 8(18)                              | 36.72(15)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 76#      | 75.921 402 6(18)                              | 7.83(7)                                                | 0+                                         | 0                                                           |                                                                |
| 33       | As            | 75       | 74.921 596 5(20)                              | 100                                                    | 3/2−                                       | +1.439 48(7)                                                | +30.0(50)**                                                    |

| <i>Z</i> | <i>Symbol</i> | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br>100 <i>x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 34       | Se            | 74       | 73.922 476 4(18)                              | 0.89(4)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 76       | 75.919 213 6(18)                              | 9.37(29)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 77       | 76.919 914 0(18)                              | 7.63(16)                                               | 1/2−                                       | +0.535 042 2(6)*                                            |                                                                |
|          |               | 78       | 77.917 309 1(18)                              | 23.77(28)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 80       | 79.916 521 3(21)                              | 49.61(41)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 82#      | 81.916 699 4(22)                              | 8.73(22)                                               | 0+                                         | 0                                                           |                                                                |
| 35       | Br            | 79       | 78.918 337 1(22)                              | 50.69(7)                                               | 3/2−                                       | +2.106 400(4)                                               | 31.8(5)**                                                      |
|          |               | 81       | 80.916 290 6(21)                              | 49.31(7)                                               | 3/2−                                       | +2.270 562(4)                                               | +26.6(4)**                                                     |
| 36       | Kr            | 78       | 77.920 364 8(12)                              | 0.355(3)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 80       | 79.916 379 0(16)                              | 2.286(10)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 82       | 81.913 483 6(19)                              | 11.593(31)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 83       | 82.914 136(3)                                 | 11.500(19)                                             | 9/2+                                       | −0.970 669(3)                                               | +25.9(1)*                                                      |
|          |               | 84       | 83.911 507(3)                                 | 56.987(15)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 86       | 85.910 610 73(11)                             | 17.279(41)                                             | 0+                                         | 0                                                           |                                                                |
| 37       | Rb            | 85       | 84.911 789 738(12)                            | 72.17(2)                                               | 5/2−                                       | +1.352 98(10)**                                             | +27.7(1)**                                                     |
|          |               | 87#      | 86.909 180 527(13)                            | 27.83(2)                                               | 3/2−                                       | +2.751 31(12)**                                             | +13.4(1)*                                                      |
| 38       | Sr            | 84       | 83.913 425(3)                                 | 0.56(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 86       | 85.909 260 2(12)                              | 9.86(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 87       | 86.908 877 1(12)                              | 7.00(1)                                                | 9/2+                                       | −1.093 603 0(13)*                                           | +33.0(20)*                                                     |
|          |               | 88       | 87.905 612 1(12)                              | 82.58(1)                                               | 0+                                         | 0                                                           |                                                                |
| 39       | Y             | 89       | 88.905 848 3(27)                              | 100                                                    | 1/2−                                       | −0.137 415 4(3)*                                            |                                                                |
| 40       | Zr            | 90       | 89.904 704 4(25)                              | 51.45(40)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 91       | 90.905 645 8(25)                              | 11.22(5)                                               | 5/2+                                       | −1.303 62(2)                                                | −17.6(3)*                                                      |
|          |               | 92       | 91.905 040 8(25)                              | 17.15(8)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 94       | 93.906 315 2(26)                              | 17.38(28)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 96#      | 95.908 273 4(30)                              | 2.80(9)                                                | 0+                                         | 0                                                           |                                                                |
| 41       | Nb            | 93       | 92.906 378 1(26)                              | 100                                                    | 9/2+                                       | +6.1705(3)                                                  | −37.0(20)*                                                     |
| 42       | Mo            | 92       | 91.906 811(4)                                 | 14.77(31)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 94       | 93.905 088 3(21)                              | 9.23(10)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 95       | 94.905 842 1(21)                              | 15.90(9)                                               | 5/2+                                       | −0.9142(1)                                                  | −2.2(1)*                                                       |
|          |               | 96       | 95.904 679 5(21)                              | 16.68(1)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 97       | 96.906 021 5(21)                              | 9.56(5)                                                | 5/2+                                       | −0.9335(1)                                                  | +25.5(13)*                                                     |
|          |               | 98       | 97.905 408 2(21)                              | 24.19(26)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 100#     | 99.907 477(6)                                 | 9.67(20)                                               | 0+                                         | 0                                                           |                                                                |
| 43       | Tc            | 98*      | 97.907 216(4)                                 |                                                        | (6)+                                       |                                                             |                                                                |
| 44       | Ru            | 96       | 95.907 598(8)                                 | 5.54(14)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 98       | 97.905 287(7)                                 | 1.87(3)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 99       | 98.905 939 3(22)                              | 12.76(14)                                              | 5/2+                                       | −0.641(5)                                                   | +7.9(4)                                                        |
|          |               | 100      | 99.904 219 5(22)                              | 12.60(7)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 101      | 100.905 582 1(22)                             | 17.06(2)                                               | 5/2+                                       | −0.719(6)*                                                  | +46.0(20)                                                      |
|          |               | 102      | 101.904 349 3(22)                             | 31.55(14)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 104      | 103.905 433(3)                                | 18.62(27)                                              | 0+                                         | 0                                                           |                                                                |
| 45       | Rh            | 103      | 102.905 504(3)                                | 100                                                    | 1/2−                                       | −0.8840(2)                                                  |                                                                |
| 46       | Pd            | 102      | 101.905 609(3)                                | 1.02(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 104      | 103.904 036(4)                                | 11.14(8)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 105      | 104.905 085(4)                                | 22.33(8)                                               | 5/2+                                       | −0.642(3)                                                   | +65.0(30)**                                                    |
|          |               | 106      | 105.903 486(4)                                | 27.33(3)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 108      | 107.903 892(4)                                | 26.46(9)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 110      | 109.905 153(12)                               | 11.72(9)                                               | 0+                                         | 0                                                           |                                                                |

| <i>Z</i> | <i>Symbol</i> | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br>100 <i>x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 47       | Ag            | 107      | 106.905 097(5)                                | 51.839(8)                                              | 1/2−                                       | −0.113 679 65(15)*                                          |                                                                |
|          |               | 109      | 108.904 752(3)                                | 48.161(8)                                              | 1/2−                                       | −0.130 690 6(2)*                                            |                                                                |
| 48       | Cd            | 106      | 105.906 459(6)                                | 1.25(6)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 108      | 107.904 184(6)                                | 0.89(3)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 110      | 109.903 002 1(29)                             | 12.49(18)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 111      | 110.904 178 1(29)                             | 12.80(12)                                              | 1/2+                                       | −0.594 886 1(8)*                                            |                                                                |
|          |               | 112      | 111.902 757 8(29)                             | 24.13(21)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 113#     | 112.904 401 7(29)                             | 12.22(12)                                              | 1/2+                                       | −0.622 300 9(9)                                             |                                                                |
|          |               | 114      | 113.903 358 5(29)                             | 28.73(42)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 116#     | 115.904 756(3)                                | 7.49(18)                                               | 0+                                         | 0                                                           |                                                                |
| 49       | In            | 113      | 112.904 058(3)                                | 4.29(5)                                                | 9/2+                                       | +5.5289(2)                                                  | +80.0(40)                                                      |
|          |               | 115#     | 114.903 878(5)                                | 95.71(5)                                               | 9/2+                                       | +5.5408(2)                                                  | +81.0(50)*                                                     |
| 50       | Sn            | 112      | 111.904 818(5)                                | 0.97(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 114      | 113.902 779(3)                                | 0.66(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 115      | 114.903 342(3)                                | 0.34(1)                                                | 1/2+                                       | −0.918 83(7)                                                |                                                                |
|          |               | 116      | 115.901 741(3)                                | 14.54(9)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 117      | 116.902 952(3)                                | 7.68(7)                                                | 1/2+                                       | −1.001 04(7)                                                |                                                                |
|          |               | 118      | 117.901 603(3)                                | 24.22(9)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 119      | 118.903 308(3)                                | 8.59(4)                                                | 1/2+                                       | −1.047 28(7)                                                |                                                                |
|          |               | 120      | 119.902 194 7(27)                             | 32.58(9)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 122      | 121.903 439 0(29)                             | 4.63(3)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 124      | 123.905 273 9(15)                             | 5.79(5)                                                | 0+                                         | 0                                                           |                                                                |
| 51       | Sb            | 121      | 120.903 815 7(24)                             | 57.21(5)                                               | 5/2+                                       | +3.3634(3)                                                  | −36.0(40)**                                                    |
|          |               | 123      | 122.904 214 0(22)                             | 42.79(5)                                               | 7/2+                                       | +2.5498(2)                                                  | −49.0(50)                                                      |
| 52       | Te            | 120      | 119.904 020(10)                               | 0.09(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 122      | 121.903 043 9(16)                             | 2.55(12)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 123#     | 122.904 270 0(16)                             | 0.89(3)                                                | 1/2+                                       | −0.736 947 8(8)                                             |                                                                |
|          |               | 124      | 123.902 817 9(16)                             | 4.74(14)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 125      | 124.904 430 7(16)                             | 7.07(15)                                               | 1/2+                                       | −0.888 450 9(10)*                                           |                                                                |
|          |               | 126      | 125.903 311 7(16)                             | 18.84(25)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 128#     | 127.904 463 1(19)                             | 31.74(8)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 130#     | 129.906 224 4(21)                             | 34.08(62)                                              | 0+                                         | 0                                                           |                                                                |
| 53       | I             | 127      | 126.904 473(4)                                | 100                                                    | 5/2+                                       | +2.813 27(8)                                                | 72.0(20)**                                                     |
| 54       | Xe            | 124      | 123.905 893 0(20)                             | 0.0952(3)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 126      | 125.904 274(7)                                | 0.0890(2)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 128      | 127.903 531 3(15)                             | 1.9102(8)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 129      | 128.904 779 4(8)                              | 26.4006(82)                                            | 1/2+                                       | −0.777 976(8)                                               |                                                                |
|          |               | 130      | 129.903 508 0(8)                              | 4.0710(13)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 131      | 130.905 082 4(10)                             | 21.2324(30)                                            | 3/2+                                       | +0.6915(2)**                                                | −11.4(1)*                                                      |
|          |               | 132      | 131.904 153 5(10)                             | 26.9086(33)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 134      | 133.905 394 5(9)                              | 10.4357(21)                                            | 0+                                         | 0                                                           |                                                                |
|          |               | 136      | 135.907 219(8)                                | 8.8573(44)                                             | 0+                                         | 0                                                           |                                                                |
| 55       | Cs            | 133      | 132.905 451 933(24)                           | 100                                                    | 7/2+                                       | +2.582 025(3)**                                             | −0.355(4)*                                                     |
| 56       | Ba            | 130      | 129.906 320(8)                                | 0.106(1)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 132      | 131.905 061(3)                                | 0.101(1)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 134      | 133.904 508 4(4)                              | 2.417(18)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 135      | 134.905 688 6(4)                              | 6.592(12)                                              | 3/2+                                       | 0.838 627(2)*                                               | +16.0(3)*                                                      |
|          |               | 136      | 135.904 575 9(4)                              | 7.854(24)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 137      | 136.905 827 4(5)                              | 11.232(24)                                             | 3/2+                                       | 0.937 34(2)*                                                | +24.5(4)*                                                      |
|          |               | 138      | 137.905 247 2(5)                              | 71.698(42)                                             | 0+                                         | 0                                                           |                                                                |

| <i>Z</i> | <i>Symbol</i> | <i>A</i>                                        | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i>                                                                                                   | <i>Isotopic</i><br><i>composition,</i><br>100 <i>x</i>                                    | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|-------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 57       | La            | 138#<br>139                                     | 137.907 112(4)<br>138.906 353 3(26)                                                                                                             | 0.090(1)<br>99.910(1)                                                                     | 5+<br>7/2+                                 | +3.713 646(7)<br>+2.783 045 5(9)                            | +45.0(20)*<br>+20.0(10)                                        |
| 58       | Ce            | 136<br>138<br>140<br>142                        | 135.907 172(14)<br>137.905 991(11)<br>139.905 438 7(26)<br>141.909 244(3)                                                                       | 0.185(2)<br>0.251(2)<br>88.450(51)<br>11.114(51)                                          | 0+<br>0+<br>0+<br>0+                       | 0<br>0<br>0<br>0                                            |                                                                |
| 59       | Pr            | 141                                             | 140.907 652 8(26)                                                                                                                               | 100                                                                                       | 5/2+                                       | +4.2754(5)                                                  | −7.7(6)*                                                       |
| 60       | Nd            | 142<br>143<br>144#<br>145<br>146<br>148<br>150# | 141.907 723 3(25)<br>142.909 814 3(25)<br>143.910 087 3(25)<br>144.912 573 6(25)<br>145.913 116 9(25)<br>147.916 893(3)<br>149.920 891(3)       | 27.2(5)<br>12.2(2)<br>23.8(3)<br>8.3(1)<br>17.2(3)<br>5.7(1)<br>5.6(2)                    | 0+<br>7/2−<br>0+<br>7/2−<br>0+<br>0+<br>0+ | 0<br>−1.065(5)<br>0<br>−0.656(4)<br>0<br>0<br>0             | −61.0(20)*<br>−31.4(12)**                                      |
| 61       | Pm            | 145*                                            | 144.912 749(3)                                                                                                                                  |                                                                                           | 5/2+                                       |                                                             |                                                                |
| 62       | Sm            | 144<br>147#<br>148#<br>149<br>150<br>152<br>154 | 143.911 999(3)<br>146.914 897 9(26)<br>147.914 822 7(26)<br>148.917 184 7(26)<br>149.917 275 5(26)<br>151.919 732 4(27)<br>153.922 209 3(27)    | 3.07(7)<br>14.99(18)<br>11.24(10)<br>13.82(7)<br>7.38(1)<br>26.75(16)<br>22.75(29)        | 0+<br>7/2−<br>0+<br>7/2−<br>0+<br>0+<br>0+ | 0<br>−0.812(2)**<br>0<br>−0.6677(11)**<br>0<br>0<br>0       | −26.1(7)*<br>+7.5(2)*                                          |
| 63       | Eu            | 151<br>153                                      | 150.919 850 2(26)<br>152.921 230 3(26)                                                                                                          | 47.81(6)<br>52.19(6)                                                                      | 5/2+<br>5/2+                               | +3.4717(6)<br>+1.5324(3)*                                   | 83.0**<br>+222.0*                                              |
| 64       | Gd            | 152#<br>154<br>155<br>156<br>157<br>158<br>160  | 151.919 791 0(27)<br>153.920 865 6(27)<br>154.922 622 0(27)<br>155.922 122 7(27)<br>156.923 960 1(27)<br>157.924 103 9(27)<br>159.927 054 1(27) | 0.20(1)<br>2.18(3)<br>14.80(12)<br>20.47(9)<br>15.65(2)<br>24.84(7)<br>21.86(19)          | 0+<br>0+<br>3/2−<br>0+<br>3/2−<br>0+<br>0+ | 0<br>0<br>−0.2572(4)*<br>0<br>−0.3373(6)*<br>0<br>0         | +127.0(50)*<br>+136.0(60)**                                    |
| 65       | Tb            | 159                                             | 158.925 346 8(27)                                                                                                                               | 100                                                                                       | 3/2+                                       | +2.014(4)                                                   | +143.2(8)                                                      |
| 66       | Dy            | 156<br>158<br>160<br>161<br>162<br>163<br>164   | 155.924 283(7)<br>157.924 409(4)<br>159.925 197 5(27)<br>160.926 933 4(27)<br>161.926 798 4(27)<br>162.928 731 2(27)<br>163.929 174 8(27)       | 0.056(3)<br>0.095(3)<br>2.329(18)<br>18.889(42)<br>25.475(36)<br>24.896(42)<br>28.260(54) | 0+<br>0+<br>0+<br>5/2+<br>0+<br>5/2−<br>0+ | 0<br>0<br>0<br>−0.480(3)*<br>0<br>+0.673(4)<br>0            | 247.7(30)**<br>+265.0(20)**                                    |
| 67       | Ho            | 165                                             | 164.930 322 1(27)                                                                                                                               | 100                                                                                       | 7/2−                                       | +4.17(3)                                                    | 358.0(20)**                                                    |
| 68       | Er            | 162<br>164<br>166<br>167<br>168<br>170          | 161.928 778(4)<br>163.929 200(3)<br>165.930 293 1(27)<br>166.932 048 2(27)<br>167.932 370 2(27)<br>169.935 464 3(30)                            | 0.139(5)<br>1.601(3)<br>33.503(36)<br>22.869(9)<br>26.978(18)<br>14.910(36)               | 0+<br>0+<br>0+<br>7/2+<br>0+<br>0+         | 0<br>0<br>0<br>−0.563 85(12)<br>0<br>0                      | +357.0(3)**                                                    |
| 69       | Tm            | 169                                             | 168.934 213 3(27)                                                                                                                               | 100                                                                                       | 1/2+                                       | −0.2310(15)*                                                |                                                                |

| <i>Z</i> | <i>Symbol</i> | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br>100 <i>x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 70       | Yb            | 168      | 167.933 897(5)                                | 0.13(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 170      | 169.934 761 8(26)                             | 3.04(15)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 171      | 170.936 325 8(26)                             | 14.28(57)                                              | 1/2−                                       | +0.493 67(1)*                                               |                                                                |
|          |               | 172      | 171.936 381 5(26)                             | 21.83(67)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 173      | 172.938 210 8(26)                             | 16.13(27)                                              | 5/2−                                       | −0.648(3)**                                                 | +280.0(40)                                                     |
|          |               | 174      | 173.938 862 1(26)                             | 31.83(92)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 176      | 175.942 571 7(28)                             | 12.76(41)                                              | 0+                                         | 0                                                           |                                                                |
| 71       | Lu            | 175      | 174.940 771 8(23)                             | 97.41(2)                                               | 7/2+                                       | +2.2323(11)**                                               | +349.0(20)*                                                    |
|          |               | 176#     | 175.942 686 3(23)                             | 2.59(2)                                                | 7−                                         | +3.162(12)**                                                | +492.0(50)*                                                    |
| 72       | Hf            | 174#     | 173.940 046(3)                                | 0.16(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 176      | 175.941 408 6(24)                             | 5.26(7)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 177      | 176.943 220 7(23)                             | 18.60(9)                                               | 7/2−                                       | +0.7935(6)                                                  | +337.0(30)*                                                    |
|          |               | 178      | 177.943 698 8(23)                             | 27.28(7)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 179      | 178.945 816 1(23)                             | 13.62(2)                                               | 9/2+                                       | −0.6409(13)                                                 | +379.0(30)*                                                    |
|          |               | 180      | 179.946 550 0(23)                             | 35.08(16)                                              | 0+                                         | 0                                                           |                                                                |
| 73       | Ta            | 180      | 179.947 464 8(24)                             | 0.012(2)                                               | 9−                                         |                                                             |                                                                |
|          |               | 181      | 180.947 995 8(19)                             | 99.988(2)                                              | 7/2+                                       | +2.3705(7)                                                  | +317.0(20)*                                                    |
| 74       | W             | 180      | 179.946 704(4)                                | 0.12(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 182      | 181.948 204 2(9)                              | 26.50(16)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 183      | 182.950 223 0(9)                              | 14.31(4)                                               | 1/2−                                       | +0.117 784 76(9)                                            |                                                                |
|          |               | 184      | 183.950 931 2(9)                              | 30.64(2)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 186      | 185.954 364 1(19)                             | 28.43(19)                                              | 0+                                         | 0                                                           |                                                                |
| 75       | Re            | 185      | 184.952 955 0(13)                             | 37.40(2)                                               | 5/2+                                       | +3.1871(3)                                                  | +218.0(20)*                                                    |
|          |               | 187#     | 186.955 753 1(15)                             | 62.60(2)                                               | 5/2+                                       | +3.2197(3)                                                  | +207.0(20)*                                                    |
| 76       | Os            | 184      | 183.952 489 1(14)                             | 0.02(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 186#     | 185.953 838 2(15)                             | 1.59(3)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 187      | 186.955 750 5(15)                             | 1.96(2)                                                | 1/2−                                       | +0.064 651 89(6)*                                           |                                                                |
|          |               | 188      | 187.955 838 2(15)                             | 13.24(8)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 189      | 188.958 147 5(16)                             | 16.15(5)                                               | 3/2−                                       | +0.659 933(4)                                               | +98.0(60)**                                                    |
|          |               | 190      | 189.958 447 0(16)                             | 26.26(2)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 192      | 191.961 480 7(27)                             | 40.78(19)                                              | 0+                                         | 0                                                           |                                                                |
| 77       | Ir            | 191      | 190.960 594 0(18)                             | 37.3(2)                                                | 3/2+                                       | +0.1507(6)*                                                 | +81.6(9)*                                                      |
|          |               | 193      | 192.962 926 4(18)                             | 62.7(2)                                                | 3/2+                                       | +0.1637(6)*                                                 | +75.1(9)*                                                      |
| 78       | Pt            | 190#     | 189.959 932(6)                                | 0.014(1)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 192      | 191.961 038 0(27)                             | 0.782(7)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 194      | 193.962 680 3(9)                              | 32.967(99)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 195      | 194.964 791 1(9)                              | 33.832(10)                                             | 1/2−                                       | +0.609 52(6)                                                |                                                                |
|          |               | 196      | 195.964 951 5(9)                              | 25.242(41)                                             | 0+                                         | 0                                                           |                                                                |
|          |               | 198      | 197.967 893(3)                                | 7.163(55)                                              | 0+                                         | 0                                                           |                                                                |
| 79       | Au            | 197      | 196.966 568 7(6)                              | 100                                                    | 3/2+                                       | +0.145 746(9)**                                             | +54.7(16)**                                                    |
| 80       | Hg            | 196      | 195.965 833(3)                                | 0.15(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 198      | 197.966 769 0(4)                              | 9.97(20)                                               | 0+                                         | 0                                                           |                                                                |
|          |               | 199      | 198.968 279 9(4)                              | 16.87(22)                                              | 1/2−                                       | +0.505 885 5(9)                                             |                                                                |
|          |               | 200      | 199.968 326 0(4)                              | 23.10(19)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 201      | 200.970 302 3(6)                              | 13.18(9)                                               | 3/2−                                       | −0.560 225 7(14)*                                           | +38.0(40)*                                                     |
|          |               | 202      | 201.970 643 0(6)                              | 29.86(26)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 204      | 203.973 493 9(4)                              | 6.87(15)                                               | 0+                                         | 0                                                           |                                                                |
| 81       | Tl            | 203      | 202.972 344 2(14)                             | 29.52(1)                                               | 1/2+                                       | +1.622 257 87(12)*                                          |                                                                |
|          |               | 205      | 204.974 427 5(14)                             | 70.48(1)                                               | 1/2+                                       | +1.638 214 61(12)                                           |                                                                |
| 82       | Pb            | 204      | 203.973 043 6(13)                             | 1.4(1)                                                 | 0+                                         | 0                                                           |                                                                |
|          |               | 206      | 205.974 465 3(13)                             | 24.1(1)                                                | 0+                                         | 0                                                           |                                                                |
|          |               | 207      | 206.975 896 9(13)                             | 22.1(1)                                                | 1/2−                                       | +0.592 583(9)*                                              |                                                                |
|          |               | 208      | 207.976 652 1(13)                             | 52.4(1)                                                | 0+                                         | 0                                                           |                                                                |

| <i>Z</i> | <i>Symbol</i> | <i>A</i> | <i>Atomic mass,</i><br><i>m<sub>a</sub>/u</i> | <i>Isotopic</i><br><i>composition,</i><br><i>100 x</i> | <i>Nuclear</i><br><i>spin,</i><br><i>I</i> | <i>Magnetic</i><br><i>moment,</i><br><i>m/μ<sub>N</sub></i> | <i>Quadrupole</i><br><i>moment,</i><br><i>Q/fm<sup>2</sup></i> |
|----------|---------------|----------|-----------------------------------------------|--------------------------------------------------------|--------------------------------------------|-------------------------------------------------------------|----------------------------------------------------------------|
| 83       | Bi            | 209#     | 208.980 398 7(16)                             | 100                                                    | 9/2−                                       | +4.1103(5)*                                                 | −51.6(15)*                                                     |
| 84       | Po            | 209*     | 208.982 430 4(20)                             |                                                        | 1/2−                                       |                                                             |                                                                |
| 85       | At            | 210*     | 209.987 148(8)                                |                                                        | (5)+                                       |                                                             |                                                                |
| 86       | Rn            | 222*     | 222.017 577 7(25)                             |                                                        | 0+                                         | 0                                                           |                                                                |
| 87       | Fr            | 223*     | 223.019 735 9(26)                             |                                                        | 3/2(−)                                     | +1.17(2)                                                    | +117.0(10)                                                     |
| 88       | Ra            | 226*     | 226.025 409 8(25)                             |                                                        | 0+                                         | 0                                                           |                                                                |
| 89       | Ac            | 227*     | 227.027 752 1(26)                             |                                                        | 3/2−                                       | +1.1(1)                                                     | +170.0(200)                                                    |
| 90       | Th            | 232#     | 232.038 055 3(21)                             | 100                                                    | 0+                                         | 0                                                           |                                                                |
| 91       | Pa            | 231*     | 231.035 884 0(24)                             | 100                                                    | 3/2−                                       | 2.01(2)                                                     |                                                                |
| 92       | U             | 233*     | 233.039 635 2(29)                             |                                                        | 5/2+                                       | 0.59(5)                                                     | 366.3(8)*                                                      |
|          |               | 234#     | 234.040 952 2(20)                             | 0.0054(5)                                              | 0+                                         | 0                                                           |                                                                |
|          |               | 235#     | 235.043 929 9(20)                             | 0.7204(6)                                              | 7/2−                                       | −0.38(3)*                                                   | 493.6(6)*                                                      |
|          |               | 238#     | 238.050 788 2(20)                             | 99.2742(10)                                            | 0+                                         | 0                                                           |                                                                |
| 93       | Np            | 237*     | 237.048 173 4(20)                             |                                                        | 5/2+                                       | +3.14(4)*                                                   | +386.6(6)                                                      |
| 94       | Pu            | 244*     | 244.064 204(5)                                |                                                        | 0+                                         |                                                             |                                                                |
| 95       | Am            | 243*     | 243.061 381 1(25)                             |                                                        | 5/2−                                       | +1.503(14)                                                  | +286.0(30)*                                                    |
| 96       | Cm            | 247*     | 247.070 354(5)                                |                                                        | 9/2−                                       | 0.36(7)                                                     |                                                                |
| 97       | Bk            | 247*     | 247.070 307(6)                                |                                                        |                                            |                                                             |                                                                |
| 98       | Cf            | 251*     | 251.079 587(5)                                |                                                        |                                            |                                                             |                                                                |
| 99       | Es            | 252*     | 252.082 980(50)                               |                                                        |                                            |                                                             |                                                                |
|          |               | 253*     | 253.084 824 7(28)                             |                                                        | 7/2+                                       | +4.10(7)                                                    | 670.0(800)                                                     |
| 100      | Fm            | 257*     | 257.095 105(7)                                |                                                        |                                            |                                                             |                                                                |
| 101      | Md            | 258*     | 258.098 431(5)                                |                                                        |                                            |                                                             |                                                                |
| 102      | No            | 259*     | 259.101 03(11)                                |                                                        |                                            |                                                             |                                                                |
| 103      | Lr            | 262*     | 262.109 63(22)                                |                                                        |                                            |                                                             |                                                                |
| 104      | Rf            | 261*     | 261.108 77(3)                                 |                                                        |                                            |                                                             |                                                                |
| 105      | Db            | 262*     | 262.114 08(20)                                |                                                        |                                            |                                                             |                                                                |
| 106      | Sg            | 263*     | 263.118 32(13)                                |                                                        |                                            |                                                             |                                                                |
| 107      | Bh            | 264*     | 264.1246(3)                                   |                                                        |                                            |                                                             |                                                                |
| 108      | Hs            | 265*     | 265.130 09(15)                                |                                                        |                                            |                                                             |                                                                |
| 109      | Mt            | 268*     | 268.138 73(34)                                |                                                        |                                            |                                                             |                                                                |
| 110      | Ds            | 271*     | 271.146 06(11)                                |                                                        |                                            |                                                             |                                                                |
| 111      | Rg            | 272*     | 272.153 62(36)                                |                                                        |                                            |                                                             |                                                                |

## 7 CONVERSION OF UNITS

Units of the SI are recommended for use throughout science and technology. However, some non-rationalized units are in use, and in a few cases they are likely to remain so for many years. Moreover, the published literature of science makes widespread use of non-SI units. It is thus often necessary to convert the values of physical quantities between SI units and other units. This chapter is concerned with facilitating this process, as well as the conversion of units in general.

Section 7.1, p. 131 gives examples illustrating the use of quantity calculus for converting the numerical values of physical quantities expressed in different units. The table in Section 7.2, p. 135 lists a variety of non-rationalized units used in chemistry, with the conversion factors to the corresponding SI units. Transformation factors for energy and energy-related units (repetency, wavenumber, frequency, temperature and molar energy), and for pressure units, are also presented in tables in the back of this manual.

Many of the difficulties in converting units between different systems are associated either with the electromagnetic units, or with atomic units and their relation to the electromagnetic units. In Sections 7.3 and 7.4, p. 143 and 146 the relations involving electromagnetic and atomic units are developed in greater detail to provide a background for the conversion factors presented in the table in Section 7.2, p. 135.



## 7.1 THE USE OF QUANTITY CALCULUS

Quantity calculus is a system of algebra in which symbols are consistently used to represent physical quantities and not their numerical values expressed in certain units. Thus we always take the values of physical quantities to be the product of a numerical value and a unit (see Section 1.1, p. 3), and we manipulate the symbols for physical quantities, numerical values, and units by the ordinary rules of algebra (see footnote <sup>1</sup>, below). This system is recommended for general use in science and technology. Quantity calculus has particular advantages in facilitating the problems of converting between different units and different systems of units. Another important advantage of quantity calculus is that equations between quantities are independent of the choice of units, and must always satisfy the rule that the dimensions must be the same for each term on either side of the equal sign. These advantages are illustrated in the examples below, where the numerical values are approximate.

*Example 1.* The wavelength  $\lambda$  of one of the yellow lines of sodium is given by

$$\lambda \approx 5.896 \times 10^{-7} \text{ m}, \quad \text{or} \quad \lambda/\text{m} \approx 5.896 \times 10^{-7}$$

The ångström is defined by the equation (see Section 7.2, “length”, p. 135)

$$1 \text{ Å} = \text{Å} := 10^{-10} \text{ m}, \quad \text{or} \quad \text{m}/\text{Å} := 10^{10}$$

Substituting in the first equation gives the value of  $\lambda$  in ångström

$$\lambda/\text{Å} = (\lambda/\text{m}) (\text{m}/\text{Å}) \approx (5.896 \times 10^{-7}) (10^{10}) = 5896$$

or

$$\lambda \approx 5896 \text{ Å}$$

*Example 2.* The vapor pressure of water at 20 °C is recorded to be

$$p(\text{H}_2\text{O}, 20 \text{ °C}) \approx 17.5 \text{ Torr}$$

The torr, the bar, and the atmosphere are given by the equations (see Section 7.2, “pressure”, p. 138)

$$\begin{aligned} 1 \text{ Torr} &\approx 133.3 \text{ Pa} \\ 1 \text{ bar} &:= 10^5 \text{ Pa} \\ 1 \text{ atm} &:= 101\,325 \text{ Pa} \end{aligned}$$

Thus

$$\begin{aligned} p(\text{H}_2\text{O}, 20 \text{ °C}) &\approx 17.5 \times 133.3 \text{ Pa} \approx 2.33 \text{ kPa} = \\ &(2.33 \times 10^3/10^5) \text{ bar} = 23.3 \text{ mbar} = \\ &(2.33 \times 10^3/101\,325) \text{ atm} \approx 2.30 \times 10^{-2} \text{ atm} \end{aligned}$$

*Example 3.* Spectroscopic measurements show that for the methylene radical,  $\text{CH}_2$ , the  $\tilde{\text{a}} \text{ } ^1\text{A}_1$  excited state lies at a repetency (wavenumber)  $3156 \text{ cm}^{-1}$  above the  $\tilde{\text{X}} \text{ } ^3\text{B}_1$  ground state

$$\tilde{\nu}(\tilde{\text{a}} - \tilde{\text{X}}) = T_0(\tilde{\text{a}}) - T_0(\tilde{\text{X}}) \approx 3156 \text{ cm}^{-1}$$

The excitation energy from the ground triplet state to the excited singlet state is thus

$$\begin{aligned} \Delta E = hc_0\tilde{\nu} &\approx (6.626 \times 10^{-34} \text{ J s}) (2.998 \times 10^8 \text{ m s}^{-1}) (3156 \text{ cm}^{-1}) \approx \\ &6.269 \times 10^{-22} \text{ J m cm}^{-1} = \\ &6.269 \times 10^{-20} \text{ J} = 6.269 \times 10^{-2} \text{ aJ} \end{aligned}$$

---

<sup>1</sup> A more appropriate name for “quantity calculus” might be “algebra of quantities”, because the principles of algebra rather than calculus (in the sense of differential and integral calculus) are involved.

where the values of  $h$  and  $c_0$  are taken from the fundamental physical constants in Chapter 5, p. 111 and we have used the relation  $1 \text{ m} = 100 \text{ cm}$ , or  $1 \text{ m} = 10^2 \text{ cm}$ . Since the electronvolt is given by the equation (Section 7.2, “energy”, p. 137)  $1 \text{ eV} \approx 1.6022 \times 10^{-19} \text{ J}$ , or  $1 \text{ aJ} \approx (1/0.16022) \text{ eV}$ ,

$$\Delta E \approx (6.269 \times 10^{-2} / 0.16022) \text{ eV} \approx 0.3913 \text{ eV}$$

Similarly the hartree is given by  $E_h = \hbar^2 / m_e a_0^2 \approx 4.3597 \text{ aJ}$ , or  $1 \text{ aJ} \approx (1/4.3597) E_h$  (Section 3.9.1, p. 94), and thus the excitation energy is given in atomic units by

$$\Delta E \approx (6.269 \times 10^{-2} / 4.3597) E_h \approx 1.4379 \times 10^{-2} E_h$$

Finally the molar excitation energy is given by

$$\Delta E_m = L \Delta E \approx (6.022 \times 10^{23} \text{ mol}^{-1})(6.269 \times 10^{-2} \text{ aJ}) \approx 37.75 \text{ kJ mol}^{-1}$$

Also, since  $1 \text{ kcal} := 4.184 \text{ kJ}$ , or  $1 \text{ kJ} := (1/4.184) \text{ kcal}$ ,

$$\Delta E_m \approx (37.75 / 4.184) \text{ kcal mol}^{-1} \approx 9.023 \text{ kcal mol}^{-1}$$

In the transformation from  $\Delta E$  to  $\Delta E_m$  the coefficient  $L$  is not a number, but has a dimension different from one. Also in this example the necessary transformation coefficient could have been taken directly from the table in the back of this manual.

*Example 4.* The molar conductivity,  $\Lambda$ , of an electrolyte is defined by the equation

$$\Lambda = \kappa / c$$

where  $\kappa$  is the conductivity of the electrolyte solution minus the conductivity of the pure solvent and  $c$  is the electrolyte concentration. Conductivities of electrolytes are usually expressed in  $\text{S cm}^{-1}$  and concentrations in  $\text{mol dm}^{-3}$ ; for example  $\kappa(\text{KCl}) \approx 7.39 \times 10^{-5} \text{ S cm}^{-1}$  for  $c(\text{KCl}) \approx 0.000500 \text{ mol dm}^{-3}$ . The molar conductivity can then be calculated as follows

$$\begin{aligned} \Lambda &\approx (7.39 \times 10^{-5} \text{ S cm}^{-1}) / (0.000500 \text{ mol dm}^{-3}) \approx \\ &0.1478 \text{ S mol}^{-1} \text{ cm}^{-1} \text{ dm}^3 = 147.8 \text{ S mol}^{-1} \text{ cm}^2 \end{aligned}$$

since  $1 \text{ dm}^3 = 1000 \text{ cm}^3$ . The above relation has previously often been, and sometimes still is, written in the not-recommended form

$$\Lambda = 1000 \kappa / c$$

However, in this form the symbols *do not* represent physical quantities, but the *numerical values* of physical quantities expressed in certain units. Specifically, the last equation is true only if  $\Lambda$  is the numerical value of the molar conductivity in  $\text{S mol}^{-1} \text{ cm}^2$ ,  $\kappa$  is the numerical value of the conductivity in  $\text{S cm}^{-1}$ , and  $c$  is the numerical value of the concentration in  $\text{mol dm}^{-3}$ . This form does not follow the rules of quantity calculus, and should be avoided. The equation  $\Lambda = \kappa / c$ , in which the symbols represent physical quantities, is true in any units. If it is desired to write the relation between numerical values it should be written in the form

$$\Lambda / (\text{S mol}^{-1} \text{ cm}^2) = \frac{1000 \kappa / (\text{S cm}^{-1})}{c / (\text{mol dm}^{-3})}$$

*Example 5.* A solution of 0.125 mol of solute B in  $m_S \approx 953$  g of solvent S has a molality  $b_B$  given by (see also footnote <sup>2</sup>, below)

$$b_B = n_B/m_S \approx (0.125/953) \text{ mol g}^{-1} \approx 0.131 \text{ mol kg}^{-1}$$

The amount-of-substance fraction of solute is approximately given by

$$x_B = n_B/(n_S + n_B) \approx n_B/n_S = b_B M_S$$

where it is assumed that  $n_B \ll n_S$ .

If the solvent is water with molar mass  $18.015 \text{ g mol}^{-1}$ , then

$$x_B \approx (0.131 \text{ mol kg}^{-1}) (18.015 \text{ g mol}^{-1}) \approx 2.36 \text{ g/kg} = 0.00236$$

These equations are sometimes quoted in the deprecated form  $b_B = 1000 n_B/M_S$ , and  $x_B \approx b_B M_S/1000$ . However, this is *not* a correct use of quantity calculus because in this form the symbols denote the *numerical values* of the physical quantities in particular units; specifically it is assumed that  $b_B$ ,  $m_S$  and  $M_S$  denote numerical values in  $\text{mol kg}^{-1}$ , g, and  $\text{g mol}^{-1}$  respectively. A correct way of writing the second equation would, for example, be

$$x_B = (b_B/\text{mol kg}^{-1})(M_S/\text{g mol}^{-1})/1000$$

*Example 6.* For paramagnetic materials the magnetic susceptibility may be measured experimentally and used to give information on the molecular magnetic dipole moment, and hence on the electronic structure of the molecules in the material. The paramagnetic contribution to the molar magnetic susceptibility of a material,  $\chi_m$ , is related to the molecular magnetic dipole moment  $m$  by the Curie relation

$$\chi_m = \chi V_m = \mu_0 N_A m^2 / 3kT$$

In the older non-rationalized esu, emu, and Gaussian systems (see Section 7.3, p. 143), this equation becomes

$$\chi_m^{(\text{ir})} = \chi^{(\text{ir})} V_m = N_A m^2 / 3kT$$

Solving for  $m$ , and expressing the result in terms of the Bohr magneton  $\mu_B$ , in the ISQ(SI)

$$m/\mu_B = (3k/\mu_0 N_A)^{1/2} \mu_B^{-1} (\chi_m T)^{1/2}$$

and in the non-rationalized emu and Gaussian systems

$$m/\mu_B = (3k/N_A)^{1/2} \mu_B^{-1} (\chi_m^{(\text{ir})} T)^{1/2}$$

Finally, using the values of the fundamental physical constants  $\mu_B$ ,  $k$ ,  $\mu_0$ , and  $N_A$  given in Chapter 5, p. 111, we obtain

$$m/\mu_B \approx 0.7977 [\chi_m / (\text{cm}^3 \text{ mol}^{-1})]^{1/2} [T/\text{K}]^{1/2} \approx 2.828 [\chi_m^{(\text{ir})} / (\text{cm}^3 \text{ mol}^{-1})]^{1/2} [T/\text{K}]^{1/2}$$

These expressions are convenient for practical calculations. The final result has frequently been expressed in the not recommended form

$$m/\mu_B \approx 2.828 (\chi_m T)^{1/2}$$

where it is assumed, contrary to the conventions of quantity calculus, that  $\chi_m$  and  $T$  denote the *numerical values* of the molar susceptibility and the temperature in the units  $\text{cm}^3 \text{ mol}^{-1}$  and K, respectively, and where it is also assumed (but rarely stated) that the susceptibility is defined using the electromagnetic equations defined within non-rationalized systems (see Section 7.3, p. 143).

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<sup>2</sup> We use  $b_B$  because of the possible confusion of notation when using  $m_B$  to denote molality, and  $m_S$  to denote the mass of S. However, the symbol  $m_B$  is frequently used to denote molality (see Section 2.10, note 14, p. 48).

*Example 7.* The esu or Gaussian unit of the electric charge is  $\sqrt{\text{erg cm}}$ . Alternatively, the franklin (Fr) is used to represent  $\sqrt{\text{erg cm}}$  (see Section 7.3, p. 143). The definition of the franklin states that two particles, each having an electric charge of one franklin and separated by a distance of one centimetre, repel each other with a force of one dyn. In order to obtain a conversion factor to the SI unit coulomb, one needs to ask what charges produce the same force in SI units at the same distance. One has thus

$$1 \text{ dyn} = 1 \text{ cm g s}^{-2} = 10^{-5} \text{ N} = \frac{1}{4\pi\epsilon_0} \frac{Q^2}{(10^{-2} \text{ m})^2}$$

It follows (see Section 7.3, p. 143) that

$$Q^2 = 4\pi\epsilon_0 \times 10^{-9} \text{ N m}^2 = \mu_0\epsilon_0 \times 10^{-2} \text{ A}^2 \text{ m}^2 = c_0^{-2} \times 10^{-2} \text{ A}^2 \text{ m}^2$$

and thus

$$Q = \frac{1}{10 c_0} \text{ A m} = \frac{10}{\zeta} \text{ C}$$

where  $\zeta$  is the exact number  $\zeta = c_0/(\text{cm s}^{-1}) = 29\,979\,245\,800$  (see Sections 7.2 and 7.3, p. 135 and 143). Thus  $1 \text{ Fr} = 1/(2.997\,924\,58 \times 10^9) \text{ C}$ .

*Example 8.* The esu or Gaussian unit of the electric field strength is  $\sqrt{\text{erg cm}} \text{ cm}^{-2}$  or, alternatively,  $\text{Fr cm}^{-2}$ . Simple insertion of the above mentioned relation between the franklin and the coulomb would yield  $1 \text{ Fr cm}^{-2} = (10^5/\zeta) \text{ C m}^{-2}$ . However,  $\text{C m}^{-2}$  is not the unit of the electric field strength in the SI. In order to obtain the conversion factor for the electric field strength in the SI, one has to set

$$1 \frac{\text{Fr}}{\text{cm}^2} = \frac{1}{4\pi\epsilon_0} \times \frac{10^5}{\zeta} \frac{\text{C}}{\text{m}^2}$$

From Example 7 one has  $4\pi\epsilon_0 \times 10^{-9} \text{ N m}^2 = (10/\zeta)^2 \text{ C}^2$ , thus

$$\frac{10^5 \text{ C}}{4\pi\epsilon_0 \zeta \text{ m}^2} = \zeta \times 10^{-6} \frac{\text{N}}{\text{C}}$$

One concludes that  $1 \text{ Fr cm}^{-2} = \zeta \times 10^{-6} \text{ V m}^{-1}$ .

*Example 9.* The esu or Gaussian unit of the polarizability is  $\text{cm}^3$ : in a system of charges with polarizability  $1 \text{ cm}^3$ , a field of  $1 \text{ Fr/cm}^2$  produces a dipole moment of  $1 \text{ Fr cm}$  (see Section 7.3, p. 143). The corresponding value of this polarizability in the SI is obtained as follows:

$$1 \text{ cm}^3 = \frac{p}{E} = \frac{(10/\zeta) \text{ C } 10^{-2} \text{ m}}{\zeta 10^{-6} \text{ V m}^{-1}} = \zeta^{-2} 10^5 \text{ m}^2 \text{ C V}^{-1}$$

One has thus  $1 \text{ cm}^3 = (10^5/\zeta^2) \text{ m}^2 \text{ C}^2 \text{ J}^{-1}$ , from the definition of the volt. This equation seems strange, at a first sight. It does not give a general relation between a volume of  $1 \text{ cm}^3$  and other units in the SI. Rather it states that a polarizability that has the value  $1 \text{ cm}^3$  in the esu or Gaussian system has the value  $(10^5/\zeta^2) \text{ m}^2 \text{ C}^2 \text{ J}^{-1}$  in the SI.

## 7.2 CONVERSION TABLES FOR UNITS

The table below gives conversion factors from a variety of units to the corresponding SI unit. Examples of the use of this table have already been given in the preceding section. For each physical quantity the name is given, followed by the recommended symbol(s). Then the SI unit is given, followed by the esu, emu, Gaussian unit, atomic unit (au), and other units in common use, with their conversion factors to SI and other units. The constant  $\zeta$  which occurs in some of the electromagnetic conversion factors is the exact number  $29\,979\,245\,800 = c_0/(\text{cm s}^{-1})$ .

The inclusion of non-SI units in this table should not be taken to imply that their use is to be encouraged. With some exceptions, SI units are always to be preferred to non-SI units. However, since many of the units below are to be found in the scientific literature, it is convenient to tabulate their relation to the SI.

For convenience units in the esu, emu and Gaussian systems are quoted in terms of the four dimensions *length*, *mass*, *time*, and *electric charge*, by including the franklin (Fr) as the electrostatic unit of charge and the biot (Bi) as the electromagnetic unit of current. This gives each physical quantity the same dimensions in all systems, so that all conversion factors are numbers. The factors  $4\pi\epsilon_0 = k_{\text{es}}^{-1}$  and the Fr may be eliminated by writing  $\text{Fr} = \text{erg}^{1/2} \text{ cm}^{1/2} = \text{cm}^{3/2} \text{ g}^{1/2} \text{ s}^{-1}$ , and  $k_{\text{es}} = 1 \text{ Fr}^{-2} \text{ erg cm} = 1$ , to recover esu expressions in terms of three base units (see Section 7.3, p. 143). The symbol Fr should be regarded as a symbol for the electrostatic unit of charge. Similarly, the factor  $\mu_0/4\pi = k_{\text{em}}$  and the Bi may be eliminated by using  $\text{Bi} = \text{dyn}^{1/2} = \text{cm}^{1/2} \text{ g}^{1/2} \text{ s}^{-1}$ , and  $k_{\text{em}} = \text{dyn Bi}^{-2} = 1$ , to recover emu expressions in terms of the three base units.

The table must be read in the following way, for example for the fermi, as a unit of length:  $\text{f} = 10^{-15} \text{ m}$  means that the symbol f of a length 13.1 f may be replaced by  $10^{-15} \text{ m}$ , saying that the length has the value  $13.1 \times 10^{-15} \text{ m}$ . A more difficult example involves for instance the polarizability  $\alpha$ . When  $\alpha$  is  $17.8 \text{ cm}^3$  in the Gaussian system, it is  $(17.8 \times 10^5/\zeta^2) \text{ m}^2 \text{ C}^2 \text{ J}^{-1} \approx 3.53 \times 10^{-9} \text{ m}^2 \text{ C}^2 \text{ J}^{-1}$  in the SI. That is, the symbol  $\text{cm}^3$  may be replaced by the expression  $(10^5/\zeta^2) \text{ m}^2 \text{ C}^2 \text{ J}^{-1}$  to obtain the value of the polarizability in the SI (see also the Example 9 in Section 7.1, p. 134). Conversion factors are either given exactly (when the = sign is used; if it is a definition, the sign := is used), or they are given to the approximation that the corresponding physical constants are known (when the sign  $\approx$  is used). In the latter case the magnitude of the uncertainty is always less than 5 in the last digit quoted, if the uncertainty is not explicitly indicated.

| <i>Name</i>           | <i>Symbol</i> | <i>Expressed in SI units</i>                                                               | <i>Notes</i> |
|-----------------------|---------------|--------------------------------------------------------------------------------------------|--------------|
| <i>length, l</i>      |               |                                                                                            |              |
| metre (SI unit)       | m             |                                                                                            |              |
| centimetre (CGS unit) | cm            | $= 10^{-2} \text{ m}$                                                                      |              |
| bohr (au)             | $a_0$         | $= 4\pi\epsilon_0\hbar^2/m_e e^2 \approx 5.291\,772\,085\,9(36) \times 10^{-11} \text{ m}$ |              |
| ångström              | Å             | $= 10^{-10} \text{ m}$                                                                     |              |
| micron                | $\mu$         | $= 1 \mu\text{m} = 10^{-6} \text{ m}$                                                      |              |
| millimicron           | m $\mu$       | $= 1 \text{ nm} = 10^{-9} \text{ m}$                                                       |              |
| x unit                | X             | $\approx 1.002 \times 10^{-13} \text{ m}$                                                  |              |
| fermi                 | f             | $= 1 \text{ fm} = 10^{-15} \text{ m}$                                                      |              |
| inch                  | in            | $= 2.54 \times 10^{-2} \text{ m}$                                                          |              |
| foot                  | ft            | $= 12 \text{ in} = 0.3048 \text{ m}$                                                       |              |
| yard                  | yd            | $= 3 \text{ ft} = 0.9144 \text{ m}$                                                        |              |
| mile                  | mi            | $= 1760 \text{ yd} = 1609.344 \text{ m}$                                                   |              |
| nautical mile         |               | $= 1852 \text{ m}$                                                                         |              |

| <i>Name</i>                  | <i>Symbol</i>         | <i>Expressed in SI units</i>                                                              | <i>Notes</i> |
|------------------------------|-----------------------|-------------------------------------------------------------------------------------------|--------------|
| <i>length, l</i> (continued) |                       |                                                                                           |              |
| astronomical unit            | ua                    | $\approx 1.495\ 978\ 706\ 91(6)\times 10^{11}\ \text{m}$                                  | 1            |
| parsec                       | pc                    | $\approx 3.085\ 678\times 10^{16}\ \text{m}$                                              |              |
| light year                   | l.y.                  | $\approx 9.460\ 730\times 10^{15}\ \text{m}$                                              |              |
| light second                 |                       | $= 299\ 792\ 458\ \text{m}$                                                               |              |
| <i>area, A</i>               |                       |                                                                                           |              |
| square metre (SI unit)       | m <sup>2</sup>        |                                                                                           |              |
| barn                         | b                     | $= 10^{-28}\ \text{m}^2$                                                                  |              |
| acre                         |                       | $\approx 4046.856\ \text{m}^2$                                                            |              |
| are                          | a                     | $= 100\ \text{m}^2$                                                                       |              |
| hectare                      | ha                    | $= 10^4\ \text{m}^2$                                                                      |              |
| <i>volume, V</i>             |                       |                                                                                           |              |
| cubic metre (SI unit)        | m <sup>3</sup>        |                                                                                           | 2            |
| litre                        | l, L                  | $= 1\ \text{dm}^3 = 10^{-3}\ \text{m}^3$                                                  |              |
| lambda                       | λ                     | $= 10^{-6}\ \text{dm}^3\ [= 1\ \mu\text{l}]$                                              |              |
| barrel (US)                  |                       | $= 158.9873\ \text{dm}^3$                                                                 |              |
| gallon (US)                  | gal (US)              | $= 3.785\ 412\ \text{dm}^3$                                                               |              |
| gallon (UK)                  | gal (UK)              | $= 4.546\ 092\ \text{dm}^3$                                                               |              |
| <i>plane angle, α</i>        |                       |                                                                                           |              |
| radian (SI unit)             | rad                   |                                                                                           |              |
| degree                       | °, deg                | $= (\pi/180)\ \text{rad} \approx (1/57.295\ 78)\ \text{rad}$                              |              |
| minute                       | '                     | $= (\pi/10\ 800)\ \text{rad}\ [= (1/60)^\circ]$                                           |              |
| second                       | "                     | $= (\pi/648\ 000)\ \text{rad}\ [= (1/3600)^\circ]$                                        |              |
| gon                          | gon                   | $= (\pi/200)\ \text{rad} \approx (1/63.661\ 98)\ \text{rad}$                              |              |
| <i>mass, m</i>               |                       |                                                                                           |              |
| kilogram (SI unit)           | kg                    |                                                                                           |              |
| gram (CGS unit)              | g                     | $= 10^{-3}\ \text{kg}$                                                                    |              |
| electron mass (au)           | <i>m</i> <sub>e</sub> | $\approx 9.109\ 382\ 15(45)\times 10^{-31}\ \text{kg}$                                    |              |
| dalton,                      | Da, u                 | $:= m_{\text{a}}(^{12}\text{C})/12 \approx 1.660\ 538\ 782(83)\times 10^{-27}\ \text{kg}$ |              |
| unified atomic mass unit     |                       |                                                                                           |              |
| gamma                        | γ                     | $= 1\ \mu\text{g}$                                                                        |              |
| tonne, (metric tonne)        | t                     | $= 1\ \text{Mg} = 10^3\ \text{kg}$                                                        |              |
| pound (avoirdupois)          | lb                    | $= 0.453\ 592\ 37\ \text{kg}$                                                             |              |
| pound, (metric pound)        |                       | $= 0.5\ \text{kg}$                                                                        |              |
| ounce (avoirdupois)          | oz                    | $= 28.349\ 52\ \text{g}$                                                                  |              |
| ounce (troy)                 | oz (troy)             | $= 31.103\ 476\ 8\ \text{g}$                                                              |              |
| grain                        | gr                    | $= 64.798\ 91\ \text{mg}$                                                                 |              |

(1) This value refers to the Julian year (1 a = 365.25 d, 1 d = 86 400 s) [152]. l.y. is not a symbol according to the syntax rules of Section 1.3, p. 5, but is an often used abbreviation.

(2) ISO and IEC only use the lower case l for the litre.

| <i>Name</i>                           | <i>Symbol</i>     | <i>Expressed in SI units</i>                                              | <i>Notes</i> |
|---------------------------------------|-------------------|---------------------------------------------------------------------------|--------------|
| <i>time, t</i>                        |                   |                                                                           |              |
| second (SI unit, CGS unit)            | s                 |                                                                           |              |
| au of time                            | $\hbar/E_h$       | $\approx 2.418\,884\,326\,505(16)\times 10^{-17}\text{ s}$                |              |
| minute                                | min               | = 60 s                                                                    |              |
| hour                                  | h                 | = 3600 s                                                                  |              |
| day                                   | d                 | = 24 h = 86 400 s                                                         | 3            |
| year                                  | a                 | $\approx 31\,556\,952\text{ s}$                                           | 4            |
| svedberg                              | Sv                | = $10^{-13}\text{ s}$                                                     |              |
| <i>acceleration, a</i>                |                   |                                                                           |              |
| SI unit                               | $\text{m s}^{-2}$ |                                                                           |              |
| standard acceleration of<br>free fall | $g_n$             | = $9.806\,65\text{ m s}^{-2}$                                             |              |
| gal (CGS unit)                        | Gal               | = $10^{-2}\text{ m s}^{-2}$                                               |              |
| <i>force, F</i>                       |                   |                                                                           |              |
| newton (SI unit)                      | N                 | = $1\text{ kg m s}^{-2}$                                                  | 5            |
| dyne (CGS unit)                       | dyn               | = $1\text{ g cm s}^{-2} = 10^{-5}\text{ N}$                               |              |
| au of force                           | $E_h/a_0$         | $\approx 8.238\,722\,06(41)\times 10^{-8}\text{ N}$                       |              |
| kilogram-force, kilopond              | kgf = kp          | = $9.806\,65\text{ N}$                                                    |              |
| <i>energy, E, U</i>                   |                   |                                                                           |              |
| joule (SI unit)                       | J                 | = $1\text{ kg m}^2\text{ s}^{-2}$                                         |              |
| erg (CGS unit)                        | erg               | = $1\text{ g cm}^2\text{ s}^{-2} = 10^{-7}\text{ J}$                      |              |
| hartree (au)                          | $E_h$             | = $\hbar^2/m_e a_0^2 \approx 4.359\,743\,94(22)\times 10^{-18}\text{ J}$  |              |
| rydberg                               | Ry                | = $E_h/2 \approx 2.179\,871\,97(11)\times 10^{-18}\text{ J}$              |              |
| electronvolt                          | eV                | = $e\cdot 1\text{ V} \approx 1.602\,176\,487(40)\times 10^{-19}\text{ J}$ |              |
| calorie, thermochemical               | cal <sub>th</sub> | = $4.184\text{ J}$                                                        |              |
| calorie, international                | cal <sub>IT</sub> | = $4.1868\text{ J}$                                                       |              |
| 15 °C calorie                         | cal <sub>15</sub> | $\approx 4.1855\text{ J}$                                                 |              |
| litre atmosphere                      | l atm             | = $101.325\text{ J}$                                                      |              |
| British thermal unit                  | Btu               | = $1055.06\text{ J}$                                                      |              |

(3) Note that the astronomical day is not exactly defined in terms of the second since so-called leap-seconds are added or subtracted from the day semiannually in order to keep the annual average occurrence of midnight at 24:00:00 on the clock. However, in reference [3] hour and day are defined in terms of the second as given.

(4) The year is not commensurable with the day and not a constant. Prior to 1967, when the atomic standard was introduced, the tropical year 1900 served as the basis for the definition of the second. For the epoch 1900.0 it amounted to  $365.242\,198\,79\text{ d} \approx 31\,556\,925.98\text{ s}$  and it decreases by  $0.530\text{ s}$  per century. The calendar years are exactly defined in terms of the day: The **Julian** year is equal to  $365.25\text{ d}$ ; the **Gregorian** year is equal to  $365.2425\text{ d}$ ; the **Mayan** year is equal to  $365.2420\text{ d}$ . The definition in the table corresponds to the Gregorian year. This is an average based on a year of length  $365\text{ d}$ , with leap years of  $366\text{ d}$ ; leap years are taken *either* when the year is divisible by 4 but is not divisible by 100, or when the year is divisible by 400. Whether the year 3200 should be a leap year is still open, but this does not have to be resolved until sometime in the middle of the 32nd century. For conversion one may use in general approximately  $1\text{ a} \approx 3.1557\times 10^7\text{ s}$ . If more accurate statements are needed, the precise definition of the year used should be stated.

| <i>Name</i>                                  | <i>Symbol</i>               | <i>Expressed in SI units</i>                                                         | <i>Notes</i> |
|----------------------------------------------|-----------------------------|--------------------------------------------------------------------------------------|--------------|
| <i>pressure, p</i>                           |                             |                                                                                      |              |
| pascal (SI unit)                             | Pa                          | $= 1 \text{ N m}^{-2} = 1 \text{ kg m}^{-1} \text{ s}^{-2}$                          |              |
| standard atmosphere                          | atm                         | $= 101\,325 \text{ Pa}$                                                              |              |
| bar                                          | bar                         | $= 10^5 \text{ Pa}$                                                                  |              |
| torr                                         | Torr                        | $= (101\,325/760) \text{ Pa} \approx 133.322 \text{ Pa}$                             |              |
| conventional millimetre<br>of mercury        | mmHg                        | $= 13.5951 \times 980.665 \times 10^{-2} \text{ Pa} \approx$<br>$133.322 \text{ Pa}$ |              |
| pounds per square inch                       | psi                         | $\approx 6.894\,758 \times 10^3 \text{ Pa}$                                          |              |
| <i>power, P</i>                              |                             |                                                                                      |              |
| watt (SI unit)                               | W                           | $= 1 \text{ kg m}^2 \text{ s}^{-3}$                                                  |              |
| Imperial horse power                         | hp                          | $\approx 745.7 \text{ W}$                                                            |              |
| metric horse power                           | hk                          | $= 735.498\,75 \text{ W}$                                                            |              |
| <i>action, angular momentum, L, J</i>        |                             |                                                                                      |              |
| SI unit                                      | J s                         | $= 1 \text{ kg m}^2 \text{ s}^{-1}$                                                  |              |
| CGS unit                                     | erg s                       | $= 10^{-7} \text{ J s}$                                                              |              |
| au of action                                 | $\hbar$                     | $:= h/2\pi \approx 1.054\,571\,628(53) \times 10^{-34} \text{ J s}$                  |              |
| <i>dynamic viscosity, <math>\eta</math></i>  |                             |                                                                                      |              |
| SI unit                                      | Pa s                        | $= 1 \text{ kg m}^{-1} \text{ s}^{-1}$                                               |              |
| poise (CGS unit)                             | P                           | $= 10^{-1} \text{ Pa s}$                                                             |              |
| centipoise                                   | cP                          | $= 1 \text{ mPa s}$                                                                  |              |
| <i>kinematic viscosity, <math>\nu</math></i> |                             |                                                                                      |              |
| SI unit                                      | $\text{m}^2 \text{ s}^{-1}$ |                                                                                      |              |
| stokes (CGS unit)                            | St                          | $= 10^{-4} \text{ m}^2 \text{ s}^{-1}$                                               |              |
| <i>thermodynamic temperature, T</i>          |                             |                                                                                      |              |
| kelvin (SI unit)                             | K                           |                                                                                      |              |
| degree Rankine                               | $^{\circ}\text{R}$          | $= (5/9) \text{ K}$                                                                  | 6            |
| <i>Celsius temperature, t</i>                |                             |                                                                                      |              |
| degree Celsius (SI unit)                     | $^{\circ}\text{C}$          | $= 1 \text{ K}$                                                                      | 6            |
| <i>entropy, S</i>                            |                             |                                                                                      |              |
| <i>heat capacity, C</i>                      |                             |                                                                                      |              |
| SI unit                                      | $\text{J K}^{-1}$           |                                                                                      |              |
| clausius                                     | Cl                          | $= 4.184 \text{ J K}^{-1} [= 1 \text{ cal}_{\text{th}} \text{ K}^{-1}]$              |              |

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(Notes continued)

(5) 1 N is approximately the force exerted by the Earth upon an apple.

(6)  $T/^{\circ}\text{R} = (9/5)T/\text{K}$ . Also, Celsius temperature  $t$  is related to thermodynamic temperature  $T$  by the equation

$$t/^{\circ}\text{C} = T/\text{K} - 273.15$$

Similarly Fahrenheit temperature  $t_{\text{F}}$  is related to Celsius temperature  $t$  by the equation

$$t_{\text{F}}/^{\circ}\text{F} = (9/5)(t/^{\circ}\text{C}) + 32$$

| <i>Name</i>                                       | <i>Symbol</i>                     | <i>Expressed in SI units</i>                                                                                    | <i>Notes</i> |
|---------------------------------------------------|-----------------------------------|-----------------------------------------------------------------------------------------------------------------|--------------|
| <i>molar entropy, <math>S_m</math></i>            |                                   |                                                                                                                 |              |
| <i>molar heat capacity, <math>C_m</math></i>      |                                   |                                                                                                                 |              |
| SI unit                                           | $\text{J K}^{-1} \text{mol}^{-1}$ | $= 1 \text{ kg m}^2 \text{s}^{-2} \text{K}^{-1} \text{mol}^{-1}$                                                |              |
| entropy unit                                      |                                   | $= 4.184 \text{ J K}^{-1} \text{mol}^{-1} =$<br>$[1 \text{ cal}_{\text{th}} \text{K}^{-1} \text{mol}^{-1}]$     |              |
| <i>molar volume, <math>V_m</math></i>             |                                   |                                                                                                                 |              |
| SI unit                                           | $\text{m}^3 \text{mol}^{-1}$      |                                                                                                                 |              |
| amagat                                            |                                   | $:= V_m$ of real gas at 1 atm<br>and $273.15 \text{ K} \approx 22.4 \times 10^{-3} \text{ m}^3 \text{mol}^{-1}$ |              |
| <i>amount density, <math>1/V_m</math></i>         |                                   |                                                                                                                 |              |
| SI unit                                           | $\text{mol m}^{-3}$               |                                                                                                                 |              |
| reciprocal amagat                                 |                                   | $:= 1/V_m$ of real gas at 1 atm<br>and $273.15 \text{ K} \approx 44.6 \text{ mol m}^{-3}$                       | 7            |
| <i>activity, <math>A</math></i>                   |                                   |                                                                                                                 |              |
| becquerel (SI unit)                               | Bq                                | $= 1 \text{ s}^{-1}$                                                                                            |              |
| curie                                             | Ci                                | $= 3.7 \times 10^{10} \text{ Bq}$                                                                               |              |
| <i>absorbed dose of radiation, <math>D</math></i> |                                   |                                                                                                                 | 8            |
| gray (SI unit)                                    | Gy                                | $= 1 \text{ J kg}^{-1} = 1 \text{ m}^2 \text{s}^{-2}$                                                           |              |
| rad                                               | rad                               | $= 0.01 \text{ Gy}$                                                                                             |              |
| <i>dose equivalent, <math>H</math></i>            |                                   |                                                                                                                 |              |
| sievert (SI unit)                                 | Sv                                | $= 1 \text{ J kg}^{-1} = 1 \text{ m}^2 \text{s}^{-2}$                                                           |              |
| rem                                               | rem                               | $= 0.01 \text{ Sv}$                                                                                             | 9            |
| <i>electric current, <math>I</math></i>           |                                   |                                                                                                                 |              |
| ampere (SI unit)                                  | A                                 |                                                                                                                 |              |
| esu, Gaussian                                     | $\text{Fr s}^{-1}$                | $:= (10/\zeta) \text{A} \approx 3.335\,64 \times 10^{-10} \text{ A}$                                            | 10           |
| biot (emu)                                        | Bi                                | $= 10 \text{ A}$                                                                                                |              |
| au                                                | $eE_h/\hbar$                      | $\approx 6.623\,617\,63(17) \times 10^{-3} \text{ A}$                                                           |              |
| <i>electric charge, <math>Q</math></i>            |                                   |                                                                                                                 |              |
| coulomb (SI unit)                                 | C                                 | $= 1 \text{ A s}$                                                                                               |              |
| franklin (esu, Gaussian)                          | Fr                                | $:= (10/\zeta) \text{C} \approx 3.335\,64 \times 10^{-10} \text{ C}$                                            | 10           |
| emu (abcoulomb)                                   | Bi s                              | $= 10 \text{ C}$                                                                                                |              |
| proton charge (au)                                | $e$                               | $\approx 1.602\,176\,487(40) \times 10^{-19} \text{ C} \approx$<br>$4.803\,21 \times 10^{-10} \text{ Fr}$       |              |

(7) The name “amagat” is unfortunately often used as a unit for both molar volume and amount density. Its value is slightly different for different gases, reflecting the deviation from ideal behaviour for the gas being considered.

(8) The unit röntgen, R, is employed to express exposure to X or  $\gamma$  radiation.  $1 \text{ R} = 2.58 \times 10^{-4} \text{ C kg}^{-1}$ .

(9) rem stands for röntgen equivalent man.

(10)  $\zeta$  is the exact number  $\zeta = c_0/(\text{cm s}^{-1}) = 29\,979\,245\,800$ .

| <i>Name</i>                                      | <i>Symbol</i>                      | <i>Expressed in SI units</i>                                                                         | <i>Notes</i> |
|--------------------------------------------------|------------------------------------|------------------------------------------------------------------------------------------------------|--------------|
| <i>charge density, <math>\rho</math></i>         |                                    |                                                                                                      |              |
| SI unit                                          | $\text{C m}^{-3}$                  | $= 1 \text{ A s m}^{-3}$                                                                             |              |
| esu, Gaussian                                    | $\text{Fr cm}^{-3}$                | $:= (10^7/\zeta) \text{ C m}^{-3} \approx$<br>$3.335\,640\,952 \times 10^{-4} \text{ C m}^{-3}$      | 10           |
| au                                               | $ea_0^{-3}$                        | $\approx 1.081\,202\,300(27) \times 10^{12} \text{ C m}^{-3}$                                        |              |
| <i>electric potential, <math>V, \phi</math></i>  |                                    |                                                                                                      |              |
| <i>electric tension, <math>U</math></i>          |                                    |                                                                                                      |              |
| volt (SI unit)                                   | V                                  | $= 1 \text{ J C}^{-1} = 1 \text{ kg m}^2 \text{ s}^{-3} \text{ A}^{-1}$                              |              |
| esu, Gaussian                                    | $\text{Fr cm}^{-1}$                | $= \zeta \times 10^{-8} \text{ V} = 299.792\,458 \text{ V}$                                          | 10           |
| “ $\text{cm}^{-1}$ ”                             | $e \text{ cm}^{-1}/4\pi\epsilon_0$ | $\approx 1.439\,964\,41(36) \times 10^{-7} \text{ V}$                                                | 11           |
| au                                               | $e/4\pi\epsilon_0 a_0$             | $= E_h/e \approx 27.211\,383\,86(68) \text{ V}$                                                      |              |
| mean international volt                          |                                    | $= 1.000\,34 \text{ V}$                                                                              |              |
| US international volt                            |                                    | $= 1.000\,330 \text{ V}$                                                                             |              |
| <i>electric resistance, <math>R</math></i>       |                                    |                                                                                                      |              |
| ohm (SI unit)                                    | $\Omega$                           | $= 1 \text{ V A}^{-1} = 1 \text{ m}^2 \text{ kg s}^{-3} \text{ A}^{-2}$                              |              |
| mean international ohm                           |                                    | $= 1.000\,49 \Omega$                                                                                 |              |
| US international ohm                             |                                    | $= 1.000\,495 \Omega$                                                                                |              |
| Gaussian                                         | $\text{s cm}^{-1}$                 | $= \zeta^2 \times 10^{-9} \Omega \approx 8.987\,551\,787 \times 10^{11} \Omega$                      | 10           |
| <i>conductivity, <math>\kappa, \sigma</math></i> |                                    |                                                                                                      |              |
| SI                                               | $\text{S m}^{-1}$                  | $= 1 \text{ kg}^{-1} \text{ m}^{-3} \text{ s}^3 \text{ A}^2$                                         |              |
| Gaussian                                         | $\text{s}^{-1}$                    | $= (10^{11}/\zeta^2) \text{ S m}^{-1} \approx$<br>$1.112\,650\,056 \times 10^{-10} \text{ S m}^{-1}$ | 10           |
| <i>capacitance, <math>C</math></i>               |                                    |                                                                                                      |              |
| farad (SI unit)                                  | F                                  | $= 1 \text{ kg}^{-1} \text{ m}^{-2} \text{ s}^4 \text{ A}^2$                                         |              |
| Gaussian                                         | cm                                 | $:= (10^9/\zeta^2) \text{ F} \approx 1.112\,650\,056 \times 10^{-12} \text{ F}$                      | 10           |
| <i>electric field strength, <math>E</math></i>   |                                    |                                                                                                      |              |
| SI unit                                          | $\text{V m}^{-1}$                  | $= 1 \text{ J C}^{-1} \text{ m}^{-1} = 1 \text{ kg m s}^{-3} \text{ A}^{-1}$                         |              |
| esu, Gaussian                                    | $\text{Fr}^{-1} \text{ cm}^{-2}$   | $:= \zeta \times 10^{-6} \text{ V m}^{-1} =$<br>$2.997\,924\,58 \times 10^4 \text{ V m}^{-1}$        | 10           |
| “ $\text{cm}^{-2}$ ”                             | $e \text{ cm}^{-2}/4\pi\epsilon_0$ | $\approx 1.439\,964\,41(36) \times 10^{-5} \text{ V m}^{-1}$                                         | 11           |
| au                                               | $e/4\pi\epsilon_0 a_0^2$           | $= E_h/ea_0 \approx$<br>$5.142\,206\,32(13) \times 10^{11} \text{ V m}^{-1}$                         |              |

(11) The units in quotation marks for electric potential through polarizability may be found in the literature, although they are strictly incorrect; the entry suggested in the column *Symbol* defines the unit in terms of physical quantities and other units, so that, for a conversion into the SI, the physical quantities only need to be replaced by their values in the SI and the units need to be interpreted as units in the SI.

| <i>Name</i>                                                                                                         | <i>Symbol</i>                                 | <i>Expressed in SI units</i>                                                                                                                                               | <i>Notes</i> |
|---------------------------------------------------------------------------------------------------------------------|-----------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------|
| <i>electric field gradient, <math>E'_{\alpha\beta}</math>, <math>q_{\alpha\beta}</math></i>                         |                                               |                                                                                                                                                                            |              |
| SI unit                                                                                                             | V m <sup>-2</sup>                             | = 1 J C <sup>-1</sup> m <sup>-2</sup> = 1 kg s <sup>-3</sup> A <sup>-1</sup>                                                                                               |              |
| esu, Gaussian                                                                                                       | Fr <sup>-1</sup> cm <sup>-3</sup>             | := $\zeta \times 10^{-4}$ V m <sup>-2</sup> =<br>2.997 924 58×10 <sup>6</sup> V m <sup>-2</sup>                                                                            | 10           |
| “cm <sup>-3</sup> ”                                                                                                 | $e$ cm <sup>-3</sup> /4 $\pi\epsilon_0$       | $\approx 1.439\,964\,41(36)\times 10^{-3}$ V m <sup>-2</sup>                                                                                                               | 11           |
| au                                                                                                                  | $e/4\pi\epsilon_0 a_0^3$                      | = $E_h/ea_0^2 \approx$<br>9.717 361 66(24)×10 <sup>21</sup> V m <sup>-2</sup>                                                                                              |              |
| <i>electric dipole moment, <math>p</math>, <math>\mu</math></i>                                                     |                                               |                                                                                                                                                                            |              |
| SI unit                                                                                                             | C m                                           | = 1 A s m                                                                                                                                                                  |              |
| esu, Gaussian                                                                                                       | Fr cm                                         | := (10 <sup>-1</sup> / $\zeta$ ) C m $\approx$<br>3.335 640 952×10 <sup>-12</sup> C m                                                                                      | 10           |
| debye                                                                                                               | D                                             | = 10 <sup>-18</sup> Fr cm $\approx$<br>3.335 640 952×10 <sup>-30</sup> C m                                                                                                 |              |
| “cm”, dipole length                                                                                                 | $e$ cm                                        | $\approx 1.602\,176\,487(40)\times 10^{-21}$ C m                                                                                                                           | 11           |
| au                                                                                                                  | $ea_0$                                        | $\approx 8.478\,352\,81(21)\times 10^{-30}$ C m                                                                                                                            |              |
| <i>electric quadrupole moment, <math>Q_{\alpha\beta}</math>, <math>\Theta_{\alpha\beta}</math>, <math>eQ</math></i> |                                               |                                                                                                                                                                            |              |
| SI unit                                                                                                             | C m <sup>2</sup>                              | = 1 A s m <sup>2</sup>                                                                                                                                                     |              |
| esu, Gaussian                                                                                                       | Fr cm <sup>2</sup>                            | := (10 <sup>-3</sup> / $\zeta$ ) C m <sup>2</sup> $\approx$<br>3.335 640 952×10 <sup>-14</sup> C m <sup>2</sup>                                                            | 10           |
| “cm <sup>2</sup> ”, quadrupole area                                                                                 | $e$ cm <sup>2</sup>                           | $\approx 1.602\,176\,487(40)\times 10^{-23}$ C m <sup>2</sup>                                                                                                              | 11           |
| au                                                                                                                  | $ea_0^2$                                      | $\approx 4.486\,551\,07(11)\times 10^{-40}$ C m <sup>2</sup>                                                                                                               |              |
| <i>polarizability, <math>\alpha</math></i>                                                                          |                                               |                                                                                                                                                                            |              |
| SI unit                                                                                                             | J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup> | = 1 F m <sup>2</sup> = 1 kg <sup>-1</sup> s <sup>4</sup> A <sup>2</sup>                                                                                                    |              |
| esu, Gaussian                                                                                                       | Fr <sup>2</sup> cm <sup>3</sup>               | := (10 <sup>5</sup> / $\zeta^2$ ) J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup> $\approx$<br>1.112 650 056×10 <sup>-16</sup> J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup> | 10           |
| “cm <sup>3</sup> ”, polarizability volume                                                                           | 4 $\pi\epsilon_0$ cm <sup>3</sup>             | $\approx 1.112\,650\,056\times 10^{-16}$ J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup>                                                                                     | 11           |
| “Å <sup>3</sup> ”                                                                                                   | 4 $\pi\epsilon_0$ Å <sup>3</sup>              | $\approx 1.112\,650\,056\times 10^{-40}$ J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup>                                                                                     | 11           |
| au                                                                                                                  | 4 $\pi\epsilon_0 a_0^3$                       | = $e^2 a_0^2/E_h \approx$<br>1.648 777 253 6(34)×10 <sup>-41</sup> J <sup>-1</sup> C <sup>2</sup> m <sup>2</sup>                                                           |              |
| <i>electric displacement, <math>D</math></i>                                                                        |                                               |                                                                                                                                                                            |              |
| <i>(volume) polarization, <math>P</math></i>                                                                        |                                               |                                                                                                                                                                            |              |
| SI unit                                                                                                             | C m <sup>-2</sup>                             | = 1 A s m <sup>-2</sup>                                                                                                                                                    |              |
| esu, Gaussian                                                                                                       | Fr cm <sup>-2</sup>                           | := (10 <sup>5</sup> / $\zeta$ ) C m <sup>-2</sup> $\approx$<br>3.335 640 952×10 <sup>-6</sup> C m <sup>-2</sup>                                                            | 10           |
| <i>magnetic flux density, <math>B</math></i>                                                                        |                                               |                                                                                                                                                                            |              |
| <i>(magnetic field)</i>                                                                                             |                                               |                                                                                                                                                                            |              |
| tesla (SI unit)                                                                                                     | T                                             | = 1 J A <sup>-1</sup> m <sup>-2</sup> = 1 V s m <sup>-2</sup> = 1 Wb m <sup>-2</sup>                                                                                       |              |
| gauss (emu, Gaussian)                                                                                               | G                                             | = 10 <sup>-4</sup> T                                                                                                                                                       |              |
| au                                                                                                                  | $\hbar/ea_0^2$                                | $\approx 2.350\,517\,382(59)\times 10^5$ T                                                                                                                                 |              |

(12) The use of the esu or Gaussian unit for electric displacement usually implies that the non-rationalized displacement is being quoted,  $D^{(\text{ir})} = 4\pi D$  (see Section 7.3, p. 143).

| <i>Name</i>                                               | <i>Symbol</i>                  | <i>Expressed in SI units</i>                                                            | <i>Notes</i> |
|-----------------------------------------------------------|--------------------------------|-----------------------------------------------------------------------------------------|--------------|
| <i>magnetic flux, <math>\Phi</math></i>                   |                                |                                                                                         |              |
| weber (SI unit)                                           | Wb                             | $= 1 \text{ J A}^{-1} = 1 \text{ V s} = 1 \text{ kg m}^2 \text{ s}^{-2} \text{ A}^{-1}$ |              |
| maxwell (emu, Gaussian)                                   | Mx                             | $= 10^{-8} \text{ Wb} [= 1 \text{ G cm}^2]$                                             |              |
| <i>magnetic field strength, <math>H</math></i>            |                                |                                                                                         |              |
| SI unit                                                   | $\text{A m}^{-1}$              |                                                                                         |              |
| oersted (emu, Gaussian)                                   | Oe                             | $= (10^3/4\pi) \text{ A m}^{-1}$                                                        | 13           |
| <i>(volume) magnetization, <math>M</math></i>             |                                |                                                                                         |              |
| SI unit                                                   | $\text{A m}^{-1}$              |                                                                                         |              |
| gauss (emu, Gaussian)                                     | G                              | $= 10^3 \text{ A m}^{-1}$                                                               | 13           |
| <i>magnetic dipole moment, <math>m, \mu</math></i>        |                                |                                                                                         |              |
| SI unit                                                   | $\text{J T}^{-1}$              | $= 1 \text{ A m}^2$                                                                     |              |
| emu, Gaussian                                             | $\text{erg G}^{-1}$            | $= 10 \text{ A cm}^2 = 10^{-3} \text{ J T}^{-1}$                                        |              |
| Bohr magneton                                             | $\mu_B$                        | $:= e\hbar/2m_e \approx$<br>$9.274\,009\,15(23) \times 10^{-24} \text{ J T}^{-1}$       | 14           |
| au                                                        | $e\hbar/m_e$                   | $:= 2\mu_B \approx 1.854\,801\,830(46) \times 10^{-23} \text{ J T}^{-1}$                |              |
| nuclear magneton                                          | $\mu_N$                        | $:= (m_e/m_p)\mu_B \approx$<br>$5.050\,783\,24(13) \times 10^{-27} \text{ J T}^{-1}$    |              |
| <i>magnetizability, <math>\xi</math></i>                  |                                |                                                                                         |              |
| SI unit                                                   | $\text{J T}^{-2}$              | $= 1 \text{ A}^2 \text{ s}^2 \text{ m}^2 \text{ kg}^{-1}$                               |              |
| Gaussian                                                  | $\text{erg G}^{-2}$            | $= 10 \text{ J T}^{-2}$                                                                 |              |
| au                                                        | $e^2 a_0^2 / m_e$              | $\approx 7.891\,036\,433(27) \times 10^{-29} \text{ J T}^{-2}$                          |              |
| <i>magnetic susceptibility, <math>\chi, \kappa</math></i> |                                |                                                                                         |              |
| SI unit                                                   | 1                              |                                                                                         |              |
| emu, Gaussian                                             | 1                              |                                                                                         | 15           |
| <i>molar magnetic susceptibility, <math>\chi_m</math></i> |                                |                                                                                         |              |
| SI unit                                                   | $\text{m}^3 \text{ mol}^{-1}$  |                                                                                         |              |
| emu, Gaussian                                             | $\text{cm}^3 \text{ mol}^{-1}$ | $= 10^{-6} \text{ m}^3 \text{ mol}^{-1}$                                                | 16           |
| <i>inductance, self-inductance, <math>L</math></i>        |                                |                                                                                         |              |
| henry (SI unit)                                           | H                              | $= 1 \text{ V s A}^{-1} = 1 \text{ kg m}^2 \text{ s}^{-2} \text{ A}^{-2}$               |              |
| Gaussian                                                  | $\text{s}^2 \text{ cm}^{-1}$   | $= \zeta^2 \times 10^{-9} \text{ H} \approx 8.987\,551\,787 \times 10^{11} \text{ H}$   | 10           |
| emu                                                       | cm                             | $= 10^{-9} \text{ H}$                                                                   |              |

(13) In practice the oersted, Oe, is only used as a unit for  $H^{(\text{ir})} = 4\pi H$ , thus when  $H^{(\text{ir})} = 1 \text{ Oe}$ ,  $H = (10^3/4\pi) \text{ A m}^{-1}$  (see Section 7.3, p. 143). In the Gaussian or emu system, gauss and oersted are equivalent units.

(14) The Bohr magneton  $\mu_B$  is sometimes denoted BM (or B.M.), but this is not recommended.

(15) In practice susceptibilities quoted in the context of emu or Gaussian units are always values for  $\chi^{(\text{ir})} = \chi/4\pi$ ; thus when  $\chi^{(\text{ir})} = 10^{-6}$ ,  $\chi = 4\pi \times 10^{-6}$  (see Section 7.3, p. 143).

(16) In practice the units  $\text{cm}^3 \text{ mol}^{-1}$  usually imply that the non-rationalized molar susceptibility is being quoted  $\chi_m^{(\text{ir})} = \chi_m/4\pi$ ; thus, for example if  $\chi_m^{(\text{ir})} = -15 \times 10^{-6} \text{ cm}^3 \text{ mol}^{-1}$ , which is often written as “ $-15 \text{ cgs ppm}$ ”, then  $\chi_m = -1.88 \times 10^{-10} \text{ m}^3 \text{ mol}^{-1}$  (see Section 7.3, p. 143).

### 7.3 THE ESU, EMU, GAUSSIAN, AND ATOMIC UNIT SYSTEMS IN RELATION TO THE SI

The ISQ (see Section 1.2, p. 4) equations of electromagnetic theory are usually used with physical quantities in SI units, in particular the four base units metre (m), kilogram (kg), second (s), and ampere (A) for length, mass, time, and electric current, respectively. The basic equations for the electrostatic force  $\mathbf{F}_{\text{es}}$  between particles with charges  $Q_1$  and  $Q_2$  in vacuum, and for the infinitesimal electromagnetic force  $d^2\mathbf{F}_{\text{em}}$  between conductor elements of length  $d\mathbf{l}_1$  and  $d\mathbf{l}_2$  and corresponding currents  $I_1$  and  $I_2$  in vacuum, are

$$\begin{aligned}\mathbf{F}_{\text{es}} &= Q_1 Q_2 \mathbf{r} / 4\pi\epsilon_0 r^3 \\ d^2\mathbf{F}_{\text{em}} &= (\mu_0/4\pi) I_1 d\mathbf{l}_1 \times (I_2 d\mathbf{l}_2 \times \mathbf{r}) / r^3\end{aligned}$$

where particles and conductor elements are separated by the vector  $\mathbf{r}$  (with  $|\mathbf{r}| = r$ ). The physical quantity  $\epsilon_0$ , the electric constant (formerly called permittivity of vacuum), is defined in the SI system to have the value

$$\epsilon_0 = (10^7/4\pi c_0^2) \text{ kg}^{-1} \text{ m}^{-1} \text{ C}^2 \approx 8.854\,187\,817 \times 10^{-12} \text{ C}^2 \text{ m}^{-1} \text{ J}^{-1}$$

Similarly, the physical quantity  $\mu_0$ , the magnetic constant (formerly called the permeability of vacuum), has the value

$$\mu_0 := 4\pi \times 10^{-7} \text{ N A}^{-2} \approx 1.256\,637\,061\,4 \times 10^{-6} \text{ N A}^{-2}$$

in the SI system. In different unit systems  $\epsilon_0$  and  $\mu_0$  may have different numerical values and units. In this book we use  $\epsilon_0$  and  $\mu_0$  as shorthands for the SI values as given above and prefer to use more general symbols  $k_{\text{es}}$  and  $k_{\text{em}}$  to describe these quantities in other systems of units and equations as discussed below. The SI value of  $\mu_0$  results from the definition of the ampere (Section 3.3, p. 87). The value of  $\epsilon_0$  then results from the Maxwell relation

$$\epsilon_0 \mu_0 = 1/c_0^2$$

where  $c_0$  is the speed of light in vacuum (see Chapter 5, p. 111).

More generally, following ideas outlined in [153] and [154], the basic equations for the electrostatic and electromagnetic forces may be written as

$$\begin{aligned}\mathbf{F}_{\text{es}} &= k_{\text{es}} \frac{Q_1 Q_2 \mathbf{r}}{r^3} \\ d^2\mathbf{F}_{\text{em}} &= \frac{k_{\text{em}}}{k^2} \frac{I_1 d\mathbf{l}_1 \times (I_2 d\mathbf{l}_2 \times \mathbf{r})}{r^3}\end{aligned}$$

where the three constants  $k_{\text{es}}$ ,  $k_{\text{em}}$  and  $k$  satisfy a more general Maxwell relation

$$k^2 \cdot k_{\text{es}}/k_{\text{em}} = c_0^2$$

The introduction of three constants is necessary in order to describe coherently the system of units and equations that are in common use in electromagnetic theory. These are, in addition to the SI, the esu (electrostatic unit) system, the emu (electromagnetic unit) system, the Gaussian system, and the system of atomic units. The constants  $k$ ,  $k_{\text{es}}$ , and  $k_{\text{em}}$  have conventionally the values given in the following table.

|                 | <i>SI</i>          | <i>esu</i>                                | <i>emu</i>                            | <i>Gaussian</i>           | <i>atomic units</i> | <i>Notes</i> |
|-----------------|--------------------|-------------------------------------------|---------------------------------------|---------------------------|---------------------|--------------|
| $k$             | 1                  | 1                                         | 1                                     | $\zeta \text{ cm s}^{-1}$ | 1                   | 1            |
| $k_{\text{es}}$ | $1/4\pi\epsilon_0$ | 1                                         | $\zeta^2 \text{ cm}^2 \text{ s}^{-2}$ | 1                         | 1                   | 2            |
| $k_{\text{em}}$ | $\mu_0/4\pi$       | $(1/\zeta^2) \text{ s}^2 \text{ cm}^{-2}$ | 1                                     | 1                         | $\alpha^2$          | 3, 4         |

(1)  $\zeta$  is the exact number  $\zeta = c_0/(\text{cm s}^{-1}) = 29\,979\,245\,800$ , see p. 135.

(2)  $1/4\pi\epsilon_0 = \zeta^2 \times 10^{-11} \text{ N A}^{-2} \text{ m}^2 \text{ s}^{-2}$

(3)  $\mu_0/4\pi = 10^{-7} \text{ N A}^{-2}$

(4)  $\alpha$  is the fine-structure constant with  $\alpha^{-1} = 137.035\,999\,679(94)$  (see Chapter 5, p. 111).

This table can be used together with Section 7.4, p. 146 to transform expressions involving electromagnetic quantities between several systems. One sees that, particularly in the Gaussian system,  $k = c_0$ . This guarantees, together with the general relation of Maxwell equations given in Section 7.4, that the speed of light in vacuum comes out to be  $c_0$  in the Gaussian system. Examples of transformations between the SI and the Gaussian system are also given in Section 7.4.

## Additional remarks

### (i) The esu system

In the esu system, the base units are the centimetre (cm), gram (g), and second (s) for length, mass and time, respectively. The franklin, symbol Fr (see footnote <sup>1</sup>, below) for the electrostatic unit of charge may be introduced alternatively as a fourth base unit. Two particles with electric charges of one franklin, one centimetre apart in a vacuum, repel each other with a force of  $1 \text{ dyn} = 1 \text{ cm g s}^{-2}$ . From this definition, one obtains the relation  $1 \text{ Fr} = (10/\zeta) \text{ C}$  (see Example 7, Section 7.1, p. 134). Since  $k_{\text{es}} = 1$ , from the general definition of the electrostatic force given above, the equation  $\text{Fr} = \text{erg}^{1/2} \text{ cm}^{1/2}$  is true in the esu system, where  $\text{erg}^{1/2} \text{ cm}^{1/2}$  is the esu of charge. In this book the franklin is thus used for convenience as a name for the esu of charge.

### (ii) The emu system

In the emu system, the base units are the centimetre (cm), gram (g), and second (s) for length, mass and time. The biot (symbol Bi) for the electromagnetic unit of electric current may be introduced alternatively as a fourth base unit. Two long wires separated by one centimetre with electric currents of one biot that flow in the same direction, repel each other in a vacuum with a lineic force (force per length) of  $1 \text{ dyn/cm}$ . From this definition one obtains the relation  $1 \text{ Bi} = 10 \text{ A}$ . Since  $k_{\text{em}} = 1$ , from the general definition of the electromagnetic force given above, the equation  $\text{Bi} = \text{dyn}^{1/2}$  is true in the emu system, where  $\text{dyn}^{1/2}$  is the emu of electric current. The biot has generally been used as a compact expression for the emu of electric current.

### (iii) The Gaussian system

In the Gaussian system, the esu and emu systems are mixed. From the relation of the franklin and the biot to the SI units one readily obtains

$$1 \text{ Bi} = \zeta \text{ Fr s}^{-1}$$

Since  $k_{\text{es}} = 1$  and  $k_{\text{em}} = 1$ , the value of the constant  $k$  is determined to be  $c_0$  by the more general Maxwell relation given above. In treatises about relativity theory  $k$  is sometimes set to 1, which corresponds to a transformation of the time axis from a coordinate  $t$  to a coordinate  $x = c_0 t$ .

<sup>1</sup> The name “franklin”, symbol Fr, for the esu of charge was suggested by Guggenheim [155], although it has not been widely adopted. The name “statcoulomb” has also been used for the esu of charge.

#### (iv) The atomic units system

In the system of atomic units [22] (see also Section 3.9.1, p. 94), charges are given in units of the elementary charge  $e$ , currents in units of  $eE_h/\hbar$ , where  $E_h$  is the hartree and  $\hbar$  is the Planck constant divided by  $2\pi$ , lengths are given in units of the bohr,  $a_0$ , the electric field strength is given in units of  $E_h/ea_0$ , the magnetic flux density in units of  $\hbar/ea_0^2$ . Conversion factors from these and other atomic units to the SI are included in Sections 3.9.1 and 7.2, p. 94 and 135. Thus, since conventionally  $k = 1$ , it follows that  $k_{\text{es}} = 1$  in this system (see footnote <sup>2</sup>) and  $k_{\text{em}} = \alpha^2$  (see footnote <sup>3</sup>).

#### (v) Non-rationalized quantities

The numerical constant  $4\pi$  is introduced into the definitions of  $\varepsilon_0$  and  $\mu_0$  because of the spherical symmetry involved in the equations defining  $\mathbf{F}_{\text{es}}$  and  $\text{d}^2\mathbf{F}_{\text{em}}$  above; in this way its appearance is avoided in later equations, i.e., in the Maxwell equations. When factors of  $4\pi$  are introduced in this way, as in the SI, the equations are also called “rationalized”.

Furthermore, it is usual to include the factor  $4\pi$  in the following quantities, when converting the electromagnetic equation from the SI system to the esu, emu, Gaussian and atomic units system:

$$\begin{aligned} D^{(\text{ir})} &= 4\pi D \\ H^{(\text{ir})} &= 4\pi H \\ \chi_e^{(\text{ir})} &= \chi_e/4\pi \\ \chi^{(\text{ir})} &= \chi/4\pi \end{aligned}$$

where the superscript (ir), for irrational, meaning non-rationalized, denotes the value of the corresponding quantity in a “non-rationalized” unit system as opposed to a system like the SI, which is “rationalized” in the sense described above.

The magnetic permeability  $\mu$  is given as  $\mu = \mu_r k_{\text{em}} 4\pi = \mu_r \mu_0$  in the SI, and as  $\mu = \mu_r k_{\text{em}}$  in the non-rationalized unit system.  $\mu_r$  is the dimensionless relative permeability and is defined in terms of the magnetic susceptibility in Section 7.4, p. 147. The electric permittivity  $\varepsilon$  is given as  $\varepsilon = \varepsilon_r 4\pi/k_{\text{es}} = \varepsilon_r \cdot \varepsilon_0$  in the SI, and as  $\varepsilon = \varepsilon_r/k_{\text{es}}$  in non-rationalized unit systems.  $\varepsilon_r$  is the dimensionless relative permittivity and is defined in terms of the electric susceptibility in Section 7.4, p. 146.

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<sup>2</sup> Since  $E_h = \frac{e^2}{4\pi\varepsilon_0 a_0}$ ,  $E = \frac{1}{4\pi\varepsilon_0} \frac{Q}{r^2} = \frac{E_h a_0}{e^2} \frac{Q}{r^2} = \frac{E_h}{ea_0} \frac{(Q/e)}{(r/a_0)^2}$ .

<sup>3</sup> Since the value of the speed of light in atomic units is the reciprocal of the fine-structure constant,  $\alpha^{-1}$ , the condition  $k^2 k_{\text{es}}/k_{\text{em}} = c_0^2$  yields  $k_{\text{em}} = \alpha^2$  in atomic units for  $k = 1$ .

## 7.4 TRANSFORMATION OF EQUATIONS OF ELECTROMAGNETIC THEORY BETWEEN THE ISQ(SI) AND GAUSSIAN FORMS

| <i>General relation</i>                                                                                                 | <i>ISQ(SI)</i>                                                                                                   | <i>Gaussian</i>                                                                                     |
|-------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------|
| Force between two localized charged particles in vacuum (Coulomb law):                                                  |                                                                                                                  |                                                                                                     |
| $\mathbf{F} = k_{\text{es}} Q_1 Q_2 \mathbf{r} / r^3$                                                                   | $\mathbf{F} = Q_1 Q_2 \mathbf{r} / 4\pi\epsilon_0 r^3$                                                           | $\mathbf{F} = Q_1 Q_2 \mathbf{r} / r^3$                                                             |
| Electrostatic potential around a localized charged particle in vacuum:                                                  |                                                                                                                  |                                                                                                     |
| $\phi = k_{\text{es}} Q / r$                                                                                            | $\phi = Q / 4\pi\epsilon_0 r$                                                                                    | $\phi = Q / r$                                                                                      |
| Relation between electric field strength and electrostatic potential:                                                   |                                                                                                                  |                                                                                                     |
| $\mathbf{E} = -\nabla\phi$                                                                                              | $\mathbf{E} = -\nabla\phi$                                                                                       | $\mathbf{E} = -\nabla\phi$                                                                          |
| Field due to a charge distribution $\rho$ in vacuum (Gauss law):                                                        |                                                                                                                  |                                                                                                     |
| $\nabla \cdot \mathbf{E} = 4\pi k_{\text{es}} \rho$                                                                     | $\nabla \cdot \mathbf{E} = \rho / \epsilon_0$                                                                    | $\nabla \cdot \mathbf{E} = 4\pi \rho$                                                               |
| Electric dipole moment of a charge distribution:                                                                        |                                                                                                                  |                                                                                                     |
| $\mathbf{p} = \int \rho \mathbf{r} dV$                                                                                  | $\mathbf{p} = \int \rho \mathbf{r} dV$                                                                           | $\mathbf{p} = \int \rho \mathbf{r} dV$                                                              |
| Potential around a dipole in vacuum:                                                                                    |                                                                                                                  |                                                                                                     |
| $\phi = k_{\text{es}} \mathbf{p} \cdot \mathbf{r} / r^3$                                                                | $\phi = \mathbf{p} \cdot \mathbf{r} / 4\pi\epsilon_0 r^3$                                                        | $\phi = \mathbf{p} \cdot \mathbf{r} / r^3$                                                          |
| Energy of a charge distribution in an electric field:                                                                   |                                                                                                                  |                                                                                                     |
| $E = Q\phi - \mathbf{p} \cdot \mathbf{E} + \dots$                                                                       | $E = Q\phi - \mathbf{p} \cdot \mathbf{E} + \dots$                                                                | $E = Q\phi - \mathbf{p} \cdot \mathbf{E} + \dots$                                                   |
| Electric dipole moment induced in an electric field:                                                                    |                                                                                                                  |                                                                                                     |
| $\mathbf{p} = \alpha \mathbf{E} + \dots$                                                                                | $\mathbf{p} = \alpha \mathbf{E} + \dots$                                                                         | $\mathbf{p} = \alpha \mathbf{E} + \dots$                                                            |
| Dielectric polarization:                                                                                                |                                                                                                                  |                                                                                                     |
| $\mathbf{P} = \chi_e \mathbf{E} / 4\pi k_{\text{es}}$                                                                   | $\mathbf{P} = \chi_e \epsilon_0 \mathbf{E}$                                                                      | $\mathbf{P} = \chi_e^{(\text{ir})} \mathbf{E}$                                                      |
| Electric susceptibility and relative permittivity:                                                                      |                                                                                                                  |                                                                                                     |
| $\epsilon_r = 1 + \chi_e$                                                                                               | $\epsilon_r = 1 + \chi_e$                                                                                        | $\epsilon_r = 1 + 4\pi \chi_e^{(\text{ir})}$                                                        |
| Electric displacement <sup>1</sup> :                                                                                    |                                                                                                                  |                                                                                                     |
| $\mathbf{D} = \mathbf{E} / 4\pi k_{\text{es}} + \mathbf{P}$                                                             | $\mathbf{D} = \epsilon_0 \mathbf{E} + \mathbf{P}$                                                                | $\mathbf{D}^{(\text{ir})} = \mathbf{E} + 4\pi \mathbf{P}$                                           |
| $\mathbf{D} = \epsilon_r \mathbf{E} / 4\pi k_{\text{es}}$                                                               | $\mathbf{D} = \epsilon_0 \epsilon_r \mathbf{E}$                                                                  | $\mathbf{D}^{(\text{ir})} = \epsilon_r \mathbf{E}$                                                  |
| Capacitance of a parallel plate condenser, area $A$ , separation $d$ :                                                  |                                                                                                                  |                                                                                                     |
| $C = \epsilon_r A / 4\pi k_{\text{es}} d$                                                                               | $C = \epsilon_0 \epsilon_r A / d$                                                                                | $C = \epsilon_r A / 4\pi d$                                                                         |
| Force between two current elements in vacuum:                                                                           |                                                                                                                  |                                                                                                     |
| $d^2 \mathbf{F} = \frac{k_{\text{em}}}{k^2} \frac{I_1 d\mathbf{l}_1 \times (I_2 d\mathbf{l}_2 \times \mathbf{r})}{r^3}$ | $d^2 \mathbf{F} = \frac{\mu_0}{4\pi} \frac{I_1 d\mathbf{l}_1 \times (I_2 d\mathbf{l}_2 \times \mathbf{r})}{r^3}$ | $d^2 \mathbf{F} = \frac{I_1 d\mathbf{l}_1 \times (I_2 d\mathbf{l}_2 \times \mathbf{r})}{c_0^2 r^3}$ |
| Magnetic vector potential due to a current element in vacuum:                                                           |                                                                                                                  |                                                                                                     |
| $d\mathbf{A} = \frac{k_{\text{em}}}{k} I d\mathbf{l} / r$                                                               | $d\mathbf{A} = \frac{\mu_0}{4\pi} I d\mathbf{l} / r$                                                             | $d\mathbf{A} = I d\mathbf{l} / c_0 r$                                                               |
| Relation between magnetic flux density and magnetic vector potential:                                                   |                                                                                                                  |                                                                                                     |
| $\mathbf{B} = \nabla \times \mathbf{A}$                                                                                 | $\mathbf{B} = \nabla \times \mathbf{A}$                                                                          | $\mathbf{B} = \nabla \times \mathbf{A}$                                                             |
| Magnetic flux density due to a current element in vacuum (Biot-Savart law):                                             |                                                                                                                  |                                                                                                     |
| $d\mathbf{B} = \frac{k_{\text{em}}}{k} I d\mathbf{l} \times \mathbf{r} / r^3$                                           | $d\mathbf{B} = \frac{\mu_0}{4\pi} I d\mathbf{l} \times \mathbf{r} / r^3$                                         | $d\mathbf{B} = I d\mathbf{l} \times \mathbf{r} / c_0 r^3$                                           |

(1) The second equation holds in isotropic media.

Magnetic flux density due to a current density  $\mathbf{j}$  in vacuum (Ampère law):

$$\nabla \times \mathbf{B} = 4\pi \frac{k_{\text{em}}}{k} \mathbf{j} \qquad \nabla \times \mathbf{B} = \mu_0 \mathbf{j} \qquad \nabla \times \mathbf{B} = 4\pi \mathbf{j}/c_0$$

Force on a current element in a magnetic flux density:

$$d\mathbf{F} = I d\mathbf{l} \times \mathbf{B}/k \qquad d\mathbf{F} = I d\mathbf{l} \times \mathbf{B} \qquad d\mathbf{F} = I d\mathbf{l} \times \mathbf{B}/c_0$$

Magnetic dipole of a current loop of area  $\mathbf{A}$ :

$$\mathbf{m} = I \mathbf{A}/k \qquad \mathbf{m} = I \mathbf{A} \qquad \mathbf{m} = I \mathbf{A}/c_0$$

Magnetic vector potential around a magnetic dipole in vacuum:

$$\mathbf{A} = k_{\text{em}} \mathbf{m} \times \mathbf{r}/r^3 \qquad \mathbf{A} = \frac{\mu_0}{4\pi} \mathbf{m} \times \mathbf{r}/r^3 \qquad \mathbf{A} = \mathbf{m} \times \mathbf{r}/r^3$$

Energy of a magnetic dipole in a magnetic flux density:

$$E = -\mathbf{m} \cdot \mathbf{B} \qquad E = -\mathbf{m} \cdot \mathbf{B} \qquad E = -\mathbf{m} \cdot \mathbf{B}$$

Magnetic dipole induced by a magnetic flux density:

$$\mathbf{m} = \xi \mathbf{B} + \dots \qquad \mathbf{m} = \xi \mathbf{B} + \dots \qquad \mathbf{m} = \xi \mathbf{B} + \dots$$

Magnetization:

$$\mathbf{M} = \chi \mathbf{H} \qquad \mathbf{M} = \chi \mathbf{H} \qquad \mathbf{M} = \chi^{(\text{ir})} \mathbf{H}^{(\text{ir})}$$

Magnetic susceptibility and relative permeability:

$$\mu_r = 1 + \chi \qquad \mu_r = 1 + \chi \qquad \mu_r = 1 + 4\pi \chi^{(\text{ir})}$$

Magnetic field strength<sup>1</sup>:

$$\begin{aligned} \mathbf{H} &= \mathbf{B}/4\pi k_{\text{em}} - \mathbf{M} & \mathbf{H} &= \mathbf{B}/\mu_0 - \mathbf{M} & \mathbf{H}^{(\text{ir})} &= \mathbf{B} - 4\pi \mathbf{M} \\ \mathbf{H} &= \mathbf{B}/4\pi \mu_r k_{\text{em}} & \mathbf{H} &= \mathbf{B}/\mu_0 \mu_r & \mathbf{H}^{(\text{ir})} &= \mathbf{B}/\mu_r \end{aligned}$$

Conductivity:

$$\mathbf{j} = \kappa \mathbf{E} \qquad \mathbf{j} = \kappa \mathbf{E} \qquad \mathbf{j} = \kappa \mathbf{E}$$

Self-inductance of a solenoid of volume  $V$  with  $n$  windings per length:

$$L = 4\pi \frac{k_{\text{em}}}{k^2} \mu_r n^2 V \qquad L = \mu_0 \mu_r n^2 V \qquad L = 4\pi \mu_r n^2 V / c_0^2$$

Faraday induction law:

$$\nabla \times \mathbf{E} + \frac{1}{k} \frac{\partial \mathbf{B}}{\partial t} = \mathbf{0} \qquad \nabla \times \mathbf{E} + \partial \mathbf{B} / \partial t = \mathbf{0} \qquad \nabla \times \mathbf{E} + \frac{1}{c_0} \frac{\partial \mathbf{B}}{\partial t} = \mathbf{0}$$

Relation between the electric field strength and electromagnetic potentials:

$$\mathbf{E} = -\nabla \phi - \frac{1}{k} \frac{\partial \mathbf{A}}{\partial t} \qquad \mathbf{E} = -\nabla \phi - \partial \mathbf{A} / \partial t \qquad \mathbf{E} = -\nabla \phi - \frac{1}{c_0} \frac{\partial \mathbf{A}}{\partial t}$$

Maxwell equations:

$$\begin{aligned} \nabla \cdot \mathbf{D} &= \rho & \nabla \cdot \mathbf{D} &= \rho & \nabla \cdot \mathbf{D}^{(\text{ir})} &= 4\pi \rho \\ k \nabla \times \mathbf{H} - \partial \mathbf{D} / \partial t &= \mathbf{j} & \nabla \times \mathbf{H} - \partial \mathbf{D} / \partial t &= \mathbf{j} & \nabla \times \mathbf{H}^{(\text{ir})} - \frac{1}{c_0} \frac{\partial \mathbf{D}^{(\text{ir})}}{\partial t} &= \frac{4\pi}{c_0} \mathbf{j} \\ k \nabla \times \mathbf{E} + \partial \mathbf{B} / \partial t &= \mathbf{0} & \nabla \times \mathbf{E} + \partial \mathbf{B} / \partial t &= \mathbf{0} & \nabla \times \mathbf{E} + \frac{1}{c_0} \frac{\partial \mathbf{B}}{\partial t} &= \mathbf{0} \\ \nabla \cdot \mathbf{B} &= 0 & \nabla \cdot \mathbf{B} &= 0 & \nabla \cdot \mathbf{B} &= 0 \end{aligned}$$

Wave equations for the electromagnetic potentials  $\phi$  and  $\mathbf{A}$  (in Lorentz gauge):

$$\Delta\phi - \frac{k_{\text{em}}}{k^2 k_{\text{es}}} \frac{\partial^2 \phi}{\partial t^2} = -4\pi k_{\text{es}} \rho$$

$$\Delta\phi - \varepsilon_0 \mu_0 \frac{\partial^2 \phi}{\partial t^2} = -\rho / \varepsilon_0$$

$$\Delta\phi - \frac{1}{c_0^2} \frac{\partial^2 \phi}{\partial t^2} = -4\pi \rho$$

$$\Delta\mathbf{A} - \frac{k_{\text{em}}}{k^2 k_{\text{es}}} \frac{\partial^2 \mathbf{A}}{\partial t^2} = -4\pi k_{\text{em}} \mathbf{j}$$

$$\Delta\mathbf{A} - \varepsilon_0 \mu_0 \frac{\partial^2 \mathbf{A}}{\partial t^2} = -\mu_0 \mathbf{j}$$

$$\Delta\mathbf{A} - \frac{1}{c_0^2} \frac{\partial^2 \mathbf{A}}{\partial t^2} = -\frac{4\pi}{c_0} \mathbf{j}$$

$$\nabla\mathbf{A} + \frac{k_{\text{em}}}{k k_{\text{es}}} \frac{\partial \phi}{\partial t} = 0$$

$$\nabla\mathbf{A} + \varepsilon_0 \mu_0 \frac{\partial \phi}{\partial t} = 0$$

$$\nabla\mathbf{A} + \frac{1}{c_0} \frac{\partial \phi}{\partial t} = 0$$

Energy density of radiation:

$$U/V = (\mathbf{E} \cdot \mathbf{D} + \mathbf{B} \cdot \mathbf{H}) / 2$$

$$U/V = (\mathbf{E} \cdot \mathbf{D} + \mathbf{B} \cdot \mathbf{H}) / 2$$

$$U/V = (\mathbf{E} \cdot \mathbf{D}^{(\text{ir})} + \mathbf{B} \cdot \mathbf{H}^{(\text{ir})}) / 8\pi$$

Rate of radiation energy flow (Poynting vector):

$$\mathbf{S} = k \mathbf{E} \times \mathbf{H}$$

$$\mathbf{S} = \mathbf{E} \times \mathbf{H}$$

$$\mathbf{S} = \frac{c_0}{4\pi} \mathbf{E} \times \mathbf{H}^{(\text{ir})}$$

Force on a moving charge  $Q$  with velocity  $\mathbf{v}$  (Lorentz force):

$$\mathbf{F} = Q (\mathbf{E} + \mathbf{v} / k \times \mathbf{B})$$

$$\mathbf{F} = Q (\mathbf{E} + \mathbf{v} \times \mathbf{B})$$

$$\mathbf{F} = Q (\mathbf{E} + \mathbf{v} \times \mathbf{B} / c_0)$$

## 8 UNCERTAINTY

It is vital to report an uncertainty estimate along with a measurement. In brief, a report of a quantitative experimental (or theoretical) result should contain a statement about what the expected “best estimate” for the true result is, as well as a statement of the probable range of possible values specifying the uncertainty. The expected dispersion of the measured values arises from a number of sources. The contribution to the uncertainty from all these sources should be estimated as part of an uncertainty budget. The tables show some of the ways in which the contributions to the uncertainty budget can be estimated and combined and the examples below show how a full statement of the resulting combined uncertainty estimate can be presented. The present summary is based on [8].



## 8.1 REPORTING UNCERTAINTY FOR A SINGLE MEASURED QUANTITY

Uncertainties for a measured quantity  $X$  can be reported in several ways by giving the expected best estimate for the quantity together with a second quantity defining an uncertainty which corresponds to some probable range for the true quantity. We use the notation  $X$  for measured quantities,  $x$  for measurement results, and  $u_c$  for uncertainty.

*Example 1.*  $m_s = 100.021\,47$  g with a combined standard uncertainty (i.e., estimated standard uncertainty) of  $u_c = 0.35$  mg. Since it can be assumed that the possible estimated values of the standard are approximately normally distributed with approximate standard deviation  $u_c$ , the unknown value of the standard is believed to lie in the interval  $m_s \pm u_c$  with a level of confidence of approximately 68 %.

*Example 2.*  $m_s = (100.021\,47 \pm 0.000\,70)$  g, where the number following the symbol  $\pm$  is the numerical value of an expanded uncertainty  $U = k u_c$ , with  $U$  determined from a combined standard uncertainty (i.e., estimated standard deviation)  $u_c = 0.35$  mg and a coverage factor of  $k = 2$ . Since it can be assumed that the possible estimated values of the standard are approximately normally distributed with approximate standard deviation  $u_c$ , the unknown value of the standard is believed to lie within the interval defined by  $U$  with a level of confidence of 95 % (see Section 8.3, p. 154).

*Example 3.*  $m_s = 100.021\,47(35)$  g, where the number in parentheses denotes the combined standard uncertainty  $u_c = 0.35$  mg and is assumed to apply to the least significant digits.

| Name                             | Symbol       | Definition                                                                                                                     | Notes |
|----------------------------------|--------------|--------------------------------------------------------------------------------------------------------------------------------|-------|
| <b>Probability distributions</b> |              |                                                                                                                                |       |
| probability distribution of $x$  | $f(x)$       | The probability density of the quantity having the value $x$ ; normal (Gaussian), rectangular, triangular, Student- $t$ , etc. |       |
| expected value of $x$            | $E[x]$       | $E[x] = \int x f(x) dx$                                                                                                        |       |
| mean                             | $\mu$        | $E[x]$                                                                                                                         |       |
| variance                         | $\sigma^2$   | $E[(x - \mu)^2]$                                                                                                               |       |
| standard deviation               | $\sigma$     | $\sigma = +\sqrt{\sigma^2}$                                                                                                    |       |
| <b>Statistics</b>                |              |                                                                                                                                |       |
| number of measurements           | $N$          |                                                                                                                                |       |
| mean                             | $\bar{x}$    | $\bar{x} = \frac{1}{N} \sum_{i=1}^N x_i$                                                                                       |       |
| variance                         | $s^2(x)$     | $s^2(x) = \frac{1}{N-1} \sum_{i=1}^N (x_i - \bar{x})^2$                                                                        | 1     |
| standard deviation               | $s(x)$       | $s(x) = +\sqrt{s^2(x)}$                                                                                                        |       |
| standard deviation of the mean   | $s(\bar{x})$ | $s(\bar{x}) = s(x)/\sqrt{N}$                                                                                                   |       |

(1) This is an unbiased estimate and takes into account the removal of one degree of freedom because the spread is measured about the mean.

| <i>Name</i>                                                 | <i>Symbol</i> | <i>Definition</i>                                                                                                                                                                                                                                                                     | <i>Notes</i> |
|-------------------------------------------------------------|---------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------|
| <b>Uncertainties</b>                                        |               |                                                                                                                                                                                                                                                                                       |              |
| standard uncertainty of $x_i$                               | $u(x_i)$      | Estimated by Type A or B approaches                                                                                                                                                                                                                                                   | 2            |
| Type A evaluation                                           |               | Type A evaluation (of standard uncertainty) is a method of evaluation of a standard uncertainty by the statistical analysis of a series of observations                                                                                                                               | 3            |
| Type B evaluation                                           |               | Type B evaluation (of standard uncertainty) is a method of evaluation of a standard uncertainty by means other than the statistical analysis of a series of observations (for instance, it is usually based on scientific judgement using all relevant information that is available) | 4            |
| relative standard uncertainty of $x$                        | $u_r(x_i)$    | $u_r(x_i) = u(x_i)/ x_i  \quad (x_i \neq 0)$                                                                                                                                                                                                                                          |              |
| combined standard uncertainty of the measurement result $y$ | $u_c(y)$      | Estimated uncertainty in $y$ from all the component measurement results<br>$y = f(x_1, x_2, \dots, x_M)$                                                                                                                                                                              | 5            |
| relative combined standard uncertainty of $y$               | $u_{c,r}(y)$  | $u_{c,r}(y) = u_c(y)/ y  \quad (y \neq 0)$                                                                                                                                                                                                                                            |              |
| expanded uncertainty                                        | $U$           | $U = k u_c(y)$                                                                                                                                                                                                                                                                        | 6            |
| coverage factor                                             | $k$           |                                                                                                                                                                                                                                                                                       | 6            |

(Notes continued)

(2) The uncertainty in the result of a measurement generally consists of several components which may be grouped into two categories according to the way in which their numerical value is estimated:

Type A: Those which are evaluated by statistical methods

Type B: Those which are evaluated by other means

A detailed report on uncertainty should consist of a complete list of the components, specifying for each the method used to obtain its numerical value. This provides the “uncertainty budget”.

(3) Examples of type A evaluations are calculating the standard deviation of the mean of a series of independent observations; using the method of least squares to fit a function to data in order to estimate the parameters of the function and their standard deviations; and carrying out an analysis of variance (ANOVA) in order to identify and quantify random effects in certain kinds of measurements. These components of the uncertainty are characterized by the estimated variances  $s_i^2$  (or the estimated “standard deviations”  $s_i$ ) and the number of degrees of freedom  $\nu_i$ . Where appropriate, the covariances should be given.

(4) This knowledge may include previous measurement data, general knowledge of the behavior and properties of relevant materials and instruments, manufacturer’s specifications, data provided in calibration and other reports. These components of uncertainty should be characterized by quantities  $u_j^2$ , which may be considered approximations to the corresponding variances, the existence of which is assumed. The quantities  $u_j^2$  may be treated like variances and the quantities  $u_j$  like standard deviations.

## 8.2 PROPAGATION OF UNCERTAINTY FOR UNCORRELATED MEASUREMENTS

For a discussion of the treatment of correlated uncertainties, see [8].

| <i>Measurement equation</i>                  | <i>Reported measurement result</i>           | <i>Equation for the combined standard uncertainty</i>             | <i>Notes</i> |
|----------------------------------------------|----------------------------------------------|-------------------------------------------------------------------|--------------|
| $Y = \sum_{i=1}^N a_i X_i$                   | $y = \sum_{i=1}^N a_i x_i$                   | $u_c(y) = \left( \sum_{i=1}^N a_i^2 u^2(x_i) \right)^{1/2}$       | 1            |
| $Y = A X_1^{a_1} X_2^{a_2} \cdots X_N^{a_N}$ | $y = A x_1^{a_1} x_2^{a_2} \cdots x_N^{a_N}$ | $u_{c,r}(y) = \left( \sum_{i=1}^N a_i^2 u_r^2(x_i) \right)^{1/2}$ | 2            |

(1) This is sometimes known as addition in quadrature. The measurement equation is represented by a sum of quantities  $X_i$  multiplied by a constant  $a_i$ . The  $a_i$  are assumed to be known with certainty.

(2) The measurement equation is represented by a product of quantities  $X_i$  raised to powers  $a_1, a_2, \dots, a_N$  and multiplied by a constant  $A$ .  $A$  and the  $a_i$  are assumed to be known with certainty.

(Notes continued)

(4) (continued)

Examples of Type B evaluations:

- (a) If it is reasonable to assume that the quantity,  $X$ , can be modeled by a normal probability distribution then lower and upper limits  $a_-$  and  $a_+$  should be estimated such that the best estimated value of the input quantity is  $x = (a_- + a_+)/2$  (i.e., the centre of the limits) and there is one chance out of two (i.e., a 50 % probability) that the value of the quantity lies in the interval  $a_-$  to  $a_+$ , then  $u \approx 1.48(a_+ - a_-)/2$ .
- (b) If however the quantity,  $X$ , is better represented by a rectangular distribution then the lower and upper limits  $a_-$  and  $a_+$  of the input quantity should be estimated such that the probability that the value lies in the interval  $a_-$  and  $a_+$  is, for all practical purposes, 100 %. Provided that there is no contradictory information, treat the quantity as if it is equally probable for its value to lie anywhere within the interval  $a_-$  to  $a_+$ ; that is, model it by a uniform (i.e., rectangular) probability distribution. The best estimate of the value of the quantity is then  $x = (a_- + a_+)/2$  with the uncertainty  $u = (a_+ - a_-)/\sqrt{3}$ .
- (5) The quantity,  $Y$  being measured, called the measurand, is not measured directly, but is determined from  $M$  other quantities  $X_1, X_2, \dots, X_M$  through a function  $f$  (sometimes called the measurement equation),  $Y = f(X_1, X_2, \dots, X_M)$ . The quantities  $X_i$  include corrections (or correction factors), as well as quantities that take into account other sources of variability, such as different observers, instruments, samples, and laboratories etc. An estimate of the measured quantity  $Y$ , denoted  $y = f(x_1, x_2, \dots, x_M)$ , is obtained from the measurement equation using *input estimates*  $x_1, x_2, \dots, x_M$  for the values of the  $M$  *input quantities*. A similar situation exists if other quantities are to be derived from measurements. The propagation of uncertainty is illustrated in the table above.
- (6) Some commercial, industrial, and regulatory applications require a measure of uncertainty that defines an interval about a measurement result  $y$  within which the value of the measurand  $Y$  can be confidently asserted to lie. In these cases the *expanded uncertainty*  $U$  is used, and is obtained by multiplying the combined standard uncertainty,  $u_c(y)$ , by a *coverage factor*  $k$ . Thus  $U = k u_c(y)$  and it is confidently believed that  $Y$  is greater than or equal to  $y - U$ , and is less than or equal to  $y + U$ , which is commonly written as  $Y = y \pm U$ . The value of the coverage factor  $k$  is chosen on the basis of the desired level of confidence to be associated with the interval defined by  $U = k u_c$ .

### 8.3 REPORTING UNCERTAINTIES IN TERMS OF CONFIDENCE INTERVALS

One can generally report the uncertainty by defining an interval, within which the quantity is expected to fall with a certain probability. This interval can be specified as  $(y - U) \leq Y \leq (y + U)$ .

*Example*     $100.021\,40\text{ g} \leq m_s \leq 100.021\,54\text{ g}$  or  $m_s = 100.021\,47(7)\text{ g}$   
or  $m_s = (100.021\,47 \pm 0.000\,07)\text{ g}$

When specifying such an interval, one should provide the confidence as the estimated probability with which the true quantity is expected to be in the given range (for example 60 %, 90 % or 95 %). This probability can be estimated using a variety of probability distributions appropriate for the measurement under consideration (Gaussian, Student-*t*, etc. [8,9]).

---

(Notes continued)

(6) (continued) Typically,  $k$  is in the range of two to three. When the normal distribution applies and  $u_c$  is a reliable estimate of the standard deviation of  $y$ ,  $U = 2u_c$  (i.e.,  $k = 2$ ) defines an interval having a level of confidence of approximately 95 %, and  $U = 3u_c$  (i.e.,  $k = 3$ ) defines an interval having a level of confidence greater than 99 %.

## 9 ABBREVIATIONS AND ACRONYMS

Abbreviations and acronyms (words formed from the initial letters of groups of words that are frequently repeated) should be used sparingly. Unless they are well established (e.g. NMR, IR) they should always be defined once in any paper, and they should generally be avoided in titles and abstracts. Some acronyms have been accepted as common words, such as “laser” from the acronym LASER. Abbreviations used to denote physical quantities should if possible be replaced by the recommended symbol for the quantity (e.g.  $E_i$  rather than IP for ionization energy, see Section 2.5, p. 22;  $\rho$  rather than dens. for mass density, see Section 2.2, p. 14). For further recommendations concerning abbreviations see [156].

A list of frequently used abbreviations and acronyms is given here to help readers, but not necessarily to encourage their universal usage. In many cases an acronym can be found written in lower case letters and in capitals. In the list which follows only the most common usage is given. More extensive lists for different spectroscopic methods have been published by IUPAC [157,158] and by Wendisch [159]; an extensive list for acronyms used in theoretical chemistry has been published by IUPAC [21].



|          |                                                                       |
|----------|-----------------------------------------------------------------------|
| A/D      | analogue-to-digital                                                   |
| AA       | atomic absorption                                                     |
| AAS      | atomic absorption spectroscopy                                        |
| ac       | alternating current                                                   |
| ACM      | adiabatic channel model                                               |
| ACT      | activated complex theory                                              |
| AD       | atom diffraction                                                      |
| ADC      | analogue-to-digital converter                                         |
| AES      | Auger electron spectroscopy                                           |
| AFM      | atomic force microscopy                                               |
| AIUPS    | angle-integrated ultraviolet photoelectron spectroscopy               |
| AM       | amplitude modulated                                                   |
| amu      | atomic mass unit                                                      |
| ANOVA    | analysis of variance                                                  |
| AO       | atomic orbital                                                        |
| APS      | appearance potential spectroscopy                                     |
| ARAES    | angle-resolved Auger electron spectroscopy                            |
| ARPEFS   | angle-resolved photoelectron fine structure                           |
| AS       | Auger spectroscopy                                                    |
| ATR      | attenuated total (internal) reflection                                |
| AU       | astronomical unit                                                     |
| au       | atomic unit                                                           |
|          |                                                                       |
| bcc      | body-centred cubic                                                    |
| BET      | Brunauer-Emmett-Teller (adsorption isotherm)                          |
| BIPM     | Bureau International des Poids et Mesures                             |
| BIS      | bremsstrahlung isochromat spectroscopy                                |
| BM       | Bohr magneton (symbol: $\mu_B$ )                                      |
| bp       | boiling point                                                         |
| Btu      | British thermal unit                                                  |
|          |                                                                       |
| CARS     | coherent anti-Stokes Raman scattering                                 |
| CAS      | complete active space                                                 |
| CAS-SCF  | complete active space - self consistent field                         |
| CAT      | computer average of transients                                        |
| CAWIA    | Commission on Atomic Weights and Isotopic Abundances (now CIAAW)      |
| CCA      | coupled cluster approximation                                         |
| CCC      | critical coagulation concentration                                    |
| CCD      | coupled charge device                                                 |
| CCL      | colour centre laser                                                   |
| CCU      | Comité Consultatif d'Unités                                           |
| ccp      | cubic close packed                                                    |
| CD       | circular dichroism                                                    |
| CEELS    | characteristic electron energy-loss spectroscopy                      |
| CELS     | characteristic energy-loss spectroscopy                               |
| CEPA     | coupled electron pair approximation                                   |
| CGPM     | Conférence Générale des Poids et Mesures                              |
| cgs, CGS | centimetre-gram-second                                                |
| CI       | chemical ionization                                                   |
| CI       | configuration interaction                                             |
| CIAAW    | Commission on Isotopic Abundances and Atomic Weights (formerly CAWIA) |

|        |                                                             |
|--------|-------------------------------------------------------------|
| CIDEP  | chemically induced dynamic electron polarization            |
| CIDNP  | chemically induced dynamic nuclear polarization             |
| CIMS   | chemical ionization mass spectroscopy                       |
| CIPM   | Comité International des Poids et Mesures                   |
| CIVR   | collision induced vibrational relaxation                    |
| CIVR   | classical intramolecular vibrational redistribution         |
| CMA    | cylindrical mass analyzer                                   |
| CME    | chemically modified electrode                               |
| CNDO   | complete neglect of differential overlap                    |
| CODATA | Committee on Data for Science and Technology                |
| COMAS  | Concentration Modulation Absorption Spectroscopy            |
| CPD    | contact-potential difference                                |
| CRDS   | cavity ring-down spectroscopy                               |
| CSRS   | coherent Stokes-Raman scattering                            |
| CT     | charge transfer                                             |
| CVD    | chemical vapor deposition                                   |
| CW     | continuous wave                                             |
|        |                                                             |
| D/A    | digital-to-analogue                                         |
| DAC    | digital-to-analogue converter                               |
| D4WM   | degenerate 4-wave mixing                                    |
| DAPS   | disappearance potential spectroscopy                        |
| dc, DC | direct current                                              |
| DFG    | difference frequency generation                             |
| DFT    | density functional theory                                   |
| DLVO   | Derjaguin-Landau-Verwey-Overbeek                            |
| DME    | dropping mercury electrode                                  |
| DQMC   | diffusion quantum Monte Carlo                               |
| DRIFTS | diffuse reflectance infrared Fourier transform spectroscopy |
| DSC    | differential scanning calorimetry                           |
| DTA    | differential thermal analysis                               |
|        |                                                             |
| E1     | elimination unimolecular                                    |
| E2     | elimination bimolecular                                     |
| EAPFS  | extended appearance potential fine structure                |
| EC     | electron capture                                            |
| ECD    | electron capture detector                                   |
| ED     | electron diffraction                                        |
| EDA    | electron donor-acceptor [complex]                           |
| EDX    | energy-dispersive X-ray analysis                            |
| EELS   | electron energy-loss spectroscopy                           |
| EH     | electron holography                                         |
| EI     | electron impact ionization                                  |
| EIS    | electron impact spectroscopy                                |
| EIS    | electrochemical impedance spectroscopy                      |
| EL     | electroluminescence                                         |
| ELDOR  | electron-electron double resonance                          |
| ELEED  | elastic low energy electron diffraction                     |
| emf    | electromotive force                                         |
| emu    | electromagnetic unit                                        |

|         |                                                                        |
|---------|------------------------------------------------------------------------|
| ENDOR   | electron-nuclear double resonance                                      |
| EPR     | electron paramagnetic resonance                                        |
| ESCA    | electron spectroscopy for chemical applications (or analysis), see XPS |
| ESD     | electron stimulated desorption                                         |
| ESDIAD  | electron stimulated desorption ion angular distribution                |
| ESR     | electron spin resonance                                                |
| esu     | electrostatic unit                                                     |
| ETS     | electron transmission spectroscopy, electron tunneling spectroscopy    |
| eu      | entropy unit                                                           |
| EXAFS   | extended X-ray absorption fine structure                               |
| EXAPS   | electron excited X-ray appearance potential spectroscopy               |
| EXELFS  | extended electron energy-loss fine structure                           |
|         |                                                                        |
| FAB(MS) | fast atom bombardment (mass spectroscopy)                              |
| fcc     | face-centred cubic                                                     |
| FD      | field desorption                                                       |
| FEESP   | field-emitted electron spin-polarization [spectroscopy]                |
| FEM     | field emission [electron] microscopy                                   |
| FES     | field emission spectroscopy                                            |
| FFT     | fast Fourier transform                                                 |
| FI      | field ionization                                                       |
| FID     | flame ionization detector                                              |
| FID     | free induction decay                                                   |
| FIM     | field-ion microscopy                                                   |
| FIMS    | field-ion mass spectroscopy                                            |
| FIR     | far-infrared                                                           |
| FM      | frequency modulated                                                    |
| FPD     | flame photometric detector                                             |
| FSR     | free spectral range                                                    |
| FT      | Fourier transform                                                      |
| FTD     | flame thermionic detector                                              |
| FTIR    | Fourier transform infrared                                             |
| FWHM    | full width at half maximum                                             |
|         |                                                                        |
| GC      | gas chromatography                                                     |
| GIXS    | grazing-incidence X-ray scattering                                     |
| GLC     | gas-liquid chromatography                                              |
| GM      | Geiger-Müller                                                          |
| GTO     | Gaussian-type orbital                                                  |
| GVB     | generalized valence bond                                               |
|         |                                                                        |
| hcp     | hexagonal close-packed                                                 |
| HEED    | high-energy electron diffraction                                       |
| HEELS   | high-energy electron energy-loss spectroscopy                          |
| HEIS    | high-energy ion scattering                                             |
| HF      | Hartree-Fock                                                           |
| hfs     | hyperfine structure (hyperfine splitting)                              |
| HMDE    | hanging mercury drop electrode                                         |
| HMO     | Hückel molecular orbital                                               |
| HOMO    | highest occupied molecular orbital                                     |

|         |                                                                         |
|---------|-------------------------------------------------------------------------|
| HPLC    | high-performance liquid chromatography                                  |
| HREELS  | high-resolution electron energy-loss spectroscopy                       |
| HTS     | Hadamard transform spectroscopy                                         |
| HWP     | half-wave potential                                                     |
| I/O     | input-output                                                            |
| IBA     | ion beam analysis                                                       |
| IC      | integrated circuit                                                      |
| ICISS   | impact-collision ion scattering spectroscopy                            |
| ICR     | ion cyclotron resonance                                                 |
| id      | inner diameter                                                          |
| IEC     | International Electrotechnical Commission                               |
| IEP     | isoelectric point                                                       |
| IEPA    | independent electron pair approximation                                 |
| IETS    | inelastic electron tunneling spectroscopy                               |
| ILEED   | inelastic low-energy electron diffraction                               |
| INDO    | incomplete neglect of differential overlap                              |
| INDOR   | internuclear double resonance                                           |
| INS     | inelastic neutron scattering                                            |
| INS     | ion neutralization spectroscopy                                         |
| IP      | ionization potential (symbol: $E_i$ )                                   |
| IPES    | inverse photoelectron spectroscopy                                      |
| IPTS    | International Practical Temperature Scale                               |
| IR      | infrared                                                                |
| IRAS    | infrared absorption spectroscopy                                        |
| IRS     | infrared spectroscopy                                                   |
| IS      | ionization spectroscopy                                                 |
| ISO     | International Organization for Standardization                          |
| ISQ     | International System of Quantities                                      |
| ISS     | ion scattering spectroscopy                                             |
| ITS     | International Temperature Scale                                         |
| IUPAC   | International Union of Pure and Applied Chemistry                       |
| IUPAP   | International Union of Pure and Applied Physics                         |
| KS      | Kohn-Sham                                                               |
| L       | ligand                                                                  |
| L2TOFMS | laser desorption laser photoionization time-of-flight mass spectroscopy |
| LASER   | light amplification by stimulated emission of radiation                 |
| LC      | liquid chromatography                                                   |
| LCAO    | linear combination of atomic orbitals                                   |
| L-CCA   | linear coupled-cluster approximation                                    |
| LCMO    | linear combination of molecular orbitals                                |
| LED     | light-emitting diode                                                    |
| LEED    | low-energy electron diffraction                                         |
| LEELS   | low-energy electron energy-loss spectroscopy                            |
| LEES    | low-energy electron scattering                                          |
| LEF     | laser excitation fluorescence                                           |
| LEIS    | low-energy ion scattering                                               |
| LEPD    | low-energy positron diffraction                                         |

|        |                                                             |
|--------|-------------------------------------------------------------|
| LET    | linear energy transfer                                      |
| LH     | Lindemann-Hinshelwood [theory]                              |
| LID    | laser induced desorption                                    |
| LIDAR  | light detection and ranging                                 |
| LIF    | laser induced fluorescence                                  |
| LIGS   | laser induced grating spectroscopy                          |
| LIMA   | laser microprobe mass analysis                              |
| LIS    | laser isotope separation                                    |
| LMR    | laser magnetic resonance                                    |
| LUMO   | lowest unoccupied molecular orbital                         |
| M      | central metal                                               |
| MAR    | magic-angle rotation                                        |
| MAS    | magic-angle spinning                                        |
| MASER  | microwave amplification by stimulated emission of radiation |
| MBE    | molecular beam epitaxy                                      |
| MBGF   | many-body Green's function                                  |
| MBPT   | many-body perturbation theory                               |
| MC     | Monte Carlo                                                 |
| MCA    | multichannel analyser                                       |
| MCD    | magnetic circular dichroism                                 |
| MCS    | multichannel scalar                                         |
| MCSCF  | multiconfiguration self-consistent field                    |
| MD     | molecular dynamics                                          |
| MDS    | metastable deexcitation spectroscopy                        |
| MEED   | medium-energy electron diffraction                          |
| MEIS   | medium-energy ion scattering                                |
| MFM    | magnetic force microscopy                                   |
| MINDO  | modified incomplete neglect of differential overlap         |
| MIR    | mid-infrared                                                |
| MKS    | metre-kilogram-second                                       |
| MKSA   | metre-kilogram-second-ampere                                |
| MM     | molecular mechanics                                         |
| MO     | molecular orbital                                           |
| MOCVD  | metal-organic chemical vapor deposition                     |
| MOMBE  | metal-organic molecular beam epitaxy                        |
| MORD   | magnetic optical rotatory dispersion                        |
| MOS    | metal oxide semiconductor                                   |
| mp     | melting point                                               |
| MPI    | multiphoton ionization                                      |
| MPPT   | Møller-Plesset perturbation theory                          |
| MP-SCF | Møller-Plesset self-consistent field                        |
| MRD    | magnetic rotatory dispersion                                |
| MRI    | magnetic resonance imaging                                  |
| MS     | mass spectroscopy                                           |
| MW     | microwave                                                   |
| MW     | molecular weight (symbol: $M_r$ )                           |
| NAA    | neutron activation analysis                                 |
| NCE    | normal calomel electrode                                    |
| Nd:YAG | Nd doped YAG                                                |

|        |                                                   |
|--------|---------------------------------------------------|
| NETD   | noise equivalent temperature difference           |
| NEXAFS | near edge X-ray absorption fine structure         |
| NIR    | near-infrared                                     |
| NIR    | non-ionizing radiation                            |
| NMA    | nuclear microanalysis                             |
| NMR    | nuclear magnetic resonance                        |
| NOE    | nuclear Overhauser effect                         |
| NQR    | nuclear quadrupole resonance                      |
| NTP    | normal temperature and pressure                   |
| od     | outside diameter                                  |
| ODMR   | optically detected magnetic resonance             |
| OGS    | opto-galvanic spectroscopy                        |
| ORD    | optical rotatory dispersion                       |
| PAS    | photoacoustic spectroscopy                        |
| PC     | paper chromatography                              |
| PD     | photoelectron diffraction                         |
| PDG    | Particle Data Group                               |
| PED    | photoelectron diffraction                         |
| PEH    | photoelectron holography                          |
| PES    | photoelectron spectroscopy                        |
| PIES   | Penning ionization electron spectroscopy, see PIS |
| PIPECO | photoion-photoelectron coincidence [spectroscopy] |
| PIPN   | periodically poled lithium niobate                |
| PIS    | Penning ionization (electron) spectroscopy        |
| PMT    | photomultiplier tube                              |
| ppb    | part per billion                                  |
| pphm   | part per hundred million                          |
| ppm    | part per million                                  |
| PPP    | Pariser-Parr-Pople                                |
| PS     | see PES                                           |
| PSD    | photon stimulated desorption                      |
| pzc    | point of zero charge                              |
| QET    | quasi equilibrium theory                          |
| QMB    | quartz microbalance                               |
| QMC    | quantum Monte Carlo                               |
| QMS    | quadrupole mass spectrometer                      |
| RADAR  | radiowave detection and ranging                   |
| RBS    | Rutherford (ion) back scattering                  |
| RD     | rotatory dispersion                               |
| RDE    | rotating disc electrode                           |
| RDF    | radial distribution function                      |
| REM    | reflection electron microscopy                    |
| REMPI  | resonance enhanced multiphoton ionization         |
| RF     | radio frequency                                   |
| RHEED  | reflection high-energy electron diffraction       |
| RHF    | restricted Hartree-Fock                           |
| RIMS   | resonant ionization mass spectroscopy             |

|                  |                                                          |
|------------------|----------------------------------------------------------|
| RKR              | Rydberg-Klein-Rees [potential]                           |
| rms              | root-mean-square                                         |
| RRK              | Rice-Ramsperger-Kassel [theory]                          |
| RRKM             | Rice-Ramsperger-Kassel-Marcus [theory]                   |
| RRS              | resonance Raman spectroscopy                             |
| RS               | Raman spectroscopy                                       |
| RSPT             | Rayleigh-Schrödinger perturbation theory                 |
| S                | singlet                                                  |
| SACM             | statistical adiabatic channel model                      |
| SAM              | scanning Auger microscopy                                |
| SBS              | stimulated Brillouin scattering                          |
| SBZ              | surface Brillouin zone                                   |
| SCE              | saturated calomel electrode                              |
| SCF              | self-consistent field                                    |
| SDCI             | singly and doubly excited configuration interaction      |
| S <sub>E</sub>   | substitution electrophilic                               |
| SEELFS           | surface extended energy-loss fine structure              |
| SEFT             | spin-echo Fourier transform                              |
| SEM              | scanning [reflection] electron microscopy                |
| SEP              | stimulated emission pumping                              |
| SERS             | surface-enhanced Raman spectroscopy                      |
| SES              | secondary electron spectroscopy                          |
| SESCA            | scanning electron spectroscopy for chemical applications |
| SEXAFS           | surface extended X-ray absorption fine structure         |
| SF               | spontaneous fission                                      |
| SFG              | sum-frequency generation                                 |
| SHE              | standard hydrogen electrode                              |
| SHG              | second-harmonic generation                               |
| SI               | Système International d'Unités                           |
| SIMS             | secondary ion mass spectroscopy                          |
| SMOKE            | surface magneto-optic Kerr effect                        |
| S <sub>N</sub> 1 | substitution nucleophilic unimolecular                   |
| S <sub>N</sub> 2 | substitution nucleophilic bimolecular                    |
| S <sub>N</sub> i | substitution nucleophilic intramolecular                 |
| SOC              | spin-orbit coupling                                      |
| SOR              | synchrotron orbital radiation                            |
| SPIES            | surface Penning ionization electron spectroscopy         |
| SPLEED           | spin-polarized low-energy electron diffraction           |
| SPM              | scanning probe microscopy                                |
| SR               | synchrotron radiation                                    |
| SRS              | synchrotron radiation source                             |
| SSIMS            | static secondary ion mass spectroscopy                   |
| STEM             | scanning transmission [electron] microscopy              |
| STM              | scanning tunnelling (electron) microscopy                |
| STO              | Slater-type orbital                                      |
| STP              | standard temperature and pressure                        |
| SVLF             | single vibronic level fluorescence                       |

|        |                                                     |
|--------|-----------------------------------------------------|
| T      | triplet                                             |
| TCC    | thermal conductivity cell                           |
| TCD    | thermal conductivity detector                       |
| TCF    | time correlation function                           |
| TDL    | tuneable diode laser                                |
| TDMS   | tandem quadrupole mass spectroscopy                 |
| TDS    | thermal desorption spectroscopy                     |
| TED    | transmission electron diffraction                   |
| TEM    | transmission electron microscopy                    |
| TG     | thermogravimetry                                    |
| TGA    | thermogravimetric analysis                          |
| THEED  | transmission high-energy electron diffraction       |
| tlc    | thin layer chromatography                           |
| TOF    | time-of-flight [analysis]                           |
| TPD    | temperature programmed desorption                   |
| TPR    | temperature programmed reaction                     |
| TR3    | time-resolved resonance Raman scattering            |
| TST    | transition state theory                             |
|        |                                                     |
| UHF    | ultra high frequency                                |
| UHF    | unrestricted Hartree-Fock                           |
| UHV    | ultra high vacuum                                   |
| UP[E]S | ultraviolet photoelectron spectroscopy              |
| UV     | ultraviolet                                         |
|        |                                                     |
| VB     | valence bond                                        |
| VCD    | vibrational circular dichroism                      |
| VEELS  | vibrational electron energy-loss spectroscopy       |
| VHF    | very high frequency                                 |
| VIS    | visible                                             |
| VLEED  | very-low-energy electron diffraction                |
| VLSI   | very large scale integration                        |
| VPC    | vapor-phase chromatography                          |
| VSEPR  | valence shell electron pair repulsion               |
| VUV    | vacuum ultraviolet                                  |
|        |                                                     |
| WFC    | work function change                                |
|        |                                                     |
| X      | halogen                                             |
| XAFS   | X-ray absorption fine structure                     |
| XANES  | X-ray absorption near-edge structure [spectroscopy] |
| XAPS   | X-ray appearance potential spectroscopy             |
| XPD    | X-ray photoelectron diffraction                     |
| XPES   | X-ray photoelectron spectroscopy                    |
| XPS    | X-ray photoelectron spectroscopy                    |
| XRD    | X-ray diffraction                                   |
| XSW    | X-ray standing wave                                 |
|        |                                                     |
| YAG    | yttrium aluminium garnet                            |
|        |                                                     |
| ZPE    | zero point energy                                   |

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## 11 GREEK ALPHABET

| Roman   | Italics | Name             | Pronunciation and<br>Latin Equivalent | Notes |
|---------|---------|------------------|---------------------------------------|-------|
| A, α    | A, α    | alpha            | A                                     |       |
| B, β    | B, β    | beta             | B                                     |       |
| Γ, γ    | Γ, γ    | gamma            | G                                     |       |
| Δ, δ    | Δ, δ    | delta            | D                                     |       |
| E, ε    | E, ε, € | epsilon          | E                                     |       |
| Z, ζ    | Z, ζ    | zeta             | Z                                     |       |
| H, η    | H, η    | eta              | Ae, Ä, Ee                             | 1     |
| Θ, θ, θ | Θ, θ, θ | theta            | Th                                    | 2     |
| I, ι    | I, ι    | iota             | I                                     |       |
| K, κ, κ | K, κ, κ | kappa            | K                                     | 2     |
| Λ, λ    | Λ, λ    | lambda           | L                                     |       |
| M, μ    | M, μ    | mu, (my)         | M                                     |       |
| N, ν    | N, ν    | nu, (ny)         | N                                     |       |
| Ξ, ξ    | Ξ, ξ    | xi               | X                                     |       |
| O, ο    | O, ο    | omikron          | O                                     |       |
| Π, π    | Π, π    | pi               | P                                     |       |
| P, ρ    | P, ρ    | rho              | R                                     |       |
| Σ, σ, ς | Σ, σ, ς | sigma            | S                                     | 2, 3  |
| T, τ    | T, τ    | tau              | T                                     |       |
| Υ, υ    | Υ, υ    | upsilon, ypsilon | U, Y                                  |       |
| Φ, φ, φ | Φ, φ, φ | phi              | Ph                                    | 2     |
| X, χ    | X, χ    | chi              | Ch                                    |       |
| Ψ, ψ    | Ψ, ψ    | psi              | Ps                                    |       |
| Ω, ω    | Ω, ω    | omega            | Oo                                    | 4     |

(1) For the Latin equivalent Ae is to be pronounced as the German Ä. The modern Greek pronunciation of the letter η is like E, long ee as in cheese, or short i as in lips. The Latin equivalent is also often called “long E”.

(2) For the lower case letters theta, kappa, sigma and phi there are two variants in each case. For instance, the second variant of the lower case theta is sometimes called “varthetaeta” in printing.

(3) The second variant for lower case sigma is used in Greek only at the end of the word.

(4) In contrast to omikron (short o) the letter omega is pronounced like a long o.



## 12 INDEX OF SYMBOLS



This index lists symbols for physical quantities, units, some selected mathematical operators, states of aggregation, processes, and particles. Symbols for chemical elements are given in Section 6.2, p. 117. Qualifying subscripts and superscripts, etc., are generally omitted from this index, so that for example  $E_p$  for potential energy, and  $E_{ea}$  for electron affinity are both indexed under  $E$  for energy. The Latin alphabet is indexed ahead of the Greek alphabet, lower case letters ahead of upper case, bold ahead of italic, ahead of Roman, and single letter symbols ahead of multiletter ones. When more than one page reference is given, bold print is used to indicate the general (defining) reference. Numerical entries for the corresponding quantities are underlined.

- |                             |                                                                                        |                             |                                                                                         |
|-----------------------------|----------------------------------------------------------------------------------------|-----------------------------|-----------------------------------------------------------------------------------------|
| <b><i>a</i></b>             | acceleration, <b>13</b> , 90, 137                                                      | $A_H$                       | Hall coefficient, 43                                                                    |
| <b><i>a</i></b>             | fundamental translation vector, <b>42</b> , 79, 80                                     | $Al$                        | Alfvén number, 82                                                                       |
| <b><i>a</i><sup>*</sup></b> | reciprocal lattice vector, <b>42</b> , 80                                              | $A_r$                       | relative atomic mass, <b>47</b> , 117                                                   |
| <i>a</i>                    | absorption coefficient, 6, 36, 37, 40, 41                                              | $A$                         | ampere (SI unit), 4, 86, <b>87</b> , 139, 143                                           |
| <i>a</i>                    | activity, <b>57–59</b> , 70, 71, 72, 74, 75                                            | $A$                         | base-centred (crystal lattice), 44                                                      |
| <i>a</i>                    | area per molecule, <b>77</b> , 78                                                      | $A$                         | symmetry label, 31–33, 51                                                               |
| <i>a</i>                    | Hermann-Mauguin symbol, 44                                                             | $\text{\AA}$                | ångström (unit of length), 24, 27, <b>131</b> , <u>135</u>                              |
| <i>a</i>                    | hyperfine coupling constant, 27                                                        |                             |                                                                                         |
| <i>a</i>                    | specific surface area, 77                                                              | <b><i>b</i></b>             | Burgers vector, 42                                                                      |
| <i>a</i>                    | thermal diffusivity, 81                                                                | <b><i>b</i></b>             | fundamental translation vector, <b>42</b> , 79, 80                                      |
| <i>a</i>                    | unit cell length, 42                                                                   | <b><i>b</i><sup>*</sup></b> | reciprocal lattice vector, <b>42</b> , 80                                               |
| <i>a</i>                    | van der Waals coefficient, 57                                                          | <i>b</i>                    | breadth, 13                                                                             |
| $a_0$                       | bohr (unit of length), <u>135</u>                                                      | <i>b</i>                    | Hermann-Mauguin symbol, 44                                                              |
| $a_0$                       | Bohr radius, 9, 18, 20, <b>22</b> , 24, 26, 95, 96, <u>111</u> , 132, <u>135</u> , 145 | <i>b</i>                    | impact parameter, 65                                                                    |
| <i>a</i>                    | adsorbed (subscript), 59                                                               | <i>b</i>                    | mobility ratio, 43                                                                      |
| <i>a</i>                    | are (unit of area), <u>136</u>                                                         | <i>b</i>                    | molality, <b>48</b> , 49, 59, <b>62</b> , 70, 75, 133                                   |
| <i>a</i>                    | atom, <b>22</b> , 24                                                                   | <i>b</i>                    | Tafel slope, 72                                                                         |
| <i>a</i>                    | atto (SI prefix), 91                                                                   | <i>b</i>                    | unit cell length, 42                                                                    |
| <i>a</i>                    | symmetry label, 32, 33                                                                 | <i>b</i>                    | van der Waals coefficient, 57                                                           |
| <i>a</i>                    | year (unit of time), 24, <u>137</u>                                                    | <i>b</i>                    | Wang asymmetry parameter, 25                                                            |
| am                          | amorphous solid, <b>54</b> , 55                                                        | <i>b</i>                    | barn (unit of area), <b>121</b> , <u>136</u>                                            |
| aq                          | aqueous solution, <b>54</b> , 55, 57, 73, 74, 97                                       | <i>b</i>                    | symmetry label, 32, 33                                                                  |
| at                          | atomization (subscript), <b>59</b> , 61                                                | bar                         | bar (unit of pressure), 48, 62, <b>131</b> , <u>138</u>                                 |
| abs                         | absorbed (subscript), 34                                                               | <b><i>B</i></b>             | magnetic flux density (magnetic induction), <b>17</b> , 27, 81, 82, 141, 146–148        |
| ads, a                      | adsorption (subscript), 54, <b>59</b>                                                  | <i>B</i>                    | Debye-Waller factor, 42                                                                 |
| app                         | apparent (superscript), 60                                                             | <i>B</i>                    | Einstein coefficient, Einstein transition probability, <b>35</b> , 37, 39               |
| atm                         | atmosphere, 40, <b>62</b> , 131, <u>138</u>                                            | <i>B</i>                    | napierian absorbance, <b>36</b> , 37                                                    |
| amagat                      | amagat unit, <u>139</u>                                                                | <i>B</i>                    | retarded van der Waals constant, 77                                                     |
| <b><i>A</i></b>             | conservation matrix, formula matrix, 53                                                | <i>B</i>                    | rotational constant, 25–27                                                              |
| <b><i>A</i></b>             | magnetic vector potential, <b>17</b> , 146–148                                         | <i>B</i>                    | second virial coefficient, 57                                                           |
| <i>A</i>                    | absorbance, 5, <b>36</b> , 37, 39, 40                                                  | <i>B</i>                    | susceptance, 17                                                                         |
| <i>A</i>                    | electron affinity, 22                                                                  | <i>B</i>                    | base-centred (crystal lattice), 44                                                      |
| <i>A</i>                    | integrated absorption coefficient, 37–41                                               | <i>B</i>                    | bel (unit of power level), <b>92</b> , 99                                               |
| <i>A</i>                    | activity (radioactive), <b>24</b> , 89, 139                                            | <i>B</i>                    | biot (unit of electric current), <b>135</b> , <u>139</u> , 144                          |
| <i>A</i> , $\mathcal{A}$    | affinity of reaction, <b>58</b> , 61                                                   | <i>Bi</i>                   | becquerel (SI unit), <b>89</b> , <u>139</u>                                             |
| <i>A</i>                    | area, <b>13</b> , 14–17, 24, 34–36, 48, 49, 56, 71, 72, 77, 78, 81, 90, 136, 146       | <i>Bq</i>                   | British thermal unit (unit of energy), <u>137</u>                                       |
| <i>A</i>                    | Einstein coefficient, Einstein transition probability, <b>35</b> , 37, 39              | <i>Btu</i>                  |                                                                                         |
| <i>A</i>                    | Helmholtz energy, <b>56</b> , 57, 78                                                   | <b><i>c</i></b>             | fundamental translation vector, 42                                                      |
| <i>A</i>                    | hyperfine coupling constant, 27                                                        | <b><i>c</i></b>             | velocity, <b>13</b> , 45, 90, 148                                                       |
| <i>A</i>                    | nucleon number, mass number, <b>22</b> , 49, 121                                       | <b><i>c</i><sup>*</sup></b> | reciprocal lattice vector, <b>42</b> , 80                                               |
| <i>A</i>                    | pre-exponential factor, 63, <b>64</b>                                                  | <i>c</i>                    | amount concentration, 4, 37, 38, <b>48</b> , 54, 58, 59, 62–67, 70, 72, 81, 82, 90, 131 |
| <i>A</i>                    | rotational constant, 25–27                                                             | <i>c</i>                    | Hermann-Mauguin symbol, 44                                                              |
| <i>A</i>                    | spin-orbit coupling constant, 25                                                       | <i>c</i>                    | specific heat capacity (constant pressure), <b>6</b> , 81, 82                           |
| <i>A</i>                    | van der Waals-Hamaker constant, 77                                                     |                             |                                                                                         |

|                    |                                                                                                                            |              |                                                                                                              |
|--------------------|----------------------------------------------------------------------------------------------------------------------------|--------------|--------------------------------------------------------------------------------------------------------------|
| $c$                | speed, <b>13</b> , 64, 81, 90, 93, 95                                                                                      | $D$          | absorbed dose of radiation, <u>139</u>                                                                       |
| $c$                | unit cell length, 42                                                                                                       | $D$          | centrifugal distortion constant, 25                                                                          |
| $c$                | speed of light in medium, 34,                                                                                              | $D$          | Debye-Waller factor, 42                                                                                      |
| $c_0$              | speed of light in vacuum, 9, 13, 16, 22,<br>23, 25, <b>34–39</b> , 41, 43, 95, <u>111</u> , 112,<br>131, 134, 143, 145–148 | $D$          | diffusion coefficient, <b>43</b> , 44, 72, 81, 90                                                            |
| $c_1$              | first radiation constant, 36, <u>112</u>                                                                                   | $D$          | dissociation energy, <b>22</b> , 96                                                                          |
| $c_2$              | second radiation constant, 36, <u>112</u>                                                                                  | $D^T$        | thermal diffusion coefficient, 81                                                                            |
| $c$                | centered superlattice, 79                                                                                                  | $D_{AB}$     | dipolar coupling constant, 28                                                                                |
| $c$                | centi (SI prefix), 91                                                                                                      | $D$          | debye (unit of electric dipole moment),<br><b>26</b> , 39, <u>141</u>                                        |
| $c$                | combustion (subscript), <b>59</b> , 61                                                                                     | Da           | dalton (unit of mass), 9, 22, 47, <b>92</b> , 94,<br><u>136</u>                                              |
| $c$                | symmetry label, 32                                                                                                         |              |                                                                                                              |
| cd                 | candela (SI unit), 34, 86, <b>88</b>                                                                                       | $e$          | unit vector, <b>13</b> , 81                                                                                  |
| cd                 | condensed phase, 40, 41, 48, <b>54</b> , 58                                                                                | $e$          | elementary charge, 9, 18–20, <b>22–24</b> , 26, 29,<br>37, 43, 70, 92, 94–96, <u>111</u> , 139, 145          |
| cr                 | crystalline, <b>54</b> , 55, 61                                                                                            | $e$          | étendue, <b>35</b> , 36                                                                                      |
| cal                | calorie (unit of energy), <b>58</b> , <u>137</u>                                                                           | $e$          | linear strain, 15                                                                                            |
| ccc                | critical coagulation concentration, 78                                                                                     | $e$          | base of natural logarithm, <b>8</b> , 99, 103, 106, 112                                                      |
| cmc                | critical miscellisation concentration, 78                                                                                  | $e$          | electron, 8, <b>50</b> , <u>115</u>                                                                          |
| $\mathbf{C}$       | spin-rotation interaction tensor, 28                                                                                       | $e$          | symmetry label, 33                                                                                           |
| $C$                | capacitance, <b>16</b> , 140, 146                                                                                          | eV           | electronvolt (unit of energy) 9, 92, <b>94</b> ,<br>132, <u>137</u>                                          |
| $C$                | heat capacity, 5, 6, 8, 43, 55–57,<br>90, 138                                                                              | erg          | erg (unit of energy), 134, 135, <u>137</u> , 138,<br>142, 144                                                |
| $C$                | integrated absorption coefficient<br>(condensed phase), 41                                                                 | e.u.         | entropy unit, <u>139</u>                                                                                     |
| $C$                | number concentration, number density,<br>43, <b>45</b> , 48, 63–65, 68, 69, 81                                             | $\mathbf{E}$ | electric field strength, 7, <b>16</b> , 24, 26, 40, 43, 73,<br>90, 104, 134, 140, 146, 148                   |
| $C$                | rotational constant, 25–27                                                                                                 | $E$          | activation energy, threshold energy, <b>64</b> , 66, 67                                                      |
| $C$                | third virial coefficient, 57                                                                                               | $E$          | cell potential, <b>71</b> , 74, 75                                                                           |
| $C$                | vibrational force constant, 27                                                                                             | $E$          | electric potential difference, <b>16</b> , 44, 74                                                            |
| $C_n$              | $n$ -fold rotation symmetry operator, 31, 32                                                                               | $E$          | electromotive force, cell potential, <b>16</b> , 89                                                          |
| $Co$               | Cowling number, 82                                                                                                         | $E$          | energy, 5, <b>14</b> , 15, 17, 19–23, 32, 33, 39, 43, 45,<br>56, 65, 67, 95, 132, 137, 140, 146,<br>147, 237 |
| $C$                | base-centred (crystal lattice), 44                                                                                         | $E$          | étendue, <b>35</b> , 36                                                                                      |
| $C$                | coulomb (SI unit), <b>89</b> , 134, 139                                                                                    | $E$          | identity symmetry operator, 31                                                                               |
| Ci                 | curie (unit of radioactivity), <u>139</u>                                                                                  | $E$          | irradiance, <b>35</b> , 90                                                                                   |
| Cl                 | clausius (unit of entropy), <u>138</u>                                                                                     | $E$          | modulus of elasticity, 15                                                                                    |
| $^{\circ}\text{C}$ | degree Celsius (SI unit), 56, <b>89</b> , 138                                                                              | $E$          | potential (electrochemistry), <b>71</b> , 72, 76                                                             |
| $d$                | centrifugal distortion constant, 25                                                                                        | $E$          | scattering amplitude, 42                                                                                     |
| $d$                | collision diameter, 64                                                                                                     | $E$          | thermoelectric force, <b>43</b> , 44                                                                         |
| $d$                | degeneracy, statistical weight, 26, 39, <b>45</b>                                                                          | $E^*$        | space-fixed inversion symmetry operator, 31                                                                  |
| $d$                | diameter, distance, thickness, <b>13</b> , 15, 146                                                                         | $E_h$        | Hartree energy, 9, 18, 20, 22, <b>95</b> , 96, <u>111</u> ,<br>132, 137, 139, 145, 147                       |
| $d$                | lattice plane spacing, 42                                                                                                  | $Eu$         | Euler number, 82                                                                                             |
| $d$                | relative density, 14                                                                                                       | $E[x]$       | expectation value of $x$ , 151                                                                               |
| d                  | day (unit of time), <b>92</b> , 136, <u>137</u>                                                                            | E            | exa (SI prefix), 91                                                                                          |
| d                  | deci (SI prefix), 91                                                                                                       | E            | excess quantity (superscript), 60                                                                            |
| d                  | deuteron, 50, <u>115</u>                                                                                                   | E            | symmetry label, 31, 32                                                                                       |
| da                 | deca (SI prefix), 91                                                                                                       | Ei           | exabinary (prefix for binary), 91                                                                            |
| dB                 | decibel, <i>see</i> bel (unit of power level), <b>92</b> ,<br>99                                                           | $f$          | activity coefficient, <b>59</b> , 61, 70, 75                                                                 |
| deg                | degree (unit of plane angle), <b>92</b> , <u>136</u>                                                                       | $f$          | atomic scattering factor, 42                                                                                 |
| dil                | dilution (subscript), 59                                                                                                   | $f$          | finesse, 36                                                                                                  |
| dpl                | displacement (subscript), 59                                                                                               | $f$          | frequency, <b>13</b> , 28, 29, 34–36, 41, 68, 81,                                                            |
| dyn                | dyne (unit of force), 27, 134, <u>137</u> , 144                                                                            |              |                                                                                                              |
| $D$                | dipolar interaction tensor, 28                                                                                             |              |                                                                                                              |
| $D$                | electric displacement, <b>16</b> , 141, 145, 146,<br>148                                                                   |              |                                                                                                              |

|                       |                                                                           |                       |                                                                                                                                                               |
|-----------------------|---------------------------------------------------------------------------|-----------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|
|                       | 82, 88, 89, 129                                                           |                       |                                                                                                                                                               |
| $f$                   | friction factor, 15                                                       | $G$                   | shear modulus, 15                                                                                                                                             |
| $f$                   | fugacity, <b>58</b> , 74                                                  | $G$                   | thermal conductance, 81                                                                                                                                       |
| $f$                   | oscillator strength, 37                                                   | $G$                   | vibrational term, 25                                                                                                                                          |
| $f$                   | vibrational force constant, 27                                            | $G$                   | weight, 14                                                                                                                                                    |
| $f(c_x)$              | velocity distribution function, 45                                        | $G_F$                 | Fermi coupling constant, <u>111</u>                                                                                                                           |
| $f$                   | femto (SI prefix), 91                                                     | $Gr$                  | Grashof number (mass transfer), 82                                                                                                                            |
| $f$                   | fermi (unit of length), <u>135</u>                                        | $G$                   | gauss (unit of magnetic flux density),<br>27, <u>141</u> , 142                                                                                                |
| $f$                   | formation reaction (subscript), <b>60</b> , 61                            | $G$                   | giga (SI prefix), 91                                                                                                                                          |
| $f$                   | fluid phase, 54                                                           | $Gi$                  | gigabinary (prefix for binary), 91                                                                                                                            |
| $ft$                  | foot (unit of length), <u>135</u>                                         | $Gy$                  | gray (SI unit), <b>89</b> , <u>139</u>                                                                                                                        |
| $fus$                 | fusion, melting (subscript), 60                                           | $Gal$                 | gal (unit of acceleration), <u>137</u>                                                                                                                        |
| <b><math>F</math></b> | Fock operator, <b>20</b> , 21                                             | $h$                   | coefficient of heat transfer, <b>81</b> , 82                                                                                                                  |
| <b><math>F</math></b> | force, 7, <b>14</b> –17, 87, 89, 95, 104, 137, 138,<br>143, 144, 146–148  | $h$                   | film thickness, 77                                                                                                                                            |
| <b><math>F</math></b> | angular momentum, 30                                                      | $h$                   | height, 13                                                                                                                                                    |
| $F$                   | Faraday constant, <b>70</b> –75, <u>111</u>                               | $h, \hbar$            | Miller index, <b>44</b> , 79, 80                                                                                                                              |
| $F$                   | fluence, 35                                                               |                       | Planck constant ( $\hbar = h/2\pi$ ), 7, 9, 18, 20,<br>22, 23, 25, 28–30, <b>34</b> –36, 38, 39, 41–43,<br>45, 65–68, 95, 96, <u>111</u> , 112, 131, 132, 145 |
| $F$                   | frequency, 28                                                             | $h$                   | hecto (SI prefix), 91                                                                                                                                         |
| $F$                   | Helmholtz energy, 56                                                      | $h$                   | helion, <b>50</b> , <u>115</u>                                                                                                                                |
| $F$                   | rotational term, 25                                                       | $h$                   | hour (unit of time), <b>92</b> , <u>137</u>                                                                                                                   |
| $F$                   | structure factor, 42                                                      | $ha$                  | hectare (unit of area), <u>136</u>                                                                                                                            |
| $F$                   | vibrational force constant, 27                                            | $hp$                  | horse power (unit of power), <u>138</u>                                                                                                                       |
| $F(c)$                | speed distribution function, 45                                           | $hk$                  | metric horse power (unit of power), <u>138</u>                                                                                                                |
| $Fo$                  | Fourier number (mass transfer), 82                                        |                       |                                                                                                                                                               |
| $Fr$                  | Froude number, 82                                                         | <b><math>H</math></b> | magnetic field strength, <b>17</b> , 90, 142, 145,<br>147, 148                                                                                                |
| $F$                   | face-centred (crystal lattice), 44                                        | $H$                   | coulomb integral, resonance integral, 19                                                                                                                      |
| $F$                   | farad (SI unit), <b>89</b> , 140                                          | $H$                   | dose equivalent, <u>139</u>                                                                                                                                   |
| $F$                   | symmetry label, 31                                                        | $H$                   | enthalpy, 5, 6, 8, 55–58, 60, 65, 66                                                                                                                          |
| $^{\circ}F$           | degree Fahrenheit (unit of temperature),<br><u>138</u>                    | $H$                   | fluence, 35                                                                                                                                                   |
| $Fr$                  | franklin (unit of electric charge), <b>134</b> , 135,<br><u>139</u> , 144 | $H$                   | Hamilton function, hamiltonian, <b>14</b> , <b>18</b> ,<br>20, 21                                                                                             |
| <b><math>g</math></b> | reciprocal lattice vector, 80                                             | $Ha$                  | Hartmann number, 82                                                                                                                                           |
| $g$                   | acceleration of free fall, 13, <b>78</b> , 81, 82,<br><u>112</u> , 137    | $H$                   | Heaviside function, 45, 66, 67, <b>107</b>                                                                                                                    |
| $g$                   | degeneracy, statistical weight, 26, 39, <b>45</b>                         | $H$                   | henry (SI unit), <b>89</b> , 142                                                                                                                              |
| $g$                   | (spectral) density of vibrational modes, 43                               | $Hz$                  | hertz (SI unit), 6, 13, 29, 88, <b>89</b>                                                                                                                     |
| $g, g_e$              | $g$ -factor, 23, <b>27</b> , 28, <u>111</u>                               | <b><math>i</math></b> | unit vector, 13                                                                                                                                               |
| $g$                   | vibrational anharmonicity constant, 25                                    | $i$                   | electric current, <b>16</b> , 72, 85, 86, 95, 139,<br>143, 144                                                                                                |
| $g$                   | gas, <b>54</b> , 61, 73                                                   | $i$                   | inversion symmetry operator, 31, 32                                                                                                                           |
| $g$                   | gram (unit of mass), 91, <u>136</u>                                       | $i$                   | square root of $-1$ , 18, 32, 42, 43, 98, 103, <b>107</b>                                                                                                     |
| $g$                   | gerade symmetry label, 32, 33                                             | $id$                  | ideal (superscript), 60                                                                                                                                       |
| $gr$                  | grain (unit of mass), <u>136</u>                                          | $in$                  | inch (unit of length), <u>135</u> , 138                                                                                                                       |
| $gal$                 | gallon (unit of volume), <u>136</u>                                       | $ir$                  | irrational (superscript), <b>145</b> , <i>see</i> non-rationalized                                                                                            |
| $gon$                 | gon (unit of angle), <u>136</u>                                           | $imm$                 | immersion (subscript), 60                                                                                                                                     |
| <b><math>G</math></b> | reciprocal lattice vector, <b>42</b> , 80                                 | $iep$                 | isoelectric point, 78                                                                                                                                         |
| $G$                   | (electric) conductance, <b>17</b> , 89                                    |                       |                                                                                                                                                               |
| $G$                   | Gibbs energy, 5, 6, <b>56</b> –58, 60, 61, 65, 66,<br>71, 74, 77          | <b><math>I</math></b> | nuclear spin angular momentum, 23, 27– <b>30</b> , 121                                                                                                        |
| $G$                   | gravitational constant, 14, <u>112</u>                                    | $I$                   | differential cross section, 65                                                                                                                                |
| $G$                   | integrated (net) absorption cross section,<br>37–40                       | $I$                   | electric current, 4, <b>16</b> , 17, 72, 85, 86, 95,<br>139, 143, 146, 147                                                                                    |
|                       |                                                                           | $I$                   | intensity, 4, <b>34</b> –37, 85, 86, 88                                                                                                                       |

|           |                                                                                  |          |                                                                                                                     |
|-----------|----------------------------------------------------------------------------------|----------|---------------------------------------------------------------------------------------------------------------------|
| $I$       | ionic strength, <b>59</b> , 70, 76                                               | $K$      | kinetic energy, <b>14</b> , 93, 94                                                                                  |
| $I$       | ionization energy, 22                                                            | $K_M$    | Michaelis constant, 66                                                                                              |
| $I$       | moment of inertia, <b>14</b> , 25                                                | $K_{AB}$ | reduced nuclear spin-spin coupling constant, 28                                                                     |
| $I$       | body-centred (crystal lattice), 44                                               | $Kn$     | Knudsen number, 82                                                                                                  |
| $j$       | internal vibrational angular momentum, 30                                        | $K$      | kelvin (SI unit), 3, 5, 40, 56, 86, <b>87</b> , 89, 138                                                             |
| $j$       | electric current density, <b>16</b> –18, 43, 72, 73, 90, 146–148                 | $Ki$     | kilobinary (prefix for binary), 91                                                                                  |
| $j$       | particle flux density, 43, <b>44</b>                                             |          |                                                                                                                     |
| $j$       | unit vector, 13                                                                  | $l$      | electron orbital angular momentum, <b>30</b> , 31                                                                   |
| $J$       | angular momentum, 30, 138                                                        | $l$      | internal vibrational angular momentum, 30                                                                           |
| $J$       | coulomb operator, <b>20</b> , 21                                                 | $l$      | cavity spacing, path length, 36, 37                                                                                 |
| $J$       | electric current density, <b>16</b> , 18, 72, 90                                 | $l$      | length, 4, <b>13</b> , 20, 42, 45, 53, 81, 82, 85–87, 103, 131, 135, 136, 143–145, 147                              |
| $J$       | heat flux vector, 43, <b>44</b>                                                  | $l$      | mean free path, 43, 44                                                                                              |
| $J$       | angular momentum quantum number (component), 31–33, 51, 66, 67, 95               | $l$      | Miller index, <b>44</b> , 79, 80                                                                                    |
| $J$       | coulomb integral, 20                                                             | $l$      | vibrational quantum number, 26, <b>30</b>                                                                           |
| $J$       | flux density, 81                                                                 | $l$      | liquid, 54, 55, 61                                                                                                  |
| $J$       | Massieu function, 56                                                             | $l$      | litre (unit of volume), 6, 92, <u>136</u>                                                                           |
| $J$       | moment of inertia, 14                                                            | $lb$     | pound (unit of mass), <u>136</u>                                                                                    |
| $J_{AB}$  | spin-spin coupling constant, 28                                                  | $lc$     | liquid crystal, 54                                                                                                  |
| $J$       | joule (SI unit), 6, 9, 29, <b>89</b> , 93, 137                                   | $lm$     | lumen (SI unit), 89                                                                                                 |
|           |                                                                                  | $lx$     | lux (SI unit), 89                                                                                                   |
| $k$       | unit vector, 13                                                                  | $l.y.$   | light year (unit of length), <u>136</u>                                                                             |
| $k$       | angular wave vector, 43                                                          | $l\ atm$ | litre atmosphere (unit of energy), <u>137</u>                                                                       |
| $k$       | absorption index, <b>37</b> , 40, 41                                             | $L$      | angular momentum, 14, 25, <b>30</b> , 32, 138                                                                       |
| $k$       | angular momentum quantum number (component), 30                                  | $L$      | Avogadro constant, 9, <b>45</b> , 47, 53, 65, 68, 78, <u>111</u> , 112, 132                                         |
| $k, k_B$  | Boltzmann constant, 36, 38, 39, <b>45</b> , 46, 64–67, 81, <u>111</u> , 112, 133 | $L$      | angular momentum quantum number, 30, 32                                                                             |
| $k$       | coefficient of heat transfer, 81                                                 | $L_D$    | Debye length, 77                                                                                                    |
| $k$       | coverage factor, 151–154                                                         | $L$      | diffusion length, 49                                                                                                |
| $k_D$     | Debye angular wavenumber, 43                                                     | $L$      | field level, power level, <b>98</b> , 99                                                                            |
| $k$       | decay constant, 24                                                               | $L$      | inductance, <b>17</b> , 89, 142, 147                                                                                |
| $k$       | Miller index, <b>44</b> , 79, 80                                                 | $L$      | Lagrangian function, Lagrangian, <b>14</b> , <b>45</b>                                                              |
| $k$       | rate coefficient, rate constant, 63–69, 72                                       | $L$      | length, 3, 4, 6, <b>13</b> , 20, 42, 45, 53, 81, 85–87, 92, 94, 95, 131, 135, 136, 143–145, 147                     |
| $k$       | thermal conductivity, <b>81</b> , 82, 90                                         | $L$      | Lorenz coefficient, 43                                                                                              |
| $k_d$     | mass transfer coefficient, <b>72</b> , 81, 82                                    | $L$      | radiance, 35                                                                                                        |
| $k_H$     | Henry's law constant, <b>58</b> , 59                                             | $L$      | term symbol, 117–120                                                                                                |
| $k_{rst}$ | vibrational force constant, 27                                                   | $Le$     | Lewis number, 82                                                                                                    |
| $k$       | kilo (SI prefix), 91                                                             | $L$      | langmuir (unit of pressure-time product), 81                                                                        |
| $kg$      | kilogram (SI unit), 4, 86, <b>87</b> , 91, <u>136</u> , 143                      | $L$      | litre (unit of volume), 6, <b>92</b> , <u>136</u>                                                                   |
| $kp$      | kilopond (unit of force), <u>137</u>                                             | $m$      | magnetic dipole moment, 17, <b>23</b> , 95, 121, 133, 142, 147                                                      |
| $kat$     | katal, 89                                                                        | $m$      | angular momentum quantum number (component), 28, <b>30</b>                                                          |
| $kgf$     | kilogram-force (unit of force), <u>137</u>                                       | $m^*$    | effective mass, 43                                                                                                  |
| $K$       | exchange operator, <b>20</b> , 21                                                | $m$      | electric mobility, 73                                                                                               |
| $K$       | rate (coefficient) matrix, <b>66</b> , 67                                        | $m$      | mass, 4, 6, 7, 9, <b>14</b> , 18, 22, 28, 45, <b>47</b> , 48, 64, 77, 81, 92, 93, 111, 115, 116, 133, 136, 143, 144 |
| $K$       | absorption coefficient, 36                                                       | $m$      | Hermann-Mauguin symbol, 44                                                                                          |
| $K$       | angular momentum quantum number (component), <b>30</b> , 31, 33                  | $m$      | molality, <b>48</b> , 49, 58, 59, 62, 70, 75, 133                                                                   |
| $K$       | bulk modulus, 15                                                                 |          |                                                                                                                     |
| $K$       | coefficient of heat transfer, 81                                                 |          |                                                                                                                     |
| $K$       | equilibrium constant, <b>58</b> , 59, 61                                         |          |                                                                                                                     |
| $K$       | exchange integral, 20                                                            |          |                                                                                                                     |

|               |                                                                                                           |            |                                                                                                 |
|---------------|-----------------------------------------------------------------------------------------------------------|------------|-------------------------------------------------------------------------------------------------|
| $m$           | order of reaction, 63, <b>66</b> , 72                                                                     | $N$        | number of entities, 6, 24, 42, <b>45</b> , 47, 77, 78, 89                                       |
| $m_u$         | atomic mass constant, <b>22</b> , 47, 94, <u>111</u> , 117                                                | $N$        | number of events, 13, 35                                                                        |
| $m_e$         | electron mass, 9, 18, 20, 22, 23, 29, 37, 94, <b>95</b> , 96, <u>111</u> , 132, 136                       | $N$        | number of measurements, 151                                                                     |
| $m_n$         | neutron mass, <u>111</u>                                                                                  | $N$        | number of states, 44, 45, <b>66</b> , 67                                                        |
| $m_p$         | proton mass, 29, <u>111</u>                                                                               | $N_A$      | Avogadro constant, 9, 17, 37, 41, 43, <b>45</b> , 47, 53, 92, <u>111</u> , 112, 117, 133        |
| $m_W$         | W-boson mass, <u>111</u>                                                                                  | $N_E$      | density of states, 43, <b>45</b> , 66, 67                                                       |
| $m_Z$         | Z-boson mass, <u>111</u>                                                                                  | $Nu$       | Nusselt number (mass transfer), 82                                                              |
| m             | metre (SI unit), 3, 6, 26, 29, 40, 53, 86, <b>87</b> , <u>135</u> , 143                                   | $N_\omega$ | (spectral) density of vibrational modes, <b>43</b> , 44                                         |
| m             | milli (SI prefix), 91                                                                                     | $N$        | newton (SI unit), <b>89</b> , <u>137</u>                                                        |
| mi            | mile (unit of length), <u>135</u>                                                                         | $N$        | nucleus, 22                                                                                     |
| min           | minute (unit of time), <b>92</b> , <u>137</u>                                                             | $Np$       | neper, 8, <b>92</b> , 98, 99                                                                    |
| mix           | mixing (subscript), 60                                                                                    |            |                                                                                                 |
| mol           | mole (SI unit), 4, 6, 40, 43, 47, 53, 59, 86, <b>88</b> , 98                                              | oz         | ounce (unit of mass), <u>136</u>                                                                |
| mon           | monomeric form, 54                                                                                        |            |                                                                                                 |
| mmHg          | millimetre of mercury (unit of pressure), <u>138</u>                                                      | Oe         | oerstedt (unit of magnetic field strength), <u>142</u>                                          |
| $M$           | magnetization, <b>17</b> , 28, 142, 147                                                                   | $p$        | electric dipole moment, 17, 23, <b>26</b> , 39, 41, 95, 134, 141, 146                           |
| $M$           | torque, 14                                                                                                | $p$        | momentum, <b>14</b> , 18, <b>45</b>                                                             |
| $M$           | transition dipole moment, 26                                                                              | $p$        | bond order, <b>19</b> , 21                                                                      |
| $M$           | angular momentum quantum number (component), 30                                                           | $p$        | fractional population, <b>38</b> , 39, 46                                                       |
| $M$           | matrix for superlattice notation, <b>79</b> , 80                                                          | $p$        | number density, number concentration, 43, 44                                                    |
| $M$           | molar mass, <b>47</b> , 54, 59, 117, 133                                                                  | $p$        | permutation symmetry operator, 31                                                               |
| $M_u$         | molar mass constant, <b>47</b> , 117                                                                      | $p$        | pressure, 3, 5–8, <b>14</b> , 15, 37–39, 48, 54, 56–58, 61–63, 72, 77, 81, 82, 89, 99, 131, 138 |
| $M$           | mutual inductance, 17                                                                                     | $p$        | probability, 45                                                                                 |
| $M$           | radiant excitance, <b>34</b> , 35                                                                         | $p^*$      | permutation-inversion symmetry operator, 31                                                     |
| $\mathcal{M}$ | Madelung constant, 43                                                                                     | p          | pico (SI prefix), 91                                                                            |
| $M_r$         | relative molecular mass, 47                                                                               | p          | primitive superlattice, 79                                                                      |
| $Ma$          | Mach number, 82                                                                                           | p          | proton, 8, 22, <b>50</b> , <u>115</u>                                                           |
| M             | mega (SI prefix), 91                                                                                      | pc         | parsec (unit of length), <u>136</u>                                                             |
| M             | nautical mile (unit of length), 92                                                                        | pH         | pH, 5, 70, <b>75</b> , 76                                                                       |
| M             | molar (unit of concentration), <b>48</b> , 49                                                             | pol        | polymeric form, 54                                                                              |
| Mi            | megabinary (prefix for binary), 91                                                                        | ppb        | part per billion, 98                                                                            |
| Mx            | maxwell (unit of magnetic flux), <u>142</u>                                                               | pph        | part per hundred, 98                                                                            |
|               |                                                                                                           | ppm        | part per million, 98                                                                            |
| $n$           | amount of substance, chemical amount, 4–7, 45, 47, 48, 51, <b>53</b> , 56, 63, 71, 77, 78, 85, 86, 88, 89 | ppq        | part per quadrillion, 98                                                                        |
| $n$           | charge number of cell reaction, <b>71</b> , 72                                                            | ppt        | part per thousand, part per trillion, 98                                                        |
| $n$           | number density, number concentration, 43– <b>45</b> , 48, 49, 53, 54, 67, 71, 81, 133                     | psi        | pounds per square inch (unit of pressure), <u>138</u>                                           |
| $n$           | number of electrons, 20                                                                                   | pzc        | point of zero charge, 78                                                                        |
| $n$           | order of (Bragg) reflection, 42                                                                           | pphm       | part per hundred million, 98                                                                    |
| $n$           | order of reaction, 63, <b>66</b> , 72                                                                     |            |                                                                                                 |
| $n$           | principal quantum number, 23                                                                              | $P$        | density matrix, 21, <b>46</b>                                                                   |
| $n$           | refractive index, <b>34</b> , 37, 40, 41                                                                  | $P$        | dielectric polarization, <b>16</b> , 40, 146                                                    |
| n             | nano (SI prefix), 91                                                                                      | $P$        | heat flux, thermal power, <b>57</b> , <b>81</b>                                                 |
| n             | nematic phase, 54                                                                                         | $P$        | permutation symmetry operator, 31                                                               |
| n             | neutron, 8, 22, <b>50</b> , <u>115</u>                                                                    | $P$        | power, <b>14</b> , 22, 98, 99, 138                                                              |
| $N$           | angular momentum, 30                                                                                      | $P$        | pressure, <b>14</b> , 39, 48, 58, 81, 89, 131, 138                                              |
| $N$           | neutron number, <b>22</b> , 24                                                                            | $P$        | probability, <b>45</b> , 46, 65, 153, 154                                                       |

|       |                                                                                     |             |                                                                                            |
|-------|-------------------------------------------------------------------------------------|-------------|--------------------------------------------------------------------------------------------|
| $P$   | probability density, <b>18</b> , <b>151</b>                                         | $R$         | particle position vector, 42                                                               |
| $P$   | radiant power, <b>34–36</b>                                                         | $R$         | transition dipole moment, 26                                                               |
| $P$   | sound energy flux, 15                                                               | $R$         | electric resistance, <b>17</b> , 89, 140                                                   |
| $P$   | transition probability, 65                                                          | $R$         | gas constant, 7, 38, <b>45</b> , 46, 54, 57–59, 61, 62, 64, 65, 71, 72, 74, 75, <b>111</b> |
| $P$   | (volume) polarization, 141                                                          | $R_H$       | Hall coefficient, 43                                                                       |
| $P$   | weight, 14                                                                          | $R$         | internal vibrational coordinate, 27                                                        |
| $P^*$ | permutation-inversion symmetry operator, 31                                         | $R$         | molar refraction, 37                                                                       |
| $Pe$  | Péclet number (mass transfer), 82                                                   | $R$         | position vector, <b>42</b> , 43                                                            |
| $Pr$  | Prandtl number, 82                                                                  | $R$         | reflectance, 36                                                                            |
| $P$   | peta (SI prefix), 91                                                                | $R$         | resolving power, 36                                                                        |
| $P$   | poise (unit of dynamic viscosity), <b>138</b>                                       | $R_\infty$  | Rydberg constant, 22, 23, <b>111</b>                                                       |
| $P$   | primitive (crystal lattice), 44                                                     | $R$         | thermal resistance, 81                                                                     |
| $P$   | symmetry label, 51                                                                  | $Ra$        | Rayleigh number, 82                                                                        |
| $Pa$  | pascal (SI unit), 40, 48, 62, <b>89</b> , <b>138</b>                                | $Re$        | Reynolds number, 5, <b>82</b>                                                              |
| $Pi$  | petabinary (prefix for binary), 91                                                  | $Rm, R_m$   | magnetic Reynolds number, 82                                                               |
| $q$   | electric field gradient, 24, <b>29</b> , 141                                        | $R$         | rhombohedral (crystal lattice), 44                                                         |
| $q$   | angular wave vector, 42, <b>43</b>                                                  | $R$         | röntgen (unit of exposure), 139                                                            |
| $q_D$ | Debye angular wavenumber, 43                                                        | $Ry$        | rydberg (unit of energy), <b>137</b>                                                       |
| $q$   | charge density, <b>16</b> , 18, 23, 43, 44, 140                                     | $^{\circ}R$ | degree Rankine (unit of thermodynamic temperature), <b>138</b>                             |
| $q$   | charge order, 19                                                                    |             |                                                                                            |
| $q$   | flux of mass, 81                                                                    | $s$         | electron spin angular momentum, 30                                                         |
| $q$   | generalized coordinate, <b>13</b> , 14, 45                                          | $s$         | length of path, length of arc, 13                                                          |
| $q$   | heat, <b>56</b> , 73, 89                                                            | $s$         | order parameter, 42                                                                        |
| $q$   | partition function, 39, <b>45</b> , 46, 66, 67                                      | $s$         | sedimentation coefficient, 13, <b>77</b>                                                   |
| $q$   | vibrational normal coordinate, 27                                                   | $s$         | solubility, <b>48</b> , 49                                                                 |
| $Q$   | quadrupole moment, 23, <b>24</b> , 28, 29, 141                                      | $s$         | standard deviation, <b>151</b> , 152                                                       |
| $Q$   | disintegration energy, 24                                                           | $s$         | symmetry number, 46                                                                        |
| $Q$   | electric charge, 4, <b>16</b> , 17, 50, 70–72, 89, 95, 134, 139, 143, 145, 146, 148 | $s^2$       | variance, <b>151</b> , 152                                                                 |
| $Q_W$ | electroweak charge, 24                                                              | $s$         | second (SI unit), 6, 29, 30, 34, 86, <b>87</b> , 92, 98, 137, 143, 144                     |
| $Q$   | heat, <b>56</b> , 57, 73, 81, 89                                                    | $s$         | solid, <b>54</b> , 73                                                                      |
| $Q$   | partition function, 39, <b>45</b> , 46, 67                                          | $sr$        | steradian (SI unit), 13, <b>89</b>                                                         |
| $Q$   | quality factor, 36                                                                  | $sln$       | solution, 54                                                                               |
| $Q$   | radiant energy, 34                                                                  | $sol$       | solution (subscript), 60                                                                   |
| $Q$   | reaction quotient, <b>58</b> , 61                                                   | $sub$       | sublimation (subscript), 60                                                                |
| $Q$   | vibrational normal coordinate, 27                                                   |             |                                                                                            |
| $Q$   | Q-branch label, 33                                                                  | $S$         | Poynting vector, <b>17</b> , 148                                                           |
| $r$   | position vector, <b>13</b> , 16, 23, 42, 45, 95, 104, 143, 145, 146                 | $S$         | probability current density, 18                                                            |
| $r$   | interatomic distance, <b>27</b> , 28, 96                                            | $S$         | scattering matrix, 65                                                                      |
| $r$   | internal vibrational coordinate, 27                                                 | $S$         | electron spin angular momentum, 25, 27, <b>30</b> , 32                                     |
| $r$   | radius, <b>13</b> , 64, 70                                                          | $S$         | nuclear spin operator, <b>28</b> , 29                                                      |
| $r$   | rate of concentration change, 63                                                    | $S$         | integrated absorption coefficient, <b>37</b> , 39, 40                                      |
| $r$   | spherical coordinate, <b>13</b> , 18, 20, 21, 96                                    | $S$         | action, 14                                                                                 |
| $r$   | reaction (subscript), 60                                                            | $S$         | area, <b>13</b> , 24, 34, 36, 78, 81, 90, 136                                              |
| rad   | rad (unit of radiation dose), <b>139</b>                                            | $S$         | entropy, 5, 55–58, 60, 61, 63, 65, 66, 78, 90, 138, 139                                    |
| rad   | radian (SI unit), 8, 13, 29, <b>89</b> , 98, <b>136</b>                             | $S$         | overlap integral, <b>19</b> , 21                                                           |
| rem   | rem (unit of dose equivalent), <b>139</b>                                           | $S$         | statistical entropy, 46                                                                    |
| refl  | reflected (subscript), 34                                                           | $S$         | vibrational symmetry coordinate, 27                                                        |
| $R$   | lattice vector, 42                                                                  | $Sc$        | Schmidt number, 82                                                                         |
| $R$   | nuclear orbital angular momentum, 30                                                | $Sh$        | Sherwood number, 82                                                                        |
|       |                                                                                     | $S_n$       | rotation-reflection symmetry operator, 31                                                  |

|                         |                                                                                                                                                      |                       |                                                                                                                         |
|-------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------|-------------------------------------------------------------------------------------------------------------------------|
| <i>Sr</i>               | Strouhal number, 82                                                                                                                                  | <i>U</i>              | expanded uncertainty, 151–154                                                                                           |
| <i>St</i>               | Stanton number (mass transfer), 82                                                                                                                   | <i>U</i>              | internal energy, 55–57, 61, 65                                                                                          |
| <i>S</i>                | siemens (SI unit), <b>89</b> , 132                                                                                                                   | <i>U</i>              | “enzyme unit”, 89                                                                                                       |
| <i>St</i>               | stokes (unit of kinematic viscosity), <u>138</u>                                                                                                     |                       |                                                                                                                         |
| <i>Sv</i>               | sievert (SI unit), <b>89</b> , <u>139</u>                                                                                                            | <i>v</i>              | velocity, <b>13</b> –16, 43, 45, 73, 90, 148                                                                            |
| <i>Sv</i>               | svedberg (unit of time), <u>137</u>                                                                                                                  | <i>v</i>              | lattice direction index, 44                                                                                             |
|                         |                                                                                                                                                      | <i>v</i>              | rate of reaction, <b>63</b> , 64, 68, 69                                                                                |
| <i>t</i>                | Celsius temperature, 38, 46, <b>56</b> , 89, 111, <u>138</u>                                                                                         | <i>v</i>              | specific volume, volume, <b>6</b> , 13, 14, 90                                                                          |
| <i>t</i>                | film thickness, thickness of layer, 77                                                                                                               | <i>v</i>              | speed, <b>13</b> , 81, 82, 93, 95                                                                                       |
| <i>t</i>                | time, 4, 8, <b>13</b> , 15, 16, 24, 29, 34–36, 57, 63, 66–69, 72, 81, 82, 85–87, 92, 95, 98, 137, 143, 144                                           | <i>v</i>              | vibrational quantum number, <b>25</b> , 26, 33, 51                                                                      |
| <i>t</i>                | transport number, 73                                                                                                                                 | vap                   | vaporization (subscript), 60                                                                                            |
| <i>t</i> <sub>1/2</sub> | half life, <b>24</b> , 64, 116                                                                                                                       | vit                   | vitreous substance, 54                                                                                                  |
| <i>t</i>                | tonne (unit of mass), 92, <u>136</u>                                                                                                                 | <i>V</i>              | electric potential, <b>16</b> , 75, 89, 140                                                                             |
| <i>t</i>                | triton, <b>50</b> , <u>115</u> , 116                                                                                                                 | <i>V</i>              | potential energy, <b>14</b> , 18, 23, 27, 29, 66, 96                                                                    |
| <i>tr</i>               | transmitted (subscript), 34                                                                                                                          | <i>V</i>              | volume, 5–8, <b>13</b> –17, 23, 34, 37, 41, 43, 45, 47, 48, 54–57, 63, 65, 66, 81, 90, 92, 111, 133, 136. 139, 146, 147 |
| <i>tr</i>               | triple point (subscript), 60                                                                                                                         |                       | volt (SI unit), 9, <b>89</b> , 94, 134, <u>140</u>                                                                      |
| <i>trs</i>              | transition (subscript), 60                                                                                                                           | <i>V</i>              |                                                                                                                         |
| <b><i>T</i></b>         | hyperfine coupling constant (tensor), 27                                                                                                             | <i>w</i>              | velocity, <b>13</b> , 90, 148                                                                                           |
| <b><i>T</i></b>         | torque, 14                                                                                                                                           | <i>w</i>              | lattice direction index, 44                                                                                             |
| <i>T</i> <sub>C</sub>   | Curie temperature, 43                                                                                                                                | <i>w</i>              | mass fraction, <b>47</b> , 49, 97, 98                                                                                   |
| <i>T</i>                | kinetic energy, <b>14</b> , 18, 93, 94                                                                                                               | <i>w</i>              | radiant energy density, 34                                                                                              |
| <i>T</i> <sub>N</sub>   | Néel temperature, 43                                                                                                                                 | <i>w</i>              | speed, <b>13</b> , 81, 90, 93, 95                                                                                       |
| <i>T</i>                | period, characteristic time interval, 13                                                                                                             | <i>w</i>              | work, 14, <b>56</b> , 70, 89                                                                                            |
| <i>T</i>                | relaxation time, <b>13</b> , 23, 29, 43                                                                                                              |                       |                                                                                                                         |
| <i>T</i>                | thermodynamic temperature, 3, 4, 6, 7, 35, 37, 39, 40, 43, 46, 54, <b>56</b> –59, 61, 62, 64–67, 71, 72, 74, 75, 77, 81, 82, 87, 89, 133, <u>138</u> | <i>W</i>              | degeneracy, statistical weight, 26, <b>45</b>                                                                           |
| <i>T</i>                | total term, electronic term, 25, 131                                                                                                                 | <i>W</i>              | number of open adiabatic channels, <b>66</b> , 67                                                                       |
| <i>T</i>                | transmittance, 36                                                                                                                                    | <i>W</i>              | number of states, <b>45</b> , 66, 67                                                                                    |
| <i>T</i> <sub>1/2</sub> | half life, <b>24</b> , 64, 116                                                                                                                       | <i>W</i>              | radiant energy, <b>34</b> , 36                                                                                          |
| <i>T</i>                | tera (SI prefix), 91                                                                                                                                 | <i>W</i>              | weight, 14                                                                                                              |
| <i>T</i>                | tesla (SI unit), 27, <b>89</b> , <u>141</u>                                                                                                          | <i>W</i> <sub>e</sub> | work, <b>14</b> , <b>56</b> , 70, 89                                                                                    |
| <i>Ti</i>               | terabinary (prefix for binary), 91                                                                                                                   | <i>W</i>              | Weber number, 82                                                                                                        |
| <i>Torr</i>             | torr (unit of pressure), 81, 131, <u>138</u>                                                                                                         | <i>W</i>              | watt (SI unit), 34, 88, <b>89</b> , <u>138</u>                                                                          |
|                         |                                                                                                                                                      | <i>W</i>              | W-boson, <b>111</b> , <u>115</u>                                                                                        |
|                         |                                                                                                                                                      | <i>Wb</i>             | weber (SI unit), <b>89</b> , <u>142</u>                                                                                 |
| <i>u</i>                | displacement vector, 42                                                                                                                              | <i>x</i>              | cartesian coordinate, <b>13</b> , 18, 21                                                                                |
| <i>u</i>                | velocity, <b>13</b> , 45, 90, 148                                                                                                                    | <i>x</i>              | energy parameter, 19                                                                                                    |
| <i>u</i>                | Bloch function, 43                                                                                                                                   | <i>x</i>              | fractional coordinate, 42                                                                                               |
| <i>u</i>                | electric mobility, 73                                                                                                                                | <i>x̄</i>             | mean, 151                                                                                                               |
| <i>u</i>                | estimated standard uncertainty, 151–154                                                                                                              | <i>x</i>              | mole fraction, amount fraction, <b>48</b> , 58, 59, 62, 81, 97, 121, 133                                                |
| <i>u</i>                | lattice direction index, 44                                                                                                                          | <i>x</i>              | vibrational anharmonicity constant, 25                                                                                  |
| <i>u</i>                | speed, <b>13</b> , 81, 90, 93, 95                                                                                                                    |                       |                                                                                                                         |
| <i>u</i>                | ungerade symmetry label, 32, 33                                                                                                                      | <i>X</i>              | reactance, 17                                                                                                           |
| <i>u</i>                | unified atomic mass unit, 9, 22, 47, <b>92</b> , 94, 111, 112, 116, 117, 121, 136                                                                    | <i>X</i>              | x unit, <u>135</u>                                                                                                      |
| <i>ua</i>               | astronomical unit (unit of length), 9, <b>92</b> , <u>94</u> , 136                                                                                   | <i>y</i>              | cartesian coordinate, <b>13</b> , 18, 21                                                                                |
| <i>U</i>                | cell potential, 71                                                                                                                                   | <i>y</i>              | fractional coordinate, 42                                                                                               |
| <i>U</i>                | electric potential difference, <b>16</b> , 17, 44, 74                                                                                                | <i>y</i>              | mole fraction (gas), <b>48</b> , 61                                                                                     |
| <i>U</i>                | electrode potential, 71                                                                                                                              | <i>y</i>              | yocto (SI prefix), 91                                                                                                   |
|                         |                                                                                                                                                      | yd                    | yard (unit of length), <u>135</u>                                                                                       |

|            |                                                            |                 |                                                       |
|------------|------------------------------------------------------------|-----------------|-------------------------------------------------------|
| $Y$        | admittance (complex), 17                                   | $\beta$         | retarded van der Waals constant, 77                   |
| $Y$        | Planck function, 56                                        | $\beta$         | spin wavefunction, <b>19</b> , 20, 29                 |
| $Y$        | spherical harmonic function, <b>18</b> , 21                | $\beta$         | unit cell length, 42                                  |
| $Y$        | yotta (SI prefix), 91                                      | $\beta$         | $\beta$ -particle, <b>50</b> , <u>115</u>             |
| $Yi$       | yottabinary (prefix for binary), 91                        | $\beta$         | phase, 55, 70, 71, 78                                 |
| $z$        | cartesian coordinate, <b>13</b> , 15, 18, 21               | $\gamma$        | hyper-polarizability, 17, <b>24</b> , 95              |
| $z$        | charge number, 20, 43, 49–51, 59, 70–74,                   | $\gamma$        | activity coefficient, <b>59</b> , 62, 70, 75, 76      |
| $z$        | collision frequency, collision                             | $\gamma$        | conductivity, <b>17</b> , 73, 90, 132, 140, 147       |
|            | frequency factor, 64, 65                                   | $\gamma$        | cubic expansion coefficient, 43, <b>56</b> , 81       |
| $z$        | cylindrical coordinate, 13                                 | $\gamma$        | Grüneisen parameter, 43                               |
| $z$        | fractional coordinate, 42                                  | $\gamma$        | gyromagnetic ratio, <b>23</b> , 27–29                 |
| $z$        | partition function, 39, <b>45</b> , 46, 67                 | $\gamma$        | mass concentration, 38, <b>48</b> , 97                |
| $z$        | zepto (SI prefix), 91                                      | $\gamma$        | plane angle, <b>13</b> , 38, 89, 92, 136              |
|            |                                                            | $\gamma$        | ratio of heat capacities, 57                          |
| $Z$        | collision density, collision number, 65                    | $\gamma$        | shear strain, 15                                      |
| $Z$        | compression factor, 57                                     | $\gamma$        | surface tension, <b>14</b> , 56, 77, 78, 81, 82, 90   |
| $Z$        | impedance, <b>17</b> , 70, <u>111</u>                      | $\gamma$        | transmission coefficient, 66                          |
| $Z$        | partition function, 39, <b>45</b> , 46, 67                 | $\gamma$        | unit cell length, 42                                  |
| $Z$        | proton number, atomic number, 20, <b>22</b> –24,           | $\gamma_p$      | proton gyromagnetic ratio, <u>112</u>                 |
|            | 49, 121                                                    | $\Upsilon$      | gamma (unit of mass), <u>136</u>                      |
| $Z$        | Z-boson, <b>111</b> , <u>115</u>                           | $\Upsilon$      | phase, 78                                             |
| $Z$        | zetta (SI prefix), 91                                      | $\Upsilon$      | photon, <b>50</b> , 67, 88, <u>115</u>                |
| $Zi$       | zettabinary (binary prefix), 91                            |                 |                                                       |
| $\alpha$   | electric polarizability, <b>24</b> , 95, 146               | $\Gamma$        | integrated absorption coefficient, <b>37</b> , 39, 40 |
| $\alpha$   | absorptance, <b>36</b> , 37                                | $\Gamma$        | Grüneisen parameter, 43                               |
| $\alpha$   | absorption coefficient, 6, 36–38, 41                       | $\Gamma$        | level width, linewidth, 24, 66, 67                    |
| $\alpha$   | acoustic factor (absorption), 15                           | $\Gamma$        | surface concentration, 48, 63, <b>77</b> , 78         |
| $\alpha$   | angle of optical rotation, <b>37</b> , 38                  | $\Gamma$        | gamma function, 107                                   |
| $\alpha$   | coefficient of heat transfer, 81                           | $\delta$        | acoustic factor (dissipation), 15                     |
| $\alpha$   | coulomb integral, <b>19</b> , 20                           | $\delta$        | centrifugal distortion constant, 25                   |
| $\alpha$   | degree of reaction, 48                                     | $\delta$        | chemical shift, 29                                    |
| $\alpha$   | electrochemical transfer coefficient, 72                   | $\delta$        | decay coefficient, 8, 98                              |
| $\alpha$   | expansion coefficient, 43, <b>56</b> , 81, 82              | $\delta$        | loss angle, 17                                        |
| $\alpha$   | fine-structure constant, <b>22</b> , 95, <u>111</u> , 144, | $\delta$        | thickness, <b>13</b> , 72, 77                         |
|            | 145                                                        | $\delta$        | Dirac delta function, Kronecker delta, 21,            |
| $\alpha$   | Madelung constant, 43                                      |                 | <b>23</b> , 42, <b>106</b>                            |
| $\alpha$   | optical rotatory power, 37                                 | $\delta$        | infinitesimal variation, 8                            |
| $\alpha$   | plane angle, <b>13</b> , 38, 89, 92, 136                   |                 |                                                       |
| $\alpha$   | polarizability, 17, <b>24</b> , 41, 95, 134, 135,          | $\Delta$        | centrifugal distortion constant, 25                   |
|            | 140, 141                                                   | $\Delta$        | inertial defect, 25                                   |
| $\alpha$   | spin wavefunction, <b>19</b> , 20, 29                      | $\Delta$        | mass excess, 22                                       |
| $\alpha$   | transfer coefficient, 72                                   | $\Delta$        | change in extensive quantity, 60                      |
| $\alpha$   | unit cell length, 42                                       | $\Delta$        | finite change, 15, 33, 106                            |
| $\alpha_p$ | relative pressure coefficient, 56                          | $\Delta$        | symmetry label, 31                                    |
| $\alpha$   | $\alpha$ -particle, 8, <b>50</b> , <u>115</u>              |                 |                                                       |
| $\alpha$   | phase, 55, 70, 71, 78                                      | $\varepsilon$   | emissivity, emittance, 35                             |
|            |                                                            | $\varepsilon_F$ | Fermi energy, 43                                      |
| $\beta$    | hyper-polarizability, 17, <b>24</b> , 95                   | $\varepsilon$   | linear strain, 15                                     |
| $\beta$    | degeneracy, statistical weight, 26, <b>45</b>              | $\varepsilon$   | molar decadic absorption coefficient, 6,              |
| $\beta$    | plane angle, <b>13</b> , 38, 89, 92, 136                   |                 | <b>36</b> , 37                                        |
| $\beta$    | pressure coefficient, 56                                   | $\varepsilon$   | orbital energy, 20                                    |
| $\beta$    | reciprocal energy parameter (to replace                    | $\varepsilon$   | permittivity, <b>16</b> , 40, 41, 90                  |
|            | temperature), 46                                           | $\varepsilon_0$ | permittivity of vacuum, electric constant,            |
| $\beta$    | resonance integral, 19                                     |                 | <b>16</b> , 17, 24, 35, 37, 39–41, 43, 70, 95,        |

|               |                                                                      |                 |                                                                                                |
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|               | <u>111</u> , 134, 143–146                                            | $\lambda$       | molar conductivity of an ion, 54, <b>73</b>                                                    |
| $\varepsilon$ | Levi-Civita symbol, 106                                              | $\lambda$       | thermal conductivity, 43, 44, <b>81</b> , 90                                                   |
| $\varepsilon$ | unit step function, Heaviside function, 45, 67, <b>107</b>           | $\lambda$       | van der Waals constant, 77                                                                     |
|               |                                                                      | $\lambda$       | wavelength, 3, 34–38, 42–44, 131                                                               |
|               |                                                                      | $\lambda$       | lambda (unit of volume), <u>136</u>                                                            |
| $\zeta$       | Coriolis $\zeta$ -constant, <b>26</b> , 30                           | $\Lambda$       | angular momentum quantum number (component), 30–32                                             |
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| $\zeta$       | magnetizability, 23                                                  |                 |                                                                                                |
| $\zeta$       | shielding parameter, 21                                              |                 |                                                                                                |
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| $\eta$        | overpotential, 72                                                    | $\mu$           | chemical potential, 5, 6, 8, 43, 44, 46, <b>57</b> , 58, 60–62                                 |
| $\eta$        | viscosity, <b>15</b> , 81, 82, 138                                   | $\mu$           | electric mobility, 73                                                                          |
| $\vartheta$   | cylindrical coordinate, 13                                           | $\tilde{\mu}$   | electrochemical potential, 71                                                                  |
| $\vartheta$   | plane angle, <b>13</b> , 38, 89, 92, 136                             | $\mu$           | dynamic friction factor, 15                                                                    |
| $\vartheta$   | volume strain, bulk strain, 15                                       | $\mu$           | Joule-Thomson coefficient, 57                                                                  |
| $\theta$      | Bragg angle, 42                                                      | $\mu$           | magnetic dipole moment, 17, <b>23</b> , 95, 115, 121, 133, 142                                 |
| $\theta$      | Celsius temperature, 38, 39, 41, 46, <b>56</b> , 61–64, 87, 129, 133 | $\mu$           | mean, 151                                                                                      |
| $\theta$      | characteristic temperature, 46                                       | $\mu$           | mobility, 43                                                                                   |
| $\theta$      | characteristic (Weiss) temperature, 43                               | $\mu$           | permeability, 5, <b>17</b> , 81, 90                                                            |
| $\theta$      | contact angle, 77                                                    | $\mu$           | reduced mass, <b>14</b> , 64–66, 95                                                            |
| $\theta$      | scattering angle, 65                                                 | $\mu$           | Thomson coefficient, 43                                                                        |
| $\theta$      | spherical polar coordinate, <b>13</b> , 18, 21, 96                   | $\mu$           | viscosity, <b>15</b> , 81                                                                      |
| $\theta$      | surface coverage, 77, 80                                             | $\tilde{\mu}$   | electrochemical potential, 71                                                                  |
| $\theta$      | internal vibrational coordinate, 27                                  | $\mu_0$         | permeability of vacuum, magnetic constant, 16, <b>17</b> , 28, <u>111</u> , 133, 143, 144, 147 |
| $\theta_W$    | characteristic (Weiss) temperature, 43                               | $\mu_B$         | Bohr magneton, <b>23</b> , 27, 29, 95, <u>111</u> , 112, 115, 133, 142                         |
| $\theta_W$    | weak mixing angle, Weinberg angle, <b>24</b> , <u>111</u> , 112      | $\mu_e$         | electron magnetic moment, <b>23</b> , <u>111</u>                                               |
| $\Theta$      | quadrupole moment, 23, <b>24</b> , 141                               | $\mu_N$         | nuclear magneton, <b>23</b> , 29, <u>111</u> , 115, 116, 121, 142                              |
| $\Theta$      | characteristic temperature, 46                                       |                 |                                                                                                |
| $\Theta_D$    | Debye temperature, <b>43</b> , 80                                    | $\mu_p$         | proton magnetic moment, <u>112</u> , 116                                                       |
| $\Theta$      | plane angle, 35                                                      | $\mu$           | micro (SI prefix), 91                                                                          |
| $\Theta$      | temperature, 56                                                      | $\mu$           | micron (unit of length), <u>135</u>                                                            |
|               |                                                                      | $\mu$           | muon, 8, <b>50</b> , <u>115</u> , 116                                                          |
| $\kappa$      | asymmetry parameter, 25                                              |                 |                                                                                                |
| $\kappa$      | bulk viscosity, 81                                                   | $\nu$           | stoichiometric number matrix, 53                                                               |
| $\kappa$      | compressibility, 43, <b>56</b>                                       | $\nu$           | charge number of cell reaction, 71                                                             |
| $\kappa$      | conductivity, 6, <b>17</b> , 73, 81, 82, 90, 132, 140, 147           | $\nu_D$         | Debye frequency, 43                                                                            |
| $\kappa$      | magnetic susceptibility, <b>17</b> , 133, 142, 145, 147              | $\nu$           | frequency, <b>13</b> , 23, 25, 28, 29, 33–36, 41, 68, 81, 88, 89, 129                          |
| $\kappa$      | molar napierian absorption coefficient, <b>36</b> , 37               | $\nu$           | kinematic viscosity, <b>15</b> , 90, 138                                                       |
| $\kappa$      | ratio of heat capacities, 57                                         | $\nu$           | stoichiometric number, <b>53</b> , 58, 61, 63, 71–74                                           |
| $\kappa$      | transmission coefficient, 65, <b>66</b>                              | $\tilde{\nu}_D$ | Debye wavenumber, 43                                                                           |
|               |                                                                      | $\tilde{\nu}$   | wavenumber (in vacuum), <b>25</b> , 26, 34–39, 131                                             |
| $\lambda$     | thermal conductivity tensor, 43                                      | $\nu$           | vibrational state symbol, 33                                                                   |
| $\lambda$     | absolute activity, 46, <b>57</b> , 58                                | $\nu_e$         | electron neutrino, <b>50</b> , <u>115</u>                                                      |
| $\lambda$     | angular momentum quantum number (component), 30                      | $\xi$           | Coriolis coupling constant, 26                                                                 |
| $\lambda$     | decay constant, 24                                                   | $\xi$           | extent of reaction, advancement, <b>48</b> , 49, <u>53</u> , 58, 60, 63, 67                    |
| $\lambda$     | mean free path, 44, <b>65</b> , 80, 81                               |                 |                                                                                                |

|            |                                                                                          |              |                                                                  |
|------------|------------------------------------------------------------------------------------------|--------------|------------------------------------------------------------------|
| $\xi$      | magnetizability, <b>23</b> , 95, 142, 147                                                | $\Sigma$     | spin quantum number (component), 30, 32                          |
| $\Xi$      | partition function (grand canonical ensemble), 46                                        | $\Sigma$     | film tension, 77                                                 |
| $\Xi$      | standardized resonance frequency, 29                                                     | $\Sigma$     | summation sign, 105                                              |
| $\pi$      | internal vibrational angular momentum, 30                                                | $\tau$       | acoustic factor (transmission), 15                               |
| $\pi$      | surface pressure, 77                                                                     | $\tau$       | characteristic time interval, relaxation time, 13, 43, <b>64</b> |
| $\pi$      | symmetry label, 32, 33                                                                   | $\tau$       | correlation time, 29                                             |
| $\pi$      | ratio of circumference to diameter, 32, 34–37, 42, 43, 45, 64, 65, 70, 95, 103, 112, 134 | $\tau$       | mean life, lifetime, <b>24</b> , 43, 44, 64, 66, 115, 116        |
| $\pi$      | pion, <u>115</u>                                                                         | $\tau$       | shear stress, <b>15</b> , 81                                     |
| $\Pi$      | osmotic pressure, 59                                                                     | $\tau$       | thickness of layer, 77                                           |
| $\Pi$      | Peltier coefficient, 43                                                                  | $\tau$       | Thomson coefficient, 43                                          |
| $\Pi$      | product sign, 105                                                                        | $\tau$       | transmittance, 36                                                |
| $\Pi$      | symmetry label, 31–33                                                                    | $\varphi$    | fluidity, 15                                                     |
| $\rho$     | density matrix, 46                                                                       | $\varphi$    | plane angle, <b>13</b> , 38, 89, 92, 136                         |
| $\rho$     | resistivity tensor, 43                                                                   | $\varphi$    | volume fraction, 97, 98                                          |
| $\rho$     | acoustic factor (reflection), 15                                                         | $\phi$       | electric potential, <b>16</b> , 75, 89, 140, 146, 147            |
| $\rho$     | charge density, <b>16</b> , 18, 23, 43, 44, 140, 146–148                                 | $\phi$       | fugacity coefficient, 58                                         |
| $\rho$     | cylindrical coordinate, 13                                                               | $\phi$       | inner electric potential, 70, 71                                 |
| $\rho$     | density of states, 43, <b>45</b> , 66, 67                                                | $\phi$       | molecular orbital, 19–21                                         |
| $\rho$     | density operator, 46                                                                     | $\phi$       | osmotic coefficient, 59                                          |
| $\rho$     | radiant energy density, <b>34</b> , 90                                                   | $\phi$       | quantum yield, <b>66</b> , 67                                    |
| $\rho$     | mass density, mass concentration, 6, <b>14</b> , 15, 38, <b>48</b> , 51, 54, 81, 82, 97  | $\phi_{rst}$ | spherical polar coordinate, <b>13</b> , 18, 21, 96               |
| $\rho$     | reflectance, 36                                                                          | $\phi$       | vibrational force constant, 27                                   |
| $\rho$     | resistivity, residual resistivity, 17, 43                                                | $\phi$       | volume fraction, <b>48</b> , 97, 98                              |
| $\rho_A$   | surface density, 14                                                                      | $\phi_W$     | wavefunction, 18, 46                                             |
| $\sigma$   | conductivity tensor, 43                                                                  | $\Phi$       | work function, 80                                                |
| $\sigma$   | shielding tensor, 29                                                                     | $\Phi$       | heat flux, thermal power, <b>57</b> , <b>81</b>                  |
| $\sigma$   | absorption cross section, 37, 38                                                         | $\Phi$       | magnetic flux, <b>17</b> , 89, 142                               |
| $\sigma$   | area per molecule, 77                                                                    | $\Phi$       | potential energy, <b>14</b> , 23                                 |
| $\sigma$   | cross section, 24, 38, 64–66                                                             | $\Phi$       | quantum yield, 66                                                |
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| $\sigma$   | density operator, 46                                                                     | $\Phi$       | work function, <b>43</b> , 80                                    |
| $\sigma$   | normal stress, 15                                                                        | $\chi$       | quadrupole interaction energy tensor, <b>24</b> , 29             |
| $\sigma$   | reflection symmetry operator, 31, 32                                                     | $\chi$       | atomic orbital, <b>19</b> , 21                                   |
| $\sigma$   | shielding constant, 29                                                                   | $\chi$       | electronegativity, 22                                            |
| $\sigma$   | order parameter, 42                                                                      | $\chi$       | magnetic susceptibility, <b>17</b> , 133, 142, 145, 147          |
| $\sigma$   | spin quantum number (component), 30                                                      | $\chi$       | surface electric potential, 70                                   |
| $\sigma$   | standard deviation, 103, <b>151</b>                                                      | $\chi_e$     | electric susceptibility, <b>16</b> , 145, 146                    |
| $\sigma$   | spin quantum number (component), 30                                                      | $\chi_m$     | molar magnetic susceptibility, <b>17</b> , 133, 142              |
| $\sigma$   | Stefan-Boltzmann constant, 35, <u>112</u>                                                | $\psi$       | outer electric potential, 70                                     |
| $\sigma$   | surface charge density, 6, 16, <b>71</b>                                                 | $\psi$       | wavefunction, 18, 20, 30, 43, 96                                 |
| $\sigma$   | surface tension, <b>14</b> , 56, 77, 78, 81, 90                                          | $\Psi$       | electric flux, 16                                                |
| $\sigma$   | symmetry number, 46                                                                      | $\Psi$       | wavefunction, 18, 46                                             |
| $\sigma^2$ | variance, 151                                                                            |              |                                                                  |
| $\sigma$   | wavenumber, 34                                                                           |              |                                                                  |
| $\sigma$   | symmetry label, 32, 33                                                                   |              |                                                                  |

|            |                                                                       |
|------------|-----------------------------------------------------------------------|
| $\omega$   | angular frequency, angular velocity, 8, <b>13</b> ,<br>23, 34, 89, 98 |
| $\omega_D$ | Debye angular frequency, 43                                           |
| $\omega$   | degeneracy, statistical weight, 26, <b>45</b>                         |
| $\omega$   | harmonic (vibrational) wavenumber, 25, 26                             |
| $\omega$   | solid angle, <b>13</b> , 34, 36, 65, 89                               |
| $\Omega$   | angular momentum quantum<br>number (component), 30, 32                |
| $\Omega$   | nutation angular frequency, 28                                        |
| $\Omega$   | partition function, 46                                                |
| $\Omega$   | solid angle, <b>13</b> , 34–36, 65, 89                                |
| $\Omega$   | volume in phase space, 45                                             |
| $\Omega$   | ohm, <b>89</b> , <u>140</u>                                           |

### Special symbols

|                           |                                                                        |
|---------------------------|------------------------------------------------------------------------|
| %                         | percent, 97, 98                                                        |
| ‰                         | permille, 97, 98                                                       |
| °                         | degree (unit of arc), 38, <b>92</b> , <u>136</u>                       |
| ⊖, °                      | standard (superscript), <b>60</b> , 71                                 |
| '                         | minute (unit of arc), <b>92</b> , <u>136</u>                           |
| "                         | second (unit of arc), <b>92</b> , <u>136</u>                           |
| *                         | complex conjugate, 107                                                 |
| *                         | excitation, 50                                                         |
| *                         | pure substance (superscript), 60                                       |
| ‡, ≠                      | activated complex (superscript),<br>transition state (superscript), 60 |
| ∞                         | infinite dilution (superscript), 60                                    |
| <sup>e</sup>              | even parity symmetry label, 32                                         |
| <sup>o</sup>              | odd parity symmetry label, 32                                          |
| [B]                       | amount concentration, concentration of B,<br><b>48</b> , 54, 69        |
| $\Delta_r$                | derivative with respect to extent<br>of reaction, <b>58</b> , 60, 74   |
| $\dim(Q)$                 | dimension of quantity $Q$ , 4                                          |
| $\nabla$                  | nabla, 16–18, 20, 43, 81, 96, <b>107</b> , 146, 147                    |
| $[\alpha]_\lambda^\theta$ | specific optical rotatory power, <b>37</b> , 38                        |
| $[Q]$                     | unit of quantity $Q$ , 4                                               |



## 13 SUBJECT INDEX



When more than one page reference is given, bold print is used to indicate the general (defining) reference. Underlining is used to indicate a numerical entry of the corresponding physical quantity. Greek letters are ordered according to their spelling and accents are ignored in alphabetical ordering. Plural form is listed as singular form, where applicable.

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## NOTES

## NOTES

## PRESSURE CONVERSION FACTORS

|        | Pa                | kPa                  | bar                                | atm                                | Torr                               | psi                                |
|--------|-------------------|----------------------|------------------------------------|------------------------------------|------------------------------------|------------------------------------|
| 1 Pa   | = 1               | = $10^{-3}$          | = $10^{-5}$                        | $\approx 9.869\,23 \times 10^{-6}$ | $\approx 7.500\,62 \times 10^{-3}$ | $\approx 1.450\,38 \times 10^{-4}$ |
| 1 kPa  | = $10^3$          | = 1                  | = $10^{-2}$                        | $\approx 9.869\,23 \times 10^{-3}$ | $\approx 7.500\,62$                | $\approx 0.145\,038$               |
| 1 bar  | = $10^5$          | = $10^2$             | = 1                                | $\approx 0.986\,923$               | $\approx 750.062$                  | $\approx 14.5038$                  |
| 1 atm  | = 101 325         | = 101.325            | = 1.013 25                         | = 1                                | = 760                              | $\approx 14.6959$                  |
| 1 Torr | $\approx 133.322$ | $\approx 0.133\,322$ | $\approx 1.333\,22 \times 10^{-3}$ | $\approx 1.315\,79 \times 10^{-3}$ | = 1                                | $\approx 1.933\,68 \times 10^{-2}$ |
| 1 psi  | $\approx 6894.76$ | $\approx 6.894\,76$  | $\approx 6.894\,76 \times 10^{-2}$ | $\approx 6.804\,60 \times 10^{-2}$ | $\approx 51.714\,94$               | = 1                                |

Examples of the use of this table: 1 bar  $\approx$  0.986 923 atm  
 1 Torr  $\approx$  133.322 Pa

*Note:* 1 mmHg = 1 Torr, to better than  $2 \times 10^{-7}$  Torr, see Section 7.2, p. 138.

NUMERICAL ENERGY CONVERSION FACTORS

$E = h\nu = hc\tilde{\nu} = kT; E_m = N_A E$

|                 | wavenumber $\tilde{\nu}$ |                                    | frequency $\nu$         |  | energy $E$                  |  |                            | molar energy $E_m$          |                            | temperature $T$            |                            |
|-----------------|--------------------------|------------------------------------|-------------------------|--|-----------------------------|--|----------------------------|-----------------------------|----------------------------|----------------------------|----------------------------|
|                 | cm <sup>-1</sup>         |                                    | MHz                     |  | aJ                          |  | eV                         | $E_h$                       | kJ mol <sup>-1</sup>       |                            | K                          |
| $\tilde{\nu}$ : | 1 cm <sup>-1</sup>       | $\cong$ 1                          | 2.997 925 $\times 10^4$ |  | 1.986 446 $\times 10^{-5}$  |  | 1.239 842 $\times 10^{-4}$ | 4.556 335 $\times 10^{-6}$  | 11.962 66 $\times 10^{-3}$ | 2.859 144 $\times 10^{-3}$ | 1.438 775                  |
| $\nu$ :         | 1 MHz                    | $\cong$ 3.335 641 $\times 10^{-5}$ | 1                       |  | 6.626 069 $\times 10^{-10}$ |  | 4.135 667 $\times 10^{-9}$ | 1.519 830 $\times 10^{-10}$ | 3.990 313 $\times 10^{-7}$ | 9.537 076 $\times 10^{-8}$ | 4.799 237 $\times 10^{-5}$ |
| $E$ :           | 1 aJ                     | $\cong$ 50 341.17                  | 1.509 190 $\times 10^9$ |  | 1                           |  | 6.241 510                  | 0.229 371 3                 | 602.2142                   | 143.9326                   | 7.242 963 $\times 10^4$    |
|                 | 1 eV                     | $\cong$ 8065.545                   | 2.417 989 $\times 10^8$ |  | 0.160 217 6                 |  | 1                          | 3.674 933 $\times 10^{-2}$  | 96.485 34                  | 23.060 55                  | 1.160 451 $\times 10^4$    |
|                 | 1 $E_h$                  | $\cong$ 219 474.6                  | 6.579 684 $\times 10^9$ |  | 4.359 744                   |  | 27.211 38                  | 1                           | 2625.500                   | 627.5095                   | 3.157 747 $\times 10^5$    |
| $E_m$ :         | 1 kJ mol <sup>-1</sup>   | $\cong$ 83.593 47                  | 2.506 069 $\times 10^6$ |  | 1.660 539 $\times 10^{-3}$  |  | 1.036 427 $\times 10^{-2}$ | 3.808 799 $\times 10^{-4}$  | 1                          | 0.239 005 7                | 120.2722                   |
|                 | 1 kcal mol <sup>-1</sup> | $\cong$ 349.7551                   | 1.048 539 $\times 10^7$ |  | 6.947 694 $\times 10^{-3}$  |  | 4.336 410 $\times 10^{-2}$ | 1.593 601 $\times 10^{-3}$  | 4.184                      | 1                          | 503.2189                   |
| $T$ :           | 1 K                      | $\cong$ 0.695 035 6                | 2.083 664 $\times 10^4$ |  | 1.380 650 $\times 10^{-5}$  |  | 8.617 343 $\times 10^{-5}$ | 3.166 815 $\times 10^{-6}$  | 8.314 472 $\times 10^{-3}$ | 1.987 207 $\times 10^{-3}$ | 1                          |

The symbol  $\cong$  should be read as meaning ‘approximately corresponding to’ or ‘is approximately equivalent to’. The conversion from kJ to kcal is exact by definition of the thermochemical kcal (see Section 2.11, note 16, p. 58). The values in this table have been obtained from the constants in Chapter 5, p. 111. The last digit is given but may not be significant.

Examples of the use of this table:

$1\text{ aJ} \cong 50\,341.17\text{ cm}^{-1}$

$1\text{ eV} \cong 96.485\,34\text{ kJ mol}^{-1}$

Examples of the derivation of the conversion factors:

$1\text{ aJ to MHz} \quad \frac{(1\text{ aJ})}{h} \cong \frac{10^{-18}\text{ J}}{6.626\,068\,96 \times 10^{-34}\text{ J s}} \cong 1.509\,190 \times 10^{15}\text{ s}^{-1} \cong 1.509\,190 \times 10^9\text{ MHz}$

$1\text{ cm}^{-1}\text{ to eV} \quad (1\text{ cm}^{-1})\,hc\left(\frac{e}{e}\right) \cong \frac{(1.986\,445\,501 \times 10^{-25}\text{ J}) \times 10^2\,e}{1.602\,176\,487 \times 10^{-19}\text{ C}} \cong 1.239\,842 \times 10^{-4}\text{ eV}$

$1\,E_h\text{ to kJ mol}^{-1} \quad (1\,E_h)\,N_A \cong (4.359\,743\,94 \times 10^{-18}\text{ J}) \times (6.022\,141\,79 \times 10^{23}\text{ mol}^{-1}) \cong 2625.500\text{ kJ mol}^{-1}$

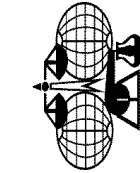
$1\text{ kcal mol}^{-1}\text{ to cm}^{-1} \quad \frac{(1\text{ kcal mol}^{-1})}{hcN_A} \cong \frac{4.184 \times (1\text{ kJ mol}^{-1})}{hcN_A} \cong \frac{4.184 \times (10^3\text{ J mol}^{-1})}{(1.986\,445\,501 \times 10^{-25}\text{ J}) \times 10^2\text{ cm} \times (6.022\,141\,79 \times 10^{23}\text{ mol}^{-1})} \cong 349.7551\text{ cm}^{-1}$



IUPAC Periodic Table of the Elements

18

|                                                |                                          |                                              |                                             |                                              |                                           |                                               |                                           |                                             |                                           |                                               |                                            |                                                 |                                          |                                               |                                           |                                           |                                         |
|------------------------------------------------|------------------------------------------|----------------------------------------------|---------------------------------------------|----------------------------------------------|-------------------------------------------|-----------------------------------------------|-------------------------------------------|---------------------------------------------|-------------------------------------------|-----------------------------------------------|--------------------------------------------|-------------------------------------------------|------------------------------------------|-----------------------------------------------|-------------------------------------------|-------------------------------------------|-----------------------------------------|
| 1                                              | 2                                        | 3                                            | 4                                           | 5                                            | 6                                         | 7                                             | 8                                         | 9                                           | 10                                        | 11                                            | 12                                         | 13                                              | 14                                       | 15                                            | 16                                        | 17                                        | 2                                       |
| 1<br><b>H</b><br>hydrogen<br>1.007 94(7)       | 2<br><b>He</b><br>helium<br>4.002 602(2) | 3<br><b>Li</b><br>lithium<br>6.941(2)        | 4<br><b>Be</b><br>beryllium<br>9.012 182(3) | 5<br><b>B</b><br>boron<br>10.811(7)          | 6<br><b>C</b><br>carbon<br>12.0107(8)     | 7<br><b>N</b><br>nitrogen<br>14.0067(2)       | 8<br><b>O</b><br>oxygen<br>15.9994(3)     | 9<br><b>F</b><br>fluorine<br>18.998 4032(5) | 10<br><b>Ne</b><br>neon<br>20.1797(6)     | 11<br><b>Na</b><br>sodium<br>22.989 769 28(2) | 12<br><b>Mg</b><br>magnesium<br>24.3050(6) | 13<br><b>Al</b><br>aluminium<br>26.981 538 6(8) | 14<br><b>Si</b><br>silicon<br>28.0855(3) | 15<br><b>P</b><br>phosphorus<br>30.973 762(2) | 16<br><b>S</b><br>sulfur<br>32.065(5)     | 17<br><b>Cl</b><br>chlorine<br>35.453(2)  | 18<br><b>Ar</b><br>argon<br>39.948(1)   |
| 19<br><b>K</b><br>potassium<br>39.0983(1)      | 20<br><b>Ca</b><br>calcium<br>40.078(4)  | 21<br><b>Sc</b><br>scandium<br>44.955 912(6) | 22<br><b>Ti</b><br>titanium<br>47.867(1)    | 23<br><b>V</b><br>vanadium<br>50.9415(1)     | 24<br><b>Cr</b><br>chromium<br>51.9961(6) | 25<br><b>Mn</b><br>manganese<br>54.938 045(5) | 26<br><b>Fe</b><br>iron<br>55.845(2)      | 27<br><b>Co</b><br>cobalt<br>58.933 195(5)  | 28<br><b>Ni</b><br>nickel<br>58.6934(2)   | 29<br><b>Cu</b><br>copper<br>63.546(3)        | 30<br><b>Zn</b><br>zinc<br>65.409(4)       | 31<br><b>Ga</b><br>gallium<br>69.723(1)         | 32<br><b>Ge</b><br>germanium<br>72.64(1) | 33<br><b>As</b><br>arsenic<br>74.921 60(2)    | 34<br><b>Se</b><br>selenium<br>78.96(3)   | 35<br><b>Br</b><br>bromine<br>79.904(1)   | 36<br><b>Kr</b><br>krypton<br>83.798(2) |
| 37<br><b>Rb</b><br>rubidium<br>85.4678(3)      | 38<br><b>Sr</b><br>strontium<br>87.62(1) | 39<br><b>Y</b><br>yttrium<br>88.905 85(2)    | 40<br><b>Zr</b><br>zirconium<br>91.224(2)   | 41<br><b>Nb</b><br>niobium<br>92.906 38(2)   | 42<br><b>Mo</b><br>molybdenum<br>95.94(2) | 43<br><b>Tc</b><br>technetium<br>[98]         | 44<br><b>Ru</b><br>ruthenium<br>101.07(2) | 45<br><b>Rh</b><br>rhodium<br>102.905 50(2) | 46<br><b>Pd</b><br>palladium<br>106.42(1) | 47<br><b>Ag</b><br>silver<br>107.8682(2)      | 48<br><b>Cd</b><br>cadmium<br>112.411(8)   | 49<br><b>In</b><br>indium<br>114.818(3)         | 50<br><b>Sn</b><br>tin<br>118.710(7)     | 51<br><b>Sb</b><br>antimony<br>121.760(1)     | 52<br><b>Te</b><br>tellurium<br>127.60(3) | 53<br><b>I</b><br>iodine<br>126.904 47(3) | 54<br><b>Xe</b><br>xenon<br>131.293(6)  |
| 55<br><b>Cs</b><br>caesium<br>132.905 451 9(2) | 56<br><b>Ba</b><br>barium<br>137.327(7)  | 57-71<br><b>lanthanoids</b>                  | 72<br><b>Hf</b><br>hafnium<br>178.49(2)     | 73<br><b>Ta</b><br>tantalum<br>180.947 88(2) | 74<br><b>W</b><br>tungsten<br>183.84(1)   | 75<br><b>Re</b><br>rhenium<br>186.207(1)      | 76<br><b>Os</b><br>osmium<br>190.23(3)    | 77<br><b>Ir</b><br>iridium<br>192.217(3)    | 78<br><b>Pt</b><br>platinum<br>195.084(9) | 79<br><b>Au</b><br>gold<br>196.966 569(4)     | 80<br><b>Hg</b><br>mercury<br>200.59(2)    | 81<br><b>Tl</b><br>thallium<br>204.3833(2)      | 82<br><b>Pb</b><br>lead<br>207.2(1)      | 83<br><b>Bi</b><br>bismuth<br>208.980 40(1)   | 84<br><b>Po</b><br>polonium<br>[209]      | 85<br><b>At</b><br>astatine<br>[210]      | 86<br><b>Rn</b><br>radon<br>[222]       |
| 87<br><b>Fr</b><br>francium<br>[223]           | 88<br><b>Ra</b><br>radium<br>[226]       | 89-103<br><b>actinoids</b>                   | 104<br><b>Rf</b><br>rutherfordium<br>[261]  | 105<br><b>Db</b><br>dubnium<br>[262]         | 106<br><b>Sg</b><br>seaborgium<br>[266]   | 107<br><b>Bh</b><br>bohrium<br>[264]          | 108<br><b>Hs</b><br>hassium<br>[277]      | 109<br><b>Mt</b><br>meitnerium<br>[268]     | 110<br><b>Ds</b><br>darmstadtium<br>[271] | 111<br><b>Rg</b><br>roentgenium<br>[272]      |                                            |                                                 |                                          |                                               |                                           |                                           |                                         |



|                                               |                                             |                                                  |                                            |                                        |                                          |                                           |                                            |                                             |                                             |                                             |                                         |                                             |                                           |                                           |
|-----------------------------------------------|---------------------------------------------|--------------------------------------------------|--------------------------------------------|----------------------------------------|------------------------------------------|-------------------------------------------|--------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|-----------------------------------------|---------------------------------------------|-------------------------------------------|-------------------------------------------|
| 57<br><b>La</b><br>lanthanum<br>138.905 47(7) | 58<br><b>Ce</b><br>cerium<br>140.116(1)     | 59<br><b>Pr</b><br>praseodymium<br>140.907 65(2) | 60<br><b>Nd</b><br>neodymium<br>144.242(3) | 61<br><b>Pm</b><br>promethium<br>[145] | 62<br><b>Sm</b><br>samarium<br>150.36(2) | 63<br><b>Eu</b><br>europium<br>151.964(1) | 64<br><b>Gd</b><br>gadolinium<br>157.25(3) | 65<br><b>Tb</b><br>terbium<br>158.925 35(2) | 66<br><b>Dy</b><br>dysprosium<br>162.500(1) | 67<br><b>Ho</b><br>holmium<br>164.930 32(2) | 68<br><b>Er</b><br>erbium<br>167.259(3) | 69<br><b>Tm</b><br>thulium<br>168.934 21(2) | 70<br><b>Yb</b><br>ytterbium<br>173.04(3) | 71<br><b>Lu</b><br>lutetium<br>174.967(1) |
| 89<br><b>Ac</b><br>actinium<br>[227]          | 90<br><b>Th</b><br>thorium<br>232.038 06(2) | 91<br><b>Pa</b><br>protactinium<br>231.035 88(2) | 92<br><b>U</b><br>uranium<br>238.028 91(3) | 93<br><b>Np</b><br>neptunium<br>[237]  | 94<br><b>Pu</b><br>plutonium<br>[244]    | 95<br><b>Am</b><br>americium<br>[243]     | 96<br><b>Cm</b><br>curium<br>[247]         | 97<br><b>Bk</b><br>berkelium<br>[247]       | 98<br><b>Cf</b><br>californium<br>[251]     | 99<br><b>Es</b><br>einsteinium<br>[252]     | 100<br><b>Fm</b><br>fermium<br>[257]    | 101<br><b>Md</b><br>mendelevium<br>[258]    | 102<br><b>No</b><br>nobelium<br>[259]     | 103<br><b>Lr</b><br>lawrencium<br>[262]   |

Notes

- 'Aluminium' and 'cesium' are commonly used alternative spellings for 'aluminium' and 'caesium'.
- IUPAC 2005 standard atomic weights (mean relative atomic masses) are listed with uncertainties in the last figure in parentheses [M. E. Wieser, *Pure Appl. Chem.* **78**, 2051 (2006)]. These values correspond to current best knowledge of the elements in natural terrestrial sources. For elements that have no stable or long-lived nuclides, the mass number of the nuclide with the longest confirmed half-life is listed between square brackets.
- Elements with atomic numbers 112 and above have been reported but not fully authenticated.