

KAYE & LABY

TABLES OF PHYSICAL AND CHEMICAL CONSTANTS

Sixteenth Edition

物理和化学常数表

第16版

G. W. C. Kaye & T. H. Laby

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Tables of Physical and Chemical Constants

16th edition

ORIGINALLY COMPILED BY

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OF AN EDITORIAL COMMITTEE

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This edition of Kaye and Laby's Tables of Physical and Chemical Constants,
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Preface to the sixteenth edition

With this new edition Kaye and Laby achieves 84 years of service. Over this period the scope of physical and chemical data required for everyday use has increased enormously and this is reflected in the size of recent editions. Successive Editorial Boards have always been at pains to ensure that the intentions of the original authors should be maintained: the primary criterion for the inclusion of material is that it should be of value not only to specialists but more generally to scientists working in a variety of fields. The present Board believes that Kaye and Laby's appeal has become more broadly based over the years and it includes material of value also to engineers and to students.

In this edition all the material has been scrutinized and revised as necessary to take account of new results. Several completely new sections have been added covering, for example, medical ultrasonics, fibre optics, high temperature superconductivity, atomic spectroscopy, infra-red and Raman spectroscopy, mass and UV-visible spectrometry, flash points and explosive limits in air and autoignition temperatures, Rutherford scattering formulae and magnetic and electrostatic bending radii. In addition, new chapters on laboratory safety and quality assurance have been added and the section on statistical methods has been rewritten as a new chapter. These three chapters are not intended to provide a complete treatment of their subjects, but rather an introduction with pointers to more definitive texts. However, the Board felt strongly that Kaye and Laby should not ignore such subjects. In addition, many of the explanatory texts that accompany the tables of data have been enlarged so as to provide easier access to the information for the non-specialist reader.

Dr G. W. C. Kaye, F.R.S., one of the original authors, was Superintendent of the Physics Department of the UK's National Physical Laboratory and after he died in 1941 other physicists at that laboratory contributed to the 9th edition, which was then in preparation. This close association has continued to the present day but with contributors drawn increasingly from other UK national laboratories and universities as well. This diversity in the backgrounds of the contributors to succeeding editions must be a source of strength for Kaye and Laby. The members of the present Board, who are working or have worked at the National Physical Laboratory, the Laboratory of the Government Chemist and the UK Atomic Energy Authority, hope that they have maintained the traditions of Kaye and Laby while bringing in a degree of freshness to this edition.

The provision of standards of measurement and of high-accuracy data is an important factor in the economic well-being of a nation and has long been accepted as a responsibility of government in industrialized countries. We may perhaps repeat the hope set out in the preface to the previous edition, and now even more urgent, that future governments will continue to support long-term programmes in national laboratories for the generation of scientific data which only those laboratories can provide.

J.N.

Extract from preface to first edition

The need for a set of up-to-date English physical and chemical tables of convenient size and moderate price has repeatedly impressed us during our teaching and laboratory experience. We have accordingly attempted in this volume to collect the more reliable and recent determinations of some of the important physical and chemical constants.

To increase the utility of the book, we have inserted, in the case of many of the sections, a brief *résumé* containing references to such books and original papers as may profitably be consulted.

Attention has been paid to the setting and accuracy of the mathematical tables; these are included merely to facilitate calculations arising out of the use of the book, and limitations of space have cut out all but a few of the more essential functions.

We began this book while at the Cavendish Laboratory, Cambridge, and Dr G. A. Carse shared in its inception. To Mr G. F. C. Searle, F.R.S., we feel we owe much for his encouragement and suggestions when the scope of the book was under consideration. . . .

G.W.C.K.

T.H.L.

September, 1911

Publisher's note

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1. Units and Fundamental Constants

1.1 Units

1.1.1 The international system of units (SI)

History

In the second half of the nineteenth century the centimetre, gram and second were in fairly general use as base units for scientific work even in such countries as the UK and the USA where the foot and the pound were employed for commerce and engineering. As a result, the units required by the rapidly emerging science of electricity were based on the centimetre, gram and second, with which they formed a coherent system known as the CGS electromagnetic system. A system of units is said to be coherent when derived units are formed from the base units without the insertion of factors of proportionality other than unity. There was also the CGS electrostatic system, but the only quantities frequently expressed in electrostatic units were electric charge, electric potential, and capacitance.

The young but fast-growing electrical industry soon found that many CGS electromagnetic units were of an extremely inconvenient size for its needs. Accordingly, in 1881, international agreement was reached to fix the practical unit of potential, to be called the volt, at 10^8 CGS units (which is approximately equal to the e.m.f. of a primary cell), and the unit of resistance, the ohm, at 10^9 CGS units (which is approximately the resistance of a column of mercury 1 m long and 1 mm^2 in cross-section). The unit of electric current, the ampere, was made a tenth of the CGS unit. A coherent system of practical electric units was thus secured which, however, was not coherent with the mechanical units based on the centimetre and gram. The practical electric units suited the needs of telegraphy, which was then the main electrical industry, and they also happen to be convenient for heavy electrical engineering and for electronics.

The magnetic units, however, were left at their CGS values, presumably because the CGS unit of magnetic flux density, subsequently called 'gauss', is of the order of the flux density of the Earth's field, and, as it was suitable for geomagnetism, there seemed no point in changing it for a unit 10^4 times larger. Coherence was thereby lost to electromagnetism as it had already been lost to the system embracing the mechanical units and the practical electric units.

Whereas the electric units, by the agreement of 1881, were chosen to be of suitable magnitude for everyday use, and whereas the centimetre and the second have acceptable sizes, the gram is too small for the practical needs of man, which are better served by a unit nearer the size of the pound or the kilogram. Moreover, the CGS unit of force, the dyne, and the unit of energy, the erg, are much too small. On the other hand, the unit of energy provided by the practical electric units, the volt-ampere-second, called the joule—which equals 10^7 ergs—is of a satisfactory size.

These considerations—the advantages of coherence and the fortuitous circumstance that a mechanical system based on the metre and the kilogram has precisely the same unit of energy as is provided by the practical electric units—led G. Giorgi in 1902 to propose a system based on the **metre**, the **kilogram**, the **second**, and one of the practical electric units. He pointed out that if magnetic field strength were expressed as amperes per metre instead of 4π times amperes per metre, which is the definition corresponding to that of the CGS unit, the number π would disappear from most electric and magnetic formulae involving rectilinear geometry, but would appear, as is to be expected, in those involving cylinders or spheres.

The International Electrotechnical Commission eventually chose the **ampere** as the fourth base unit of the MKSA or 'Giorgi' system, and in 1948 the 9th General Conference of Weights and Measures[†] recommended it for science and technology, as well as for commerce and industry. This system admirably

[†] The General Conference of Weights and Measures (CGPM) is the authority set up by the Metre Convention of 1875 to promote and improve the metric system, and to secure international uniformity in metric units and standards of measurement. It consists of delegations from the member nations (of which there were 46, including the UK, in 1982), which meet every few years, the 15th, 16th and 17th Conferences having been held in 1975, 1979, and 1983. The International Bureau of Weights and Measures (BIPM), Sèvres (near Paris) is the central office and laboratory of the organization, and is managed, under the authority of the General Conference, by the International Committee of Weights and Measures (CIPM) consisting of 18 members, each from a different nation. The International Committee meets yearly and is responsible for recommending proposals for approval by the General Conference. Eight specialist advisory committees assist the International Committee in planning co-operative programmes of research, and in the preparation of recommendations on units of measurement, on length (definition of the metre), mass, time (definition of the second), temperature, electricity, photometry and radiometry, and ionizing radiations.

covers mechanics and electromagnetism, but it does not provide for other branches of science such as heat. In 1960, in the hope of securing world-wide uniformity in the units employed in natural science, the 11th CGPM added to the units metre, kilogram, second and ampere, the **kelvin** for thermodynamic temperature, the **candela** for luminous intensity, and the radian and steradian for plane and solid angle. The first two joined the original four in being called 'base' units, and the last two were called 'supplementary' units. Any unit formed from two or more base units is called 'derived'. The radian and steradian are regarded as derived units. The MKSA system thus broadened is called the International System of Units, often abbreviated to SI, and is the most satisfactory system of units we have had so far, in that it caters for the commercial and industrial activities of man as well as for the needs of science. In 1971, the 14th CGPM added the **mole**, the unit of amount of substance used in chemistry, to the list of base units, thus making them seven in all.

Definitions of some SI units

The seven base quantities, each with its unit and unit symbol, are listed below.

SI base quantities and units

Quantity	Name of unit	Unit symbol
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

The SI base units are defined as follows:

The **metre** is the length of the path travelled by light in vacuum during a time interval of $1/299\,792\,458$ of a second.

The **kilogram** is the unit of mass; it is equal to the mass of the international prototype of the kilogram.

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.

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 2×10^{-7} newton per metre of length.

The **kelvin**, unit of thermodynamic temperature, is the fraction $1/273.16$ of the thermodynamic temperature of the triple point of water.

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.

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.

The **candela** is the luminous intensity, in a given direction, of a source that emits monochromatic radiation of frequency 540×10^{12} hertz and that has a radiant intensity in that direction of $(1/683)$ watt per steradian.

The SI supplementary units are defined thus:

The **radian** is the plane angle between two radii of a circle which cut off on the circumference an arc equal in length to the radius.

The **steradian** is the solid angle which, having its vertex in the centre of a sphere, cuts off an area of the surface of the sphere equal to that of a square with sides of length equal to the radius of the sphere.

Derived units. The table below lists some of the more common SI derived quantities, each with its unit and unit symbol. The composite symbols in the last column are to some extent indicative of the definition of the quantity.

Quantity	Unit	Symbol
Supplementary		
Plane angle	radian	rad
Solid angle	steradian	sr
Derived		
Area	square metre	m^2
Volume	cubic metre	m^3
Frequency	hertz	Hz s^{-1}
Density	kilogram per cubic metre	kg m^{-3}
Concentration	mole per cubic metre	mol m^{-3}
Velocity	metre per second	m s^{-1}
Angular velocity	radian per second	rad s^{-1}
Acceleration	metre per second squared	m s^{-2}
Angular acceleration	radian per second squared	rad s^{-2}
Force	newton	N m kg s^{-2}
Pressure, stress	pascal	Pa N m^{-2}
Viscosity (dynamic)	pascal second	Pa s
Viscosity (kinematic)	metre squared per second	$\text{m}^2 \text{s}^{-1}$
Energy, work, quantity of heat	joule	J N m
Power, radiant flux	watt	W J s^{-1}
Quantity of electricity	coulomb	C A s
Potential difference, electromotive force	volt	V W A^{-1}
Electric field strength	volt per metre	V m^{-1}
Electric resistance	ohm	Ω V A^{-1}
Electric conductance	siemens	S Ω^{-1}
Capacitance	farad	F C V^{-1}
Magnetic flux	weber	Wb V s
Magnetic flux density	tesla	T Wb m^{-2}
Inductance	henry	H Ωs
Magnetic field strength	ampere per metre	A m^{-1}
Magnetomotive force	ampere	A
Wave number*	1 per metre	m^{-1}
Activity (of a radionuclide)	becquerel	Bq s^{-1}
Absorbed dose	gray	Gy J kg^{-1}
Dose equivalent	sievert	Sv J kg^{-1}
Luminous flux	lumen	lm cd sr
Luminance	candela per square metre	cd m^{-2}
Illuminance	lux	lx lm m^{-2}
Heat flux density, irradiance	watt per square metre	W m^{-2}
Heat capacity, entropy	joule per kelvin	J K^{-1}
Specific heat capacity, specific entropy	joule per kilogram kelvin	$\text{J kg}^{-1} \text{K}^{-1}$
Thermal conductivity	watt per metre kelvin	$\text{W m}^{-1} \text{K}^{-1}$
Molar energy	joule per mole	J mol^{-1}
Molar entropy, molar heat capacity	joule per mole kelvin	$\text{J mol}^{-1} \text{K}^{-1}$

* Wave numbers in the infra-red are still often expressed in cm^{-1} .