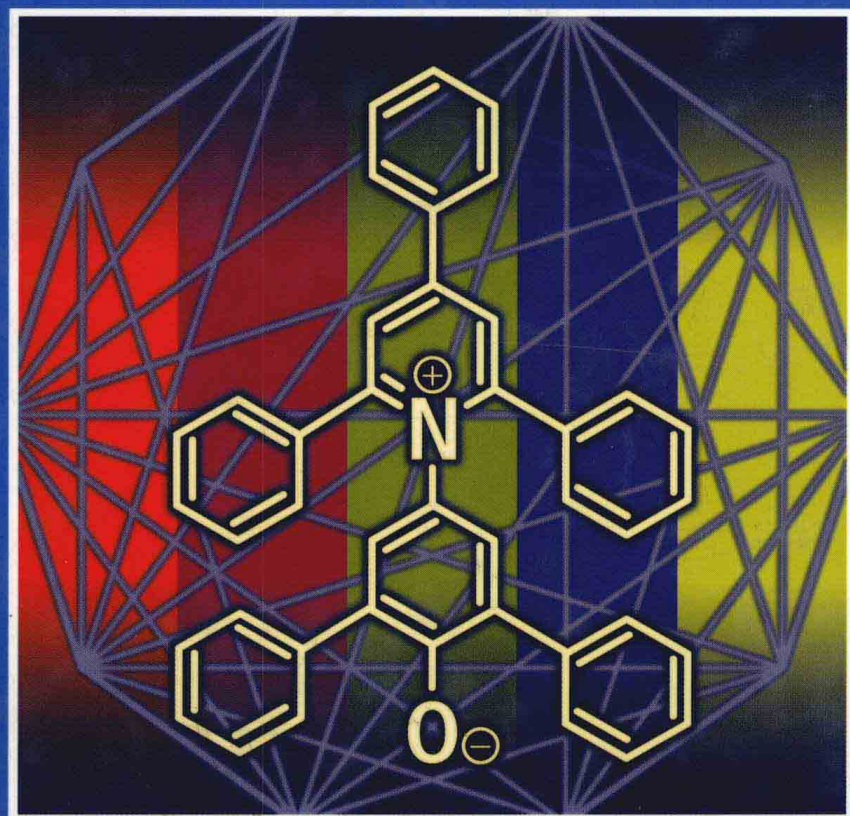


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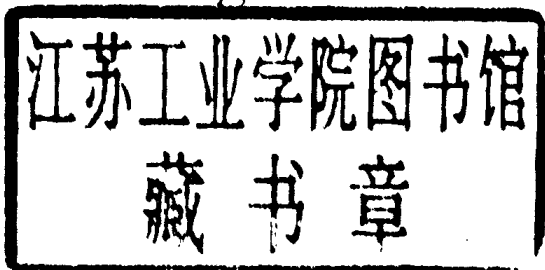
Third, Updated and Enlarged Edition



Christian Reichardt

Solvents and Solvent Effects in Organic Chemistry

Third, Updated and Enlarged Edition



 **WILEY-VCH**

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*To Maria
and in memory of my parents*

Preface to the Third Edition

Meeting the demand for the second edition of this book, which is – despite a reprint in 1990 – no longer available, and considering the progress that has been made during the last decade in the study of solvent effects in experimental and theoretical organic chemistry, this improved third edition is presented to the interested reader.

Following the same layout as in the second edition, all topics retained have been brought up to date, with smaller and larger changes and additions on nearly every page. Two Sections (4.4.7 and 5.5.13) are completely new, dealing with solvent effects on host/guest complexation equilibria and reactions in biphasic solvent systems and neoteric solvents, respectively. More than 900 new references have been added, giving preference to review articles, and many older ones have been deleted. New references either replace older ones or are added to the end of the respective reference list of each chapter. The references cover the literature up to the end of 2001.

From the vast number of published papers dealing with solvent effects in all areas of organic chemistry, only some illustrative examples from the didactic and systematic point of view could be selected. This book is not a monograph covering *all* relevant literature in this field of research. The author, responsible for this subjective selection, apologizes in advance to all chemists whose valuable work on solvent effects is not mentioned in this book. However, using the reviews cited, the reader will find easy access to the full range of papers published in a certain field of research on solvent effects.

Great progress has been made during the last decade in theoretical treatments of solvent effects by various quantum-chemical methods and computational strategies. When indicated, relevant references are given to the respective solution reactions or absorptions. However, a critical evaluation of all the theoretical models and methods used to calculate the differential solvation of educts, activated complexes, products, ground and excited states, is outside the expertise of the present author. Thus, a book on all kinds of theoretical calculations of solvent influences on chemical reactions and physical absorptions has still to be written by someone else.

Consistent use of the nomenclature,^{a)} symbols,^{b)} terms,^{c)} and SI units^{d)} recommended by the IUPAC commissions has also been made in this third edition.

For comments and valuable suggestions I have to thank many colleagues, in particular Prof. E. M. Kosower, Tel Aviv/Israel, Prof. R. G. Makitra, Lviv/Ukraine, Prof. N. O. Mchedlov-Petrossyan, Kharkiv/Ukraine, and Prof. K. Möckel, Mühlhausen/Germany. For their assistance in drawing formulae, preparing the indices, and providing me with difficult to obtain literature, I thank Mr. G. Schäfer (technician), Mrs. S. Schellenberg (secretary), and Mrs. B. Becht-Schröder (librarian), all at the Department

^{a)} G. J. Leigh, H. A. Favre, and W. V. Metanomski: *Principles of Chemical Nomenclature – A Guide to IUPAC Recommendations*, Blackwell Science Publications, London, 1998.

^{b)} I. Mills, T. Cvitas, K. Homann, N. Kallay, and K. Kuchitsu: *Quantities, Units and Symbols in Physical Chemistry*, 2nd ed., Blackwell Science Publications, London, 1993.

^{c)} P. Müller: *Glossary of Terms used in Physical Organic Chemistry – IUPAC Recommendations 1994*, Pure Appl. Chem. 66, 1077 (1994).

^{d)} G. H. Aylward and T. J. V. Tristan: *SI Chemical Data*, 4th ed., Wiley, Chichester, 1999; *Datensammlung Chemie in SI-Einheiten*, 3rd ed., Wiley-VCH, Weinheim/Germany, 1999.

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of Chemistry, Philipps University, Marburg/Germany. Special thanks are due to the staff of Wiley-VCH Verlag GmbH, Weinheim/Germany, particularly to Dr. Elke Westermann, for their fine work in turning the manuscript into the final book. Lastly, my biggest debt is to my wife Maria, not only for her assistance in the preparation of the manuscript, but also for her constant encouragement and support during the writing of this book.

Marburg (Lahn), Spring 2002

Christian Reichardt

Preface to the Second Edition

The response to the first English edition of this book, published in 1979, has been both gratifying and encouraging. Its mixed character, lying between that of a monograph and a textbook, has obviously made it attractive to both the industrial and academic chemist as well as the advanced student of chemistry.

During the last eight years the study of solvent effects on both chemical reactions and absorption spectra has made much progress, and numerous interesting and fascinating examples have been described in the literature. In particular, the study of ionic reactions in the gas phase – now possible due to new experimental techniques – has allowed direct comparisons between gas-phase and solution reactions. This has led to a greater understanding of solution reactions. Consequently, Chapters 4 and 5 have been enlarged to include a description of ionic gas-phase reactions compared to their solution counterparts.

The number of well-studied solvent-dependent processes, *i.e.* reactions and absorptions in solution, has increased greatly since 1979. Only a representative selection of the more instructive, recently studied examples could be included in this second edition.

The search for empirical parameters of solvent polarity and their applications in multiparameter equations has recently been intensified, thus making it necessary to rewrite large parts of Chapter 7.

Special attention has been given to the chemical and physical properties of organic solvents commonly used in daily laboratory work. Therefore, all Appendix Tables have been improved; some have been completely replaced by new ones. A new well-referenced table on solvent-drying has been added (Table A-3). Chapter 3 has been enlarged, in particular by the inclusion of solvent classifications using multivariate statistical methods (Section 3.5). All these amendments justify the change in the title of the book to *Solvents and Solvent Effects in Organic Chemistry*.

The references have been up-dated to cover literature appearing up to the first part of 1987. New references were added to the end of the respective reference list of each chapter from the first edition.

Consistent use of the nomenclature, symbols, terms, and SI units recommended by the IUPAC commissions has also been made in the second edition.*)

I am very indebted to many colleagues for corrections, comments, and valuable suggestions. Especially helpful suggestions came from Professors H.-D. Försterling, Marburg, J. Shorter, Hull/England, and R. I. Zalewski, Poznań/Poland, to whom I am very grateful. For critical reading of the whole manuscript and the improvement of my English I again thank Dr. Edeline Wentrup-Byrne, now living in Brisbane/Australia. Dr. P.-V. Rinze, Marburg, and his son Lars helped me with the author index. Finally, I would like to thank my wife Maria for her sympathetic assistance during the preparation of this edition and for her help with the indices.

Marburg (Lahn), Spring 1988

Christian Reichardt

* Cf. Pure Appl. Chem. 51, 1 (1979); *ibid.* 53, 753 (1981); *ibid.* 55, 1281 (1983); *ibid.* 57, 105 (1985).

Preface to the First Edition

The organic chemist usually works with compounds which possess labile covalent bonds and are relatively involatile, thereby often rendering the gas-phase unsuitable as a reaction medium. Of the thousands of reactions known to occur in solution only few have been studied in the gas-phase, even though a description of reaction mechanisms is much simpler for the gas-phase. The frequent necessity of carrying out reactions in the presence of a more or less inert solvent results in two main obstacles: The reaction depends on a larger number of parameters than in the gas-phase. Consequently, the experimental results can often be only qualitatively interpreted because the state of aggregation in the liquid phase has so far been insufficiently studied. On the other hand, the fact that the interaction forces in solution are much stronger and more varied than in the gas-phase, permits to affect the properties and reactivities of the solute in manifold modes.

Thus, whenever a chemist wishes to carry out a chemical reaction he not only has to take into consideration the right reaction partners, the proper reaction vessels, and the appropriate reaction temperature. One of the most important features for the success of the planned reaction is the selection of a suitable solvent. Since solvent effects on chemical reactivity have been known for more than a century, most chemists are now familiar with the fact that solvents may have a strong influence on reaction rates and equilibria. Today, there are about three hundred common solvents available, nothing to say of the infinite number of solvent mixtures. Hence the chemist needs, in addition to his intuition, some general rules and guiding-principles for this often difficult choice.

The present book is based on an earlier paperback "Lösungsmittleffekte in der organischen Chemie" [1], which, though following the same layout, has been completely rewritten, greatly expanded, and brought up to date. The book is directed both toward the industrial and academic chemist and particularly the advanced student of chemistry, who on the one hand needs objective criteria for the proper choice of solvent but on the other hand wishes to draw conclusions about reaction mechanisms from the observed solvent effects.

A knowledge of the physico-chemical principles of solvent effects is required for proper bench-work. Therefore, a description of the intermolecular interactions between dissolved molecules and solvent is presented first, followed by a classification of solvents derived therefrom. Then follows a detailed description of the influence of solvents on chemical equilibria, reaction rates, and spectral properties of solutes. Finally, empirical parameters of solvent polarity are given, and in an appendix guidelines to the everyday choice of solvents are given in a series of Tables and Figures.

The number of solvent systems and their associated solvent effects examined is so enormous that a complete description of all aspects would fill several volumes. For example, in Chemical Abstracts, volume 85 (1976), approximately eleven articles per week were quoted in which the words "Solvent effects on ..." appeared in the title. In the present book only a few important and relatively well-defined areas of general importance have been selected. The book has been written from the point of view of practical use for the organic chemist rather than from a completely theoretical one.

In the selection of the literature more recent reviews were taken into account mainly. Original papers were cited in particular from the didactic point of view rather

than priority, importance or completeness. This book, therefore, does not only have the character of a monograph but also to some extent that of a textbook. In order to help the reader in his use of the literature cited, complete titles of the review articles quoted are given. The literature up until December 1977 has been considered together with a few papers from 1978. The use of symbols follows the recommendations of the Symbols Committee of the Royal Society, London, 1971 [2].

I am very grateful to Professor Karl Dimroth, Marburg, who first stimulated my interest in solvent effects in organic chemistry. I am indebted to Professors W. H. Pirkle, Urbana/Illinois, D. Seebach, Zürich/Switzerland, J. Shorter, Hull/England, and numerous other colleagues for helpful advice and information. Thanks are also due to the authors and publishers of copyrighted materials reproduced with their permission (cf. Figure and Table credits on page 495). For the careful translation and improvement of the English manuscript I thank Dr. Edeline Wentrup-Byrne, Marburg. Without the assistance and patience of my wife Maria, this book would not have been written.

Marburg (Lahn), Summer 1978

Christian Reichardt

References

[1] C. Reichardt: *Lösungsmittelleffekte in der organischen Chemie*. 2nd edition. Verlag Chemie, Weinheim 1973;

Effets de solvant en chimie organique (translation of the first-mentioned title into French, by I. Tkatchenko), Flammarion, Paris 1971;

Rastvoriteli v organicheskoi khimii (translation of the first-mentioned title into Russian, by E. R. Zakhisa), Izdatel'stvo Khimiya, Leningrad 1973.

[2] *Quantities, Units, and Symbols*, issued by The Symbols Committee of the Royal Society, London, in 1971.

List of Abbreviations

Abbreviations and Recommended Values of Some Fundamental Constants and Numbers^{a, b)}

N_A	Avogadro constant	$6.0221 \cdot 10^{23} \text{ mol}^{-1}$
c_0	speed of light in vacuum	$2.9979 \cdot 10^8 \text{ m} \cdot \text{s}^{-1}$
ϵ_0	absolute permittivity of vacuum [$= 1/(\mu_0 \cdot c_0^2)$; μ_0 = permeability of vacuum]	$8.8542 \cdot 10^{-12} \text{ C}^2 \cdot \text{J}^{-1} \cdot \text{m}^{-1}$
e	elementary charge	$1.6022 \cdot 10^{-19} \text{ C}$
h	Planck constant	$6.6261 \cdot 10^{-34} \text{ J} \cdot \text{s}$
R	gas constant	$8.3145 \text{ J} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$ (or 0.08206 $\text{L} \cdot \text{atm} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$)
k_B	Boltzmann constant ($= R/N_A$)	$1.3807 \cdot 10^{-23} \text{ J} \cdot \text{K}^{-1}$
V_m	standard molar volume of an ideal gas (at $t = 0^\circ\text{C}$ and $p = 100 \text{ kPa}$)	$22.711 \text{ L} \cdot \text{mol}^{-1}$
T_0	zero of the Celsius scale	273.15 K
π	ratio of the circumference to the diameter of a circle	3.1416
e	exponential number and base of natural logarithms (ln)	2.7183
$\ln 10$	natural logarithm of ten ($\ln x = \ln 10 \cdot \lg x$; \lg = decadic logarithm)	2.303

Abbreviations and Symbols for Units^{a, b)}

bar	bar ($= 10^5 \text{ Pa} = 10^5 \text{ N} \cdot \text{m}^{-2}$)	pressure
cg/g	centigram/gram	weight percent
cL/L, cl/l	centilitre/litre	volume percent
cmol/mol	centimol/mol	mole percent
cm	centimetre (10^{-2} m)	length
cm^3	cubic centimetre (millilitre mL; 10^{-6} m^3)	volume
C	coulomb	electric charge

^{a)} I. Mills, T. Cvitaš, K. Homann, N. Kallay, and K. Kuchitsu: *Quantities, Units and Symbols in Physical Chemistry*. 2nd ed., Blackwell Scientific Publications, London, 1993.

^{b)} G. H. Aylward and T. J. V. Tristan: *SI Chemical Data*. 4th ed., Wiley, Chichester, 1999; *Datensammlung Chemie in SI-Einheiten*. 3rd ed., Wiley-VCH, Weinheim/Germany, 1999.

XVIII List of Abbreviations

°C	degrees centigrade (Celsius)	temperature
dm ³	cubic decimetre (litre L; 10 ⁻³ m ³)	volume
J	joule	energy
kJ	kilojoule (10 ³ J)	energy
K	kelvin	temperature
L, l	litre (1 dm ³ ; 10 ⁻³ m ³)	volume
m	metre	length
min	minute	time
mol	mole	amount of substance
MPa	megapascal (10 ⁶ Pa)	pressure
mT	millitesla (10 ⁻³ T)	magnetic flux density (magnetic field)
nm	nanometre (10 ⁻⁹ m)	length
Pa	pascal (1 N · m ⁻² = 10 ⁻⁵ bar)	pressure
percent (%)	part per hundred (10 ⁻²)	dimensionless fraction
ppm	part per million (10 ⁻⁶)	dimensionless fraction
s	second	time

Abbreviations and Symbols for Properties^{e)}

a_i	activity of solute i	
$a(^1\text{H})$	ESR hyperfine coupling constant (coupling with ^1H)	mT (= 10 ⁻³ T)
A_j	the solvent's anion-solvating tendency or 'acity' (Swain)	
AN	solvent acceptor number, based on ^{31}P NMR chemical shift of Et ₃ PO (Gutmann and Meyer)	
α	electric polarizability of a molecule, polarizability volume	C ² · m ² · J ⁻¹ or 4 $\pi\epsilon_0$ · cm ³
α	empirical parameter of solvent hydrogen-bond donor acidity (Taft and Kamlet)	
B	empirical parameter of solvent Lewis basicity (Palm and Koppel)	
B_{MeOD}	IR based empirical parameter of solvent Lewis basicity (Palm and Koppel)	

^{e)} P. Müller: *Glossary of Terms used in Physical Organic Chemistry – IUPAC Recommendations 1994*. Pure Appl. Chem. 66, 1077 (1994).

B_{PhOH}	IR based empirical parameter of solvent Lewis basicity (Koppel and Paju; Makitra)	
B_j	the solvent's cation-solvating tendency or 'basity' (Swain)	
β	empirical parameter of solvent hydrogen-bond acceptor basicity (Taft and Kamlet)	
c	cohesive pressure (cohesive energy density) of a solvent	MPa (= 10^6 Pa)
$c_i, c(i)$	molar concentration of solute i	$\text{mol} \cdot \text{L}^{-1}$
C_A, C_B	Lewis acidity and Lewis basicity parameter (Drago)	
cmc	critical micelle concentration	$\text{mol} \cdot \text{L}^{-1}$
D_{HA}	molar bond-dissociation energy of the bond between H and A	$\text{kJ} \cdot \text{mol}^{-1}$
D_π	empirical parameter of solvent Lewis basicity, based on a 1,3-dipolar cycloaddition reaction (Nagai <i>et al.</i>)	
DN	solvent donor number (Gutmann) [$= -\Delta H(\text{D}-\text{SbCl}_5)$]	$\text{kcal} \cdot \text{mol}^{-1}$
DN^{N}	normalized solvent donor number (Marcus)	
$\delta, \delta_{\text{H}}$	Hildebrand's solubility parameter	$\text{MPa}^{1/2}$
δ	chemical shift of NMR signals	ppm
δ	solvent polarizability correction term (Taft and Kamlet)	
E	energy, molar energy	$\text{kJ} \cdot \text{mol}^{-1}$
E	electric field strength	$\text{V} \cdot \text{m}^{-1}$
E	enol constant (K. H. Meyer)	
E	empirical parameter of solvent Lewis acidity (Palm and Koppel)	
E_A, E_a	Arrhenius activation energy	$\text{kJ} \cdot \text{mol}^{-1}$
E_A, E_B	Lewis acidity and Lewis basicity parameter (Drago)	
EA	electron affinity	$\text{kJ} \cdot \text{mol}^{-1}$
E_{B}^{N}	empirical solvent Lewis basicity parameter, based on the $n \rightarrow \pi^*$ absorption of an aminyloxide radical (Mukerjee; Wrona)	
E_{K}	empirical solvent polarity parameter, based on the $d \rightarrow \pi^*$ absorption of a molybdenum complex (Walther)	$\text{kcal} \cdot \text{mol}^{-1}$

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E_{MLCT}^*	empirical solvent polarity parameter, based on the $d \rightarrow \pi^*$ absorption of a tungsten complex (Lees)	
E_{T}	molar electronic transition energy, molar electronic excitation energy	$\text{kJ} \cdot \text{mol}^{-1}$ or $\text{kcal} \cdot \text{mol}^{-1}$
$E_{\text{T}}(30)$	empirical solvent polarity parameter, based on the intramolecular CT absorption of a pyridinium- <i>N</i> -phenolate betaine dye (Dimroth and Reichardt)	$\text{kcal} \cdot \text{mol}^{-1}$
E_{T}^{N}	normalized $E_{\text{T}}(30)$ solvent polarity parameter (Reichardt)	
E_{T}^{SO}	empirical solvent polarity parameter, based on the $n \rightarrow \pi^*$ absorption of an <i>S</i> -oxide (Walter)	$\text{kcal} \cdot \text{mol}^{-1}$
EPA	electron-pair acceptor	
EPD	electron-pair donor	
ϵ_{r}	relative permittivity ($=\epsilon/\epsilon_0$) (“dielectric constant”)	
Φ	empirical solvent polarity parameter, based on the $n \rightarrow \pi^*$ absorption of ketones (Dubois)	
G	IR based empirical solvent polarity parameter (Schleyer and Allerhand)	
ΔG°	standard molar Gibbs energy change	$\text{kJ} \cdot \text{mol}^{-1}$
ΔG^\ddagger	standard molar Gibbs energy of activation	$\text{kJ} \cdot \text{mol}^{-1}$
$\Delta G_{\text{solv}}^\circ$	standard molar Gibbs energy of solvation	$\text{kJ} \cdot \text{mol}^{-1}$
$\Delta G_{\text{hydr}}^\circ$	standard molar Gibbs energy of hydration	$\text{kJ} \cdot \text{mol}^{-1}$
$\Delta G_{\text{t}}^\circ(\text{X}, \text{O} \rightarrow \text{S}),$ $\Delta G_{\text{t}}^\circ(\text{X}, \text{W} \rightarrow \text{S})$	standard molar Gibbs energy of transfer of solute X from a reference solvent (O) or water (W) to another solvent (S)	$\text{kJ} \cdot \text{mol}^{-1}$
γ_i	activity coefficient of solute i	
ΔH°	standard molar enthalpy change	$\text{kJ} \cdot \text{mol}^{-1}$
ΔH^\ddagger	standard molar enthalpy of activation	$\text{kJ} \cdot \text{mol}^{-1}$
ΔH_{v}	molar enthalpy (heat) of vapourization	$\text{kJ} \cdot \text{mol}^{-1}$
H_0	acidity function (Hammett)	
HBA	hydrogen-bond acceptor	

HBD	hydrogen-bond donor	
HOMO	highest occupied molecular orbital	
E_i, I, IP	ionization energy	$\text{kJ} \cdot \text{mol}^{-1}$
I	gas-chromatographic retention index (Kováts)	
J	NMR spin-spin coupling constant	Hz
k	rate constant for monomolecular ($n = 1$) and bimolecular ($n = 2$) reactions	$(\text{L} \cdot \text{mol}^{-1})^{n-1} \cdot \text{s}^{-1}$
k_0	rate constant in a reference solvent or in the gas phase for monomolecular ($n = 1$) and bimolecular reactions ($n = 2$)	$(\text{L} \cdot \text{mol}^{-1})^{n-1} \cdot \text{s}^{-1}$
k_0	in Hammett equations the rate constant of unsubstituted substrates	$(\text{L} \cdot \text{mol}^{-1})^{n-1} \cdot \text{s}^{-1}$ with $n = 1$ or 2
K, K_c	equilibrium constant (concentration basis; $v =$ stoichiometric number)	$(\text{mol} \cdot \text{L}^{-1})^{\Sigma v}$
K_a, K_b	acid and base ionization constants	$(\text{mol} \cdot \text{L}^{-1})^{\Sigma v}$
K_{auto}	autoionization ion product, autoprotolysis constant	$\text{mol}^2 \cdot \text{L}^{-2}$
$K_{\text{Assoc}}, K_{\text{Dissoc}}, K_{\text{ion}}, K_{\text{T}}$	equilibrium constants of association, dissociation, ionization, resp. tautomerization reactions	$(\text{mol} \cdot \text{L}^{-1})^{\Sigma v}$
$K_{\text{O/w}}$	1-octanol/water partition constant (Hansch and Leo)	
KB	kauri-butanol number	
L	desmotropic constant (K. H. Meyer)	
LUMO	lowest unoccupied molecular orbital	
λ	wavelength	$\text{nm} (= 10^{-9} \text{ m})$
m	mass of a particle	g
M_r	relative molecular mass of a substance ("molecular weight")	
M	miscibility number (Godfrey)	
MH	microscopic hydrophobicity parameter of substituents (Menger)	
μ	empirical solvent softness parameter (Marcus)	
μ	permanent electric dipole moment of a molecule	$\text{C} \cdot \text{m}$ (or D)
μ_{ind}	induced electric dipole moment of a molecule	$\text{C} \cdot \text{m}$ (or D)

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μ_i°	standard chemical potential of solute i	$\text{kJ} \cdot \text{mol}^{-1}$
μ_i^∞	standard chemical potential of solute i at infinite dilution	$\text{kJ} \cdot \text{mol}^{-1}$
n, n_D	refractive index (at sodium D line) ($= c_0/c$)	
N	empirical parameter of solvent nucleophilicity (Winstein and Grunwald)	
N_+	nucleophilicity parameter for (nucleophile + solvent)-systems (Ritchie)	
ν	frequency	Hz, s^{-1}
ν°	frequency in the gas phase or in an inert reference solvent	Hz, s^{-1}
$\tilde{\nu}$	wavenumber ($= 1/\lambda$)	cm^{-1}
Ω	empirical solvent polarity parameter, based on a Diels-Alder reaction (Berson)	
p	pressure	$\text{Pa} (= 1\text{N} \cdot \text{m}^{-2}),$ $\text{bar} (= 10^5 \text{Pa})$
P	measure of solvent polarizability (Palm and Koppel)	
P	empirical solvent polarity parameter, based on ^{19}F NMR measurements (Taft)	
PA	proton affinity	$\text{kJ} \cdot \text{mol}^{-1}$
P_y	empirical solvent polarity parameter, based on the $\pi^* \rightarrow \pi$ emission of pyrene (Winnik)	
$P_{o/w}$	1-octanol/water partition coefficient (Hansch and Leo)	
pH	$-\lg[\text{H}_3\text{O}^+], -\lg c(\text{H}_3\text{O}^+)$ (abbreviation of <i>potentia hydrogenii</i> or <i>puissance d'hydrogène</i> (Sørensen 1909)	
pK	$-\lg K$	
π	internal pressure of a solvent	$\text{MPa} (= 10^6 \text{Pa})$
π^*	empirical solvent dipolarity/ polarizability parameter, based on the $\pi \rightarrow \pi^*$ absorption of substituted aromatics (Taft and Kamlet)	

π_{azo}^*	empirical solvent dipolarity/ polarizability parameter, based on the $\pi \rightarrow \pi^*$ absorption of azo merocyanine dyes (Buncel)	
π_x	hydrophobicity parameter of substituent X in $\text{H}_5\text{C}_6\text{-X}$ (Hansch)	
r	radius of sphere representing an ion or a cavity	cm ($= 10^{-2}$ m)
r	distance between centres of two ions or molecules	cm ($= 10^{-2}$ m)
ρ	density (mass divided by volume)	$\text{g} \cdot \text{cm}^{-3}$
ρ, ρ_A	Hammett reaction resp. absorption constants	
S	generalized for solvent	
S	empirical solvent polarity parameter, based on the Z-values (Brownstein)	
\mathcal{P}	$\lg k_2$ for the Menshutkin reaction of tri- <i>n</i> -propylamine with iodomethane (Drougard and Decroocq)	
ΔS°	standard molar entropy change	$\text{J} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$
ΔS^\ddagger	standard molar entropy of activation	$\text{J} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$
S_p	solvophobic power of a solvent (Abraham)	
SA	empirical parameter of solvent hydrogen-bond donor acidity (Catalán)	
SB	empirical parameter of solvent hydrogen-bond acceptor basicity (Catalán)	
SPP	empirical parameter of solvent dipolarity/polarizability, based on the $\pi \rightarrow \pi^*$ absorption of substituted 7- nitrofluorenes (Catalán)	
σ	Hammett substituent constant	
σ	NMR screening constant	
t	Celsius temperature	$^\circ\text{C}$
T	thermodynamic temperature	K
t_{mp}	melting point	$^\circ\text{C}$
t_{bp}	boiling point	$^\circ\text{C}$
U	internal energy	kJ
ΔU_v	molar energy of vapourization	$\text{kJ} \cdot \text{mol}^{-1}$

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$V_m, V_{m,i}$	molar volume (of i)	$\text{cm}^3 \cdot \text{mol}^{-1}$
ΔV^\ddagger	molar volume of activation	$\text{cm}^3 \cdot \text{mol}^{-1}$
$x_i, x(i)$	mole fraction of i ($x_i = n_i / \sum n$)	
X	empirical solvent polarity parameter, based on an S_E2 reaction (Gielen and Nasielski)	
χ_R, χ_B	empirical solvent polarity parameters, based on the $\pi \rightarrow \pi^*$ absorption of merocyanine dyes (Brooker)	$\text{kcal} \cdot \text{mol}^{-1}$
${}^O y_X^S, {}^W y_X^S$	solvent-transfer activity coefficient of a solute X from a reference solvent (O) or water (W) to another solvent (S)	
Y	empirical parameter of solvent ionizing power, based on <i>t</i> -butyl chloride solvolysis (Winstein and Grunwald)	
Y_{OTs}	empirical parameter of solvent ionizing power, based on 2-adamantyl tosylate solvolysis (Schleyer and Bentley)	
Y	measure of solvent polarization (Palm and Koppel)	
z_i	charge number of an ion i	positive for cations, negative for anions
Z	empirical solvent polarity parameter, based on the intermolecular CT absorption of a substituted pyridinium iodide (Kosower)	$\text{kcal} \cdot \text{mol}^{-1}$