### INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY

ANALYTICAL CHEMISTRY DIVISION COMMISSION ON SOLUBILITY DATA

## SOLUBILITY DATA SERIES

## Volume 60

## HALOGENATED METHANES WITH WATER

## SOLUBILITY DATA SERIES

Editor-in-Chief J.W. Lorimer The University of Western Ontario London, Ontario, Canada

F.W. Getzen Sub-editor Liquid/Liquid Systems C.L. Young Sub-editor Indexes

## EDITORIAL BOARD

P.G.T. Fogg (UK)
H. Gamsjäger (Austria)
H.J.M. Grünbauer (The Netherlands)
G.T. Hefter (Australia)
S. Krause (USA)
A. Maczynski (Poland)
K.N. Marsh (Denmark)
V.V. Myasoedova (Russia)

M. Salomon (USA) M.-T. Saugier-Cohen Adad (France) V.P. Sazonov (Russia) D.G. Shaw (USA) A. Skrzecz (Poland) R.P.T. Tomkins (USA) R.G. Wissinger (USA) Yu. Yampol'skii (Russia)

Managing Editor

P.D. Gujral

IUPAC Secretariat, Oxford, UK

INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY IUPAC SECRETARIAT: Bank Court Chambers, 2-3 Pound Way Templars Square, Cowley, Oxford OX4 3YF, UK

## SOLUBILITY DATA SERIES

## Volume 60

## HALOGENATED METHANES WITH WATER

Volume Editors

Ari L. Horváth Imperial Chemical Industries PLC North Carolina State University Runcorn, UK

Forrest W. Getzen Raleigh, NC, USA

Contributors

Z. Maczynska Institute of Physical Chemistry Warsaw, Poland

**OXFORD UNIVERSITY PRESS** 

## **IUPAC Solubility Data Series Rates for 1995**

Subscriptions:	UK and Europe	<b>Rest of World</b>
Full subscription	£265	\$395
Single volume	£80	\$120
Reduced rates are available to	members of IUPAC. Please appl	y directly to the publisher for

Prices include postage by surface mail or, for subscribers in the USA and Canada by Airfreight or in Japan, India, Australia and New Zealand by Air Speeded Post. Airmail rates are available on request.

The IUPAC Solubility Data Series (ISSN 0191-5622) is published quarterly in March, June, September and December by Oxford University Press, Oxford, UK. Annual subscription price is US\$395. The IUPAC Solubility Data Series is distributed by Virgin Mailing & Distribution, 10 Camptown Road, Irvington, New Jersey 07111-1105, USA. Second Class postage paid at Newark, New Jersey, USA and additional entry points.

US POSTMASTER: send address corrections to IUPAC Solubility Data Series, c/o Virgin Mailing & Distribution, 10 Camptown Road, Irvington, New Jersey 07111-1105, USA.

New subscriptions may be registered at any time during the year but will be reckoned as running from January 1st of the year in which the subscription is received. Claims regarding non-receipt of issues must be received within 4 months of publication or date of order, whichever is later. Back issues are available—for information contact Journals Customer Services Department, Oxford University Press, Walton Street, Oxford OX2 6DP.

#### **Order Information**

details.

Payment is required with all orders and may be made by the following methods: Cheque (made payable to Oxford University Press) National Girobank (Account 500 1056) Credit Card (Access, Visa, American Express, Diners Club) Direct Debit: Please send for further details UNESCO Coupons Bankers: Barclays Bank plc, PO Box 333, Oxford Code 20-65-18. Account 00715654 Please send orders to: Journals Subscription Department, Oxford University Press, Walton Street, Oxford OX2 6DP. Tel: +44(0)1865 56767, Fax: +44(0)1865 267773

Any enquires regarding this publication should be addressed to: USA and Canada: Oxford University Press, 2001 Evans Road, Cary, NC 27513, USA. Rest of World: Journals Customer Services Department, Oxford University Press, Walton Street, Oxford OX2 6DP, UK.

©1995 International Union of Pure and Applied Chemistry. All rights reserved. No part of this publication may be reproduced, stored in a retrieval system, or transmitted in any form or by any means, electronic, mechanical, photocopying, recording or otherwise, without the prior written permission of the Copyright owner.

Printed and bound in Great Britain by Antony Rowe Ltd, Chippenham, Wiltshire

CONTENTS

	Introduction to the Solubility Data Series	
	Solubility of Liquids in Liquids	vii
	Preface	xiv
	Halogenated methanes solubilities	
	Note: an asterisk(*) indicates presence of a Critical Evaluation	
	<ul> <li>I. Bromotrichloromethane with water</li> </ul>	1
	2. Tribromofluoromethane with water	1
	-3. Tetrabromomethane with water	2
	4. Trichlorofluoromethane with water	3*
	/ 5. Trichlorofluoromethane with seawater	13
	· 6. Tetrachloromethane with water	17*
	7. Tetrachloromethane with MQ-Water	58
	8. Tetrachloromethane with municipal tap water (PASE)	58
	9. Tetrachloromethane with seawater	59
	~10. Tetrachloromethane with calcium chloride and water	61
	11. Tetrachloromethane with hydrogen chloride and water	62
	12. Tetrachloromethane with lithium chloride and water	63
	13. Tetrachloromethane with magnesium sulfate and water	64
l	,14. Tetrachloromethane with potassium chloride and water	65
	15. Tetrachloromethane with potassium hydroxide and water	66
	- 16. Tetrachloromethane with potassium sulfate and water	66
	- 17. Tetrachloromethane with sodium chloride and water	67
	<sup>18</sup> . Tetrachloromethane with sodium sulfate and water	68
	19. Tetrachloromethane with water-d <sub>2</sub>	69
	✓20. Bromodichloromethane with water	71 <b>•</b>
	21. Bromodichloromethane with Milang water	74
.	· 22. Bromodichloromethane with Myponga water +	75
	23. Bromodichloromethane with Myponga water humic acid	75
	.24. Dibromochloromethane with water	76*
	$\checkmark$ 25. Dibromochloromethane with Milang water	80
	$_{\nu}$ 26. Dibromochloromethane with Myponga water	80
	27. Dibromochloromethane with Myponga water + humic acid;	81
"	$^{\prime\prime}$ 28. Tribromomethane with water	82*
L	29. Tribromomethane with Milang water	90
1	30. Tribromomethane with Myponga water	90
1	31. Tribromomethane with Myponga water + humic acid	91
	32. Trichloromethane with water	92*
	/33. Trichloromethane with Milang water	132
J	/ 34. Trichloromethane with MQ-water	132

/ 35. Trichloromethane with Municipal tap water (PASE)	133
· 36. Trichloromethane with Myponga water	133
<sup>4</sup> 37. Trichloromethane with Myponga water + humic acid	134
38. Trichloromethane with seawater	135
39. Trichloromethane with cesium chloride and water	136
40. Trichloromethane with hydrogen chloride and water	137
41. Trichloromethane with lithium chloride and water	138
- 42. Trichloromethane with potassium chloride and water	139
43. Trichloromethane with rubidium chloride and water	139
44. Trichloromethane with sodium chloride and water	140
45. Trichloromethane with water-d <sub>2</sub>	140
46. Triiodomethane with water	141
/47.Bromochloromethane with water	143 <b>*</b>
48.Dibromomethane with water	146*
49.Dichloromethane with water	151 <b>*</b>
.50. Dichloromethane with hydrogen chloride and water	176
51.Diiodomethane with water	177*
/ 52.Diiodomethane with water-d <sub>2</sub>	180
53.Iodomethane; with water	181*
54. Iodomethane; with seawater	186
55. Iodomethane; with water-d <sub>2</sub>	187
System Index	188
Registry Number Index	190
Author Index	191
Solubility Data Series: Published and Forthcoming Volumes	196

Γ

#### INTRODUCTION TO THE SOLUBILITY DATA SERIES

#### SOLUBILITY OF LIQUIDS IN LIQUIDS

#### NATURE OF THE PROJECT

The Solubility Data project (SDP) has as its aim a comprehensive review of published data for solubilities of gases, liquids and solids in liquids or solids. Data of suitable precision are compiled for each publication on data sheets in a uniform format. The data for each system are evaluated and, where data from different sources agree sufficiently, recommended values are proposed. The evaluation sheets, recommended values, and compiled data sheets are published on consecutive pages.

This series is concerned primarily with liquid-liquid systems, but a limited number or related solid-liquid, fluid-fluid and multicomponent (organic-water-salt) systems are included where it is considered logical and appropriate. Solubilities at elevated and low temperatures and at elevated pressures have also been included, as it is considered inappropriate to establish artificial limits on the data presented if they are considered relevant or useful.

For some systems, the two components may be miscible in all proportions at certain temperatures and pressures. Data on reported miscibility gaps and upper and lower critical solution temperatures are included where appropriate and when available.

#### COMPILATIONS AND EVALUATIONS

The formats for the compilations and critical evaluations have been standardized for all volumes. A description of these formats follows.

#### Compilations

The format used for the compilations is, for the most part, self-explanatory. A compilation sheet is divided into boxes, with detailed contents described below.

*Components*: Each component is listed according to IUPAC name, formula, and Chemical Abstracts (CA) Registry Number. The Chemical Abstracts name is also included if this differs from the IUPAC name, as are trivial names if appropriate. IUPAC and common names are cross-referenced to Chemical Abstracts names in the System Index.

The formula is given either in terms of the IUPAC or Hill (1) system and the choice of formula is governed by what is usual for most current users: i.e., IUPAC for inorganic compounds, and Hill system for organic compounds. Components are ordered on a given compilation sheet according to:

- (a) saturating components:
- (b) non-saturating components in alphanumerical order;
- (c) solvents in alphanumerical order.

The saturating components are arranged in order according to the IUPAC 18-column periodic table with two additional rows:

H, alkali elements, ammonium, alkaline earth elements
transition elements
boron, carbon, nitrogen groups; chalcogenides, halogens
noble gases
Ce to Lu
Th to the end of the known elements, in order of atomic number

Organic compounds within each Hill formula are ordered in the following succession:

- (a) by degree of unsaturation
- (b) by order of increasing chain length in the parent hydrocarbon
- (c) by order of increasing chain length of hydrocarbon branches
- (d) numerically by position of unsaturation
- (e) numerically by position by substitution
- (f) alphabetically by IUPAC name.

Example:	
C <sub>5</sub> H <sub>8</sub>	cyclopentane
	2-methyl-1.3-butadiene
	1.4-pentadiene
	1-pentyne
CeHin	cvclopentane
- 310	3-methyl-1-butene
	2-methyl-2-butene
	1-pentene
	2-pentene
Cellin	2.2-dimethylpropane
- 512	2-methylbutane
	pentane
C <sub>5</sub> H <sub>12</sub> O	2,2-dimethyl-1-propanol
5 12	2-methyl-1-butanol
	2-methyl-2-butanol
	3-methyl-1-butanol
	3-methyl-2-butanol
	1-pentanol
	2-pentanol
	3-pentanol
C <sub>6</sub> H <sub>12</sub> O	cyclohexanol
• • • •	4-methyl-1-penten-3-ol
	1-hexen-3-ol
	4-hexen-3-ol
Deuterated ( <sup>2</sup> H) con	npounds follow immediately the corresponding H compounds.

Original Measurements: References are abbreviated in the forms given by Chemical Abstracts Service Source Index (CASSI). Names originally in other than Roman alphabets are given as transliterated by Chemical Abstracts. In the case of multiple entries (for example, translations) an

asterisk indicates the publication used for compilation of the data.

Variables: Ranges of temperature, pressure, etc. are indicated here.

Prepared by: The names of all compilers are given here.

*Experimental Values*: Components are described as (1), (2), etc., as defined in the "Components" box. Data are reported in the units used in the original publication, with the exception that modern names for units and quantities are used; e.g., mass per cent for weight per cent; mol dnt<sup>3</sup> for molar; etc. In most cases, both mass and molar values are given. Usually, only one type of value (e.g., mass per cent) is found in the original paper, and the compiler has added the other type of value (e.g., mole per cent) from computer calculations based on 1989 atomic weights (2). Temperatures are expressed as  $t/^{\circ}$ C,  $t/^{\circ}$ F or T/K as in the original; if necessary, conversions to T/K are made, sometimes in the compilations, and always in the critical evaluation. However, the author's units are expressed according to IUPAC recommendations (3) as far as possible.

Errors in calculations, fitting equations, etc. are noted, and where possible corrected. Material inserted by the compiler is identified by the word "compiler" or by the compiler's name in parentheses or in a footnote. In addition, compiler-calculated values of mole or mass fractions are included if the original data do not use these units. If densities are reported in the original paper, conversions from concentrations to mole fractions are included, but otherwise this is done in the evaluation, with the values and sources of the densities being quoted and referenced.

Details of smoothing equations (with limits) are included if they are present in the original publication and if the temperature or pressure ranges are wide enough to justify this procedure and if the compiler finds that the equations are consistent with the data.

The precision of the original data is preserved when derived quantities are calculated, if necessary by the inclusion of one additional significant figure. In some cases graphs have been included, either to illustrate presented data more clearly, or if this is the only information in the original. Full grids are not usually inserted as it is not intended that users should read data from the graphs.

*Method*: The apparatus and procedure are mentioned briefly. Abbreviations used in Chemical Abstracts are often used here to save space, reference being made to sources of further detail if these are cited in the original paper.

Source and Purity of Materials: For each component, referred to as (1), (2), etc., the

following information (in this order and in abbreviated form) is provided if available in the original paper: source and specified method of preparation; properties; degree of purity.

*Estimated Error*: If estimated errors were omitted by the original authors, and if relevant information is available, the compilers have attempted to estimate errors (identified by "compiler" or the compiler's name in parentheses or in a footnote) from the internal consistency of data and type of apparatus used. Methods used by the compilers for estimating and reporting errors are based on Ku and Eisenhart (4).

Comments and/or Additional Data: Many compilations include this section which provides short comments relevant to the general nature of the work or additional experimental and thermodynamic data which are judged by the compiler to be of value to the reader.

*References*: The format for these follows the format for the Original Measurements box, except that final page numbers are omitted. References (usually cited in the original paper) are given where relevant to interpretation of the compiled data, or where cross-reference can be made to other compilations.

#### Evaluations

The evaluator's task is to assess the reliability and quality of the data, to estimate errors where necessary, and to recommend "best" values. The evaluation takes the form of a summary in which all the data supplied by the compiler have been critically reviewed. There are only three boxes on a typical evaluation sheet, and these are described below.

Components: The format is the same as on the Compilation sheets.

*Evaluator*: The name and affiliation of the evaluator(s) and date up to which the literature was checked.

Critical Evaluation:

(a) Critical text. The evaluator checks that the compiled data are correct, assesses their reliability and quality, estimates errors where necessary, and recommends numerical values based on all the published data (including theses, reports and patents) for each given system. Thus, the evaluator reviews the merits or shortcomings of the various data. Only published data are considered. Documented rejection of some published data may occur at this stage, and the corresponding compilations may be removed.

The solubility of comparatively few systems is known with sufficient accuracy to enable a set of recommended values to be presented. Although many systems have been studied by at least two workers, the range of temperatures is often sufficiently different to make meaningful comparison impossible.

Occasionally, it is not clear why two-groups of workers obtained very different but internally consistent sets of results at the same temperature, although both sets of results were obtained by reliable methods. In such cases, a definitive assessment may not be possible. In some cases, two or more sets of data have been classified as tentative even though the sets are mutually inconsistent.

(b) Fitting equations. If the use of a smoothing equation is justifiable the evaluator may provide an equation representing the solubility as a function of the variables reported on all the compilation sheets, stating the limits within which it should be used.

(c) Graphical summary. In addition to (b) above, graphical summaries are often given.

(d) Recommended values. Data are recommended if the results of at least two independent groups are available and they are in good agreement, and if the evaluator has no doubt as to the adequacy and reliability of the applied experimental and computational procedures. Data are reported as tentative if only one set of measurements is available, or if the evaluator considers some aspect of the computational or experimental method as mildly undesirable but estimates that it should cause only minor errors. Data are considered as doubtful if the evaluator considers some aspect of the computational or experimental method as undesirable but still considers the data to have some value where the order of magnitude of the solubility is needed. Data determined by an inadequate method or under ill-defined conditions are rejected. However, references to these data are included in the evaluation together with a comment by the evaluator as to the reason for their rejection.

(e) References. All pertinent references are given here, including all those publications appearing in the accompanying compilation sheets and those which, by virtue of their poor precision, have been rejected and not compiled.

(f) Units. While the original data may be reported in the units used by the investigators, the final recommended values are reported in SI units (3) when the data can be converted accurately.

# QUANTITIES AND UNITS USED IN COMPILATION AND EVALUATION OF SOLUBILITY DATA

#### Mixtures, Solutions and Solubilities

A mixture (5, 6) describes a gaseous, liquid or solid phase containing more than one substance, where the substances are all treated in the same way.

A solution (5, 6) describes a liquid or solid phase containing more than one substance, when for convenience one of the substances, which is called the *solvent*, and may itself be a mixture, is treated differently than the other substances, which are called *solutes*. If the sum of the mole fractions of the solutes is small compared to unity, the solution is called a *dilute solution*.

The solubility of a solute 1 (solid, liquid or gas) is the analytical composition of a saturated solution, expressed in terms of the proportion of the designated solute in a designated solvent (7).

"Saturated" implies equilibrium with respect to the processes of dissolution and demixing; the equilibrium may be stable or metastable. The solubility of a substance in metastable equilibrium is usually greater than that of the same substance in stable equilibrium. (Strictly speaking, it is the activity of the substance in metastable equilibrium that is greater.) Care must be taken to distinguish true metastability from supersaturation, where equilibrium does not exist.

Either point of view, mixture or solution, may be taken in describing solubility. The two points of view find their expression in the reference states used for definition of activities, activity coefficients and osmotic coefficients. Note that the composition of a saturated mixture (or solution) can be described in terms of any suitable set of thermodynamic components.

#### **Physicochemical Quantities and Units**

Solubilities of solids have been the subject of research for a long time, and have been expressed in a great many ways, as described below. In each case, specification of the temperature and either partial or total pressure of the saturating gaseous component is necessary. The nomenclature and units follow, where possible, ref. (3)

A note on nomenclature. The nomenclature of the IUPAC Green Book (3) calls the solute cpmponent B and the solvent component A. In compilations and evaluations, the first-named component (component 1) is the solute, and the second (component 2 for a two-component system) is the solvent. The reader should bear these distinctions in nomenclature in mind when comparing equations given here with those in the Green Book.

1. Mole fraction of substance 1,  $x_1$  or x(1):

$$x_1 = n_1 / \sum_{s=1}^{c} n_s$$
 [1]

where  $n_s$  is the amount of substance of s, and c is the number of distinct substances present (often the number of thermodynamic components in the system). Mole per cent of substance 1 is 100  $x_1$ .

2. Ionic mole fractions of salt i,  $x_{i+}$ ,  $x_{i-}$ :

For a mixture of s binary salts i, each of which ionizes completely into  $v_{s+}$  cations and  $v_{s-}$  anions, with  $v_s = v_{s+} + v_{s-}$  and a mixture of p non-electrolytes j, of which some may be solvent components, a generalization of the definition in (8) gives:

$$x_{i+} = \frac{v_{i+}x_i}{1 + \sum_{i=1}^{s} (v_i - 1)x_s}, \quad x_{i-} = \frac{v_{i-}x_{i+}}{v_{i+}} \quad i = 1...s$$
[2]

$$x'_{j} = \frac{x_{j}}{1 + \sum_{l=1}^{s} (v_{l} - 1)x_{l}}, \quad j = (s+1), \dots, p$$
[3]

The sum of these mole fractions is unity, so that, with c = s + p,

$$\sum_{i=1}^{s} (x_{i+} + x_{i-}) + \sum_{i=s+1}^{c} x_{i}' = 1$$
[4]

General conversions to other units in multicomponent systems are complicated. For a threecomponent system containing non-electrolyte 1, electrolyte 2 and solvent 3,

$$x_{1} = \frac{v_{2+}x_{1}'}{v_{2+}(v_{2}-1)x_{2+}} \qquad x_{2} = \frac{x_{2+}}{v_{2+}(v_{2}-1)x_{2+}}$$
[5]

These relations are used in solubility equations for salts, and for tabulation of salt effects on solubilities of gases (see below).

3. Mass fraction of substance 1,  $w_1$  or w(1):

$$w_1 = g_1 / \sum_{s=1}^{c} g_s$$
 [6]

where  $g_s$  is the mass of substance s. Mass per cent of substance 1 is 100  $w_1$ . The equivalent terms weight fraction, weight per cent and g (1)/100 g solution are no longer used.

4. Molality of solute 1 in a solvent 2,  $m_1$ :

$$m_1 = n_1/n_2 M_2$$
 (7)

[7]

SI base units: mol kg<sup>-1</sup>. Here,  $M_2$  is the molar mass of the solvent.

5. Aquamolality, Solvomolality of substance 1 in a mixed solvent with components 2, 3 (9),  $m_1^{(3)}$ :

$$m_1^{(3)} = m_1 \overline{M} / M_3$$
 [8]

SI base units: mol kg<sup>-1</sup>. Here, the average molar mass of the solvent is

$$\overline{M} = x_2' M_2 + (1 - x_2') M_3$$
<sup>[9]</sup>

and  $x_2$ ' is the solvent mole fraction of component 2. This term is used most frequently in discussing comparative solubilities in water (component 2) and heavy water (component 3) and in their mixtures.

6. Amount concentration of solute 1 in a solution of volume V,  $c_1$ :

 $c_1 = [\text{formula of solute}] = n_1/V$  [10]

SI base units: mol  $m^{-3}$ . The symbol  $c_1$  is preferred to [formula of solute], but both are used. The old terms *molarity*, *molar* and *moles per unit volume* are no longer used.

7. Mass concentration of solute 1 in a solution of volume V,  $\rho_1$ :SI base units: kg m<sup>-3</sup>.

ρ

7

$$f_1 = g_1/V \tag{11}$$

8. Mole ratio,  $r_{n,12}$  (dimensionless)

$$r_{n,12} = n_1/n_2$$
 [12]

Mass ratio, symbol  $r_{g,12}$ , may be defined analogously.

Mole and mass fractions are appropriate to either the mixture or the solution point of view. The other quantities are appropriate to the solution point of view only. Conversions between pairs of these quantities can be carried out using the equations given in Table 1 at the end of this Introduction. Other useful quantities will be defined in the prefaces to individual volumes or on specific data sheets.

9. Density, ρ:

 $\rho = g/V$ 

[13]

SI base units: kg m<sup>-3</sup>. Here g is the total mass of the system.

10. Relative density,  $d = \rho/\rho^{\circ}$ : the ratio of the density of a mixture at temperature *t*, pressure *p* to the density of a reference substance at temperature *t'*, pressure *p'*. For liquid solutions, the reference substance is often water at 4°C, 1 bar. (In some cases 1 atm is used instead of 1 bar.) The term specific gravity is no longer used.

#### Thermodynamics of Solubility

Thermodynamic analysis of solubility phenomena provides a rational basis for the construction of functions to represent solubility data, and thus aids in evaluation, and sometimes enables thermodynamic quantities to be extracted. Both these are often difficult to achieve because of a lack of experimental or theoretical activity coefficients. Where thermodynamic quantities can be found, they are not evaluated critically, since this task would involve examination of a large body of data that is not directly relevant to solubility. Where possible, procedures for evaluation are based on established thermodynamic methods. Specific procedures used in a particular volume will ybe described in the Preface to that volume.

#### REFERENCES

- 1. Hill, E.A. J. Am. Chem. Soc. <u>1900</u>, 22, 478.
- 2. IUPAC Commission on Atomic Weights and Isotopic Abundances. Pure Appl. Chem. <u>1989</u>, 63, 975.
- 3. Mills, I.; et al., eds. Quantities, Units and Symbols in Physical Chemistry (the Green Book). Blackwell Scientific Publications. Oxford, UK. <u>1993</u>.
- 4. Ku, H.H., p. 73; Eisenhart, C., p. 69; in Ku, H.H., ed. Precision Measurement and Calibration. *NBS Special Publication 300.* Vol. 1. Washington. <u>1969</u>.
- 5. Regaudy, J.; Klesney, S. P. Nomenclature of Organic Chemistry (IUPAC), ("The Blue Book"), Pergamon, Oxford, <u>1979.</u>
- 6. Gold, V.; et al., eds. Compendium of Chemical Technology (the Gold Book). Blackwell Scientific Publications. Oxford, UK. <u>1987</u>.
- 7. Freiser, H.; Nancollas, G.H., eds. Compendium of Analytical Nomenclature (the Orange Book). Blackwell Scientific Publications. Oxford, UK. <u>1987</u>. Sect. 9.1.8.
- 8. Robinson, R.A.; Stokes, R.H. *Electrolyte Solutions*. Butterworths. London. <u>1959</u>. 2nd ed.
- Lorimer, J.W. in Cohen-Adad, R.; Lorimer, J.W. Alkali Metal and Ammonium Chlorides in Water and Heavy Water (Binary Systems). IUPAC Solubility Data Series, Vol. 47. Pergamon Press. Oxford, UK, <u>1991</u>. p. 495.

A.F.M. Barton Perth, WA, Australia

F.W. Getzen Raleigh, NC, USA

G.T. Hefter Perth, WA, Australia

D.G. Shaw Fairbanks, AK, USA

January, 1994

	x <sub>i</sub>	w <sub>i</sub>	m <sub>i</sub>	c <sub>i</sub>
<i>x<sub>i</sub></i> =	x <sub>i</sub>	$\frac{1}{1 + \frac{M_i}{M_c} \left(\frac{1}{w_i} - 1 + \sum_{j \neq i}^{c-1} \left(\frac{M_c}{M_j} - 1\right) \frac{w_j}{w_i}\right)}$	$\frac{1}{1 + \frac{1}{m_i M_c} + \sum_{j \neq i}^{c-1} \frac{m_j}{m_i}}$	$\frac{1}{1 + \frac{1}{M_c} \left(\frac{\rho}{c_i} - M_i\right) + \sum_{j \neq i}^{c-1} \frac{c_j}{c_i} \left(1 - \frac{M_j}{M_c}\right)}$
w <sub>i</sub> =	$\frac{1}{1 + \frac{M_c}{M_i} \left(\frac{1}{x_i} - 1 + \sum_{j \neq i}^{c-1} \left(\frac{M_j}{M_c} - 1\right) \frac{x_j}{x_i}\right)}$	w <sub>i</sub>	$\frac{1}{1 + \frac{1}{m_i M_i} \left(1 + \sum_{j \neq i}^{c-1} m_j M_j\right)}$	<u> <i>c<sub>i</sub>M<sub>i</sub></i></u> ρ
<i>m<sub>i</sub></i> =	$\frac{1}{M_c \left(\frac{1}{x_i} - 1 - \sum_{j \neq i}^{c-1} \frac{x_j}{x_i}\right)}$	$\frac{1}{M_i\left(\frac{1}{w_i}-1-\sum_{j\neq i}^{c-1}\frac{w_j}{w_i}\right)}$	m <sub>i</sub>	$\frac{1}{\frac{1}{c_i} \left( \rho - \sum_{j \neq i}^{c-1} c_j M_j \right) - M_i}$
<i>c<sub>i</sub></i> =	$\frac{\rho}{M_i + M_c \left(\frac{1}{x_i} - 1 + \sum_{j \neq i}^{c-1} \left(\frac{M_j}{M_c} - 1\right) \frac{x_j}{x_i}\right)}$	$\frac{\rho w_i}{M_i}$	$\frac{\rho}{\frac{1}{m_i} \left(1 + \sum_{j \neq i}^{c-1} M_j m_j\right) + M_i}$	c <sub>i</sub>

Table 1. Interconversions between Quantities Used as Measures of Solubility c-component Systems Containing c - 1 Solutes i and Single Solvent c

 $\rho$  - density of solution;  $M_i$  - molar masses of i. For relations for 2-component systems, set summations to 0.

Ě

## PREFACE

This volume contains a comprehensive collection and critical evaluation of solubility data published prior to 1993 for halogenated methanes with water, seawater, heavy water, and aqueous electrolyte solutions. The systems are ordered on the basis of chemical formula according to the Hill system.

A critical evaluation of the solubility data is not satisfactory without a comprehensive literature search followed by compilation of all the available information reported in journals, patents, pamphlets, brochures, books, etc. The difficulties and problems in retrieval of all solubility data are a well recognized fact. Some of the difficulties which arose during the retrieval of the original publications for this volume were:

- i. The articles, reports, etc. were not cited in the Chemical Abstracts
- ii. The British Library did not hold the publications
- iii. The Slavonic Section of the British Library was unable to obtain some publications from abroad
- iv. Manufacturers' reports, leaflets, brochures, etc. were not available from industrial source
- v. University reports from experimental stations were not available from university libraries.

In summary, there were several cases where the original sources were not obtainable and the information had to be taken from secondary sources.

The referenced literature sources can be classified into the following groups:

- A. Bibliographies
- B. Secondary sources (books, reviews)
- C. Databases (computer softwares)
- D. Primary sources (journals, reports, etc.)

A list of the various sources of solubility data following the above classification is summarized in Table 1.

#### Table 1. Sources of Solubility Data

#### A. Bibliographies:

- 1. Wisniak, J.; Tamir, A. Liquid-Liquid Equilibrium and Extraction. A Literature Source Book, Elsevier, Amsterdam, <u>1980</u>, 1252 pp.
- 2. Wisniak, J.; Herskowitz, M. Solubility of Gases and Solids. A Literature Source Book, Parts A and B, Elsevier, Amsterdam, <u>1984</u>, 2070 pp.

#### **B. Secondary Sources:**

- 1. Ware, G. W. Review of Environmental Contamination and Toxicology, Vol. 116, Springer-Verlag, New York, <u>1990</u>, 200 pp.
- 2. Freier, R. K. Aqueous Solutions, Vol. 1, Walter de Gruyter, Berlin, 1976, 477 pp.
- 3. Landolt-Börnstein Zahlenwerte und Funktionen aus Physik, Chemie, Astronomie, Geophysik und Technik, Gleichgewichte der Absorption von Gasen in Flüssigkeiten, Vol. 4a/c1, Springer-Verlag, Berlin, <u>1976</u>, 479 pp.
- Landolt-Börnstein Zahlenwerte und Funktionen aus Physik, Chemie, Astronomie, Geophysik und Technik, (6<sup>th</sup> ed.), Vol. 1, Eigenschaften der Materie in Ihren Aggregationstäden, Part 2b, Lösungsgleichgewichte I, Springer-Verlag, Berlin, <u>1962</u>.
- Landolt-Börnstein Zahlenwerte und Funktionen aus Physik, Chemie, Astronomie, Geophysik und Technik, (6<sup>th</sup> ed.), Vol. 2, Eigenschaften der Materie in Ihren Aggregatzustäden, Part 2c, Lösungsgleichgewichte II, Springer-Verlag, Berlin, <u>1964</u>.
- 6. Horvath, A. L. Halogenated Hydrocarbons. Solubility-Miscibility with Water, Marcel Dekker, New York, <u>1982</u>, 889 pp. 7. Washburn, E. W., Ed. International Critical Tables of Numerical Data, Vol. 3, McGraw-Hill, New York, <u>1928</u>.
- 7. Washburn, E. W., Ed. International Critical Tables of Numerical Data, Vol. 3, McGraw-Hill, New York, <u>1928</u>.

Table 1.	Sources	of Solubility	Data	(continued)
				(/

- 8. Beilsteins Handbuch der Organischen Chemie, Beilstein Institut für Literatur der Organischen Chemie, Springer-Verlag, Berlin, Vol. 1, <u>1918-</u>
- 9. Kirk-Othmer Encyclopedia of Chemical Technology, 1<sup>st</sup>, 2<sup>nd</sup>, 3<sup>rd</sup>, and 4<sup>th</sup> eds., Wiley, New York.
- 10. Seidell, A. Solubility of Organic Compounds, 3rd ed., Van Nostrand, New York, 1963, 926 pp.
- 11. Linke, W. F. Solubilities of Inorganic and Metal-Organic Compounds, 4th Ed., van Nostrand, Princeton, N. J., Vol. 1, <u>1958</u>, Vol. 2, <u>1965</u>.
- 12. Stephen, H.; Stephen, T. Solubilities of Inorganic and Organic Compounds, Vols. 1 and 2, Pergamon Press, Oxford, <u>1963</u>.
- 13. Riddich, J. A.; Bunger, W. B. Organic Solvents, 3<sup>rd</sup> ed., Vol. 2, Wiley-Interscience, New York, <u>1970</u>, 603 pp.
- 14. DECHEMA Data Serires, Gmeling, J.; Onken, U. Vapor-Liquid Equilibrium Data Collection. Aqueous-Organic Systems, Vol. 1, <u>1977</u>.
- 15. Dreisbach, R. R. Physical Properties of Chemical Componds, Am. Chem. Soc. Adv. in Chem. Ser. 15 (1955, 536 pp.), 22 (1959, 491 pp.), and 29(1961, 489 pp.), Washington, D. C.

#### C. Databases:

1. AQUASOL database of Aqueous Solubility, 5<sup>th</sup>., S. H. Yalkowsky, Ed., University of Arizona, Tucson, Arizona, <u>1991</u>.

#### D. Primary Sources:

- 1. Journals
- 2. Reports
- 3. Brochures, leaflets
- 4. Patents
- 5. Private communications

Once a copy of the original document - usually a photocopy - was obtained, the first step was to read it carefully. Often the original article was in a foreign language which required a translation. If the description was not focused on the solubility aspect, then often some relevant details, e.g., source and purity of the solute and solvent, or both, were not included in the text. In many cases, the method, apparatus, and procedure were simply referred to other articles. Furthermore, when the experimental errors were not stated, the compiler sometimes introduced a subjective judgement for the accuracy of the measured solubility and tem-perature. The estimated accuracy of the measured quantities is quite subjective and the readers must take this fact into consideration.

The summaries of the procedures and abstracts were based on the available description of the method, apparatus, and procedures. Often a short statement "Details are not available," had to be used when the main contents of the articles focused not on solubility measurements but some other aspects of chemistry.

The conversion of the published solubility data into conventional units often presented difficulties, particularly when the dimensions were not expressed clearly. For example, the Henry's law constants, separation coefficients, distribution ratio, partition coefficients, etc. were not specified explicitly. Some of the more relevant conversion formulas are given in Table 2.

To convert solubility data from one unit to another, often the density and vapor pressure of the pure components and/or of the mixture were needed. The physical properties of the pure compounds were taken from the DIPPR (ref. 1) and TRCVP (ref. 2) databases. To find or derive the physical properties of mixtures required a more elaborate effort. Experimental data, e.g., density and vapor pressure for aqueous solutions of halogenated hydrocarbons are extremely rare in the literature. Consequently, approximations were obtained using calculations recommended in standard handbooks, e.g., Reid et al. (ref. 2). An illustration of the

Table 2. Conversion Formulas for Solubility Units.  
a. Henry's Law Constant (*H*):  

$$Mass \% = \frac{\left[\frac{P_{up}}{H(\dim ensionless)}\right]}{H(\dim ensionless)} \times 100$$

$$Mass \% = \frac{\left[\frac{P_{up}}{(M + 1)}(\frac{dim}{d \mod 1})\right]}{d_{udwar}} \times 100$$

$$Mass \% = \frac{\left[\frac{P_{up}}{(M + 1)}(\frac{dim}{d \mod 1})\right]}{d_{udwar}} \times 10^{-4}$$

$$H(\dim ensionless) = \frac{C_{ur}}{C_{udwar}} = \frac{16.04}{T(K)} \frac{P_{up}}{S_{udwar}} (\operatorname{mm}Hg) M_{udwar}}$$
b. Air/Water Partition Coefficient (*K*<sub>a</sub>m):  

$$Mass \% = \frac{100}{82.054} \frac{K_{up}P_{up}}{T(K)} \frac{M_{udwar}}{M_{udwar}}$$
c. Distribution Ratio (*p/c*):  

$$(p/c) = \frac{\text{solute vapor pressure}}{\text{solute concentration}} \frac{(\operatorname{mm}Hg)}{(\operatorname{mmole}/dm^3)}$$
d. Partition Coefficient (*K*):  

$$K = \frac{1}{\text{Ostwald coefficient}} = \frac{1}{L}$$
e. Distribution Coefficient (*K*):  

$$K_{i} = \text{Ostwald coefficient} = L = \frac{\gamma_{i}}{x_{i}}$$
difficulty of obtaining accurate conversions is shown by the following typical case.  
The solubility of earbon tetrachloride (CCL) in water (H\_{i}O) is 0.10 cm<sup>3</sup>100 cm<sup>3</sup> water at 25 °C. The solubility in mass per cent (100 w<sub>i</sub>) is required. The density of pure carbon tetrachloride is 1.5834 g CCL/cm<sup>3</sup> at 25 °C. Older Marken at 25 °C. The solubility in mass per cent (100 w<sub>i</sub>) is required. The density of pure carbon tetrachloride is 1.5834 g CCL/cm<sup>4</sup> at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. The solution of earbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, the density of the aqueous solution of carbon tetrachloride at 25 °C. However, th

nal. As a consequence, the errors obtained upon the determination of solubility are considerably greater than the possible difference between the solubility in pure water or in its aqueous solutions at the same temperature. It follows, in this case, one must use the approximation

$$g CCl_4 / 100 g H_2 O \approx 100 w_1$$
.

On the other hand, for solubility values larger than, say, 2 - 3 mass per cent, the differences become significant, as the following example illustrates. The solubility in water of dichloromethane (CH<sub>2</sub>Cl<sub>2</sub>) is 2.363 g/100 g water at 0°C. The mass per cent solubility, 100  $w_1$ , at the same temperature is required.

If 100 g water dissolves 2.363 g  $CH_2Cl_2$ , then the total mass of the solution becomes 102.363 g. This means that 100 g aqueous solution will dissolve less  $CH_2Cl_2$  than 100 g pure water, that is,

100 
$$w_1 = \frac{2.363 \text{ g CH}_2\text{Cl}_2}{102.363 \text{ g soln}}$$
 100 = 2.3084

The resulting mass per cent is 2.3084 at 0°C. Consequently, there is a 2.31 per cent reduction in the solubility relative to 100 g pure water. This is in comparison to the reduction of only 0.10 per cent in the case of the solubility of CCl<sub>4</sub> in 100 g water or 100 g aqueous solution at 25°C.

All critical evaluations of the solubility data are presented on the evaluation sheets. The information available on the relevant compilation sheets served as the fundamental source for the critical evaluations. The following main aspects of the solubility data have been considered:

Purity of the components (solute and solvents) Date of the experiment Accuracy of the method Reproducibility of the experiment Experience of the investigator(s), previous publications Consistency of the data, trends in series

Once these aspects of the works had been examined, efforts were made to weight the reliability of the reported data. As much as possible, the differences in the quality of the data have been taken into account.

The available solubility data for binary systems may be very scarce, medial, or abundant, depending on the commercial application of the systems. Whenever data from different sources exist, they often disagree. For evaluation of the agreement or disagreement among the data, graphical presentations usually provide the clearest picture. The figures normally contain the original data from all references for a given system. A plot representing the solubility data (expressed in mole fraction or mass per cent) can reveal whether or not a trend exists among the data in addition to highlighting any discrepancies of the experimental solubility measurements.

The final data can be classified as:

obscure or dubious - rejected data tentative - not recommended data recommended - good agreement among the published results.

A graphical presentation of the solubility data may be performed on linear, semi-logarithmic, etc. scales. The unit of solubility may be given as specific or molar scales. For practical reasons, the mass per cent unit is often used, whereas the mole fraction is another common unit of solubility. In this volume most figures are plotted using one of two different scales. For the solubilities in water, linear graphs are used and for the solubilities of water in halogenated hydrocarbons, the so called Cox chart is applied. In the Cox charts, the solubility of water is given as  $\log_{10} x_2$  versus 1/T. A straight line on a Cox plot can be very useful for revealing consistent data covering a limited temperature range. However, the straight line does not apply for solubility data over a very wide temperature interval, e.g., from the triple point to the critical point. Therefore, in some cases (e.g., solubility of water in chloroform), the normal polynomial equations were used for smoothing the solubility data.

To fit experimental data into the various equations, the individual data points are often weighted. The smoothed data may be influenced somewhat by such procedures of weighting individual data points. Data analysis is a process of mathematical and/or graphical treatment of experimental data through the use of statitical or parametric procedures. Whenever the raw data are fragmentary, the aim is to generate an extended range of internally consistent values by synthesis. In any case, the temperature dependence of solubility must be examined and any unusual irregularities must be investigated carefully.

The temperature dependence of the solubility of halogenated hydrocarbons in water has been correlated using normal polynomial equations over limited temperature intervals. These equations should be used only in the temperature range indicated. The equations play two major roles; to provide data users with interpolated data with good accuracy and to provide data users with acceptable data for use in computer programs, if desired. The deviation of the various experimental values from the fitted data are given as standard or average deviations. If the experimental data cover a reasonable temperature interval, then the data and the smoothing equation are often illustrated in a figure. The graphical presentation of the reported and fitted data clearly indicates the risks involved when one attempts to extrapolate the raw experimental data. In some cases, an extrapolation is quite safe for the desired temperature (see for example, the solubility of water in halogenated hydrocarbons over moderate temperature intervals); whereas, in other cases, the risk is very high and it is not recommended. The shape of the solubility curves suggests the extrapolation and interpolation possibilities.

Whether the mutual solubility between two liquids is partial or complete depends on the similarities or differences between the molecules of the two chemical compounds. The halogenated hydrocarbon and water systems are only partially miscible without exception. Furthermore, the miscibility or immiscibility of water with other liquids is also very dependent upon temperature. At ambient temperature and atmospheric pressure, the miscibility is very limited between water and halogenated hydrocarbons. For most systems, the mutual solubility is less than 2 mass per cent at ambient room temperatures.

In a liquid-liquid system the variation of solubility with temperature depends on the sign of the heat of mixing or solution. The minimum dissolution temperature occurs for most halogenated hydrocarbon - water systems between 270 and 310 K. The interactions between the solute and solvent molecules provide the explanation for the phenomena. The temperature dependence of solubility is a result of temperature dependent structural modifications of water. A different energy is required for cavity formation and for the changes in rigidity of the lattice as temperature is changed.

The appearance of these minima resembles the dependence of the critical micelle concentration upon temperature. Here, the iceberg formation of water molecules around the monodisperse solute enhances the critical micelle concentration, the logarithm of which would be decreased linearly with 1/T provided the solution behaves regularly, and the iceberg formation increases with a temperature reduction.

It is well recognised that the rare gases and hydrocarbon gases form iceberg-like structures when dissolved in cold water, followed by an iceberg melting as the temperature is raised. The solubility of most gases in water shows a minimum when plotted against temperature. The shape of the solubility curves for molecules with alkyl groups is explained by considering that the ice-like structure of water is also formed around the nonpolar alkyl group of molecules in water. This structure tends to break down as the temperature of the solution is increased.

The minimum solubility temperature can be calculated from a calorimetric measurement of the enthalpy of solution at infinite dilution ( $\Delta_{sole}H^{\infty}$ ). This value is very large and varies from negative to positive values within a small temperature range. Consequently, very large errors may be caused by an erroneous interpretation. In addition to the enthalpy of solution, the change in the heat capacity at infinite dilute aqueous solution ( $\Delta_{sole}C_{p}^{\infty}$ ) is required for the calculation of the minimum solubility temperature ( $T_{min}$ );

$$T_{\min} = 298.15 \text{ K} - \frac{\Delta_{\text{soln}} H^{\circ}(298.15 \text{ K})}{\Delta_{\text{soln}} C_{p}^{\circ}}$$

A more detailed description of the minimum solubility phenomenon is given by Horvath (ref. 4), Shinoda (ref. 5), Nishino and Nakamura (ref. 6), and Privalov and Gill (ref. 7).

The relations between the solubility and some of the physical properties of the solute or solvent have been investigated from early times in solution studies. The various experiments showed that in very diluted solutions the influence of the solute molecules does not extend to all solvent molecules in the solution. Consequently, the large fraction of solvent molecules, which are not affected by solute molecules, must exist more or less in the same state of aggregation as in the pure state. A change of solvent molar volume in dilute solutions is very small. However, in more concentrated solutions the effects of the presence of the solute molecules are not negligible. The amount of molar volume change of the solvent depends on the nature of the solutes. The dissolution process is related to the disintegration of attraction of the solute in the solvent, as compared to its pure state molecular order, as a result of the mutual action of attraction of the solute and solvent.

In general, solubility depends on many parameters and is of a very complex nature. There is no simple relation established between solubility and the other properties of the pure components (solute and solvent). Despite the lack of a simple relationship, the solubility of halogenated benzenes in water at 25°C has been correlated with a reasonable success using the molar volume of the solutes (ref. 8). Similarly, the solubility of halogenated methanes in water shows a linear relationship (ref. 9). The correlation is based on the assumption that the molar volumes in saturated solutions are not greatly different from those in the pure state. The simple linear relation at 25°C is illustrated graphically in Figure 1. This is a very useful and practical way to show the consistency of the solubility data. Any significant deviation from such a linear relationhip suggests some inconsistency and introduces doubt concerning the reliability of the data.



Figure 1. Logarithm of Saturated Molarity versus Solute Molar Volume at 25 °C for Halogenated Methanes in Water.

Correlation equation:

 $\log_{10} (S/\text{mole } \text{dm}^{-3}) = 1.8995 - 4.3707 \times 10^{-2} (V_m/\text{cm}^3 \text{ mole}^{-1})$ 

**REFERENCES:** 

- 1. DIPPR, Design Institute for Physical Property Data, Am. Inst. Chem. Eng., New York, Software package for IBM PC, <u>1985</u>.
- 2. TRCVP, Thermodynamic Research Center, Texas A & M University System, College Station, TX., 1989.
- 3. Reid, R. C.; Prausnitz, J. M.; Poling, B. E. The Properties of Gases and Liquids, 4<sup>th</sup> ed., McGraw-Hill, New York, <u>1987</u>, 742 pp.
- 4. Horvath, A. L. Halogenated Hydrocarbons: Solubility-Miscibility with Water, Marcel Dekker, New York, <u>1982</u>, 889 pp.
- 5. Shinoda, K. J. Phys. Chem. 1977, 81, 1300.
- 6. Nishino, N.; Nakamura, M. Bull. Chem. Soc. Japan <u>1978</u>, 51, 1617.
- 7. Pivalov, P. L.; Gill, S. J. Pure Appl. Chem. 1989, 61, 1097.
- 8. Horvath, A. L.; Getzen, F. W. Halogenated Benzenes, Toluenes and Phenols with Water, IUPAC Solubility Data Series, Vol. 20, Pergamon Press, Oxford, <u>1985</u>, 266 pp.
- 9. Horiba, S. Memoires of the College of Science and Engineering, Kyoto Imperial University, <u>1917</u>, Vol. 2, p. 1-43.

	1	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	O'Connell, W. L.	
(2) Bromotrichloromethane; CBrCl <sub>3</sub> ; [75-62-7]	Trans. Am. Inst. Mech. Eng. <u>1963</u> , 226, 126-32.	
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 g <sub>1</sub> /g <sub>2</sub>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & (compiler) \end{array}$	
20 $6.0 \times 10^{-3}$	$6.0 \times 10^3$ $6.6$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	(1) Distilled (compiler).	
	(2) Dow Chemicals Co., used as received.	
	ESTIMATED ERRORS:	
	Solubility: Not specified.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromofluoromethane; CBr <sub>3</sub> F; [353-54-8]	O'Connell, W. L.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Trans. Am. Inst. Mech. Eng. <u>1963</u> , 226, 126-32.	
VARIABLES:	PREPARED BY:	
T/K = 293 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 g <sub>1</sub> /g <sub>2</sub> 10 <sup>5</sup> (com	$x_1 = \frac{100 g_2/g_1}{(compiler)}$	
20	$2.0 \times 10^{-2}$ 3.0	
25 $4.0 \times 10^{-2}$ 2	.66	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ol> <li>Dow Chemicals Co., used as received.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: + 0.5 K (compiler).	

2	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrabromomethane (Carbon tetrabromide);	Gross, P. M.; Saylor, J. H.
$CDI_4$ ; [336-13-4] (2) Water: H.O. [7732-18-5]	J. Am. Soc. Soc. <u>1931</u> , 53, 1744-51.
(2) walki, 1120, (7752-10-5)	
VARIABLES:	PREPARED BY:
T/K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	•
$t/^{\circ}C$ 1000 $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
30 0.24	$2.4 \times 10^{-2}$ 1.30
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS
An excess of tetrabromomethane in 500 g water was shaken for 12 hours in a thermostat bath. Sam- ples were then withdrawn and read against water in an interferometer made by Zeiss (ref. 1). A detailed description of the complete procedure is given in a	<ol> <li>Eastman Kodak Co., recrystallized from ethyl alcohol and petroleum ether before use.</li> <li>Distilled.</li> </ol>
Ph. D. thesis (ref. 2).	ESTIMATED ERRORS:
	Solubility: $\pm$ 8.0%. Temperature: $\pm$ 0.02 K.
	REFERENCES:
	<ol> <li>Gross, P. M. J. Am. Chem. Soc. <u>1929</u>, 51, 2362.</li> <li>Saylor, J. H. Ph. D. thesis, Duke University, Durham, <u>1930</u>.</li> </ol>
COMPONENTS	ODICINAL MEASUDEMENTS.
(1) Tetrahromomethane (Carbon tetrahromide):	Van Arkel A. E.: Vies S. E.
(1) retrationentiale (Carbon Etrationide), CBr <sub>4</sub> ; [558-13-4]	Recl. Tray. Chim. Pays-Bas 1936, 55, 407-11.
(2) Water; H <sub>2</sub> O; [7732-18-5]	
VARIABLES:	PREPARED BY:
T/K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C m <sub>i</sub> /mol kg <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$30 7.0 \times 10^{-4}$	$2.32 \times 10^{-2}$ 1.26
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
No details are available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).

COMPONENTS:	EVALUATOR:
<ol> <li>Trichlorofluoromethane; CCl<sub>3</sub>F; [75-69-4]; R-11</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	January 1993.

#### **CRITICAL EVALUATION:**

The trichlorofluoromethane (1) and water (2) binary system is discussed in two parts; part 1 is trichlorofluoromethane (1) in water (2) and part 2 is water (2) in trichlorofluoromethane (1).

**Part 1.** The solubility of trichlorofluoromethane (1) in water (2) has been studied by at least eleven different work groups with only partial agreement of results. The data of Howe *et al.* (ref. 1) are substantially lower than other measurements and are therefore rejected. The solubility data reported by du Pont (refs. 2 and 3), Rauws *et al.* (ref. 4), McConnell *et al.* (ref. 5), Pearson and McConnell (ref. 6), and Sukornick (ref. 7) show some consistency, while the measurements reported by Park *et al.* (ref. 8), Warner and Weiss (ref. 9), Wisegarver and Cline (ref. 10), Warner et al. (ref. 11), Balls (ref. 14), and Zeininger (ref. 15) are relatively high in value.

It may be assumed that the solubility data reported in the du Pont bulletins were produced in the same laboratories with similar methods and accuracy at 298.15 and 304.15 K, respectively. There appears to be no significant difference between the two data points despite the 6 K difference in temperature. Similarly, the two measurements reported by McConnell et al. and Pearson and McConnell in 1975 most likely were made in the same laboratories under much the same conditions.

There is no obvious reason to explain the difference between the lower (refs. 2, 3, 5, 6, and 7) and higher (refs. 8, 9, 10, 11, 14, and 15) sets of data. In the opinion of the evaluator, there is not enough evidence to justify rejection of measurements from either of the two data sets. Although both data sets are classified as tentative, the smoothed solubility values in mass per cent (1) for the 273 through 313 K temperature interval (Table 1) are based upon a regression equation for all data shown in Figure 1:

Solubility  $[100 w_l] = 25.0094 - 0.16263 (T/K) + 2.6547 \times 10^4 (T/K)^2$ 

with a standard error of 5.0 x  $10^{-2}$  about the regression line.

A minimum solubility occurs for most halogenated hydrocarbons in water systems between 270 and 310 K (refs. 12 and 13). A minimum solubility temperature can be calculated from an enthalpy of solution at infinite dilution and a change in solute heat capacity in going from a pure liquid to an infinitely dilute aqueous solution. However, such thermodynamic data are rarely available in the literature.

**Part 2.** The solubility of water (2) in trichlorofluoromethane (1) has been measured by three research groups (refs. 2, 3, and 7) with only partially consistent results with the exception of the measurements of Sukornick (ref. 7). Generally, the solubility behavior of water in halogenated hydrocarbons shows a definite increase with increasing temperature. This trend has been demonstrated in a comprehensive compilation of solubility data (ref. 12). The two measurements at 294.15 and 298.15 K from the du Pont laboratories compare favorably with the recommended values, whereas the reported value of Sukornick (ref. 7) at 304.15 K is low and classified as doubtful. More accurate solubility measurements are required for more reliable values.

(continued next page)



	co	MPONENTS:	EVALUATOR:
	(1)	Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
	(2)	Water; H <sub>2</sub> O; [7732-18-5]	January 1993.
	CRI	TICAL EVALUATION: (continued)	
	REI	FERENCES:	
1	1.	<ol> <li>Howe, G. B.; Mullins, M. E.; Rogers, T. N. AFESC Tyndall Air Force Base, Report ESL-TR-86-66, Vol. 1, Florida, <u>Sept. 1987</u>, 86 pp. (AD-A188 517).</li> </ol>	
1	2.	du Pont de Nemours & Company, Solubility Rela Tech. Bull. B-7, Wilmington, Del., <u>1966</u> , 16 pp.	ntionship of the Freon Fluorocarbon Compounds,
-	3.	du Pont de Nemours & Company, <i>Solubility Rela</i> Bull. B-43, Wilmington, Del., <u>1966</u> , 17 pp.	ationships between Fluorocarbons and Water, Tech.
i,	4.	Rauws, A. G.; Olling, M.; Wibowo, A. E. J. P.	harm. Pharmac. <u>1973</u> , 25, 718.
7	5.	McConnell, G.; Ferguson, D. M.; Pearson, C. R	. Endeavour <u>1975</u> , 34, 13.
15	6.	Pearson, C. R.; McConnell, G. Proc. Roy. Soc.	B. <u>1975</u> , <i>189</i> , 305.
	7.	Sukornick, B. Inter. J. Thermophys. 1989, 10, 52	53.
,	8.	Park, T.; Rettich, T. R.; Battino, R.; Peterson, D 324.	.; Wilhelm, E. J. Chem. Eng. Data <u>1982</u> , 27,
* 4	9.	Warner, M. J.; Weiss, R. F. Deep-Sea Res. 1983	5, 32, 1485.
12 3	10.	Wisegarver, D. P.; Cline, J. D. Deep-Sea Res. 1	<u>985,</u> <i>32</i> , 97.
2	11.	Warner, H. P.; Cohen, J. M.; Ireland, J. C. Det Priority Pollutants, EPA Technical Report PB87-2	ermination of Henry's Law Constants of Selected 12684, Cincinnati, OH., July 1987.
	12.	Horvath, A. L. Halogenated Hydrocarbons: Solu. New York, <u>1982</u> , 889 pp.	bility-Miscibility with Water. Marcel Dekker, Inc.,
	13.	Privalov, P. L.; Gill, S. J. Pure & Appl. Chem.	<u>1989, 61, 1097.</u>
۲. ۲	14.	Balls, P. W. Ph. D. Thesis, Univ. of East Anglia	, Norwich, U. K., <u>July 1980</u> .
, ·	15.	Zeininger, H. Hoechst A. G., Internat Report, 19	<u>75</u> .

<u>(</u> +

' (٫۱

5

: : \*

6	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	du Pont de Nemours & Company,
(2) Water; $H_2O$ ; [7732-18-5]	Solubility Relationship of the Freon Fluorocarbon Compounds, Tech. Bull. B-7, Wilmington, Del., <u>1966</u> , 16 pp.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
<i>t/</i> °C 100 <i>w<sub>i</sub></i> 10 (con	$x_1^{4} x_1 = 100 w_2 = 10^{4} x_2$ npiler) (compiler)
25 0.11 1.	44 $1.1 \times 10^{-2}$ 8.38
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available. The original report is not obtainable; therefore, the solubility data were taken from a secondary	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
source (ref. 1).	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).
	REFERENCES:
	<ol> <li>Saunders, R. A. Handbook of Aerosol Technology, 2<sup>nd</sup> ed., Van Nostrand Reinhold Co., New York, <u>1979</u>, p. 184.</li> </ol>
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	du Pont de Nemours & Company,
(2) Water; $H_2O$ ; [7732-18-5]	Solubility Relationships between Fluorocarbons and Water, Tech. Bull. B-43, Wilmington, Del., <u>1966</u> , 17 pp.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 294 - 304	A. L. Horvath
EVDEDIMENTAL VALUES.	
$\frac{t^{\circ}C}{t^{\circ}} = 100 w_{i} \qquad 10$	$x_1 = 100 w_2 = 10^4 x_2$ piler) (compiler)
21 - 31 0.106 1.	9.0 × 10 <sup>-3</sup> 6.86
AUXILIARY INFORMATION	
AUXILIARY I	NFORMATION
AUXILIARY I METHOD/APPARATUS/PROCEDURE:	NFORMATION SOURCE AND PURITY OF MATERIALS:
AUXILIARY I METHOD/APPARATUS/PROCEDURE: Details are not available. The original report is not obtainable; therefore,	NFORMATION SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Distilled (compiler).
AUXILIARY I METHOD/APPARATUS/PROCEDURE: Details are not available. The original report is not obtainable; therefore, the solubility data were taken from a secondary source (ref. 1).	NFORMATION SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Distilled (compiler). ESTIMATED ERRORS:
AUXILIARY I METHOD/APPARATUS/PROCEDURE: Details are not available. The original report is not obtainable; therefore, the solubility data were taken from a secondary source (ref. 1).	NFORMATION SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Distilled (compiler). ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 1 K (compiler).
AUXILIARY I METHOD/APPARATUS/PROCEDURE: Details are not available. The original report is not obtainable; therefore, the solubility data were taken from a secondary source (ref. 1).	NFORMATION SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Distilled (compiler). ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 1 K (compiler). REFERENCES:

•

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Rauws, A. G.; Olling, M.; Wibowo, A. E.	
[75-69-5]; R-11	J. Pharm. Pharmac. 1973. 25. 718-22.	
(2) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXDEDIMENTAL VALUES.		
	104 x 100 w <i>M</i> -//mol g-1	
	(compiler) (compiler)	
25 0.11	1.44 $8.0 \times 10^{-4}$	
AUXILIARY I	NFORMATION	
	SOLIDCE AND DUDITY OF MATERIALS.	
A calibrated bottle was filled with nitrogen	(1) Source and purity not given	
measured quantity of water, and the organic vapor. After mixing and equilibration, the concentrations of	(2) Distilled (compiler).	
injection of the head space samples into a gas	ESTIMATED ERRORS:	
by means of a tritium foil electron capture detector.	Solubility: Not specified.	
	Temperature: $\pm 0.5 \text{ K}$ (complet).	
-		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F;	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R.	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5]	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, <u>1975</u> , 34, 13-8.	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES:	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, <u>1975</u> , 34, 13-8. PREPARED BY:	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 293	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8. PREPARED BY: A. L. Horvath	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: 77K = 293 EXDEDIMENTAL VALUES:	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8. PREPARED BY: A. L. Horvath	
COMPONENTS: (1) Trichlorofluoromethane; $CCl_3F$ ; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: 7/K = 293 EXPERIMENTAL VALUES: $t^{0}C$ $10^6 c_1/c_2$	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. <i>Endeavour</i> , <u>1975</u> , <i>34</i> , 13-8. PREPARED BY: A. L. Horvath	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $10^{6} g_{1}/g_{2}$	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8. PREPARED BY: A. L. Horvath $100 w_i$ $10^4 x_i$ (compiler) $10^4 x_i$	
COMPONENTS:         (1) Trichlorofluoromethane; CCl <sub>3</sub> F;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t/^{\circ}$ C $10^{6} g_{1}/g_{2}$ 20       1100	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 0.1099 $1.443$	
COMPONENTS:         (1) Trichlorofluoromethane; $CCl_3F$ ;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $10^{6} g_{1}/g_{2}$ 20       1100	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.PREPARED BY: A. L. Horvath $100 w,$ (compiler) 0.1099 $1.443$	
COMPONENTS:         (1) Trichlor of luoromethane; CCl <sub>3</sub> F;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}$ C         10 <sup>6</sup> g <sub>1</sub> /g <sub>2</sub> 20	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 0.1099 $104 x_i$ (compiler) 1.443	
COMPONENTS:         (1) Trichlorofluoromethane; CCl <sub>3</sub> F;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}$ C         10 <sup>6</sup> g <sub>1</sub> /g <sub>2</sub> 20	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.PREPARED BY: A. L. Horvath $100 w,$ (compiler) 0.1099 $1.443$	
COMPONENTS:         (1) Trichlorofluoromethane; CCl <sub>3</sub> F;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}$ C         10 <sup>6</sup> $g_1/g_2$ 20         1100	ORIGINAL MEASUREMENTS: McConnell, G.; Ferguson, D. M.; Pearson, C. R. Endeavour, 1975, 34, 13-8.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 0.1099 $104 x_i$ (compiler) 0.1099 $1.443$	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>6</sup> g <sub>1</sub> /g <sub>2</sub> 20 1100 AUXILIARY II METHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS:         McConnell, G.; Ferguson, D. M.; Pearson, C. R.         Endeavour, 1975, 34, 13-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         0.1099         1.443	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: 7/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $10^{6} g_{1}/g_{2}$ 20 1100 AUXILIARY II METHOD/APPARATUS/PROCEDURE: The solubility of the trichlorofluoromethane in water was determined by gas-liquid chromatography (GLC) using an electron canture detector. Where non-	ORIGINAL MEASUREMENTS:         McConnell, G.; Ferguson, D. M.; Pearson, C. R.         Endeavour, 1975, 34, 13-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         0.1099         1.443         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) Distilled (compiler).	
COMPONENTS:         (1) Trichlorofluoromethane; CCl <sub>3</sub> F;         [75-69-4]; R-11         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}$ C         10 <sup>6</sup> g <sub>1</sub> /g <sub>2</sub> 20         1100         AUXILIARY II         METHOD/APPARATUS/PROCEDURE:         The solubility of the trichlorofluoromethane in water was determined by gas-liquid chromatography (GLC) using an electron capture detector. Where possible, identification was confirmed by a linked mass-spectrometer (MS).	ORIGINAL MEASUREMENTS:         McConnell, G.; Ferguson, D. M.; Pearson, C. R.         Endeavour, 1975, 34, 13-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         0.1099         1.443         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) Distilled (compiler).	
COMPONENTS: (1) Trichlorofluoromethane; CCl <sub>3</sub> F; [75-69-4]; R-11 (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>6</sup> g <sub>1</sub> /g <sub>2</sub> 20 1100 AUXILIARY II METHOD/APPARATUS/PROCEDURE: The solubility of the trichlorofluoromethane in water was determined by gas-liquid chromatography (GLC) using an electron capture detector. Where pos- sible, identification was confirmed by a linked mass- spectrometer (MS).	ORIGINAL MEASUREMENTS:         McConnell, G.; Ferguson, D. M.; Pearson, C. R.         Endeavour, 1975, 34, 13-8.         PREPARED BY:         A. L. Horvath         100 w, 10 <sup>4</sup> x, (compiler)         0.1099         1.443         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) Distilled (compiler).         ESTIMATED ERRORS:         Solubility:	

Ĭ

8		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Pearson, C. R.; McConnell, G.	
[/3-09-4]; K-11 (2) Water: H O: [7722.18_5]	Proc. Roy. Soc. B <u>1975</u> , 189, 305-32.	
(2) water, n <sub>2</sub> O, [//32-16-5]		
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 10 <sup>6</sup> ₩,	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 1100	0.110 1.44	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Saturated solutions were prepared in a constant temperature thermostat bath. Water samples were ex- tracted with n-pentane and an aliquot of the extract	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
taken for gas-liquid chromatographic analysis. The gas chromatograph was fitted with a <sup>53</sup> Ni electron	ESTIMATED ERRORS:	
capture detector.	Solubility: Not specified.	
	Temperature. $\pm 0.1 \text{ K}$ (complet).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Zeininger, H.	
[75-69-4]; R-11	Hoechst A. G., Internat Report, 1975.	
(2) water, $n_20$ ; [7732-16-5]		
VARIABLES:	PREPARED BY:	
T/K = 273 - 313	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^{4} x_{i} & 100 \ w_{i} M_{j}^{-i} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
0 0.37 10 0.25	$\begin{array}{ccc} 4.87 & 2.69 \times 10^{-3} \\ 3.29 & 1.82 \times 10^{-3} \end{array}$	
20 0.16 40 0.070	$\begin{array}{cccc} 2.10 & 1.17 \times 10^{-3} \\ 0.919 & 5.10 \times 10^{-4} \end{array}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE	SOURCE AND PURITY OF MATERIALS	
The organic compound was condensed into a	(1) Source and purity not given.	
degassed liquid water sample up to saturation at which two phases appeared. After shaking the mix-	(2) Distilled (compiler).	
ture for several hours, equilibrium was achieved and the respective phases were analysed in a gas chro-	ESTIMATED ERRORS:	
matograph. The original report is not obtainable; therefore,	Solubility: Not specified.	
the details were taken from junge (ref. 1).	1 emperature: ± 1 K (compiler).	
	REFERENCES:	
	(1) Junge, C. Z. Naturforsch. <u>1976</u> , 31a, 482.	

	· · · · ·	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Balls, P. W.	
(2) Water: H <sub>-</sub> O: [7732-18-5]	Ph. D. Thesis, Univ. of East Anglia, Norwich, U. K. July 1980, 375 pp	
(-)	····, <u>zoij 1200</u> , 575 pp.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:	I	
t/°C Henry's law constant.	$100 w$ , $10^4 x$ .	
H/dimensionless	(compiler) (compiler)	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.137 1.79 0.136 1.79	
$20$ $3.40 \pm 0.21$	0.147 1.93	
$25  3.60 \pm 0.22$	0.163 2.14	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The method involved repeated equilibrations of	(1) Source and purity not given.	
nitrogen with a liquid water sample containing the trichlorofluoromethane. Each sample was shaken	(2) Distilled.	
temperature bath for 30 minutes. The equilibrated gas	ESTIMATED EPROPS	
phase was then injected into a gas chromatograph. The experiments were performed at least three times		
at 10, 15, 20, and 25 °C; each experiment involved	Solubility: See above. Temperature: + 0.5 K (compiler).	
	• • • • •	
COMPONENTS	OPIGINAL MEASUREMENTS.	
(1) Trichlorofluoromethane: CCLF:	Park T $\cdot$ Rettich T R $\cdot$ Ratting R $\cdot$ Peterson D $\cdot$	
[75-69-4]; R-11	Wilhelm, E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chem. Eng. Data <u>1982</u> , 27, 324-6.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Bunsen coefficient.	$100 w_1$ $10^4 x_1$	
a/dimensionless	(compiler) (compiler)	
10 0.2491	0.1602 2.104	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The Bunsen coefficient was measured in a	(1) Matheson Gas Co., minimum purity 99.9 mole	
Scholander microgasometric apparatus which was contained in an air bath. After the distilled water was decased, the trichlorodiucromethane sample was in-	(2) Distilled (reverse osmosis), passed through ion-	
troduced into the apparatus. About 2 hours were re-	exemunge commina.	
	ESTIMATED ERRORS	
ure for the calculation of the Bunsen coefficient is	ESTIMATED ERRORS:	
ure for the calculation of the Bunsen coefficient is described by Douglas (ref. 1).	ESTIMATED ERRORS: Solubility: ± 1.7 std. dev. Temperature: ± 0.05 K.	
described by Douglas (ref. 1).	ESTIMATED ERRORS: Solubility: ± 1.7 std. dev. Temperature: ± 0.05 K.	
ure for the calculation of the Bunsen coefficient is described by Douglas (ref. 1).	ESTIMATED ERRORS: Solubility: ± 1.7 std. dev. Temperature: ± 0.05 K. REFERENCES: (1) Develop E. J. Diver Cham. 10(4, 69, 100)	

ŀ

10		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F; 175-69-21: R-11	Wisegarver, D. P.; Cline, J. D.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Deep-Sea Res. <u>1985</u> , 32, 97-106.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	<u> </u>	
t/°C Bunsen coefficient, α/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
25 0.246	0.158 2.08	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibration took place in a 2 liter flask rotating in a constant temperature bath for 20 hours. The analysis of CCI,F was carried out using a	<ol> <li>AIRCO, Inc., used as received.</li> <li>Distilled.</li> </ol>	
mounted in a gas chromatograph. Peak areas were cupatified by a HP 3388A integrator and compared to	ESTIMATED ERRORS:	
the detector response of a standard sample.	Solubility: $\pm 2.0$ %. Temperature: $\pm 0.05$ K.	
(1) Tricklosoftworomethanet, CCI Fr	ORIGINAL MEASUREMENTS:	
[175-69-4]; R-11	Deen-Sea Res. 1985. 32. 1485-97.	
(2) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 274 - 313.95	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 10 <sup>2</sup> c <sub>1</sub> P <sub>1</sub> <sup>-1</sup> /mol dm <sup>-3</sup> atm <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
0.85 3.7221 0.85 3.6555	0.511 6.73 0.502 6.61	
0.85 3.6636 0.85 3.6818	0.503 6.60 0.506 6.66	
	(continued)	
	(continues)	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibration of a dilute mixture of tri- chlorofluoromethane gas with water took place in an	<ol> <li>Source and purity not given.</li> <li>Deionized.</li> </ol>	
equilibration chamber at various temperatures. The	ESTIMATED ERRORS:	
were measured by gas chromatography using a	ESTIMATED ERRORS:	

				1
COMPONENTS:		ORIGINAL MEAS	UREMENTS:	
(1) Trichlorofluoromethane; [75-69-4]: R-11	CCl <sub>3F</sub> ;	Warner, M. J.; We	eiss, R. F.	
(2) Water: $H_2O$ : [7732-18-5]		Deep-Sea Res. <u>198</u>	5, 32, 1485-97.	
VARIABLES:		PREPARED BY:		
T/K = 274 - 313.95		A. L. Horvath		
EXPERIMENTAL VALUES	: (continued)			
t/°C	10 <sup>2</sup> c, P, <sup>-1</sup> /mol dm <sup>-3</sup> atm <sup>-1</sup>	100 w, (compiler)	$10^4 x_1$ (compiler)	
$\begin{array}{c} 0.85\\ 0.85\\ 9.58\\ 9.59\\ 9.59\\ 9.59\\ 9.59\\ 9.59\\ 19.82\\ 19.87\\ 19.89\\ 19.89\\ 19.93\\ 29.87\\ 29.87\\ 29.87\\ 29.87\\ 29.87\\ 29.87\\ 29.87\\ 31.79\\ 31.79\\ 31.79\\ 31.79\\ 31.89\\ 31.$	$\begin{array}{c} 3.6561\\ 3.6371\\ 2.1092\\ 2.1600\\ 2.1532\\ 2.1498\\ 2.1760\\ 2.1647\\ 1.2561\\ 1.2646\\ 1.2756\\ 1.2707\\ 1.2785\\ 1.2785\\ 0.8697\\ 0.8783\\ 0.8715\\ 0.8712\\ 0.8645\\ 0.8223\\ 0.8314\\ 0.8543\\ 0.8222\\ 0.8129\\ 0.8240\\ 0.8263\\ 0.6180\\ 0.6282\\ 0.5979\\ 0.6130\\ 0.6048\\ 0.6060\\ \end{array}$	0.502 0.500 0.290 0.297 0.296 0.295 0.295 0.174 0.171 0.176 0.175 0.176 0.176 0.120 0.120 0.121 0.120 0.120 0.120 0.120 0.120 0.120 0.120 0.120 0.120 0.121 0.120 0.120 0.120 0.121 0.120 0.121 0.120 0.121 0.120 0.121 0.120 0.121 0.120 0.121 0.120 0.120 0.120 0.120 0.120 0.120 0.114 0.114 0.114 0.114 0.0856 0.0870 0.0828 0.0837 0.0839	$\begin{array}{c} 6.62\\ 6.58\\ 3.81\\ 3.90\\ 3.89\\ 3.88\\ 3.93\\ 3.88\\ 2.28\\ 2.25\\ 2.30\\ 2.30\\ 2.30\\ 2.31\\ 2.31\\ 1.58\\ 1.59\\ 1.58\\ 1.58\\ 1.58\\ 1.58\\ 1.58\\ 1.50\\ 1.52\\ 1.51\\ 1.55\\ 1.49\\ 1.48\\ 1.50\\ 1.50\\ 1.50\\ 1.50\\ 1.50\\ 1.50\\ 1.12\\ 1.14\\ 1.09\\ 1.11\\ 1.10\\$	

Ŧ

12	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Warner, H. P.; Cohen, J. M.; Ireland, J. C.
(2) Water; $H_2O$ ; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants, EPA Tech. Rept. PB87- 212684, Cincinatti, OH., July 1987.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	1
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ (\text{compiler}) & (\text{compiler}) \end{array}$
24.85 5.83 $\times$ 10 <sup>-2</sup>	0.1969 2.586
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The original method and apparatus for the de- termination of Henry's law constants are described in Mackay <i>et al.</i> (ref. 1). The general procedure was to	<ol> <li>Purest quality available and used without further purification; stated purity &gt;99 %.</li> <li>Distilled deionized.</li> </ol>
deionized water and stir the mixture overnight. A portion of this solution was returned to a stripping	ESTIMATED ERRORS:
vessel. The solute was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from a log of the concentra-	Solubility: $\pm 6 \%$ std. dev. Temperature: $\pm 0.05$ K.
tion versus time plot. The experimental data points are averages of two or more replicates.	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Techn. <u>1979</u> , 13, 333.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Howe, G. B.; Mullins, M. E.; Rogers, T. N.
(2) Water; $H_2O$ ; [7732-18-5]	AFESC Tyndall Air Force Base, Rept. ESL-TR-86- 66, Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp. (AD- A188 571).
VARIABLES:	PREPARED BY:
T/K = 283 - 303	A. L. Horvath
EXPERIMENTAL VALUES:	1
$t/^{\circ}$ C 10 <sup>6</sup> $w_{1}$	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
10 541 20 341 30 262	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
250 ml bottles were filled with distilled de- ionized water and sealed. Measured volumes of tri- chlorofluoromethane were injected into the bottles through each bottle septum using a microliter syr- inge. The solute was in excess of the anticipated	<ol> <li>Probably a commercial reagent, at least 99 % purity, used as received.</li> <li>Distilled deionized.</li> </ol>
solubility limit. The bottles were shaken for 1 hour with a wrist-action shaker and allowed to equilibrate for about 3 weeks. Samples were then injected into a gas chromatograph equipped with a Carbopack col- umn and a FID detector. The GC responses were compared with a calibration plot to establish concen- trations.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Sukornick, B.	
[/3-69-4]; K-11 (2) Water: H O: (7722-18-5)	Inter. J. Thermophys. <u>1989</u> , 10, 553-61.	
(2) water, H <sub>2</sub> O; [//32-16-3]		
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub> 10 (con	$4x_1$ 100 $w_2$ 104 $x_2$ ppiler) (compiler)	
25 9.50 $\times$ 10 <sup>-2</sup> 1.1	$25 7.0 \times 10^{-3} 5.34$	
	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ul> <li>(1) Amed-Signal Inc., Burlato, N. T., purity not given.</li> <li>(2) Distilled (compiler)</li> </ul>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: ± 1 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Zeininger, H.	
(2) Seawater	Hoechst A. G., Internat Report, 1975.	
VARIABLES:	PREPARED BY:	
77K = 276 - 313	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^4 x_1 & 100 \ w_l M_l^{-l} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
3 0.19 10 0.12	$\begin{array}{cccc} 2.50 & 1.38 \times 10^{-3} \\ 1.58 & 8.74 \times 10^{-4} \end{array}$	
20 0.09 40 0.05	$\begin{array}{cccc} 1.18 & 6.55 \times 10^{4} \\ 0.656 & 3.64 \times 10^{4} \end{array}$	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Degassed liquid trichlorofluoromethane samples were introduced into degassed seawater samples up to saturation at which point two phases appeared. After	<ol> <li>Source and purity not given.</li> <li>North Atlantic surface water, salinity not given.</li> </ol>	
snaking a sample for several nours equilibrium was achieved and the respective phases were analysed with a gas liquid chromatograph	ESTIMATED ERRORS:	
The original report is not obtainable; therefore, the details were taken from Junge (ref. 1).	Solubility: Not specified. Temperature: $\pm 1$ K (compiler).	
	REFERENCES:	
	(1) Junge, C. Z. Naturforsch. <u>1976</u> , 31a, 482.	

14		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Balls, P. W.	
(2) Seawater	Ph. D. Thesis, Univ. of East Anglia, Norwich, U. K., July 1980, 375 pp.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.1301.710.1291.700.1301.710.1451.90	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The method involved repeated equilibrations of nitrogen with a water sample containing trichloro- fluoromrthane. The sample was shaken vigorously at the required temperature in a constant temperature	<ol> <li>Source and purity not given.</li> <li>Salinity = 20 ‰.</li> </ol>	
bath for 30 minutes. The equilibrated gas phase was then injected into a gas chromatograph. The experi-	ESTIMATED ERRORS:	
20, and 25 °C with each experiment involving about six equilibrations.	Solubility: See above. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS	ODICINAL MEASUDEMENTS.	
(1) Trichlorofluoromethane: CCLE:	Wisegarver D. P. Cline I. D.	
[75-69-4]; R-11	Deep-Sea Res. 1985, 32, 97-106.	
(2) Seawater		
VARIABLES:	PREPARED BY:	
T/K = 273 - 305	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Bunsen coefficient, α/dimensionless	t/°C Bunsen coefficient, α/dimensionless	
$\begin{array}{cccccc} 0.0 & 0.603 \pm 0.024 & (33) \\ 2.6 & 0.517 \pm 0.010 & (8) \\ 3.5 & 0.493 \pm 0.008 & (7) \\ 5.2 & 0.451 \pm 0.007 & (6) \end{array}$	5.3 0.448 $\pm$ 0.005 (11) 5.4 0.460 $\pm$ 0.010 (7) 7.0 0.402 $\pm$ 0.004 (14) 7.2 0.415 $\pm$ 0.012 (9)	
	(continued)	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibration occurred in a 2 liter flask ro- tating in a constant temperature water bath for 20 hours. The analysis of CCl <sub>3</sub> F was carried out using a Howlett Packace <sup>62</sup> Ni electron conture detector	<ol> <li>AIRCO, Inc., used as received.</li> <li>Collected from Puget Sound, Washington with salinity of 35.0x10<sup>-3</sup>.</li> </ol>	
mounted in a gas chromatograph. Peak areas were ulantified by a HP 3388A integrator and compared to	ESTIMATED ERRORS:	
the detector response of a standard sample.	Solubility: $\pm 2.0 \%$ . Temperature: $\pm 0.5 K$ .	

	15	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Wisegarver, D. P.; Cline, J. D.	
(2) Seawater	Deep-Sea Res. <u>1985</u> , 32, 97-106.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 305	A. L. Horvath	
EXPERIMENTAL VALUES: (continued)	I	
t/°C Bunsen coefficient, α/dimensionless	t/°C Bunsen coefficient, α/dimensionless	
$\begin{array}{ccccccc} 7.4 & 0.411 \pm 0.001 & (6) \\ 7.9 & 0.391 \pm 0.003 & (5) \\ 10.0 & 0.331 \pm 0.005 & (6) \\ 10.2 & 0.343 \pm 0.004 & (10) \\ 10.4 & 0.344 \pm 0.002 & (7) \\ 10.5 & 0.352 \pm 0.004 & (10) \\ 12.3 & 0.313 \pm 0.003 & (9) \\ 15.0 & 0.276 \pm 0.007 & (10) \\ 15.1 & 0.274 \pm 0.003 & (9) \\ 15.2 & 0.274 \pm 0.001 & (8) \end{array}$	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	
The standard deviations are followed by the number	of observations in parantheses.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichlorofluoromethane; CCl <sub>3</sub> F;	Warner, M. J.; Weiss, R. F.	
(2) Seawater	Deep-Sea Res. <u>1985</u> , 32, 1485-97.	
VARIABLES:	PREPARED BY:	
T/K = 272 - 313	A. L. Horvath	
Concentration		
EXPERIMENTAL VALUES:		
t/°C 10 <sup>2</sup> c, P, - <sup>1</sup> /mol dm <sup>-3'</sup> atm <sup>-1</sup>	$t^{\circ}C$ 10 <sup>2</sup> c, $P_{i}^{-1}$ /mol dm <sup>-3</sup> ath <sup>-1</sup> a t m	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	4.79 1.9601 4.79 1.9607 4.81 1.8770 4.82 1.9773 4.84 1.9318 9.19 1.5302 9.19 1.5293 9.19 1.5330 9.19 1.5372 9.19 1.5727 (continued)	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Equilibration of a dilute mixture of trichloro- fluoromethane gas with seawater occurred in an equilibration chamber at various temperatures. Both gas and liquid phase concentrations were measured by gas chromatography using a Shimadzu <sup>63</sup> Ni electron	<ol> <li>Source and purity not given.</li> <li>Surface water collected at La Jolla. Salinity = 35.807 ± 0.006 × 10<sup>-3</sup>.</li> </ol>	
capture detector operating in a constant current mode.	ESTIMATED ERRORS:	
	Solubility: $\pm 1.5 \%$ . Temperature: $\pm 0.02 K$ .	

COMPONENTS:	ORIGINAL MEASUREMENTS:
<ol> <li>Trichlorofluoromethane; CCl<sub>3</sub>F; [75-69-4]; R-11</li> <li>Seawater</li> </ol>	Warner, M. J.; Weiss, R. F. Deep-Sea Res. <u>1985</u> , 32, 1485-97.
VARIABLES:	PREPARED BY:
T/K = 272 - 313	A. L. Horvath
Concentration	

EXPERIMENTAL VALUES: (continued)

t/⁰C	$10^2 c_1 P_1^{-1}$ /atm mol dm <sup>-3</sup>	t/°C	$10^2 c_1 P_1^{-1}$ /atm mol dm <sup>-3</sup>
9.19	1.5404	29.55	1.6182
20.43	0.8648	29.55	0.6042
20.44	0.8868	29.55	0.6037
20.44	0.8792	29.55	0.6020
20.44	0.8749	39.90	0.4280
20.44	0.8788	39.91	0.4301
20.44	0.8780	39.93	0.4299
29 53	0.6085	39.93	0.4357
29.53	0.6031	39.93	0.4281

COMPONENTS:	EVALUATOR:
(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	January 1993.

#### CRITICAL EVALUATION:

The tetrachloromethane (1) and water (2) binary system is discussed in two parts; part 1 is tetrachloromethane (1) in water (2) and part 2 is water (2) in tetrachloromethane (1).

**Part 1.** The solubility of tetrachloromethane (1) in water (2) has been studied by 41 workers. The experimental data of most investigators in the 273 to 303 K temperature interval are sufficiently reliable to use in a smoothing equation. Even though a number of measurements above 308 K were reported, the solubility values in this temperature range are rather uncertain and are classified as doubtful.

The experimental work of several workers was not used for the smoothing equation for a variety of reasons. The measured solubility of Booth and Everson (refs. 1 and 2) and of Karger *et al.* (ref. 3) are markedly higher than the solubility values calculated from the smoothing equation. The measurements by Powell (ref. 4), Tettamanti *et al.* (ref. 5), Svetlanov *et al.* (ref. 6), Antropov *et al.* (ref. 7), Simonov *et al.* (ref. 8), Sato and Nakijima (ref. 9), Coca *et al.* (ref. 10), Balls (ref. 11), Yoshioka *et al.* (ref. 12), Howe *et al.* (ref. 13), and Wright *et al.* (ref. 14) are all several per cents lower than the smoothed solubility values and have also been rejected. The solubilities calculated from the distribution coefficients of Prosyanov *et al.* (ref. 15) are in very poor agreement, giving little confidence in their values which were regarded as dubious.

The remaining data from 26 laboratories were compiled or used for the smoothing equation. The data are also shown in Figure 1. The fitting equation obtained for mass per cent (1) is given by:

Solubility  $[100 w_1] = 3.4653 - 0.0230285 (T/K) + 3.91621 \times 10^{-5} (T/K)^2$ 

The equation represents the combined data points which yield a standard deviation of  $5.8 \times 10^{-3}$  in the 273 to 308 K temperature range.

The curve obtained from the smoothing equation shows a distinct minimum at 294 K, as seen in Figure 1. The existence of the minimum solubility is discussed in the Preface.

The recommended solubility values at 5 K intervals for tetrachloromethane (1) in water (2) are presented in Table 1.

Table 1. Tentative Solubility of Tetrachloromethane (1) in Water (2)

Temperature		Solubility	
°C	K	100 w <sub>1</sub>	$10^4 x_l$
0	273.15	0.0970	1.137
5	278.15	0.0898	1.053
10	283.15	0.0846	0.9916
15	288.15	0.0813	0.9529
20	293.15	0.0800	0.9376
25	298.15	0.0813	0.9529
30	303.15	0.0832	0.9752
35	308.15	0.0878	1.029

(continued next page)


Part 2. The solubility of water (2) in tetrachloromethane (1) has been reported by 37 laboratories since 1921. The solubility measurements cover the 273 to 373 K temperature range, see Figure 2. However, the data available above 323 K originate from only two investigators (refs. 16 and 17). The recommended solubility values are limited to the 283 to 307 K temperature interval.

Although data from all the workers show a general increase in solubility with temperature, the extent of the increase is variable above 313 K.

The data of Bell (ref. 18), Niini (ref. 19), and Fox and Martin (ref. 20) are substantially lower than all other studies and they have been rejected. The data of Eberius (ref. 21), Zielinski (ref. 22), Grigsby (ref. 23), Goldman (ref. 24), Ohtsuka and Kazama (ref. 25), and Kleeberg et al. (ref. 26) are significantly higher than the likely solubility and were also rejected. The remaining data mainly due to Clifford (ref. 27), Rosenbaum and Walton (ref. 28), Staverman (ref. 29), Hutchison and Lyon (ref. 30), McGovern (ref. 31), Donahue and Bartell (ref. 32), Rotariu et al. (ref. 33), Greinacher et al. (ref. 34), Desnoyer (ref. 35), Tettamanti et al. (ref. 5), Johnson et al. (refs. 36 and 37), Johnson (ref. 38), Christian et al. (ref. 39), Högfeldt and Fredlund (ref. 41), Simonov et al. (refs. 8, 41 and 42), Glasoe and Schultz (ref. 43), Antropov et al. (ref. 7), Kirchnerova (ref. 44), Kirchnerova and Cave (ref. 45), Coca et al. (ref. 10), Ksiazczak and Buchowski (ref. 46), Wu (ref. 17), and Orlandini et al. (ref. 47) are in good agreement, especially the values given in (refs. 31, 43, and 17).

The correlating equation for the combined mass per cent solubilities of water in tetrachloromethane between 283 and 308 K is as follows:

Solubility  $[100 w_2] = 0.301104 - 2.348078 \times 10^{-3} (T/K) + 4.607143 \times 10^{-6} (T/K)^2$ 

This equation yielded a standard deviation of  $1.5 \times 10^3$  in the 273 to 308 K temperature range.

(continued next page)

COMPONENTS:		EVALUATOR:
(1) Tetrachloromethane (car CCl <sub>4</sub> ; [56-23-5]	bon tetrachloride);	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; $H_2O$ ; [7732-18-:	5]	January 1993.
CRITICAL EVALUATION:	(continued)	
The recommended mass per operation of the presented in the Table 2.	cent solubility values at 5	K intervals for water in tetrachloromethane are
Table 2. Re	commended Solubility of	Water (2) in Tetrachloromethane (1)
Ten	nperature S	olubility
°C	K $100 w_2$	$10^4 x_2$
10 15 20 25 30 35	283.15         0.00558           288.15         0.00699           293.15         0.00864           298.15         0.01051           303.15         0.01261           308.15         0.01495	4.762 5.965 7.372 8.967 10.757 12.750
0 045	· · · ·	<del>, , , ,</del>
0 040	-	À
+ 0 035	-	<b>/</b>
U 0 030	-	0 0 -
	_	
	_	
	0	80
<b>3</b> 0 0 13	88	
- 0010		-
0 005	200	-
0 000 2	70 280 290 300 3	i 10 320 330 340 350
	Т	/ К
Figu	re 2. Solubility of Water	(2) in Tetrachloromethane (1).
REFERENCES:		
1. Booth, H. S.; Everson,	H. E. Ind. Eng. Chem.	<u>1948,</u> <i>40</i> , 1491.
2. Booth, H. S.; Everson,	H. E. Ind. Eng. Chem. ]	<u>949</u> , <i>41</i> , 2627.
3. Karger, B. L.; Chatterje Northeastern Univ., Bos	e, A. K.; King, J. W. <i>T</i> ton, Mass., <u>May 10, 197</u>	ech. Rept. No: 3, Dept. of Chemistry, 1.
4. Powell, J. F. Brit. J. In	ndustr. Med. <u>1947</u> , 4, 233	
5. Tettamanti, K.; Nogradi	, M.; Sawinsky, J. Peric	d. Polytech, Chem. Eng. <u>1960</u> , 4, 201.

رد لار

59

36

41

(continued next page)

.

Committee L

19

	20			
	СО	MPONENTS:	EVALUATOR:	
	(1)	Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.	
	(2)	Water; H <sub>2</sub> O; [7732-18-5]	January 1993.	
	CR	ITICAL EVALUATION: (continued)		Ì
	RE	FERENCES:		
51	6.	Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. <i>Phys. Chem.</i> <u>1971</u> , 45, 488.	I.; Treger, Yu. A.; Flid, R. M. Russ. J.	
()	7.	Antropov, L. I.; Pogulyai, V. E.; Simonov, V. E. <u>1971</u> , <i>46</i> (2), 311-312 ( <i>VINIT1</i> No. 3739-71).	D.; Shamsutdinov, T. M. Russ. J. Phys. Chem.	
$\mathbf{G}$	8.	Simonov, V. D.; Shamsutdinov, T. M.; Pogulyai <u>1974</u> , <i>48</i> , 1573.	, V. E.; Popova, L. N. Russ. J. Phys. Chem.	
1.9	9.	Sato, A.; Nakijima, T. Arch. Environ. Health 19	<u>79,</u> <i>34</i> , 69.	
1)	10.	Coca, J.; Diaz, R. M.; Pazos, C. Fluid Phase Ed	quilibr. <u>1980,</u> 4, 125.	
71	11.	Balls, P. W. Ph. D. Thesis, Univ. of East Anglia	a, Norwich, U. K., <u>1980,</u> 375 pp.	
84	12.	Yoshioka, Y.; Ose, Y.; Sato, T. Ecotoxicol. Env	iron. Saf. <u>1986</u> , 12, 15.	
<b>२१</b>	13.	Howe, G. B.; Mullins, M. E.; Rogers, T. N. AF 66, Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp. (AD-A188	ESC Tyndall Air Force Base, Report ESL-TR-86- 8 571).	
91	14.	Wright, D. A.; Sandler, S. I.; DeVoll, D. Envir	on, Sci. Technol. <u>1992</u> , 26, 1828.	
C,	15. Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. KhimTekhnol. Inst. 1973, 183.		670ª CA	
$()^{+}$	16. Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. KhimTekhnol. Inst. 1974, 55.			2004 6
11	17.	Wu, X. Huaxue Shiji <u>1981</u> , 221.		
29	18.	Bell, R. P. J. Chem. Soc. 1932, 2905.		
31	19.	Niini, A. Suomen Kemistilehte 1938, 11a, 19.		
·} <b>`</b>	20.	Fox, J. J.; Martin, A. K. Proc. Roy. Soc. Londo	n, Ser. A, <u>1940</u> , <i>174</i> , 234.	
44	21. Eberius, E. Wasserbestimmung mit Karl-Fischer-Lösung, Verlag Chemie, GMBH, Weinheim, <u>1954</u> , p. 67.			
40	22.	Zielinski, A. Z. Chem. Stosowana 1959, 3, 377.		
72	<ol> <li>Grigsby, R. D. Self-Association and Hydration of N-Methylacetamide in Carbon Tetrachloride, <i>Ph. D. Thesis</i>, Univ. of Oklahoma, Norman, Oklahoma, <u>1966</u>.</li> </ol>			
ر. در	24.	Goldman, S. Can. J. Chem. 1974, 52, 1668.		
30	25.	Ohtsuka, K.; Kazama, K. Sen'i Seihin Shohi Kag	aku Kaishi <u>1982,</u> 22, 197.	
<sub>2</sub> 1	26.	Kleeberg, H.; Klein, D.; Luck, W. A. P. Chem.	-IngTech. <u>1987</u> , 59, 409.	
<b>ว</b> ร	27.	Clifford, C. W. J. Ind. Eng. Chem. 1921, 13, 63	BI. L Convection.	
			(continued next page)	

•

ſ	CO	MPONENTS:	EVALUATOR:
	(1)	Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
	(2)	Water; H <sub>2</sub> O; [7732-18-5]	January 1993.
Ī	CRI	TICAL EVALUATION: (continued)	
	REF	FERENCES:	
	28.	Rosenbaum, C. K.; Walton, J. H. J. Am. Chem.	Soc. <u>1930</u> , 52, 3568.
3	29.	Staverman, A. J. Recl. Trav. Chim. Pays-Bas 19	<u>41,</u> <i>60</i> , 836.
,	30.	Hutchison, C. A.; Lyon, A. M. Columbia Unive	rsity Report A-745, July 1, 1943.
	31.	McGovern, E. W. Ind. Eng. Chem. 1943, 35, 12	230.
·	32.	Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1	<u>952,</u> <i>56</i> , 480.
	33.	Rotariu, G. J.; Fraga, D. W.; Hildebrand, J. H.	J. Am. Chem. Soc. <u>1952</u> , 74, 5783.
3	34.	Greinacher, E.; Lüttke, W.; Mecke, R. Z. Elektr	rochem. <u>1955</u> , 59, 23.
	35.	Desnoyer, M. Dosage de l'Eau dans les Solvants Mesure des Constantes Dielectriques, Centre l'Etu Saclay, June 23, 1959.	or Organiques par Absorption Infra Rouge et ades Nucleaires de Saclay, Report No. 1254,
;	36.	Johnson, J. R.; Christian, S. D.; Affsprung, H. E	3. J. Chem. Soc. <u>1965</u> , 1.
,	37.	Ibid., <u>1966</u> , 77.	
/	38.	Johnson, J. R. Ph. D. Thesis, Univ. of Oklahom	a, Norman, Oklahoma, <u>1966</u> .
	39.	Christian, S. D.; Affsprung, H. E.; Hunter, W. J Properties of Water, U. S. Office Saline Water Ro p. 71, 79-81.	A.; Gillam, W. S.; McCoy, W. H. Solute es. Develop. Progr. Report, No. 301, <u>1968</u> ,
	40.	Högfeldt, E.; Fredlund, F. Acta Chem. Scand. 1	<u>970,</u> 24, 1858.
	41.	Simonov, V. D.; Pogulya, V. E.; Shamsutdinov,	T. M. Russ. J. Phys. Chem. <u>1970</u> , 44, 1755.
	42.	Simonov, V. D.; Pogulya, V. E.; Shamsutdinov, Respub. Pravl. Vses. Khim. Obshchest. <u>1971</u> , 346	T. M. et al. Dokl. Neftekim. Sekt. Bashkir. j.
7	43.	Glasoe, P. K.; Schultz, S. D. J. Chem. Eng. Dat	ta <u>1972</u> , 17, 66.
	44.	Kirchnerova, J. Ph. D. Thesis, McGill Univ., Me	ontreal, <u>April 1975,</u> 280 pp.
	45.	Kirchnerova, J.; Cave, G. C. B. Can. J. Chem.	<u>1976,</u> <i>54</i> , 3909.
,	46.	Ksiazczak, A.; Buchowski, H. Fluid Phase Equil	<i>ibr.</i> <u>1980</u> , <i>5</i> , 131.
	47.	Orlandini, M.; Fermeglia, M.; Kikic, I.; Alessi, I	P. Chem. Eng. J. <u>1983</u> , 26, 245.

•

22			
COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) Tetrachlorometh	ane (carbon tetr	achloride);	Rex, A.
(2) Water; $H_2O$ ; [7	732-18-5]		Z. Phys. Chem. <u>1906</u> , 55, 355-70.
VARIABLES:			PREPARED BY:
T/K = 273 - 303			A. L. Horvath
EXPERIMENTAL	VALUES:		
	t/°C	$100 g_1/g_2$	$\begin{array}{ccc} 100 \ w_{i} & 10^{4} \ x_{i} \\ (compiler) & (compiler) \end{array}$
	0	$9.7 \times 10^{-2}$	$9.7 \times 10^{-2}$ 1.14
	20	$8.0 \times 10^{-2}$	$8.0 \times 10^2$ 0.938 $8.0 \times 10^2$ 0.938
	50	0.5 × 10	0.5 × 10 0.550
		AUXILIARY I	NFORMATION
METHOD/APPARA	TUS/PROCEDU	JRE:	SOURCE AND PURITY OF MATERIALS:
The solubility was determined usin	of tetrachlorom	ethane in water aned flasks with	(1) Kahlbaum, redistilled and washed before use; b, p, = 76.7°C.
calibrated capillary ( equilibrated in a the	cylinders. After mostatic bath, t	samples were heir volumes and	(2) Distilled (compiler).
weights were detern calculated.	nined and the sol	lubility was	ESTIMATED ERRORS:
			Solubility: Not specified.
			Temperature: $\pm 0.5$ K.
COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) Water: $H_{-}O$ : [7732-18-5]			Clifford, C. W.
(2) Tetrachlorometh	ane (carbon tetr	achloride);	J. Ind. Eng. Chem. 1921, 13, 631-2.
CCl₄; [56-23-5]			
VARIABLES:			PREPARED BY:
T/K = 297 - 302			A. L. Horvath
EXPERIMENTAL V	ALUES:		
	t/℃	100 w <sub>1</sub>	$10^4 x_1$ 100 $w_1 M_1^{-1} / \text{mol g}^{-1}$
	24.0	$1.0 \times 10^{-2}$	(compiler) (compiler) 8.52 $5.55 \times 10^{-1}$
	28.5	$1.0 \times 10^{-2}$ $1.3 \times 10^{-2}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
		AUXILIARY II	NFORMATION
METHOD/APPARA	TUS/PROCEDU	JRE:	SOURCE AND PURITY OF MATERIALS:
The calcium c determination of wat was abstracted from	the samples by	was used for the es. The moisture air and absorbed	<ol> <li>Distilled (compiler).</li> <li>Technical grade, used after distilling over CaCl<sub>2</sub>.</li> </ol>
in two calcium chloride tubes. The two tubes were weighted before and after desorption. For further		wo tubes were For further	ESTIMATED ERRORS:
details, see (ref. 1).			Solubility: about 1 mg. Temperature: ± 0.5 K (compiler).
			REFERENCES:
			(1) Clifford, C. W. J. Ind. Eng. Chem. 1921,
			13, 628.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Gross, P. M.	
(2) Water; $H_2O$ ; [7732-18-5]	J. Am. Chem. Soc. <u>1929</u> , 51, 2362-6.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
t/°C 100 g <sub>1</sub> /g <sub>2</sub>	$100 w_1$ $10^5 x_1$	
25 $7.7 \times 10^{-2}$	$7.7 \times 10^{-2}$ 9.02	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A weighted amount of tetrachloromethane was mixed with 1000 g water in a bottle and immersed in a water thermostat. The bottle was placed on a shaker	<ol> <li>Kahlbaum's sulfur-free analytical reagent, used as received.</li> <li>Distilled.</li> </ol>	
until no more droplets of the liquid remained un- dissolved. The concentration of the tetrachloro-	ESTIMATED ERRORS:	
water interferometer. One or more samples were syphoned out for the determination of the solubility. The German translation is also published	Solubility: $\pm 5 \%$ . Temperature: $\pm 0.01$ K.	
(ref. 1).	REFERENCES:	
	(1) Gross, P. M. Z. Phys. Chem. <u>1929</u> , 6B, 215.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Rosenbaum, C. K.; Walton, J. H.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	J. Am. Chem. Soc. <u>1930</u> , 52, 3568-73.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 323	A. L. Horvath	
EXPERIMENTAL VALUES:	r	
$t/^{\circ}C$ 100 $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$\begin{array}{cccc} 10 & 7.11 \times 10^{-3} \\ 20 & 8.44 \times 10^{-3} \end{array}$	$\begin{array}{ccc} 7.11 \times 10^{\cdot3} & 0.607 \\ 8.44 \times 10^{\cdot3} & 0.720 \end{array}$	
	(continued)	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Tetrachloromethane was saturated with water in a flask using continuous shaking, followed by standing for a day or more to establish equilibrium.	<ol> <li>Distilled.</li> <li>Reagent grade, free from impurities, b. p. = 76.8°C.</li> </ol>	
troduced into a bulb containing CaH <sub>2</sub> . The complete apparatus was placed in a water bath. The evolved	ESTIMATED ERRORS:	
hydrogen was measured volummetrically after three days. Two to four determinations were carried out at each temperature. For more details, see (ref. 1).	Solubility: $\pm 0.001 \%$ H <sub>2</sub> O. Temperature: $\pm 0.2$ K (compiler).	
	REFERENCES:	
	(1) Rosenbaum, C. K. Ph. D. Thesis, Univ. of Wisconsin, Wisconsin, <u>1930</u> .	

Ϊ

Ł

24					
COMPONENTS:			ORIC	GINAL MEAS	UREMENTS:
(1) Water; H <sub>2</sub> O; [77	/32-18-5]		Rosenbaum, C. K.; Walton, J. H.		
(2) Tetrachlorometha CCl <sub>4</sub> ; [56-23-5]	ane (carbon	tetrachloride);	J. Am. Chem. Soc. <u>1930</u> , 52, 3568-73.		
VARIABLES:			PRE	PARED BY:	
T/K = 283 - 323			A. I	. Horvath	
EXPERIMENTAL V	ALUES: (co	ontinued)			
	t/°C	100 g <sub>1</sub> /g <sub>2</sub>	(	100 w <sub>i</sub> compiler)	10 <sup>s</sup> x, (compiler)
	30 40 50	$\begin{array}{c} 1.09 \times 10^{-2} \\ 1.52 \times 10^{-2} \\ 2.37 \times 10^{-2} \end{array}$	1. 1. 2.	$\begin{array}{r} 09 \times 10^{-2} \\ 52 \times 10^{-2} \\ 37 \times 10^{-2} \end{array}$	0.930 1.30 2.02
COMPONENTS:			ORIC	GINAL MEAS	UREMENTS:
(1) Tetrachlorometha	ane (carbon	tetrachloride);	Gross, P. M.; Saylor, J. H.		
$CCl_4; [56-23-5]$	22 19 51		J. A	m. Chem. Soc.	<u>1931</u> , <i>53</i> , 1744-51.
(2) water; $H_2O$ ; [77	32-18-3]				
VARIABLES:			PREI	PARED BY:	
T/K = 288 - 303			A. I	. Horvath	
EXPERIMENTAL V	ALUES:				
	t/℃	$1000 g_1/g_2$	(	100 w <sub>i</sub> compiler)	$10^5 x_i$ (compiler)
	15 30	0.77 0.81	7. 8.	$7 \times 10^{-2}$ 1 × 10^{-2}	9.02 9.49
		AUXILIARY I	NFORM	MATION	
METHOD/APPARA	TUS/PROCE	EDURE:	SOU	RCE AND PU	RITY OF MATERIALS:
An excess of t water was shaken for Samples were then w using an interferomet detailed description of	etrachlorome 12 hours in withdrawn and ter made by of the comple	thane in 500 g a thermostat. d read against water Zeiss (ref. 1). A ete procedure is	(1) (2)	Eastman Kod NaOH solutic use. Distilled.	ak Co., shaken with $H_2SO_4$ and on, dried and fractionated before
given in a Ph. D. the	esis (ref. 2).		ESTI	MATED ERR	ORS:
			Solu Tem	bility: perature:	± 2.5 %. ± 0.02 K.
			REF	ERENCES:	
			(1) (2)	Gross, P. M. 2362. Saylor, J. H. sity, Durham	J. Am. Chem. Soc. <u>1929</u> , 51, Ph. D. Thesis, Duke Univer- , <u>1930</u> .

	25
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]	Bell, R. P.
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	J. Chem. Soc. <u>1932</u> , 2905-11.
VARIABLES:	PREPARED BY:
T/K = 288 - 298	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/kg m^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
AUXILIARY I	NFORMATION
	SOURCE AND BURITY OF MATERIALS
An about 1 to 5 ratio of water/tetrachloro- methane mixture was rotated in a thermostat for 12 hours. After the equilibrium was attained, samples were taken and filtered through cottonwool. The	<ol> <li>(1) Distilled (compiler).</li> <li>(2) Merck reagent, analytical grade, redistilled before use.</li> </ol>
the reaction with $\alpha$ -naphthoxydichlorophosphine. The	ESTIMATED ERRORS:
NaOH solution (ref. 1). Two to three successive de- terminations were carried out with the tetrachloro- methane.	Solubility: $\pm$ 4.4 % mean. Temperature: $\pm$ 0.02 K.
	REFERENCES:
	(1) Bell, R. P. J. Chem. Soc. <u>1932</u> , 2903.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	van Arkel, A. E.; Vles, S. E.
(2) Water; $H_2O$ ; [7732-18-5]	Recl. Trav. Chim. Pays-Bas, <u>1936</u> , 55, 407-11.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}$ C 10 <sup>4</sup> $x_1$	$\begin{array}{ccc} 100 \ w_i & 100 \ w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & \text{(compiler)} \end{array}$
30 0.955	$8.15 \times 10^{-2}$ $5.3 \times 10^{-4}$
	NEODMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Niini, A.	
$CC_{4}$ ; [30-23-3]	Suomen Kemistilehti <u>1938</u> , 11a, 19-20.	
(2) water; $H_2O$ ; [//32-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EVDEDIMENTAL VALUES.		
EXPERIMENTAL VALUES:	5 ··· 100 ··· 104 ··	
1/°C 100 W <sub>1</sub> 10 (com	$\begin{array}{ccc} x_1 & 100 \ w_2 & 10^{\circ} \ x_2 \\ \text{piler} & (\text{compiler}) \end{array}$	
20 $7.1 \times 10^{-2}$ 8.3	$5.0 \times 10^{-3}$ 4.27	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The change caused by the addition of tetra-	(1) Kahlbaum, analytical grade, $d_4 = 1.59413$ at	
and refractive index methods. The measurements of	(2) Distilled.	
were carried out by the conventional methods, that is by the use of a sefective and a dilatemeter		
respectively.	ESTIMATED ERRORS:	
	Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Fox, J. J.; Martin, A. K.	
(2) Tetrachloromethane (carbon tetrachloride);	Proc. Roy. Soc. London, Ser. A, <u>1940</u> , 174, 234-62	
0014, [50 25 5]	204 02.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 291	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 100 $g_{1}/g_{2}$	$100 w_1$ $10^4 x_1$	
	(compiler) (compiler)	
18.3 $6.9 \times 10^{-3}$	$6.9 \times 10^{-3}$ 5.89	
AUXILIARY IN	IFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A grating spectrometer was used to observe the	(1) Distilled.	
intrared absorption spectrum of water in the saturated tetrachloromethane solution. The absorption cells	(2) Source and purity not given.	
removal of water was revealed by the diminution of	ESTIMATED ERRORS:	
using the measured weight of water collected.	Solubility: Not specified.	

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]	Staverman, A. J.
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Recl. Trav. Chim. Pays-Bas, <u>1941</u> , 60, 836-41.
VARIABLES:	PREPARED BY:
T/K = 273 - 303	A. L. Horvath
EXPERIMENTAL VALUES: t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^{5} x_{i} & 100 \ w_{i} M_{i}^{-i} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$
$\begin{array}{cccc} 0 & 0.50 \times 10^{-2} \\ 25 & 1.16 \times 10^{-2} \\ 30 & 1.58 \times 10^{-2} \end{array}$	$\begin{array}{cccc} 42.8 & 2.77 \times 10^{-4} \\ 99.1 & 6.44 \times 10^{-4} \\ 135.0 & 8.77 \times 10^{-4} \end{array}$
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Tetrachloromethane was shaken with water for about 12 hours in a liquid paraffin thermostat bath.	<ol> <li>Distilled.</li> <li>Source and purity not given.</li> </ol>
The water content of the organic phase was determin- ed by the Karl Fischer titration method. All measure-	ESTIMATED ERRORS:
ments were done in duplicates. The results were averages of two measurements at each temperature. A full description of the method used is given in a Ph. D. thesis (ref. 1).	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiled).
	REFERENCES:
	(1) Staverman, A. J. Ph. D. Thesis, Univ. of Leiden, Leiden, Belgium, <u>1938</u> .
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]	Hutchison, C. A.; Lyon, A. M.
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Columbia University Report A-745, July 1, 1943.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
<i>t/</i> °C 100 <i>w<sub>1</sub>M<sub>1</sub><sup>-1</sup></i> /mol g <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_{i} & 10^{4} \ x_{i} \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$25   5.43  imes 10^4$	9.78 × 10 <sup>-3</sup> 8.34
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A 1 to 15 volume ratio mixture of water and tetrachloromethane was introduced into an equilibra- tion flask and then lowered into a waterbath thermo-	<ol> <li>Distilled.</li> <li>Source is not known. Purified and dried before use.</li> </ol>
90 minutes at constant temperature. The amount of water in the organic phase was determined by a modified Karl Fischer titration. The determination	ESTIMATED ERRORS: Solubility: $\pm 0.1 \times 10^4$ avg. dev.
was taken from a secondary source (ref. 1). The original university report is no longer available.	1 emperature: ± 0.05 K.
	REFERENCES:
	<ol> <li>Eidinoff, M. L.; Joris, G. G.; Taylor, H. S.; Urey, H. C., Eds., <i>Production of Heavy</i> <i>Water</i>, McGraw-Hill, New York, <u>1955</u>, p. 129.</li> </ol>

, J

28	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	McGovern, E. W.
(2) Water; $H_2O$ ; [7732-18-5]	Ind. Eng. Chem. <u>1943</u> , 35, 1230-9.
VARIABLES:	PREPARED BY:
T/K = 273 - 303	A. L. Horvath
EXPERIMENTAL VALUES:	I
t/°C 100 w, 10 (com	$x_{1}^{4} x_{1}^{2} = 100 w_{2}^{2} = 10^{4} x_{2}^{2}$ appiler) (compiler)
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Solubility data as a function of temperat tabulated data point at $25^{\circ}$ C.	ure were presented in graphical form except the
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	(1) Source not given, commercial grade.
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	Powell, J. F.
(2) Water; $H_2O$ ; [7732-18-5]	Brit. J. Industr. Med. <u>1945</u> , 2, 212-6.
VARIABLES:	PREPARED BY:
T/K = 293 - 310	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Ostwald coefficient, L/dimensionless	$\begin{array}{ccc} 100 & w_i & 10^s & x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$
20 1.05 37 0.45	$\begin{array}{cccc} 8.05 \times 10^{-2} & 9.44 \\ 6.82 \times 10^{-2} & 7.99 \end{array}$
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A tetrachloromethane - air mixture was bub- bled through 50 cm <sup>3</sup> distilled water contained in a gas	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>
Equilibrium was reached in 3/4 hour with a gas flow of 50 cm <sup>3</sup> /mim. The saturated solution was trans-	ESTIMATED ERRORS:
ferred to a measuring cylinder and shaken with tolu- ene. The toluene extraction was allowed to settle and then analyzed for tetrachloromethane by the addition	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiled).
ot pyridine and alkali; see (ref. 1).	REFERENCES:
	(1) Habgood, S.; Powell, J. F. Brit. J. Industr. Med. 1944, J. 39.

	29
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	Booth, H. S.; Everson, H. E.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	I
t/°C 100 V <sub>1</sub> /V <sub>2</sub>	$\begin{array}{ccc} 100 \ w_{1} & 10^{4} \ x_{1} \\ \text{(compiler)} & \text{(compiler)} \end{array}$
25.0 0.10	0.159 1.86
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The equilibrium was established through re- peated shaking and centrifuging of a tetrachloro- methane and water mixture in a stoppered Goetz tube	<ol> <li>Commercial reagent, C. P. grade, used as received.</li> <li>Distilled.</li> </ol>
and returning the tube to a constant temperature water bath. The difference between the total amount of	ESTIMATED ERRORS:
which remained in excess was taken as the amount of solute dissolved in the known volume of water. The determination of the excess amount of solute added is	Solubility: Not specified. Temperature: ± 1 K (compiler).
described by Hanaslick in his Ph. D. thesis (ref. 1).	REFERENCES:
	(1) Hanslick, R. S. Ph. D. Thesis, Columbia University, <u>1935</u> .
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	Booth, H. S.; Everson, H. E.
(2) Water; $H_2O$ ; [7732-18-5]	Ind. Eng. Chem. <u>1949</u> , 41, 2627-8.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298 - 333	A. L. Horvath
EXPERIMENTAL VALUES:	1
$t/^{\circ}C$ 100 $V_{1}/V_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
25.0 0.10 60 0.10	0.159 1.86 0.159 1.86
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The solute-solvent mixture in a solubility tube was gently rotated while the tube was immersed in a constant temperature bath. The difference between the	<ol> <li>Commercial reagent, C. P. grade, used as received.</li> <li>Distilled</li> </ol>
total amount of solute and the amount remaining in excess was taken as the amount of solute dissolved in	ESTIMATED ERRORS:
the known volume of solvent. The volumetric deter- mination is described elsewhere (ref. 1).	Solubility: Not specified. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Hanslick, R. S. <i>Ph. D. Thesis</i> , Columbia University, <u>1935</u> .

30	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	Chitwood, B. G.
(2) Water; $H_2O$ ; [7732-18-5]	Adv. in Chem. Ser., Am. Chem. Soc., <u>1952</u> , 7, 91-9.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	,
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^{5} x_{i} & 10^{6} w_{i} M_{j}^{-1} / \text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$
25 $8.0 \times 10^{2}$	9,38 5.20
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).
COMPONENTS:	ORIGINAL MEASUREMENTS:
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride);	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E.
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5]	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. <u>1952</u> , 56, 480-4.
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES:	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. <u>1952</u> , 56, 480-4. PREPARED BY:
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 298	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. <u>1952</u> , 56, 480-4. PREPARED BY: A. L. Horvath
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T/</i> K = 298 EXPERIMENTAL VALUES:	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. <u>1952</u> , 56, 480-4. PREPARED BY: A. L. Horvath
COMPONENTS:(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5](2) Water; H <sub>2</sub> O; [7732-18-5]VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>5</sup> x <sub>1</sub> 10 (corr	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. <u>1952</u> , 56, 480-4. PREPARED BY: A. L. Horvath $0.w_1$ $10^4 x_2$ $100 w_2$ piler) $10^4 x_2$ (compiler)
COMPONENTS:(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5](2) Water; H <sub>2</sub> O; [7732-18-5]VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>5</sup> x <sub>1</sub> 10 (corr 259.78.28	ORIGINAL MEASUREMENTS:         Donahue, D. J.; Bartell, F. E.         J. Phys. Chem. 1952, 56, 480-4.         PREPARED BY:         A. L. Horvath $0.w_1$ $10^4 x_2$ $0.0 w_2$ piler)         (compiler)         × $10^{-2}$ 8.3 $9.73 \times 10^{-3}$
COMPONENTS:(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5](2) Water; H <sub>2</sub> O; [7732-18-5]VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>5</sup> x <sub>1</sub> 10 (corr 259.78.28	ORIGINAL MEASUREMENTS:         Donahue, D. J.; Bartell, F. E.         J. Phys. Chem. 1952, 56, 480-4.         PREPARED BY:         A. L. Horvath $0w_1$ $10^4 x_2$ $00w_2$ (compiler)         × $10^{-2}$ 8.3         9.73 × $10^{-3}$
COMPONENTS:(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5](2) Water; H <sub>2</sub> O; [7732-18-5]VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>5</sup> x <sub>1</sub> 259.78.28	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1952, 56, 480-4.PREPARED BY: A. L. Horvath $0.w_1$ $10^4 x_2$ $100 w_2$ piler) $(compiler)$ $\times 10^{-2}$ $8.3$ $9.73 \times 10^{-3}$ NFORMATION
COMPONENTS:(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5](2) Water; H <sub>2</sub> O; [7732-18-5]VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ 10 <sup>5</sup> $x_1$ 10 (corr 25 9.7 8.28AUXILIARY IIMETHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS: Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1952, 56, 480-4.PREPARED BY: A. L. Horvath $0. w_1$ $10^4 x_2$ $0. w_2$ piler) × $10^{-2}$ $8.3$ $9.73 \times 10^{-3}$ NFORMATIONSOURCE AND PURITY OF MATERIALS:
COMPONENTS:         (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $10^{3} x_{1}$ (corr         25       9.7         8.28         METHOD/APPARATUS/PROCEDURE:         A mixture of water and tetrachloromethane         was placed in a glass stoppered flask and was shaken         intermittently for at least three days in a water bath         held at constant temperature. The organic phase was	ORIGINAL MEASUREMENTS:         Donahue, D. J.; Bartell, F. E.         J. Phys. Chem. 1952, 56, 480-4.         PREPARED BY:         A. L. Horvath $0 w_1$ $10^4 x_2$ $00 w_2$ piler)         (compiler)         × $10^{-2}$ 8.3 $9.73 \times 10^{-3}$ NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1)       Reagent grade, purified by fractional distillation.         (2)       Purified by distillation.
COMPONENTS:         (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $10^{5} x_{1}$ 10         (corr         25       9.7         8.28         METHOD/APPARATUS/PROCEDURE:         A mixture of water and tetrachloromethane         was placed in a glass stoppered flask and was shaken         intermittently for at least three days in a water bath         held at constant temperature. The organic phase was         analyzed for water content by the Karl Fischer meth-         of and the aqueous phase was analyzed by using an	ORIGINAL MEASUREMENTS:         Donahue, D. J.; Bartell, F. E.         J. Phys. Chem. 1952, 56, 480-4.         PREPARED BY:         A. L. Horvath $0 w_1$ $10^4 x_2$ $00 w_2$ piler)         (compiler)         × $10^{-2}$ 8.3 $9.73 \times 10^{-3}$ NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1)       Reagent grade, purified by fractional distillation.         (2)       Purified by distillation.         ESTIMATED ERRORS:

	31		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Rotariu, G. J.; Fraga, D. W.; Hildebrand, J. H.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	J. Am. Chem. Soc. <u>1952</u> , 74, 5783.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 100 \ V_{j}/V_{j} & 10^{4} \ x_{j} \\ (\text{compiler}) \end{array}$		
25 $1.0 \times 10^{-2}$	$1.6 \times 10^{-2}$ 8.53		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The solute-solvent mixture was rocked for 24 hours in a water bath. Samples were removed from the organic-rich phase with a special, turned-up tip	<ol> <li>Distilled (compiler).</li> <li>Source and purity not given.</li> </ol>		
bipet. The liquid was exposed only to dry nitrogen during its removal and analysis with a Karl Fischer	ESTIMATED ERRORS:		
terminations.	Solubility: Not specified.		
	Temperature: $\pm 0.01$ K.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Eberius, E.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	Wasserbestimmung mit Karl-Fischer-Lösung, Verlag Chemie, GMBH, Weinheim, <u>1954</u> , p. 67.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C 100 $w_{I}$	$\begin{array}{ccc} 10^3 x_i & 10^6 w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
20 $0.012 \pm 0.003$	1.02 6.66		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A 50 g tetrachloromethane sample was shaken with pure water in a 100 ml Erlenmeyer flask. The two phases were separated when equilibrium had	<ol> <li>Distilled.</li> <li>Source and purity not given.</li> </ol>		
been established. The organic phase was analyzed for water using the Karl Fischer titration method. Five	ESTIMATED ERRORS:		
determinations were performed.	Solubility: See above.		
	remperature: Not specified.		

~~			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Greinacher, E.; Lüttke, W.; Mecke, R.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	Z. Elektrochem. <u>1955</u> , 59, 23-31.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
$t/^{\circ}C$ $n_{1}/V_{2}/mol m^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
20 7.5	$8.48 \times 10^{-3}$ 7.24		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A saturated solution of water and tetrachloro- methane was prepared and the water content of the organic phase was determined by the Karl Fischer titration method.	<ol> <li>Spectroscopic grade.</li> <li>Source not given, purified and dried before use.</li> </ol>		
	ESTIMATED ERRORS:		
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).		
1			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	ORIGINAL MEASUREMENTS: Fitzgerald, M. E.; Griffing, V.; Sullivan, J.		
<ul> <li>COMPONENTS:</li> <li>(1) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ul>	ORIGINAL MEASUREMENTS: Fitzgerald, M. E.; Griffing, V.; Sullivan, J. J. Chem. Phys. <u>1956</u> , 25, 926-33.		
<ul> <li>COMPONENTS:</li> <li>(1) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> <li>VARIABLES:</li> </ul>	ORIGINAL MEASUREMENTS: Fitzgerald, M. E.; Griffing, V.; Sullivan, J. J. Chem. Phys. <u>1956</u> , 25, 926-33. PREPARED BY:		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T/K</i> = 293	ORIGINAL MEASUREMENTS: Fitzgerald, M. E.; Griffing, V.; Sullivan, J. J. Chem. Phys. <u>1956</u> , 25, 926-33. PREPARED BY: A. L. Horvath		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T/K</i> = 293 EXPERIMENTAL VALUES:	<ul> <li>ORIGINAL MEASUREMENTS:</li> <li>Fitzgerald, M. E.; Griffing, V.; Sullivan, J.</li> <li>J. Chem. Phys. <u>1956</u>, 25, 926-33.</li> <li>PREPARED BY:</li> <li>A. L. Horvath</li> </ul>		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/mol m^{-3}$	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $100 w_i$ $(compiler)$		
COMPONENTS:         (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t/^{\circ}$ C $n_1/V_2/mol m^{-3}$ 20       5.8	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $100 w_i$ $(compiler)$ $8.9 \times 10^{-2}$ $1.04$		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/\text{mol m}^{-3}$ 20 5.8	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $100 w_i$ $(compiler)$ $8.9 \times 10^{-2}$ $1.04$		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/mol m^{-3}$ 20 5.8 AUXILIARY II	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $100 w_i$ $(compiler)$ $8.9 \times 10^{-2}$ 1.04		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/mol m^{-3}$ 20 5.8 AUXILIARY II METHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $100 w_i$ $(compiler)$ $8.9 \times 10^{-2}$ 1.04NFORMATIONSOURCE AND PURITY OF MATERIALS:		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/mol m^{-3}$ 20 5.8 AUXILIARY II METHOD/APPARATUS/PROCEDURE: Details are not available.	ORIGINAL MEASUREMENTS:         Fitzgerald, M. E.; Griffing, V.; Sullivan, J.         J. Chem. Phys. <u>1956</u> , 25, 926-33.         PREPARED BY:         A. L. Horvath $100 w_i$ (compiler) $(compiler)$ (compiler) $8.9 \times 10^{-2}$ 1.04         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) Double distilled.		
COMPONENTS: (1) Tetrachloromethane (carbon tetrachloride); (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ $n_1/V_2/mol m^{-3}$ 20 5.8 AUXILIARY II METHOD/APPARATUS/PROCEDURE: Details are not available.	ORIGINAL MEASUREMENTS:Fitzgerald, M. E.; Griffing, V.; Sullivan, J.J. Chem. Phys. 1956, 25, 926-33.PREPARED BY:A. L. Horvath $100 w_i$ $10^4 x_i$ (compiler)8.9 × 10 <sup>-2</sup> 1.04NFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Double distilled.ESTIMATED ERRORS:		

	33		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Desnoyer, M.		
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Dosage de l'Eau dans les Solvants Organiques par Absorption Infra-Rouge et Mesure des Constantes Dielectriques, Centre D'Etudes Nucleaires de Saclay, Report No. 1254, Saclay, France, June 23, 1959.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 299	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^4 x_i & 10^6 w_i M_j^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
$26   1.01 \times 10^{-2}$	8.62 5.61		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The saturation of tetrachloromethane with water took place at room temperature under vigorous stirring. The concentration of water in the organic-	<ol> <li>Distilled.</li> <li>Commercial reagent, washed and dried before use.</li> </ol>		
rich phase was determined by infrared spectrometry.	ESTIMATED ERRORS:		
	Solubility: $\pm 4 \%$ . Temperature: $\pm 0.5 $ K (compiler).		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Zielinski, A. Z.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	Chem. Stosowana <u>1959</u> , 3, 377-84.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}C$ $\rho_1/kg m^{-3}$	$10^3 x$ , $100 w$ , (compiler) (compiler)		
25 1.54	8.24 $9.72 \times 10^2$		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The concentration of water in the organic phase was determined by the Karl Fischer titration method (caf. 1)	<ol> <li>Distilled (compiler).</li> <li>Gliwice Chemicals, pure grade, used as received</li> </ol>		
	ESTIMATED ERRORS:		
	Solubility: Not specified. Temperature: $\pm 1$ K (compiler).		
	(1) Mitchell I. Smith D. M. Acuamatou A		
	<i>Treatise on Metrhods for the Determination of Water</i> , Wiley, New York, <u>1952</u> , pp. 73 and 260.		

34			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Tettamanti, K.; Nogradi, M.; Sawinsky, J.		
(2) Water; $H_2O$ ; [7732-18-5]	Period. Polytech., Chem. Eng. <u>1960</u> , 4, 201-18.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	Z. Maczynska		
EXPERIMENTAL VALUES:			
t/°C 100 w₁ 10 (con	$\begin{array}{ccccc} s & x_1 & 100 & w_2 & 10^4 & x_2 \\ \text{npiler} & & & (\text{compiler}) \end{array}$		
20 $7.0 \times 10^{-2}$ 8.	$1.0 \times 10^2$ 8.53		
The reported values were graphically sn	noothed.		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Turbidimetric titrations were used for both the aqueous and the organic phases. Tetrachloromethane or water was weighted into a stoppered flask with an accuracy of 0.01 g. The temperature was maintained	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
constant in an ultrathermostat. First, the organic solvent was titrated with water from a microburet	ESTIMATED ERRORS:		
until the appearance of turbidity. Then, the water was titrated with the organic solvent until the appearance of turbidity.	Solubility: $\pm 10 \%$ . Temperature: $\pm 0.1 $ K.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Liu, JL.; Huang, TC.		
(2) Water; $H_2O$ ; [7732-18-5]	Scientia Sinica (Peking) <u>1961</u> , 10, 700-10.		
VARIABLES:	PREPARED BY:		
T/K = 288 - 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 1000 g₁/g₂	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
15 0.755 20 0.748 25 0.762	$\begin{array}{cccc} 7.54 \times 10^{2} & 8.837 \\ 7.47 \times 10^{2} & 8.755 \\ 7.61 \times 10^{2} & 8.919 \end{array}$		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
An accurate amount of tetrachloromethane was transferred into an Erlenmeyer flask containing a weighed amount of distilled water. The flask was rotated in a thermostat bath until the equilibrium was established. The optical densities of different con- centrations of tetrachloromethane in water water	<ol> <li>Peking Chemical Factory, further purified by washing with H<sub>2</sub>SO<sub>4</sub>, NaOH, and water, dried over CaCl<sub>2</sub>, and fractionated.</li> <li>Distilled.</li> </ol>		
measured using water as the reference liquid. Dupli- cate experiments were made and the results averag- ed. The absorption spectrum of the tetrachloro- methane in water was determined by using a Hilger spectrophotometer.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).		
speen sphotomotori			

	35		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Johnson, J. R.; Christian, S. D.; Affsprung, H. E.		
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	J. Chem. Soc. <u>1965</u> , 1-4.		
VARIABLES:	PREPARED BY:		
T/K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C $c_1/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x, \\ (compiler) & (compiler) \end{array}$		
25.0 8.7	9.89 × 10 <sup>-3</sup> 8.44		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The method of establishing the water and tetrachloromethane solution equilibrium in a thermo- stat has been reported elsewhere (ref. 1). The con-	<ol> <li>Distilled (compiler).</li> <li>Source not given. Purified by distillation in an Oldershaw column before use.</li> </ol>		
centration of water in the tetrachloromethane phase was determined using a Beckman KF-3 aquameter.	ESTIMATED ERRORS:		
For all measurements, 50 ml samples were analyzed.	Solubility: Not specified. Temperature: $\pm 0.1$ K.		
	REFERENCES:		
	(1) Christian, S. D.; Affsprung, H. E.; Johnson,		
	J. R.; Worley, J. D. J. Chem. Educ. <u>1963</u> , 40, 419.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Johnson, J. R.; Cristian, S. D.; Affsprung, H. E.		
<ul><li>(2) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li></ul>	J. Chem. Soc. A, <u>1966</u> , 77-8.		
VARIABLES:	PREPARED BY:		
T/K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}C$ $c_{1}/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
25.0 8.7	$9.89 \times 10^{-3}$ 8.44		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Samples of water and tetrachloromethane mix- tures were equilibrated in a constant temperature waterbath. The solute isopiestic apparatus has been	<ol> <li>Distilled (compiler).</li> <li>Source not given, reagent grade, distilled in an Oldershaw column before use.</li> </ol>		
described elsewhere (ref. 1). The water solubilities were determined using a Beckman Model KF-3	ESTIMATED ERRORS:		
Aquameter.	Solubility: $\pm 0.3 \times 10^{-3}$ . Temperature: $\pm 0.1^{\circ}$ C.		
	REFERENCES:		
	(1) Christisn, S. D.; Affsprung, H. E.; Johnson, J. R.; Worley, J. D. J. Chem. Educ, 1963,		

36		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Johnson, J. R.	
(2) Water; $H_2O$ ; [7732-18-5]	Self-Association and Hydration of Phenol in Several Organic Solvents, Ph. D. Thesis, Univ. of Oklahoma, Norman, Oklahoma, <u>1966</u> .	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C <i>c<sub>1</sub></i> /mol m <sup>-3</sup> 10 (con	$\begin{array}{llllllllllllllllllllllllllllllllllll$	
25 5.1 7.8 >	$\times 10^{-2}$ 8.7 9.89 $\times 10^{-3}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibrium between tetrachloromethane and water took place in a well-stirred water bath. The experimental samples were allowed to equilibrate for two or more days. Water analyses were made with a	<ol> <li>Allied Chemical Corp., purified by distillation before use.</li> <li>Distilled.</li> </ol>	
Beckman Aquameter using a Karl Fischer reagent. The solubility of tetrachloromethane in water was	ESTIMATED ERRORS:	
vice. Full details are available in the thesis.	Solubility: $\pm 4 \%$ . Temperature: $\pm 0.1$ K.	
	ODICINAL MEASUDEMENTS.	
(1) Water: $H_0$ : [7732-18-5]	Grigshy R D	
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ul>	Self-Association and Hydration of N-Methyl- acetamide in Carbon Tetrachloride, Ph. D. Thesis, Univ. of Oklahoma, Norman, Oklahoma, <u>1966</u> .	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C <i>c<sub>1</sub></i> /mol m <sup>-3</sup> 10 (com	0 $w_i$ 10 <sup>3</sup> $x_i$ Method used: piler) (compiler)	
25 9.8 1.8 × 25 8.8 1.6 ×	$\begin{array}{cccc} 10^{-2} & 1.53 & \text{coulometric} \\ 10^{-2} & 1.36 & \text{total pressure} \end{array}$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Two methods were used for the water concen- tration in tetrachloromethane determination. After equilibration in a thermostated closed	<ol> <li>Distilled.</li> <li>Fischer reagent, refluxed with Hg and redistilled before use.</li> </ol>	
total water content using a coulometric Karl Fischer	ESTIMATED ERRORS:	
the saturated solution was measured in a liquid-vapor equilibrium still apparatus. Full details are described in the Ph. D. thesis; see also (ref. 1).	Solubility: $\pm 12$ %. Temperature: $\pm 0.1$ K.	
	REFERENCES:	
	<ol> <li>Grigsby R. D.; Christian, S. D.; Affsprung, H. R. J. Phys. Chem. <u>1968</u>, 72, 2465.</li> </ol>	

	37		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Christian, S. D.; Affsprung, H. E.; Hunter, W. J.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	A.; Gillam, W. S.; MCCOY, W. H. Solute Properties of Water, U. S. Office of Saline Water Research and Development Program, Report No. 301 <u>1968</u> , p. 71, 79-81.		
VARIABLES:	PREPARED BY:		
T/K = 288 - 298	A. L. Horvath		
EXPERIMENTAL VALUES:	1		
$t/^{\circ}$ C $n_1/V_2/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$		
$\begin{array}{cccc} 15 & 5.9 \pm 0.2 \\ 25 & 8.7 \pm 0.3 \end{array}$	$\begin{array}{cccccccc} 6.64 \times 10^{-3} & 5.66 \\ 9.89 \times 10^{-3} & 8.44 \end{array}$		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The solute isopiestic method was used without modification. The apparatus employed consisted of a closed vessel divided into two compartments, one for water and the other for the organic solvent. Equilibri- um was attained often for a period of 6 to 8 hours. The solution water with the Back	<ol> <li>Distilled.</li> <li>Source not given, certified or reagent grade, distilled through a 30 plate Oldershaw column before use.</li> </ol>		
man KF-3 Aquameter. The Karl Fisher reagent was standardized against crystalling sodium tartarate	ESTIMATED ERRORS:		
dihydrate.	Solubility: See above. Temperature: $\pm 0.1$ K.		
COMPONENTS	ORIGINAL MEASUREMENTS		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Högfeldt, E.; Fredlund, F.		
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ul>	Acta Chem. Scand. <u>1970</u> , 24, 1858-60.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
$t/^{\circ}$ C $c_1/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$		
25 10	9.7 × 10 <sup>-3</sup> 8.28		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Samples of tetrachloromethane were shaken to equilibrium with pure water. The liquid mixture was centrifuged and the organic phase analyzed for water following a modification of the Karl Fischer titration method.	<ol> <li>Distilled (compiler).</li> <li>Source not given, 99.95 % pure, checked by VPC.</li> </ol>		
memou.	ESTIMATED ERRORS:		
	Solubility:Not specified.Temperature: $\pm$ 0.5 K (compiler).		

38		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Simonov, V. D.; Pogulyai, V. E.; Shamsutdinov,	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	1. M. Russ. J. Phys. Chem. <u>1970</u> , 44, 1755-7.	
VARIABLES:	PREPARED BY:	
7/K = 303 - 323	A. L. Horvath	
EXPERIMENTAL VALUES		
t/°C 100 w <sub>1</sub>	$\begin{array}{cccc} 10^{3} x_{i} & 100 \ w_{i} M_{i}^{-1} / \text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
$\begin{array}{cccc} 30 & 1.35 \times 10^{-2} \\ 50 & 3.05 \times 10^{-2} \end{array}$	$\begin{array}{cccc} 1.15 & 7.49 \times 10^{4} \\ 2.60 & 1.69 \times 10^{-3} \end{array}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The Alexejew's synthetic method of solubility determination was used (ref. 1). A fixed weight of water and tetrachloromethane was sealed in a tube	<ol> <li>Distilled (compiler).</li> <li>Source not given. Chemically pure grade, dried and redistilled before use.</li> </ol>	
and subjected to a gradually increasing temperature with constant agitation. The appearence of an	ESTIMATED ERRORS:	
tion temperature. The temperature was then allowed to fall and an observation made, while the tube was constantly agitated, to establish the temperature of the	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
first appearance of opalescence. The observation was repeated several times.	REFERENCES:	
	(1) Alexejew, W. Ann. Phys. Chem. <u>1886</u> , 28, 305.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Simonov, V. D.; et al.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Dokl. Neftekim. Sekt., Bashkir. Respub. Pravl. Vses. Khim. Obshchest. <u>1971</u> , 346-51.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 288 - 299	A. L. Horvath	
EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 w, 10 <sup>4</sup> x,		
Alexeev Fischer Spectroscopy Alexeev Fischer Spectros		
14.5 $6.33 \times 10^{-3}$ - 6.	$30 \times 10^{-3}$ 5.402 - 5.377	
	(continued)	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The determination of the solubility of water in tetrachloromethane was based on the infrared spectro- scopic method. A UR-20 spectrophotometer was used	<ol> <li>Distilled.</li> <li>"KhCh" brand, treated with Na<sub>2</sub>CO<sub>3</sub>, dried with CaCl<sub>2</sub> and P<sub>2</sub>O<sub>5</sub> and distilled before use.</li> </ol>	
The spectra were recorded at 3600-810 cm <sup>-1</sup> , and the absorbance at 3702 cm <sup>-1</sup> was found by the baseline	ESTIMATED ERRORS:	
method. Standard mixtures of tetrachloromethane and water were used for the calibration graphs. The results obtained were compared with measurements	Solubility: Not specified. Temperature: $\pm$ 0.5 K.	
Fischer titration.	REFERENCES:	
	(1) Alexeev, V. F. Wied. Ann. <u>1886</u> , 28, 305.	

•

_						39
COMPONENTS:			ORIGINAL	MEASURE	MENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]		Simonov, V. D.; et al.				
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]		Dokl. Neftekim. Sekt., Bashkir. Respub. Pravl. Vses. Khim. Obshchest. <u>1971</u> , 346-51.		Respub. Pravl. 346-51.		
VARIABLES:		PREPAREI	D BY:			
T/K = 288 - 299			A. L. Horv	/ath		
EXPERIMENTAL	VALUES: (con	tinued)	=			
t/°C		100 w <sub>1</sub>			$10^4 x_1$	
	Alexeev	Fischer Spec	troscopy	Alexeev	Fischer	Spectroscopy
18 22 24 26	$\begin{array}{c} 7.00 \times 10^{-3} \\ 8.85 \times 10^{-3} \\ 9.12 \times 10^{-3} \\ 10.70 \times 10^{-3} \end{array}$	$\begin{array}{c} 7.20 \times 10^{-3} & 7 \\ 8.90 \times 10^{-3} & 8 \\ 10.60 \times 10^{-3} \end{array}$	$10 \times 10^{-3}$ $90 \times 10^{-3}$ $10 \times 10^{-3}$	5.974 7.551 7.782 9.129	6.144 7.594 9.043	6.059 7.594 8.617 -
COMPONENTS:			ORIGINAL	MEASURE	MENTS:	
(1) Tetrachlorometh CCl <sub>4</sub> ; [56-23-5]	hane (carbon te	rachloride);	Svetlanov, I.; Treger	Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. I.; Treger, Yu. A.; Flid, R. M.		
(2) Water; H <sub>2</sub> O; [7	732-18-5]		Russ. J. Phys. Chem. <u>1971</u> , 45, 488-90.			
VARIABLES:			PREPARED BY:			
T/K = 288 - 333		A. L. Horvath				
EXPERIMENTAL	VALUES:					
	t/°C	1000 g <sub>1</sub> /g <sub>2</sub>	100 w (compile	er)	10 <sup>s</sup> x (compi	iler)
	15 30 45 60	0.81 0.45 0.25 0.14	$8.1 \times 10$ $4.5 \times 10$ $2.5 \times 10$ $1.4 \times 10$	)-2 )-2 )-2 )-2	9.4 5.2 2.9 1.6	9 7 3 4
		AUAILIANT		//\		
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:				
The dynamic method used for the solubility de- termination consisted of the saturation of nitrogen with tetrachloromethane vapor which was then bub-		(1) Sourc (2) Distil	e and purity led (compile	not given. r).	•	
with a definite volument	me of water. Af	ter 2 - 3 hours, amples were tak-	ESTIMATED ERRORS:			
en periodicallty (ref tetrachloromethane i Some 3 - 5 measure	1). The conce in water was de ments were tak	ntration of the termined by GLC. en at each temper-	Solubility: Temperatur	re: ±	15 %. 0.05 K.	
ature.		•	REFERENC	CES:		
			(1) Trege Russ.	er, Yu. A.; F J. Phys. Ch	Flid, R. M 1964, 1964,	.; Spektor, S. S. 38, 253.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ol> <li>Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ol>	Karger, B. L.; Chatterjee, A. K.; King, J. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Tech. Report No. 3, Dept. of Chem., Northeastern Univ., Boston, Mass., <u>May 10, 1971</u> .	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 286	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
$t^{\circ}C$ Partition coefficient <sup>1</sup> , $K_L$ /dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_j \\ (compiler) & (compiler) \end{array}$	
12.5 1.9	0.106 1.24	
<sup>1</sup> Gas-liquid chromatographic parameter, The solubility was calculated from the p	, from instrument calibration. artition coefficient and the vapor pressure.	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Gas-liquid chromatography was used for the determination of the partition of tetrachloromethane with thin layers of water coated on Porasil D. Helium	<ol> <li>J. T. Baker Chemical Co., reagent grade, used as received.</li> <li>Distilled (compiler).</li> </ol>	
carrier. The gas was presaturated with water at the column temperature. The eluents were detected with	ESTIMATED ERRORS:	
an F. & N. flame ionization detector. The partition coefficient $(K_l)$ was obtained from the slope of the straight line plot of $V_N/A_L$ vs. $V_L/A_L$ .	Solubility: Not specified. Temperature: $\pm$ 0.05 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Glasoe, P. K.; Schultz, S. D.	
(2) Totrachlagomathana (asshan tatrachlagida)		
CCl <sub>4</sub> ; [56-23-5]	J. Chem. Eng. Data <u>1972</u> , 17, 66-8.	
VARIABLES:	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY:	
VARIABLES: T/K = 288 - 318	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY: A. L. Horvath	
VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES:	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY: A. L. Horvath	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]VARIABLES: $T/K = 288 - 318$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/\text{mol m}^{-3}$	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $100 w_i$ $(compiler)$ $10^4 x_i$ $(compiler)$	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $\frac{15}{25}$ $6.2 \pm 0.1$ $8.6 \pm 0.3$	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY: A. L. Horvath $100 w_i$ $10^4 x_i$ (compiler) (compiler) $6.97 \times 10^3$ 5.95 $9.78 \times 10^3$ 8.35	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $15   6.2 \pm 0.1$ $25   8.6 \pm 0.3$ $30   9.3 \pm 0.3$ $35   11.4 \pm 0.3$	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY: A. L. Horvath $100 w_i$ 10 <sup>4</sup> $x_i$ (compiler) (compiler) $6.97 \times 10^3$ 5.95 $9.78 \times 10^3$ 8.35 $1.06 \times 10^{-2}$ 9.08 $1.31 \times 10^{-2}$ 11.2	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $15   6.2 \pm 0.1$ $25   8.6 \pm 0.3$ $30   9.3 \pm 0.3$ $35   11.4 \pm 0.3$ $45   15.6 \pm 0.8$	J. Chem. Eng. Data <u>1972</u> , 17, 66-8. PREPARED BY: A. L. Horvath $100 w_i$ 10 <sup>4</sup> $x_i$ (compiler) (compiler) $6.97 \times 10^3$ 5.95 $9.78 \times 10^3$ 8.35 $1.06 \times 10^2$ 9.08 $1.31 \times 10^2$ 11.2 $1.82 \times 10^2$ 15.5	
(2) Tetrachloromethane (carbon tetrachlorode); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $15$ $6.2 \pm 0.1$ $25$ $8.6 \pm 0.3$ $30$ $9.3 \pm 0.3$ $35$ $11.4 \pm 0.3$ $45$ $15.6 \pm 0.8$ AUXILIARY II	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \end{pmatrix}$ $6.97 \times 10^3$ $5.95$ $9.78 \times 10^3$ $1.06 \times 10^{-2}$ $9.08$ $1.31 \times 10^2$ $11.2$ $1.82 \times 10^2$ 15.5	
(2) Tetrachloromethane (carbon tetrachlorode); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_{I}/mol m^{-3}$ $15$ $6.2 \pm 0.1$ $25$ $8.6 \pm 0.3$ $30$ $9.3 \pm 0.3$ $35$ $11.4 \pm 0.3$ $45$ $15.6 \pm 0.8$ AUXILIARY II METHOD/APPARATUS/PROCEDURE:	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \end{pmatrix}$ $6.97 \times 10^3$ $5.95$ $9.78 \times 10^3$ $1.06 \times 10^{-2}$ $9.08$ $1.31 \times 10^{-2}$ $1.82 \times 10^{-2}$ 15.5         NFORMATION         SOURCE AND PURITY OF MATERIALS:	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $15$ $6.2 \pm 0.1$ $25$ $8.6 \pm 0.3$ $30$ $9.3 \pm 0.3$ $35$ $11.4 \pm 0.3$ $45$ $15.6 \pm 0.8$ AUXILIARY II METHOD/APPARATUS/PROCEDURE: The saturation of tetrachloromethane with water occurred in a closed system, protected from atmospheric moisture; the saturating vessel was	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \end{pmatrix}$ $6.97 \times 10^3$ $5.95$ $9.78 \times 10^3$ $8.35$ $1.06 \times 10^2$ $9.08$ $1.31 \times 10^2$ $1.82 \times 10^2$ $15.5$ NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Distilled.         (2) Source not given. Reagent grade, purified b distillation before use.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_{I}$ /mol m <sup>-3</sup> 15 6.2 ± 0.1 25 8.6 ± 0.3 30 9.3 ± 0.3 35 11.4 ± 0.3 45 15.6 ± 0.8 AUXILIARY II METHOD/APPARATUS/PROCEDURE: The saturation of tetrachloromethane with water occurred in a closed system, protected from atmospheric moisture; the saturating vessel was placed in a constant temperature water bath. The concentration of water in tetrachloromethane was placed in a constant temperature water bath. The	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \end{pmatrix}$ $(compiler) \end{pmatrix}$ $6.97 \times 10^3$ $5.95$ $9.78 \times 10^{-1}$ $8.35$ $1.06 \times 10^{-2}$ $9.08$ $1.31 \times 10^{-2}$ $11.2$ $1.82 \times 10^{-2}$ $15.5$ NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Distilled.         (2) Source not given. Reagent grade, purified b         distillation before use.         ESTIMATED ERRORS:	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ $15$ $6.2 \pm 0.1$ $25$ $8.6 \pm 0.3$ $30$ $9.3 \pm 0.3$ $35$ $11.4 \pm 0.3$ $45$ $15.6 \pm 0.8$ AUXILIARY II METHOD/APPARATUS/PROCEDURE: The saturation of tetrachloromethane with water occurred in a closed system, protected from atmospheric moisture; the saturating vessel was placed in a constant temperature water bath. The concentration of water in tetrachloromethane was measured using the Karl Fischer titration method with a "dead stop" end-point determination (ref. 1). Fifteen observations were made at each temperature.	J. Chem. Eng. Data 1972, 17, 66-8.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \end{pmatrix}$ $6.97 \times 10^3$ $5.95$ $9.78 \times 10^3$ $1.06 \times 10^{-2}$ $9.08$ $1.31 \times 10^2$ $1.82 \times 10^2$ $15.5$ NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Distilled.         (2) Source not given. Reagent grade, purified by distillation before use.         ESTIMATED ERRORS:         Solubility:       See above.         Temperature: $\pm$ 0.1 K (compiler).	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5] VARIABLES: T/K = 288 - 318 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_{i}$ /mol m <sup>-3</sup> 15 $6.2 \pm 0.1$ 25 $8.6 \pm 0.3$ 30 $9.3 \pm 0.3$ 35 $11.4 \pm 0.3$ 45 $15.6 \pm 0.8$ AUXILIARY II METHOD/APPARATUS/PROCEDURE: The saturation of tetrachloromethane with water occurred in a closed system, protected from atmospheric moisture; the saturating vessel was placed in a constant temperature water bath. The concentration of water in tetrachloromethane was measured using the Karl Fischer titration method with a "dead stop" end-point determination (ref. 1). Fifteen observations were made at each temperature.	J. Chem. Eng. Data 1972, 17, 66-8. PREPARED BY: A. L. Horvath $\begin{pmatrix} 100 & w_i & 10^4 & x_i \\ (compiler) & (compiler) \\ 6.97 \times 10^3 & 5.95 \\ 9.78 \times 10^3 & 8.35 \\ 1.06 \times 10^2 & 9.08 \\ 1.31 \times 10^2 & 11.2 \\ 1.82 \times 10^2 & 15.5 \\ \hline NFORMATION \\ \hline SOURCE AND PURITY OF MATERIALS: (1) Distilled. (2) Source not given. Reagent grade, purified by distillation before use. ESTIMATED ERRORS: Solubility: See above. Temperature: \pm 0.1 K (compiler).REFERENCES:$	

	4		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Antropov, L. I.; Pogulyai, V. E.; Simonov, V. D.; Shamsutdinov, T. M.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Russ. J. Phys. Chem. <u>1972</u> , 46, 311-2 (VINITI No. 3739-71).		
VARIABLES:	PREPARED BY:		
T/K = 288 - 299	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 100 w <sub>1</sub> 1 (co	$\begin{array}{ccccc} 0^4 x_1 & 100 \ w_2 & 10^4 \ x_2 \\ mpiler ) & (compiler) \end{array}$		
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$\frac{92}{7.10 \times 10^{-3}}$ 6.12		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$31$ $ 8.30 \times 10^{-3}$ $7.08$		
$26   9.76 \times 10^{-3}   11$	43 $1.20 \times 10^{-2}$ 10.24		
AUXILIARY	INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Alexejev's synthetic solubility determination method was used (ref. 1). A fixed weight of tetra-	<ol> <li>Source not given. Chemically pure grade.</li> <li>Distilled.</li> </ol>		
mixture was subjected to gradually increasing temper-	ESTIMATED ERRORS:		
opalescence or clouding was the indication of the saturation temperature. The temperature was then	Solubility: Not specified.		
allowed to fall and an observation made, while the tube was constantly apitated, of the temperature of			
the first appearance of opalescence. The observation was repeated several times.	REFERENCES:		
•	(1) Alexejew, W. Ann. Phys. Chem. <u>1886</u> , 28, 305.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
<ol> <li>Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ol>	Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Tr. Mosk. KhimTekhnol. Inst. <u>1972</u> , 183-6.		
VARIABLES:	PREPARED BY:		
T/K = 298 - 373	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
t/°C Distribution coefficient	$1, 100 w_i$ $10^5 x_i$		
25.0 29200	0.00439 0.514		
<sup>1</sup> Gas-liquid system analysis parameter, from calibration measurements. The solubility was calculated from the distribution coefficient and the vapor pressure.			
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The vapor-liquid equilibrium was measured by using a modification of the Rayleigh distillation	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
method. An inert carrier gas was used for the re- moval of the vapor phase from the equilibration	ESTIMATED ERRORS:		
vessel. The analysis for tetrachloromethane in the water phase was accomplished radiometrically by	Solubility: Not specified.		
using "CI labeled compounds.	Temperature: ± 1.0 K (compiler)		

42			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Tr. Mosk. KhimTekhnol. Inst. <u>1972</u> , 183-6.		
VARIABLES:	PREPARED BY:		
T/K = 298 - 373	A. L. Horvath		
EXPERIMENTAL VALUES: (continued)	·		
t/°C Distribution coefficient <sup>1</sup> D <sub>t</sub> /dimensionless	, $100 w_i$ , $10^s x_i$ (compiler) (compiler)		
50.1 9670 70.0 4100 90.0 1960 96.2 1500	0.0364 4.26 0.1687 19.79 0.637 75.01 0.983 116.09		
100.0 1380 1.179 139.52 <sup>1</sup> Gas-liquid system analysis parameter, from calibration measurements. The solubility was calculated from the distribution coefficient and the vapor pressure.			
COMPONENTS	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii,		
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Ya. D. Tr. Mosk. KhimTekhnol. Inst. <u>1974</u> , 55-6.		
VARIABLES:	PREPARED BY:		
T/K = 298 - 343	A. L. Horvath		
EXPERIMENTAL VALUES:			
$\log_{10} \alpha = \frac{733.92}{T(K)} - 0.0161$			
where $\alpha$ = Distribution Coefficient			
At the normal boiling point of CCL, 349.79 K, $\alpha = 120$ , the activity coefficient.			
$\gamma_1 = 296$ , and the mole fraction of (1) at saturation, $x_1 = 3.378 \times 10^{-3}$ (compiler).			
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The water concentration in the organic-rich phase was determined by a radiometric method using tritium labeled water.	<ol> <li>Distilled (compiler).</li> <li>Source and purity not given.</li> </ol>		
The experimental procedure is described in more detail elsewhere (ref. 1).	ESTIMATED ERRORS:		
	Solubility: Not specified. Temperature: ± 1.0 K (compiler).		
	REFERENCES:		
	<ol> <li>Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. Khim Tekhnol. Inst. <u>1973</u>, 100.</li> </ol>		

	43	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Goldman, S.	
<ul><li>(2) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li></ul>	Can. J. Chem. <u>1974</u> , 52, 1668-80.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 313	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $c_{i}/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{ccccc} 0.942 \times 10^{-2} & 8.04 \\ 1.09 \times 10^{-2} & 9.30 \\ 1.34 \times 10^{-2} & 11.4 \\ 1.60 \times 10^{-2} & 13.6 \\ 1.86 \times 10^{-2} & 15.8 \\ 2.23 \times 10^{-2} & 19.0 \\ 2.58 \times 10^{-2} & 22.0 \end{array}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Tetrachloromethane was equilibrated with an excess of water in a vessel under rotation in a water bath for 1 hour. Aliquots of the water saturated organic solvent were taken for analysis with cali- brated Hamilton syringes. The analyses were per-	<ol> <li>Distilled.</li> <li>Source not given. Certified grade, dried with CaCl<sub>2</sub> and distilled under dry nitrogen, density = 1.5826 g/cc at 25°C.</li> </ol>	
formed with an Aquatest II automatic Karl Fischer titrator. During the analyses, the coulometer current generated by the instrument was checked with an accurate milliammeter.	ESTIMATED ERRORS: Solubility: $\pm$ 0.18 std. dev. Temperature: $\pm$ 0.02 K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ol> <li>Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ol>	Simonov, V. D.; Shamsutdinov, T. M; Pogulyai, V. E.; Popova, L. N.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Russ. J. Phys. Chem. <u>1974</u> , 48, 1573-5.	
VARIABLES:	PREPARED BY:	
T/K = 313	A. L. Horvath	
EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $w_1$ 10 <sup>5</sup> $x_1$ 100 $w_2$ 10 <sup>3</sup> $x_2$ (compiler) (compiler) 40 2.50 × 10 <sup>-2</sup> 2.93 1.97 × 10 <sup>-2</sup> 1.68		
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS		
The synthetic method of Alexejew was used. Details are found elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
	REFERENCES:	
	<ol> <li>Simonov, V. D.; Popova, L. N.; Shamsutdinov, T. M.; Pogulyai, V. I.; Memina, F. A. Symp. Doklay Neftekhimicheskoi Sektsii, Ufa, <u>1971</u>, No. 6.</li> </ol>	

COMPONEN	TS:		ORIGINAL MEASUREMENTS:	
<ul> <li>(1) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ul>		McConnell, G.; Ferguson, D. M.;	Pears	
		Endeavour <u>1975</u> , 34, 13-8.		
			VARIABLES:	:
T/K = 293			A. L. Horvath	
EXPERIMEN	TAL VALUES:		۱ <u> </u>	
	t/℃	$10^4 g_l/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	er)
	20	7.85	$7.85 \times 10^{-2}$ 9.19	
		AUXILIARY I	NFORMATION	
			SOURCE AND DUBITY OF MATE	
	ubility of tetrachk	promethane in water	(1) Source and purity not given	INIAL
was determine using an elect	ed by gas-liquid ch ron capture detect	romatography (GLC) or. Where possible,	(2) Distilled (compiler).	
identification spectrometer (	was confirmed by (MS).	a linked mass-	ESTIMATED ERRORS:	
			Solubility: Not specifie Temperature: ± 0.3 K (com	:d. piler)
COMPONEN	 TS:		ORIGINAL MEASUREMENTS:	<u> </u>
(1) Tetrachlor	romethane (carbon	tetrachloride);	Pearson, C. R.; McConnell, G.	
(2) Water; H <sub>2</sub>	20; [7732-18-5]		Proc. Roy. Soc. B, <u>1975</u> , 189, 305	-32.
VARIABLES:	· · · · · · · · · · · · · · · · · · ·		PREPARED BY:	
<i>T</i> /K = 293			A. L. Horvath	
EXPERIMEN	TAL VALUES:			
	<b>///</b> C	10° w <sub>1</sub>	$100 w_1$ $10^5 x_1$	:г)
	<i>"</i> C		(compiler) (compile	-,
	20	785	(compiler) (compile 7.85 $\times$ 10 <sup>-2</sup> 9.20	
	20	785	(compiler) (compile 7.85 × 10 <sup>-2</sup> 9.20	
	20	785 AUXILIARY II	(compiler) (compile 7.85 × 10 <sup>-2</sup> 9.20 NFORMATION	
METHOD/AP	20 20 PARATUS/PROC	785 AUXILIARY II EDURE:	(compiler) (compile 7.85 × 10 <sup>-2</sup> 9.20 NFORMATION SOURCE AND PURITY OF MATE	RIAL
METHOD/AP Saturate temperature th tracted with n- taken for GI C	20 20 PARATUS/PROC d solutions were p ermostat bath. Wa pentane and an ali analysis. The gas	785 AUXILIARY II EDURE: prepared in a constant ter samples were ex- quot of the extract chromatograph was	(compiler) (compile 7.85 × 10 <sup>-2</sup> 9.20 NFORMATION SOURCE AND PURITY OF MATE (1) Source and purity not given. (2) Distilled.	RIAL
METHOD/AP Saturate temperature the tracted with n- taken for GLC fitted with a <sup>31</sup>	20 20 PARATUS/PROC d solutions were p ermostat bath. Wa pentane and an ali analysis. The gas Ni electron capture	785 AUXILIARY II EDURE: prepared in a constant ter samples were ex- quot of the extract chromatograph was e detector.	(compiler) (compile 7.85 × 10 <sup>-2</sup> 9.20 NFORMATION SOURCE AND PURITY OF MATE (1) Source and purity not given. (2) Distilled. ESTIMATED ERRORS:	RIAL

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Kirchnerova, J.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Ph. D. Thesis, McGill Univ., Montreal, Quebec, April 1975, 280 pp.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/°</i> C <i>c<sub>1</sub>/mol</i> m <sup>-3</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
25 8.7	9.96 × 10 <sup>-3</sup> 8.50	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Binary mixtures containing 50 ml tetra- chloromethane were isopiestically equilibrated with water in an equilibration vessel which was submerg- ed in a waterbath. The equilibrium concentration of water in the binary mixtures was then determined by a Keal Eiseber distribution method. The operative is	<ol> <li>Distilled and deionized.</li> <li>Fisher Scientific Co., spectroanalyzed grade, further purified by extraction with H<sub>2</sub>SO<sub>4</sub>, dried and redistilled, 00.78 % pure.</li> </ol>	
cluded a Zeromatic II potentiometer with the dead-	ESTIMATED ERRORS:	
made at least in duplicate.	Solubility: $\pm 1 \%$ . Temperature: $\pm 0.1 $ K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Kirchnerova, J.; Cave, G. C. B.	
<ul> <li>(2) Tetrachloromethane (carbon tetrachloride);</li> <li>CCl<sub>4</sub>; [56-23-5]</li> </ul>	Can. J. Chem. <u>1976</u> , 54, 3909-16.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/℃ c₁/mol m <sup>-3</sup>	$\begin{array}{ccc} 10^4 x_i & 100 w_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25.0 8.7	8.5 $9.96 \times 10^3$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A bottle containing a mixture of 50 ml tetra- chloromethane and 6 ml water was submerged into a water thermostat bath for two days. The concentra- tion of water in the organic phase was determined by a conventional Karl Fischer deadstop backtitration. The determination was carried out in triplicate. A determined description of the complete exceeding in	<ol> <li>Distilled and deionized.</li> <li>Fisher - C199, washed with H<sub>2</sub>SO<sub>4</sub> and K<sub>2</sub>CO<sub>3</sub> solutions and finally with water. Dried over silica gel and fractionally distilled. Purity was better than 99.8 %.</li> </ol>	
given in a Ph. D. thesis (ref. 1).	ESTIMATED ERRORS:	
	Solubility: $\pm 0.2 c_r$ std. dev. Temperature: $\pm 0.1 K$ .	
	REFERENCES:	
	(1) Kirchnerova, J. Ph. D. Thesis, McGill Univ., Montreal, Quebec, <u>1975</u> .	

46		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Sato, A.; Nakijima, T.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Arch. Envir. Health <u>1979</u> , 34, 69-75.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 310	A. L. Horvath	
EXPERIMENTAL VALUES	1	
$t/^{\circ}C$ Distribution coefficient <sup>1</sup> $D_t$ /dimensionless	, $100 w_i$ , $10^5 x_i$ (compiler) (compiler)	
37 0.25	$3.78 \times 10^{-2}$ 4.43	
<sup>1</sup> Water-air system analysis parameter,	from calibration measurements.	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Tetrachloromethane vapor was equilibrated in an airtight vial with water and the overlying air. When equilibrium was reached, a portion of the	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
equilibrated air in the vessel was withdrawn using an air-tight syringe and this sample was injected into a	ESTIMATED ERRORS:	
peak height was used to calculate the distribution	Solubility: $\pm 0.04$ std. dev.	
	Temperature: ± 0.5 K (complet).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Freed, V. H.; Chiou, C. T.; Schmedding, D.;	
(2) Water; $H_2O$ ; [7732-18-5]	Environ. Health Perspect. <u>1979</u> , 30, 75-80.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/°</i> C 10 <sup>6</sup> w,	$100 w_1$ $10^5 x_1$	
	(compiler) (compiler)	
25 800	8.U × 10 <sup>-</sup> 9.38	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A known volume of water was mixed with tetrachloromethane in Erlenmeyer flasks. The flasks, with their contents magnetically stirred, were im- mersed in a water bath. Samples were removed for	<ol> <li>Source not given. Analytical standard grade, purity is greater than 95 %.</li> <li>Distilled and run through a resin column.</li> </ol>	
analyses at regular intervals. All samples were ana- lyzed by gas-liquid chromatography using an electron	ESTIMATED ERRORS:	
capture detector. The reported solubility is the average of five consecutive samples with less than 5% variation	Solubility: Not specified.	
570 variation.		

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Aref'eva, R. P.; Korenman, I. M.; Gorokhov, A. A.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	USSR Patent 672 548, <u>July 5</u> , <u>1979</u> , 3 pp. (CA. 91:113256k).		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES	1		
$t/^{\circ}C$ $g_1V_2^{-1}/kg m^{-3}$	$\begin{array}{ccc} 100 \ w_{i} & 10^{4} \ x_{i} \\ (compiler) & (compiler) \end{array}$		
20 0.87 ± 0.03	$8.72 \times 10^{-2}$ 1.02		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Equal quantities of diphenylthiocarbazone were added to aqueous solutions with different amounts of tetrachloromethane in several flasks. The contents of the flasks were shaken for 5 minutes and left to stand	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
for 1 hour. The mixtures were then filtered through a naper filter and then concentration measurements	ESTIMATED ERRORS:		
were made by spectrophotometry. The turning point on the graph of optical density versus tetrachloro- methane concentration was taken as the measurement.	Solubility: See above. Temperature: $\pm 1$ K (compiler).		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachioromethane (carbon tetrachioride); CCl <sub>4</sub> ; [56-23-5]	Coca, J.; Diaz, R. M.; Pazos, C.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Fiula Phase Equilo. <u>1980</u> , 4, 125-36.		
VARIABLES: PREPARED BY:			
<i>T/</i> K = 298 A. L. Horvath			
EXPERIMENTAL VALUES:			
t/°C 100 w <sub>1</sub> 10 (com	$x_1 = 100 w_2 = 10^4 x_2$ (compiler)		
$25 7.0 \times 10^{-2} 8.2$	$20   1.0 \times 10^{-2}   8.53$		
AUXILIARY INFORMATION			
	SOLIDCE AND DIDITY OF MATERIALS		
The mutual as while data must determine the	(1) Brokus socost further surified by distillation		
the method described by Othmer <i>et al.</i> (ref. 1). Ten cm <sup>3</sup> of tetrachloromethane was added to water from a buret and agitated until the solution became turbid.	<ol> <li>(1) Froous reagent, further purified by distillation in a heli-packing column.</li> <li>(2) Distilled.</li> </ol>		
appearance of the turbidity indicated the for- mation of a second phase. The solubility was calcu-	ESTIMATED ERRORS:		
lateu nom known densities and volumes.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).		
	REFERENCES:		
	(1) Othmer, D. F.; White, R. E.; Trueges, E. <i>Ind. Eng. Chem.</i> <u>1941</u> , 33, 1240.		

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Ksiazczak, A.; Buchowski, H.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Fluid Phase Equilib. <u>1980</u> , 5, 131–40.	
VARIABLES:	PREPARED BY:	
T/K = 323	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 10 <sup>4</sup> $x_1$	$\begin{array}{ccc} \cdot 100 \ w_i & 100 \ w_i M_j^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
50 21	$2.46 \times 10^{-2}$ $1.365 \times 10^{-3}$	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The difference between the vapor pressures of the aqueous solution and the dry tetrachloromethane as a function of the ratio of vapor and liquid vol- umes was measured in an apparatus consisting of 3	<ol> <li>Purified by repeated distillation.</li> <li>Source not given. Purified and dried before use.</li> </ol>	
tubes with mercury and a pressure adjusting device. The difference between the mercury levels in the	ESTIMATED ERRORS:	
middle tube and the reference tube gave the vapor pressure of the reference liquid. The solubility was calculated from the close of 1/AP versus V(V	Solubility: Not specified.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Balls, P. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ph. D. Thesis, Univ. of East Anglia, Norwich, U. K., July 1980, 375 pp.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ (compiler) & (compiler) \end{array}$	
$\begin{array}{cccc} 10 & 0.75 \pm 0.06 \\ 15 & 0.90 \pm 0.07 \\ 20 & 1.05 \pm 0.07 \end{array}$	$\begin{array}{cccc} 6.54 \times 10^2 & 7.66 \\ 6.84 \times 10^2 & 8.02 \\ 7.28 \times 10^2 & 8.53 \end{array}$	
25 1.30 ± 0.10	$7.24 \times 10^{-2}$ 8.48	
AUXILIARY IN	IFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The measurements involved repeated equilibra- tions of nitrogen with a water sample containing tetrachloromethane. The mixture was shaken vigor- ously at the required temperature in a constant	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
phase was then injected into a gas chromatograph for analysis. The experiments were performed at least	ESTIMATED ERRORS:	
three times at 10, 15, 20, and 25°C. Each run at constant temperature involved about six equilibra- tions.	Solubility: See above. Temperature: ± 0.5 K (compiler).	

----

	49	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Techn. <u>1980</u> , 14, 1227-9.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C <i>c</i> <sub>1</sub> /mol m <sup>-3</sup>	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ (compiler) & (compiler) \end{array}$	
25 4.92	$7.58 \times 10^2$ 8.88	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of tetrachloromethane was added to a stainless steel tube containing water and then the tube was sealed. It was allowed to equilibrate during	<ol> <li>New England Nuclear, used as received.</li> <li>Distilled.</li> </ol>	
shaking for a week at 25°C. After centrifugation the concentration was obtained from a measurement of	ESTIMATED ERRORS:	
solubility procedure was carried out at least twice for each sample and the analysis was done in duplicate.	Solubility: $\pm 2.5$ % std. dev. Temperature: $\pm 0.3$ K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ul> <li>(1) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> </ul>	Veith, G. D.; Macek, K. J.; Petrucelli, S. R.; Carroll, J.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Proc. 3 <sup>rd</sup> Ann. Symp. on Aquatic Toxicology, ASTM Publ. 707, Philadelphia, <u>1980</u> , 116-29.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
$t/^{\circ}$ C $n_1 V_2^{-1}/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
20 5.01	$7.71 \times 10^{-2}$ 9.04	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of tetrachloromethane was added to $10 \text{ cm}^3$ distilled water in a 50 cm <sup>3</sup> flask. The mixture was magnetically stirred in a constant temperature	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
water bath. The mixture was analyzed by using radio- active techniques. The ${}^{14}C$ activity of the water	ESTIMATED ERRORS:	
samples was measured in a liquid scintillation spec- trometer by recording the number of counts per minute.	Solubility: Not specified. Temperature: $\pm 1$ K.	

50			
COMPONENTS:		ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]		Wu, X.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]		Huaxue Shiji <u>1981</u> , 221-4.	
VARIABLES:		PREPARED BY:	
<i>T</i> /K = 278 - 348		A. L. Horvath	
EXPERIMENTAL V	ALUES:		
	t/°C 10 <sup>6</sup> w <sub>1</sub>	$\begin{array}{ccc} 100 \ w_i & 10 \\ (compiler) & (compiler) \end{array}$	⁴ <i>x,</i> npiler)
	5       42         10       46         15       67         20       92         25       109         30       123         35       144         40       190         45       201         50       232         55       250         60       299         65       324         70       380         75       408	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	8.59 8.93 5.72 7.85 9.30 9.5 2.3 5.2 7.1 9.8 1.3 5.5 7.6 2.4 8.7
	AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF M	ATERIALS:
A known amount of water was added to dried tetrachloromethane to make a series of standard sam- ples for preparing calibration curves and obtaining the saturated solutions at each temperature. The deter- mination of the water content of the tetrachloro- methane was made with an RM-250 NMR spectrome- ter (radiation frequency 250 MHz). The water peak amplitude was the mean of four readings taken at a particular temperature.		<ol> <li>Distilled.</li> <li>Source not given. Grade 2 with KOH and H<sub>2</sub>SO<sub>4</sub> solu water before dried on mol</li> </ol>	purity, washed tions and distilled ecular sieve.
		ESTIMATED ERRORS: Solubility: $\pm$ 28 std. Temperature: $\pm$ 1 - 2 K.	dev.

-	51	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Leighton, D. T.; Calo, J. M.	
(2) Water; $H_2O$ ; [7732-18-5]	J. Chem. Eng. Data <u>1981</u> , 26, 382-5.	
VARIABLES:	PREPARED BY:	
T/K = 274 - 300	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
t/°C Distribution coefficient <sup>1</sup> D <sub>L</sub> /dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
<sup>1</sup> Gas-liquid system analysis parameter,	from calibration measurements.	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A 5 $\mu$ L tetrachloromethane sample was inject- ed into a 2.3 liter equilibration cell containing distilled water. After the cell was shaken vigorously for about 5 minutes, the homogeneity of the liquid sample was maintained with a magnetic stirrer. Compressed air was passed through the cell and the gas flow was measured with a soap film flowmeter. The tetrachloromethane content was extracted and analyzed using a dual flame ionization detector gas chromatograph.	<ul> <li>(1) Source and purity not given.</li> <li>(2) Distilled.</li> <li>ESTIMATED ERRORS:</li> <li>Solubility: ± 3.4 %.</li> <li>Temperature: ± 0.5 K.</li> </ul>	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ol> <li>(1) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	Mackay, D.; et al. Volatilization of Organic Pollutants from Water, U. S. EPA Report 600/3-82-019, Athens, Georgia, <u>1982</u> (PB 82-230939).	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t^{\circ}C$ $c_1$ /mol m <sup>-3</sup> 100 $w_1$ 10 <sup>5</sup> $x_1$ 100 $w_2$ 10 <sup>4</sup> $x_2$ (compiler) (compiler) (compiler)		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	9.376 9.200 $8.20 \times 10^{-3}$ 7.0	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An aqueous solution with an excess amount of tetrachloromethane was stirred for one day before being introduced into a 1 liter glass vessel through a sintered glass disk. The exit gas flow rate was meas- ured by a soap bubble flow meter. The concentration of tetrachloromethane in water was determined by gas chromatography. The GC was accurated with beth of	<ol> <li>Source and purity not given.</li> <li>Doubly distilled.</li> <li>ESTIMATED ERRORS:</li> </ol>	
dual flame ionization detector and an electron capture detector.	Solubility: Not specified. Temperature: ± 1.0 K.	

52		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Ohtsuka, K.; Kazama, K.	
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Sen'i Seihin Shohi Kagaku Kaishi <u>1982</u> , 22, 197-201.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
25 0.22	$1.4 \times 10^{-2}$ 1.19	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Water was added gradually to 50 ml tetra- chloromethane in a flask which was then lowered into a thermostat bath. The flask was shaken vigorously	<ol> <li>Distilled (compiler).</li> <li>Commercial JTS extra pure reagent. Further purified by conventional methods.</li> </ol>	
until the first cloud (turbidity) appeared. The sample water content was determined by the Karl Fischer ti-	ESTIMATED ERRORS:	
tration method.	Solubility: Not specified.	
	Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S.	
CCI <sub>4</sub> ; [56-23-5]	Tellus <u>1983</u> , 35B, 170-6.	
(2) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 278 - 306	A. L. Horvath	
EXPERIMENTAL VALUES: <i>t</i> /°C Henry's law constant, <i>H</i> /dimensionless	$\begin{array}{ccc} 100 \ w, & 10^5 \ x, \\ (compiler) & (compiler) \end{array}$	
5 0.4864	$8.006 \times 10^2$ 9.383	
10 0.5854 15 0.7000	$\begin{array}{cccc} 8.420 \times 10^2 & 9.869 \\ 8.823 \times 10^2 & 10.34 \end{array}$	
20 0.8320 25 0.9831	$\begin{array}{cccc} 9.206 \times 10^2 & 10.79 \\ 9.584 \times 10^2 & 11.23 \end{array}$	
33 1.2696	$1.0554 \times 10^{-1}$ 12.37	
Ine Henry's law constant (H) was derived from the best-fit lines of van't Hoff plots: $\log_2 H = -\frac{2918}{100} + 9.77$		
$\overline{T(K)} = \overline{T(K)}$		
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
to measure the Henry's law constant as described by McAuliffe (ref. 1). A known volume of water in a	<ol> <li>Source and purity not specified.</li> <li>Distilled.</li> </ol>	
syringe was shaken with a known volume of pure tetrachloromethane. After equilibration of the sample.	ESTIMATED ERRORS:	
the headspace was separated and analyzed for the compound of interest. A gas chromatograph fitted with an electron capture detector was used for the	Solubility: $\pm$ 5.5 % std. dev. Temperature: $\pm$ 0.2 K.	
analysis.	REFERENCES:	
	(1) McAuliffe, C. D. Chem. Techn. 1971. 1, 46.	

	53		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Orlandini, M.; Fermeglia, M.; Kikic, I.; Alessi, P.		
(2) Water; $H_2O$ ; [7732-18-5]	Chem. Eng. J. <u>1983</u> , 26, 245-50.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:	1		
t/°C 10 <sup>4</sup> x <sub>1</sub> 10 (con	$\begin{array}{cccc} 0 & w_1 & 10^4 & x_2 & 100 & w_2 \\ \text{apiler} & & & (\text{compiler}) \end{array}$		
20 0.9 7.68	$\times 10^{-2}$ 7.1 8.322 $\times 10^{-3}$		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE: A mixture of water and tetrachloromethane was agitated overnight and left at equilibrium for 24 hours. Three samples were withdrawn from each phase with a syringe and analyzed by gas chro- matography. The gas chromatograph (C. Erba ATc/t)	<ul> <li>SOURCE AND PURITY OF MATERIALS:</li> <li>(1) Fluka reagent, purified by distillation before use.</li> <li>(2) Distilled (compiler).</li> </ul>		
was equipped with a Shimadzu Chromatopac E-1A integrator. A stainless column packed with 100-120 mesh Chromosorb W DMCD coated with hepta- decane nitrile was used for analysis.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.1 K.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Munz, C. D.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ph. D. Thesis, Stanford Univ., Stanford, CA., <u>1985</u> , 306 pp.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 283 - 303	A. L. Horvath		
EXPERIMENTAL VALUES:	<u> </u>		
t/°C Henry's Law Constant, $\rho_1/kg \text{ m}^{-3}$ 100 $w_1$ 10 <sup>5</sup> $x_1$ , H/dimensionless (compiler) (compiler)			
10         0.609         0.810           20         0.980         0.78           30         1.530         0.75	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
The multiple equilibrium technique with direct	(1) Aldrich Chemicals Co. WI > 99.5 % pure		
aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law constant. A known volume of liquid and gas was introduced	(2) Distilled and deionized.		
into a syringe and allowed to equilibrate. The ratio of gas to liquid volume was maintained constant. While	ESTIMATED ERRORS:		
tograph, the liquid samples were pre-concentrated by liquid-liquid extraction before analysis. The ex- periments were carried out in duplicates.	Solubility: $\pm 1.28 \times 10^{-2}$ std. dev. Temperature: $\pm 0.5$ K.		

B
54				
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Tetrachloromethane (carbon tetrachloride);	Yoshioka, Y.; Ose, Y.; Sato, T.			
(2) Water; $H_2O$ ; [7732-18-5]	Ecotoxicol. Envir. Saf. <u>1986</u> , 12, 15-21.			
VARIABLES:	PREPARED BY:			
<i>T</i> /K = 293	A. L. Horvath			
EXPERIMENTAL VALUES				
$t/^{\circ}C$ $\rho_{i}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^7 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
20 $2.6 \times 10^{-3}$ <sup>1</sup>	2.6 × 10 <sup>-4</sup> 3.04			
<sup>1</sup> Cited value is in error (compiler).				
	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A supersaturated solution of tetrachloro- methane in water was agitated for two hours at 20°C. After storing for 2 hours the water phase was filtered through Whatman no 2 filter paper before analysis	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>			
The concentration of tetrachloromethane in water was determined by gas chromatography or by ultraviolet	ESTIMATED ERRORS:			
spectroscopy.	Solubility: Not specified. Temperature: $\pm 1.0$ K.			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Tetrachloromethane (carbon tetrachloride);	Lo, J. M.; Tseng, C. L.; Yang, J. Y.			
(2) Water: $H_{2}O$ : [7732-18-5]	Anal. Chem. <u>1986</u> , 58, 1596-7.			
VARIABLES:	PREPARED BY:			
T/K = 298	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C 100 w <sub>1</sub>	$10^{5} x_{1}$ $100 w_{1} M_{1}^{-1} / \text{mol g}^{-1}$ (compiler)			
25 $8.0 \times 10^{-2}$	9.38 5.20 × 10 <sup>-4</sup>			
AUXILIARY I	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A 5 ml tetrachloromethane sample containing a labeled complex, cobalt(III) pyrrolidinecarbodithio- ate, was placed together with 1000 ml water into a separation funnel. After rigorous shaking of the	<ol> <li>E. Merck Co., used as received.</li> <li>Demineralized and then boiled at reduced pressure.</li> </ol>			
minutes before separation of the two phases. Sam- oles were withdrawn for activity measurement with	ESTIMATED ERRORS:			
NaI (TI) scintillation detector. Several factors were adjusted to obtain accurate measurements.	Solubility: $\pm$ 0.7 × 10-2 std. dev.Temperature: $\pm$ 1.0 K (compiler).			

	55		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Gossett, J. M.		
(2) Water; $H_2O$ ; [7732-18-5]	Environ. Sci. Tehn. <u>1977</u> , 21, 202-8.		
VARIABLES:	PREPARED BY:		
T/K = 283 - 308	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
t/°C Henry's Law Constant, CV <sup>1</sup> H/m <sup>3</sup> atm mol <sup>-1</sup> %	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$		
$\begin{array}{ccccccc} 10.0 & 1.32 \times 10^2 & 5.15 \\ 17.5 & 2.11 \times 10^2 & 4.20 \\ 24.8 & 3.04 \times 10^2 & 3.92 \\ 34.6 & 4.60 \times 10^{-2} & 3.42 \end{array}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
$^{1}$ CV = coefficient of variation (= 1	00 S.D./mean).		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A modification of the EPICS procedure was used for measuring Henry's law constants. A precise quantity of tetrachloromethane was injected into serum bottles which contained distilled water. The bottles were incubated for 18 - 24 hours at four	<ol> <li>J. T. Baker reagent, analytical grade. Used as received.</li> <li>Distilled.</li> </ol>		
desired temperatures in a reciprocating shaker bath. The headspace concentrations of the EPICS bottles were measured by a gas chromatograph which was equipped with a flame ionization detector. The mean of the coefficient of variation values was approxi- mately 4.3 %.	ESTIMATED ERRORS: Solubility: See above. Temperature: ± 0.1 K.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Kleeberg, H.; Klein, D.; Luck, W. A. P.		
<ul><li>(2) Tetrachloromethane (carbon tetrachloride); CCl<sub>4</sub>; [56-23-5]</li></ul>	ChemIngTechn. <u>1987</u> , 58, 409-11.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C $c_1/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
25 9.3	$1.68 \times 10^{-2}$ 1.43		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Samples of solutions of different composition were prepared by mixing the pure components at room temperature. The samples were incubated for	<ol> <li>(1) Distilled.</li> <li>(2) Source and purity not specified.</li> </ol>		
was determined by Karl Fischer titration.	ESTIMATED ERRORS:		
	Solubility: Not specified. Temperature: $\pm 1.0$ K.		

56			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Warner, H. P.; Cohen, J. M.; Ireland, J. C.		
(2) Water; $H_2O$ ; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants, EPA Technical Report, PB 87-212684, Cincinnati, OH., July 1987.		
VARIABLES:	PREPARED BY:		
<i>TI</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ (compiler) & (compiler) \end{array}$		
24.85 $3.02 \times 10^{-2}$	$7.61 \times 10^{-2}$ 8.92		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The original method and apparatus for the determination of Henry's law constant, as described by Mackay <i>et al.</i> (ref. 1), was used. The general procedure was to add an excess quantity of tetra- chloromethane to distilled deionized water, place the	<ol> <li>Source not given. Purest quality available was used without further purification. Stated purity &gt; 99 %.</li> <li>Distilled and deionized.</li> </ol>		
sample in a thermostat bath, and provide overnight mixing. A portion of this solution was returned to the stripping vessel. Tetrachloromethane was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from the log of the concentration versus time plot. The	ESTIMATED ERRORS: Solubility: ± 6 % std. dev. Temperature: ± 0.05 K.		
experimental data values are averages of two or more replicates.	<ul> <li>REFERENCES:</li> <li>(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Techn. <u>1979</u>, 13, 333.</li> </ul>		
(1) Tetrachloromethane (carbon tetrachloride);	Howe, G. B.; Mullins, M. E.; Rogers, T. N.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	AFESC Tyndall Air Force Base, Report ESL-TR- 86-66, Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp (AD-A188 571).		
VARIABLES:	PREPARED BY:		
T/K = 283 - 303	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C 10 <sup>6</sup> $w_{1}$	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
10         104           20         146           30         105	$\begin{array}{cccccccc} 1.04 \ \times \ 10^{-2} & 1.218 \\ 1.46 \ \times \ 10^{-2} & 1.710 \\ 1.05 \ \times \ 10^{-2} & 1.230 \end{array}$		
AUXILIARY II	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
250 ml bottles were filled with distilled and deionized water and sealed. A measured volume of tetrachloromethane was injected into the bottles through each septum using a microliter syringe. Tetrachloromethane was in excess of the anticipated	<ol> <li>Source not given. Probably a commercial reagent, at least 99 % purity. Used as received.</li> <li>Distilled and deionized.</li> </ol>		
solubility limit. The bottles were shaken for one hour with a wristaction shaker and allowed to equilibrate for about 3 weeks. Samples were then injected into a gas chromatograph equipped with a Carbopack col- umn and a FID detector. The GC response was com- pared with calibration plots.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).		

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Vogel, A. I., rev. by Furniss, B. S.; et al.		
(2) Water; $H_2O$ ; [7732-18-5]	Vogel's Textbook of Practical Organic Chemistry, 5 <sup>th</sup> ed., Longman, London, <u>1989</u> , p. 1442.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 100 w <sub>1</sub>	$10^{5} x_{j}$ $100 w_{j} M_{j}^{-j} / \text{mol g}^{-1}$		
$25.0 8.0 \times 10^{-2}$	(compiler) (compiler) 9.38 $5.2 \times 10^{-4}$		
AUXILIARY II	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The solubility determination was carried out in small test tubes (e. g. 100x12 mm) to permit vigorous shaking of the tetrachloromethane and water mix-tures. During the experiments 0.20 ml tetrachloro-	<ol> <li>Source not given. Analytical reagent, treated with KOH solution and dried by distillation.</li> <li>Distilled.</li> </ol>		
methane was added to 3.0 ml of water and agitated. The concentration of tetrachloromethane in water was	ESTIMATED ERRORS:		
graphic column.	Solubility: Not specified. Temperature: $\pm$ 1.0 K (compiler).		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Wright, D. A.; Sandler, S. I.; DeVoll, D.		
(2) Water; $H_2O$ ; [7732-18-5]	Environ. Sci. Techn. <u>1992</u> , 26, 1828-31.		
VARIABLES:	PREPARED BY:		
<i>TT</i> K = 293 - 313	A. L. Horvath		
EXPERIMENTAL VALUES: $t/^{\circ}C \qquad \gamma^{\circ} \qquad 10^{5} x_{i}$	100 w, 100 w, $M_{1}^{-1}$ /mole g <sup>-1</sup> (compiler) (compiler)		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$		
AUXILIARY IN	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A differential static cell equilibrium apparatus was used to measure the infinite dilution activity coefficient of tetrachloromethane ( $\gamma^{\infty}$ ) in water. Cells containing degassed water were submerged in a ther-	<ol> <li>Source and purity not given.</li> <li>Distilled, filtered and deionized.</li> </ol>		
mostated water bath. Tetrachloromethane was injected into the mixture cells and a magnetic stirrer was	ESTIMATED ERRORS:		
turned on. The cells were allowed to equilibrate. The stirrer was then turned of f and the differential pressure was recorded. The experiment was repeated at least three times at each temperature.	Solubility: See above. Temperature: ± 0.05 K.		

58			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Munz, C. D.		
(2) MQ-Water	Ph. D. Thesis, Stanford University, Stanford, CA., <u>1985</u> , 306 pp.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:	·		
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ (compiler) & (compiler) \end{array}$		
20 0.980	$7.80 \times 10^{-2}$ 9.145		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The multiple equilibrium technique with direct aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law constant. A known volume of liquid and gas was introduced in-	<ol> <li>Aldrich Chemicals Co., WI, &gt; 99.5 % pure.</li> <li>0.25 mg total organic carbon per liter and 0.75 mg COD per liter.</li> </ol>		
to a syringe and allowed to equilibrate. The ratio of gas to liquid volume was maintained constant. While	ESTIMATED ERRORS:		
vapor samples were injected direct into a gas chro- matograph, the liquid samples were pre-concentrated by liquid-liquid extraction before analysis. The experiments were carried out in duplicates.	Solubility: $\pm 0.1 \times 10^{-3}$ std. dev. Temperature: $\pm 0.5$ K.		
COMPONENTS	ORIGINAL MEASUREMENTS		
(1) Tetrachloromethane (carbon tetrachloride):	Munz, C. D.		
CCl₄; [56-23-5]	Ph. D. Thesis, Stanford University, Stanford, CA.,		
(2) Municipal tap water (PASE)	<u>1985</u> , 306 pp.		
VARIABLES:	PREPARED BY:		
T/K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
20 0.958	7.996 × 10 <sup>-2</sup> 9.372		
AUXILIARY II	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The multiple equilibrium technique with direct aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law constant. A known volume of liquid and gas was introduced in-	<ol> <li>Aldrich Chemicals Co., WI, &gt; 99.5 % pure.</li> <li>10 mg total organic carbon per liter and 40 mg COD per liter.</li> </ol>		
to a syringe and allowed to equilibrate. The ratio of gas to liquid volume was maintained constant. While vapor samples were instand direct into a case above	ESTIMATED ERRORS:		
matograph, the liquid samples were pre-concentrated by liquid-liquid extraction before analysis. The experiments were carried out in duplicates.	Solubility: $\pm 1.7 \times 10^{-2}$ std. dev. Temperature: $\pm 0.5$ K.		

	59		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Liss, P. S.; Slater, P. G.		
(2) Seawater	Nature <u>1974</u> , 247, 181-4.		
(-)	1		
VARIABLES:	PREPARED BY:		
T/K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant <sup>1</sup> , H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
20 1.08	$7.22 \times 10^{-2}$ 8.46		
<sup>1</sup> Obtained from [mass(1) cm <sup>-3</sup> (air)/mass	s(1) cm <sup>-3</sup> (seawater)].		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Details are not available except that the data originate elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Samples from Atlantic.</li> </ol>		
	ESTIMATED ERRORS:		
	Solubility: Not specified.		
	Temperature: $\pm 2 \text{ K}$ (compiler).		
	REFERENCES:		
	(1) Lovelock, J. E. Personal communication.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Balls, P. W.		
(2) Seawater	Ph. D. Thesis, Univ. of East Anglia, Norwich, U. K., July <u>1980</u> , 375 pp.		
T/K = 283 - 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_1 & 10^3 \ x_1 \\ (compiler) & (compiler) \end{array}$		
$\begin{array}{cccc} 10 & 0.82 \pm 0.01 \\ 15 & 0.95 \pm 0.10 \end{array}$	$6.19 \times 10^{-2}$ 7.25 $6.68 \times 10^{-2}$ 7.83		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
_			
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The method involved repeated equilibrations of nitrogen with a water sample containing tetrachloro- methane. The sample was shaken vigorously at the required temperature in a constant temperature both	<ol> <li>Source and purity not given.</li> <li>Salinity = 20 %.</li> </ol>		
for 30 minutes. The equilibrated gas phase was then injected into a gas chromatograph. The experiments	ESTIMATED ERRORS:		
were performed at least three times at 10, 15, 20, and 25°C, each experiment involving about six equili- brations.	Solubility: See above. Temperature: $\pm 0.5$ K (compiler)		

60			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S.		
(2) Securities	Tellus <u>1983</u> , 35B, 170-6.		
(2) Seawater			
VARIABLES:	PREPARED BY:		
T/K = 278 - 306	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_1 & 10^5 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
5 0.7100	$5.596 \times 10^{-2}$ 6.557		
10 0.8716 15 1.0624	$5.759 \times 10^{-2} \qquad 6.748$ 5.911 $\times 10^{-2} \qquad 6.927$		
20 1.2007 25 1.5473 23 2.0536	$6.180 \times 10^{-2}$ 7.066 $6.180 \times 10^{-2}$ 7.242 $6.625 \times 10^{-2}$ 7.53		
35 2.050	0.025 × 10 7.705		
The Henry's law constants (H) were der	ived from the best-fit lines of van't Hoff plots:		
$\log_e H = -\frac{3230}{777} + 12$	1.27		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The multiple equilibrium technique was used to measure the Henry's law constant ( $H$ ) as described by McAuliffe (ref. 1). A known volume of seawater in a surface was that a backen with a known volume of the seawater.	<ol> <li>Source and purity not given.</li> <li>Samples from 48° N to 65° S in the Atlantic in late 1981.</li> </ol>		
chloromethane. After equilibration, the headspace was separated and analyzed for tetrachloromethane. A	ESTIMATED ERRORS:		
gas chromatograph with an electron capture detector was used for the analysis.	Solubility: $\pm$ 5.5 % std. dev.		
	Temperature: $\pm$ 0.2 K.		
	REFERENCES:		
	(1) McAuliffe C D <i>Chem Techn</i> 1971 1 46		
	(-,		

							61
COMPONENTS:				ORIC	GINAL MEASURE	EMENTS:	
(1) Tetrachloromethane (carbon tetrachloride); CCL: [56-23-5]			Liu,	JL.; Huang, T	С.		
(2) Calcium chloride: CaCl <sub>2</sub> ; [10043-52-4]			Scier	utia Sinica (Peking	) <u>1961</u> , <i>10</i> , 700-10.		
(3) Water; $H_2O$ ;	[7732-18-5]						
VARIABLES:				PREF	ARED BY:		
1/K = 298 Concentration				A. L	. Horvath		
EXPERIMENTAL	VALUES:						
	t/⁰C	<i>c</i> <sub>2</sub> /mol dm <sup>-3</sup>	ρ <sub>l</sub> /kg	; m <sup>-3</sup>	100 w	$10^{5} x_{1}$	
	25	0.005	0.4		(compiler)	(compiler)	
	25	0.205 0.280	0.0	552 598	$6.41 \times 10^{-2}$ $5.85 \times 10^{-2}$ $5.40 \times 10^{-2}$	7.66 7.03	
		0.581	0.2	452	$4.29 \times 10^{-2}$	5.31	
		AUXIL	IARY IN	IFORM	IATION		
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:					
An accurate amount of tetrachloromethane was transferred into an Erlenmeyer flask containing a weighed amount of aqueous solution. The flask was rotated in a thermostat bath until equilibrium was established. The optical densities of different concentration of tetrachloromethane in the aqueous solutions were measured using water as the reference liquid. Duplicate experiments were made and the results averaged. The absorption spectra of tetra- chloromethane in aqueous calcium chloride solutions were determined using a Hilger spectrophotometer.		(1) (2) (3)	Peking Chemical washing with H <sub>2</sub> S over CaCl <sub>2</sub> and fr Peking Chemical before use. Distilled.	Factory, further purified b O <sub>4</sub> , NaOH and water; drie actionated. Factory, recrystallized twie	y ce		
		ra- utions eter.	ESTII Solut Temp	MATED ERRORS bility: perature: ±	: Not specified. 0.5 K (compiler).		

62		
COMPONENTS:		ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbor CCl <sub>4</sub> ; [56-23-5]	n tetrachloride);	Svetlanov, E. B.; Velichko, S. M. Levinskii, M. I.; Treger, Yu. A.; Flid, R. M.
(2) Hydrogen chloride; HCl; [	7647-01-0]	Russ. J. Phys. Chem. <u>1971</u> , 45, 488-90.
(3) Water; H <sub>2</sub> O; [7732-18-5]		
VARIABLES:		PREPARED BY:
T/K = 288 - 333 Concentration		A. L. Horvath
EXPERIMENTAL VALUES:		
Concentration of HCl, $100 w_2$	t/℃ 1000 w	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
10	15         1.00           30         0.55           45         0.30           60         0.18	$\begin{array}{ccccc} 1.00 \times 10^{-1} & 12.35 \\ 5.5 \times 10^{-2} & 6.79 \\ 3.0 \times 10^{-2} & 3.70 \\ 1.8 \times 10^{-2} & 2.22 \end{array}$
20	15       1.25         30       0.74         45       0.40         60       0.23	$\begin{array}{ccccc} 1.25 \times 10^{-1} & 16.31 \\ 7.4 \times 10^{-2} & 9.65 \\ 4.0 \times 10^{-2} & 5.21 \\ 2.3 \times 10^{-2} & 3.00 \end{array}$
	AUXILIARY	INFORMATION
METHOD/APPARATUS/PROC		SOURCE AND PURITY OF MATERIALS
The dynamic method was	used for the determi-	(1) Source and purity not given.
nation of solubility. It involved nitrogen with terachloromethane then bubbled through a thermos	the saturation of vapor which was tatically controlled	<ul> <li>Source and purity not given.</li> <li>Distilled (compiler)</li> </ul>
solution. After 2 - 3 hours the s	solution became	ESTIMATED ERRORS:
saturated and samples were taken periodically (ref. 1). The concentration of tetrachloromethane in the aqueous solution was determined by gas chromatogra- nhy. On average $3 - 5$ measurements were taken at		Solubility: $\pm 15 \%$ . Temperature: $\pm 0.05 $ K.
each temperature.		REFERENCES:
		(1) Treger, Yu. A.; Flid, R. M.; Spector, S. S. Russ. J. Phys. Chem. <u>1964</u> , 38, 253.

							63
COMPONENTS:			ORIC	SINAL MEASUR	EMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);			Liu,	JL.; Huang, T	-C.		
(2) Lithium chloride: LiCl: $[7447-41-8]$			Scier	utia Sinica (Peking	g) <u>1961,</u> <i>10</i> , 700-10.		
(2) Eltinum ellie (3) Water: H <sub>2</sub> O:	[7732-18-5]	[141 41 0]					
(0) ** 100, 02,00,							
VARIABLES:				PREP	ARED BY:		
T/K = 293 Concentration				A. L	. Horvath		
EXPERIMENTA	L VALUES						
	t/⁰C	c <sub>2</sub> /mol dm <sup>-3</sup>	ρ <sub>l</sub> /kg	m <sup>-3</sup>	100 w, (compiler)	10 <sup>s</sup> x, (compiler)	
	20	0.1867 0.3496 0.4352 0.7015 0.9765	0.0 0.6 0.6 0.6	717 596 576 544 509	$\begin{array}{rrrr} 7.13 \ \times \ 10^{-2} \\ 6.86 \ \times \ 10^{-2} \\ 6.69 \ \times \ 10^{-2} \\ 6.33 \ \times \ 10^{-2} \\ 5.94 \ \times \ 10^{-2} \end{array}$	8.389 8.108 7.924 7.541 7.130	
		AUXIL	IARY IN	IFORM	IATION		
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:					
An accurate amount of tetrachloromethane was transferred into an Erlenmeyer flask containing a weighed amount of aqueous solution. The flask was rotated in a thermostat bath until the equilibrium was established. The optical densities of different con- centrations of tetrachloromethane in the aqueous solution were measured using water as the reference liquid. Duplicate experiments were done and the re- sults averaged. The absorption spectra of tetra- chloromethane in aqueous lithium chloride solutions were determined using a Hilger spectrophotometer.		(1) (2) (3)	Peking Chemical further purified b NaOH and water fractionated. Merck reagent, r crystallized. Distilled.	Factory, by washing with $H_2SO_4$ , ; dried over CaCl <sub>2</sub> and nixed with HCl and re-			
		ESTIN Solut Temp	MATED ERRORS bility: perature: ±	S: Not specified. - 0.5 K (compiler).			
							_

64		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Gross, P. M.	
(2) Magnesium sulfate: $MgSQ \cdot [7487-88-9]$	Z. phys. Chem. <u>1929</u> , 6B, 215-20.	
(2) Water: H <sub>2</sub> O: [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
One concentration		
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ 100	$g_i/g_j$ 100 $w_i$ 10 <sup>5</sup> $x_i$	
25 0.5 4.8 ×	$10^2$ 4.8 × $10^2$ 5.93	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A mixture of 10 to 20 ml tetrachloromethane in 500 ml of an aqueous solution of $MgSO_4$ was placed in a water thermostat bath. The sample con- tainer was placed on a shaker and agitated until no	<ol> <li>Source not given. Commercial reagent, fractionally distilled before use.</li> <li>Kahlbaum chemical, certified grade.</li> <li>Distilled.</li> </ol>	
more liquid droplets remained undissolved. One or more samples were syphoned out and the concentra-	ESTIMATED ERRORS:	
tion of tetrachloromethane was determined using a Zeiss water interferometer.	Solubility: $\pm 5\%$ .	
	Temperature: $\pm 0.01$ K.	
COMPONENTS	ORIGINAL MEASUREMENTS	
(1) Tetrachloromethane (carbon tetrachloride):	Liu. JL.: Huang. TC.	
CCI <sub>4</sub> ; [56-23-5]	Sciencia Sinica (Peking) <u>1961</u> , 10, 700-10.	
(2) Magnesium sulfate; MgSO <sub>4</sub> ; [7487-88-9]		
(3) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 298 Concentration	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ $\rho_1/k_2$	$g m^{-3}$ 100 $w_i$ 10 <sup>5</sup> $x_i$ (compiler) (compiler)	
25 0.199 0.	$537   6.21 \times 10^2   7.429$	
0.318 0. 0.433 0.	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
0.537 0.4	487 4.58 × 10 <sup>-2</sup> 5.679	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An accurate amount of tetrachloromethane was transferred into an Erlenmeyer flask containing a	(1) Peking Chemical Factory, further purified by	
weighed amount of aqueous solution. The flask was	over CaCl <sub>2</sub> and fractionated.	
established. The optical densities of different con- centration of tetrachloromethane in the aqueous	(3) Distilled.	
solutions were measured using water as the reference liquid. Duplicate experiments were made and the	ESTIMATED ERRORS:	
results averaged. The absorption spectra of tetra- chloromethane in aqueous magnesium sulfate solu-	Solubility: Not specified.	
tions were determined using a Hilger spectro- photometer.	Temperature: $\pm 0.5$ K (compiler).	

,

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride);	Gross, P. M.		
(2) Potassium chloride; KCl; [7447-40-7]	Z. phys. Chem. <u>1929</u> , 6B, 215-20.		
(3) Water; H <sub>2</sub> O; [7732-18-5]			
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 298	A. L. Horvath		
One concentration			
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ 100	$g_1/g_3$ 100 w, 10 <sup>5</sup> x <sub>1</sub> (compiler) (compiler)		
25 0.5 6.5 ×	$10^{-2}$ 6.5 × $10^{-2}$ 7.84		
AUXILIARY I	NFORMATION		
	SOURCE AND DUDITY OF MATERIALS.		
A mixture of 10 to 20 ml tetrachloromethane	(1) Source not given Commercial reagent		
in 500 ml of an aqueous solution of KCl was placed in a water thermostat bath. The sample container was placed on a shaker and agitated until no more liquid	<ul> <li>(1) Source hot given. Commercial reagent, fractionally distilled before use.</li> <li>(2) Kahlbaum chemical, certified grade.</li> <li>(3) Distilled.</li> </ul>		
droplets remained undissolved. One or more samples were syphoned out and the concentration of tetra-	ESTIMATED ERRORS:		
chloromethane was determined using a Zeiss water interferometer.	Solubility: $\pm 5\%$ .		
	Temperature: $\pm$ 0.01 K.		
COMPONENTS	ORIGINAL MEASUREMENTS:		
(1) Tetrachloromethane (carbon tetrachloride):	Lin, L-L: Huang, TC.		
CCl <sub>4</sub> ; [56-23-5]	Scientia Sinica (Peking) 1961, 10, 700-10.		
(2) Potassium chloride; KCl; [7447-40-7]			
(3) Water; H <sub>2</sub> O; [7732-18-5]			
VARIABLES:	PREPARED BY:		
T/K = 298 Concentration	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ $\rho_1/\text{kg}$	$m^{-3}$ 100 $w_i$ 10 <sup>5</sup> $x_i$ (compiler) (compiler)		
25 0.3032 0.	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
0.7232 0.0	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
AUXILIARY II	VFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
An accurate amount of tetrachloromethane was transferred into an Erlenmeyer flask containing a	<ol> <li>Peking Chemical Factory, further purified by washing with H<sub>2</sub>SO<sub>4</sub>, NaOH and water; dried</li> </ol>		
weighed amount of aqueous solution. The flask was rotated in a thermostat bath until equilibrium was	<ul> <li>over CaCl<sub>2</sub> and fractionated.</li> <li>(2) Peking Chemical Factory,</li> </ul>		
established. The optical densities of different concentration of tetrachloromethane in the aqueous	<ul><li>recrystallized twice before use.</li><li>(3) Distilled.</li></ul>		
solutions were measured using water as the reference liquid. Duplicate experiments were made and the	ESTIMATED ERRORS:		
chloromethane in aqueous potassium chloride solu- tions were determined using a Hiltor spectra	Solubility: Not specified.		
photometer.	remperature. T 0.5 K (complice).		

66		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Treadwell, W. D.; Köhl, A.	
(2) Potentium hydroxida: KOU: [1210.58.2]	Helv. Chim. Acta <u>1926</u> , 9, 681-91.	
(2) Polassium nydroxide; $KOR; [1510-38-5]$		
VARIABLES.		
T/K = 333	A. L. Horvath	
One concentration		
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ $\rho_1/k_2$	$g m^{-3}$ 100 $w_l$ 10 <sup>5</sup> $x_l$	
	(compiler) (compiler)	
60 1.0 0.	2 1.98 × 10 <sup>-2</sup> 2.41	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibrium between a known volume of tetrachloromethane gas and KOH solution was estab-	(1) Prepared in the authors' laboratory; purified by fractional distillation before use	
lished in an container under continuous shaking. The volume of tetrachloromethane not dissolved in the	<ul><li>(2) Source and purity not given.</li><li>(3) Distilled.</li></ul>	
KOH solution was measured and the amount of dis- solved gas was determined at the same time and		
conditions. A single measurement was done at each temperature.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tetrachloromethane (carbon tetrachloride);	Liu, JL.; Huang, TC.	
(2) Potassium sulfate: K-SQ : [7778-80-5]	Scientia Sinica (Peking) <u>1961</u> , 10, 700-10.	
(3) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
Concentration		
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ $\rho_1/\text{kg}$	$m^{-3}$ 100 w, 10 <sup>5</sup> x, (compiler) (compiler)	
25 0.1323 0.6 0.3150 0.5	$60  6.49 \times 10^{-2}  7.766$ $44  5.22 \times 10^{-2}  6.433$	
0.3518 0.3	$507$ $4.84 \times 10^{-2}$ $6.001$ (continued)	
AUXILIARY II	VFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An accurate amount of tetrachloromethane was	(1) Peking Chemical Factory, further purified by	
transferred into an Erlenmeyer flask containing a weighed amount of aqueous solution. The flask was	washing with H <sub>2</sub> SO <sub>4</sub> , NaOH and water; dried over CaCl <sub>2</sub> and fractionated.	
rotated in a thermostat bath until equilibrium was established. The optical densities of different	(2) Peking Chemical Factory, recrystallized twice before use.	
concentration of tetrachloromethane in the aqueous solutions were measured using water as the reference	(3) Distilled.	
results averaged. The absorption spectra of tetra-	ESTIMATED ERRORS:	
were determined using a Hilger spectrophotometer.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	

-						67
COMPONENTS:				ORIG	INAL MEASURE	MENTS:
(1) Tetrachloromet	hane (carbo	on tetrachloride);		Liu,	JL.; Huang, T(	С.
(2) Potassium sulfa	ate: KaSO4:	[7778-80-5]		Scien	ntia Sinica (Peking	) <u>1961</u> , <i>10</i> , 700-10.
(3) Water; H <sub>2</sub> O; [7	732-18-5]	[///0/00/0]				
VARIABLES:				PREP	ARED BY:	
T/K = 298 Concentration				A. L	. Horvath	
EXPERIMENTAL	VALUES:	(continued)				
1	°/°C	c <sub>2</sub> /mol dm <sup>-3</sup>	ρ <sub>ι</sub> /kg	g m <sup>-3</sup>	100 w, (compiler)	10 <sup>s</sup> x, (compiler)
2	25	0.4516 0.5177	0.4 0.4	464 430	$4.37 \times 10^{-2}$ $4.02 \times 10^{-2}$	5.509 5.124
		0.6034	0.:	388	$3.60 \times 10^{-2}$	4.657
COMPONENTS:				ORIG	INAL MEASURE	MENTS:
(1) Tetrachloromethane (carbon tetrachloride);		Liu, JL.; Huang, TC.				
(2) Sodium chloride; NaCl; [7647-14-5]		Scien	tia Sinica (Peking,	) <u>1961</u> , <i>10</i> , 700-10.		
(3) Water; H <sub>2</sub> O; [7	732-18-5]					
VARIABLES:				PREP	ARED BY:	
T/K = 298				A. L.	. Horvath	
Concentration						
EXPERIMENTAL	VALUES:					
t	/°C	c₂/mol dm⁻³	ρ <sub>ı</sub> /kg	m <sup>-3</sup>	100 w, (compiler)	10 <sup>s</sup> x <sub>1</sub> (compiler)
2	5	0.2549	0.6 0.6	590 542	$6.84 \times 10^{-2}$ $6.30 \times 10^{-2}$	8.099 7.536
		0.6375	0.6	516	$6.01 \times 10^{-2}$ $5.58 \times 10^{-2}$	7.229
		0.9623	0.5 0.5	553 541	$5.32 \times 10^{-2}$ 5.19 × 10^{-2}	6.486 6.344
		AUXILI	ARY IN	NFORM	IATION	
METHOD/APPARA	TUS/PRO	CEDURE:		SOUR	CE AND PURITY	Y OF MATERIALS:
An accurate a	amount of t	etrachloromethan	e was	(1)	Peking Chemical	Factory, further purified by
transferred into an I weighed amount of rotated in a thermos established. The opt	Erlenmeyer aqueous sol stat bath unit tical densition	flask containing lution. The flask til equilibrium wa es of different	a was as	(2) (3)	washing with H <sub>2</sub> S over CaCl <sub>2</sub> and fr Peking Chemical Distilled.	O4, NaOH and water; dried actionated. Factory, analytical grade.
solutions were meas	sured using	water as the refe	erence	FCTIN		
results averaged. The	e absorptio	n spectra of tetra	i- tions	Soluk		Not specified
were determined us	ing a Hilger	spectrophotome	ter.	Temp	berature: ±	0.5 K (compiler).

68	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tetrachloromethane (carbon tetrachloride);	Liu, JL.; Huang, TC.
(2) Sodium sulfate: Na $_{3}$ SO <sub>4</sub> : [7757-82-6]	Scientia Sinica (Peking) <u>1961</u> , 10, 700-10.
(3) Water; $H_2O$ ; [7732-18-5]	
VARIABLES:	PREPARED BY:
T/K = 298 Concentration	A. L. Horvath
EXPERIMENTAL VALUES:	•
$t/^{\circ}C$ $c_2/\text{mol dm}^{-3}$ $\rho$	$1/\text{kg m}^{-3}$ 100 $w_i$ 10 <sup>5</sup> $x_i$ (compiler) (compiler)
25 0.130 0.230 0.380 0.464 0.564	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
AUXILIAR	Y INFORMATION
METHOD/APPARATUS/PROCEDURE	SOURCE AND PUBITY OF MATERIALS
An accurate amount of tetrachloromethane w transferred into an Erlenmeyer flask containing a weighed amount of aqueous solution. The flask was rotated in a thermostat bath until equilibrium was established. The optical densities of different concentration of tetrachloromethane in the aqueous solutions were measured using water as the referent	<ul> <li>(1) Peking Chemical Factory, further purified by washing with H<sub>2</sub>SO<sub>4</sub>, NaOH and water; dried over CaCl<sub>2</sub> and fractionated.</li> <li>(2) Peking Chemical Factory, recrystallized twice before use.</li> <li>(3) Distilled.</li> </ul>
results averaged. The absorption spectra of tetra- chloromethane in aqueous sodium sulfate solutions were determined using a Hilger spectrophotometer.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water-d <sub>2</sub> ; D <sub>2</sub> O; [7789-20-0]	Hutchison, C. A.; Lyon, A. M.
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	Columbia University Report A-745, July 1, 1943.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}$ C 100 $w_{I}M_{I}^{-1}/\text{mol g}^{-1}$	$\begin{array}{c} 100 \ w_{i} \\ (compiler) \\ (compiler) \\ \end{array}$
$25 4.78 \times 10^{4}$	$9.57 \times 10^{-3}$ 7.34
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A 1 to 15 volume ratio mixture of heavy water and tetrachloromethane was introduced into an equilibration flask and then lowered into the water thermostat bath. The assembly was shaken mechan-	<ol> <li>Source and purity not given.</li> <li>Source not given. Purified and dried before use.</li> </ol>
The amount of heavy water in the organic phase was	ESTIMATED ERRORS:
determined by a modified Karl Fischer titration method. The determinations were done in triplicate. The description was taken from a secondary source (ref. 1). The original report is not available from Columbia University.	Solubility: $\pm 0.1 \times 10^{-5}$ avg. dev. Temperature: $\pm 0.05$ K.
	REFERENCES:
	(1) Eidinoff, M. L., Joris, G. G., Taylor, H. S., Urey, H. C., Eds. <i>Production of Heavy Wa-</i> <i>ter</i> , McGraw-Hill, New York, <u>1955</u> , p. 129.
$(1) Water-d: D \cap (7789, 20, 0)$	Enherre D. Poth E
(2) Tetrachloromethane (carbon tetrachloride):	Bull Soc. Chim. Fr. 1968, 2291-2.
CCl <sub>4</sub> ; [56-23-5]	Dam. Doc. China 17. <u>1700</u> , 2271-2.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	1
$t/^{\circ}$ C 10 <sup>5</sup> $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ (compiler) & (compiler) \end{array}$
25 9	$0.9 \times 10^{-2}$ 6.9
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The isotope dilution technique was used. It	(1) Received from pile EL 3 of Saclay, contained
involved the mixing of about 100 cm <sup>3</sup> tetrachloro- methane with 1 cm <sup>3</sup> heavy water in a volumetric flask which was then inserted into a constant temperature bath. The mixture was acitated for 15 minutes to	<ul> <li>traces of tritium.</li> <li>(2) Source not given, commercial grade.</li> </ul>
ensure the equilibrium between the two phases. The	ESTIMATED ERRORS:
liquid scintillation counting procedure.	Solubility: $\pm 0.4 \times 10^{-5}$ g. Temperature: $\pm 1$ K (compiler).

•

200

4 .....

70	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water-d <sub>2</sub> ; D <sub>2</sub> O; [7789-20-0]	Glasoe, P. K.; Schultz, S. D.
(2) Tetrachloromethane (carbon tetrachloride); CCl <sub>4</sub> ; [56-23-5]	J. Chem. Eng. Data <u>1972</u> , 17, 66-8.
VARIABLES:	PREPARED BY:
T/K = 288 - 308	A. L. Horvath
EXPERIMENTAL VALUES:	100
	(compiler) (compiler)
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{ccccc} 0.713 \times 10^{-2} & 5.47 \\ 0.949 \times 10^{-2} & 7.28 \\ 1.07 \times 10^{-2} & 8.21 \end{array}$
$35  10.3 \pm 0.3$	$1.32 \times 10^{-2}$ 10.13
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturation of heavy water with tetrachloro- methane was done in a closed system, protected from atmospheric moisture, and placed in a constant tem- perature water bath. The solubility of heavy water in tetrachloromethane was measured using the Karl	<ol> <li>Source not given, 99.8 % pure, used as received.</li> <li>Source not given. Reagent grade, purified by distillation before use.</li> </ol>
rischer titration method with a "dead stop" end-point determination (ref. 1). 15 Trials were done at each	ESTIMATED ERRORS:
temperature.	Solubility: See above. Temperature: $\pm 0.1$ K (compiler).
	REFERENCES:
	(1) Wernimont, G.; Hopkinson, F. J. Ind. Eng. Chem. <u>1943</u> , 12, 272.

COMPONENTS:	EVALUATOR:
<ol> <li>Bromodichloromethane; CHBrCl<sub>2</sub>; [75-27-4]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	January 1993.

1

ļ

# CRITICAL EVALUATION:

All the available solubility data for bromodichloromethane (1) in water (2) are summarized in Table 1.

The data are in very poor agreement and in the absence of other independent studies it is not possible to determine which values are the more reliable. Further studies will be necessary before even tentative values can be assigned. Even so, it may be observed that the solubility trend for halogenated hydrocarbon - water systems, as presented in the Preface, provided some support for the mass per cent (1) value, 100  $w_1$ , of 1.618 at 298.15 K (ref. 1).

Table 1. Reported Solubility of Bromodichloromethane (1) in Water (2)

Tem	Temperature Reporte		lity
°C	К	100 w <sub>1</sub>	$10^3 x_1$
20	293.15	2.099 (ref. 4)	2.352 (ref. 4)
24.85	298.00	1.821 (ref. 5)	2.035 (ref. 5)
25	298.15	1.618 (ref. 1)	1.805 (ref. 1)
30	303.15	0.3023 (ref. 2),	0.3333 (ref. 2),
		0.2972 (ref. 3)	0.3277 (ref. 3)

#### **REFERENCES:**

- 1. Symons, J. M. et al. Treatment Techniques for Controlling Trihalomethanes in Drinking Water, U. S. EPA Report 600/2-81-156, Cincinnati, OH., <u>1981</u>.
- 2. McNally, M. E.; Grob, R. L. J. Chromatogr. 1983, 260, 23.
- 3. Ibid., <u>1984</u>, 284, 105.
- 4. Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518.
- 5. Warner, H. P.; Cohen, J. M.; Ireland, J. C. Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB87-212684, Cincinnati, OH., July 1987.

12	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	Symons, J. M.; et al.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Treatment Techniques for Controlling Trihalo- methanes in Drinking Water, U. S. EPA Report 600/2-81-156, Cincinnati, OH., <u>1981</u> .
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	•
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$
$25    9.5 \times 10^{-2}$	1.6179 1.805
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are given elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Cincinnati tap water.</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm 0.5$ K.
	REFERENCES:
	(1) Werner, U. S. Environmental Protection Agency, Cincinnati, OH., <i>Personal</i> <i>Comm nication</i> , <u>1980</u> .
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	McNally, M. E.; Grob, R. L.
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chromatogr. <u>1983</u> , 260, 23-32.
VARIABLES:	PREPARED BY:
T/K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	1
t/°C 10 <sup>6</sup> ₩,	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
30 3031.9	0.3023 3.333
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Standard solutions were prepared in 100 ml flasks. They were shaken for 5 minutes and then allowed to equilibrate in a constant temperature bath held at 30°C for 24 hours. Samples were pipetted in headspace vials, sealed, thermostated and then exam-	<ol> <li>Chemical Service, West Chester, PA., purest grade available.</li> <li>Distilled and run through two Barnstead purification cartridges.</li> </ol>
ined by the gas chromatographic head space analysis technique. An F. & M. Model 402 gas chromato- graph equipped with dual columns and dual flame- ionization detectors was used.	ESTIMATED ERRORS: Solubility: ± 150 ppm std. dev. Temperature: ± 0.5 K (compiler).

	73
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	McNally, M. E.; Grob, R. L.
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chromatogr. <u>1984</u> , 284, 105-16.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	•
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/kg m^{-3}$	$100 w_i$ $10^4 x_i$
30 2.968	0.2972 3.276
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Standard bromodichloromethane solutions of increasing concentration were prepared in volumetric flasks as described elsewhere (ref. 1). The aqueous organic liquid phase was 25 ml and the vapor phase above the mixture was 35 ml. The system was allow-	<ol> <li>Chemical Service, West Chester, PA., purest grade available.</li> <li>Distilled and run through two Barnstead purification cartridges.</li> </ol>
ed to reach equilibrium in a constant temperature bath and then a sample was injected into the gas chromato-	ESTIMATED ERRORS:
graph. Values reported are an average of at least 5 measurements.	Solubility: $\pm$ 7.12 % std. dev. Temperature: $\pm$ 0.5 K (compiler).
	REFERENCES:
	(1) McNally, M. E.; Grob, R. L. J. Chromatogr. <u>1983</u> , 260, 23.
COMPONENTS	ODIGINAL MEASUDEMENTS.
(1) Bromodichloromethane: CHBrCl <sub>2</sub> : [75-27-4]	Nicholson, B. C.: Maguire, B. P.: Bursill, D. B.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$
20 $1.6 \times 10^{-3}$	2.099 2.352
AUXILIARY IN	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow was	<ol> <li>Source and purity not given.</li> <li>Distilled and deionized.</li> </ol>
	ESTIMATED ERRORS:
maintained. Samples of the solution being stripped were taken for analysis. After extraction of the sample, the extracts were analyzed by a gas chroma- tograph equipped with a capillary column.	Solubility: $\pm$ 0.2 × 10-3 std. dev.Temperature: $\pm$ 0.1 K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.

74	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	Warner, H. P.; Cohen, J. M.; Ireland, J. C.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB87-212684, Cincinnati, OH., July 1987.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	·
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$
24.85 $2.12 \times 10^{-3}$	1.821 2.036
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The original method and apparatus for the determination of Henry's law constants, described by Mackay <i>et al.</i> (ref. 1), were used. The general procedure was to add an excess quantity of bromo-	<ol> <li>Source not given. Purest quality available and was used without further purification, stated purity &gt; 99 %.</li> <li>Distilled and deionized.</li> </ol>
dichloromethane to distilled deionized water and mix the system overnight. A portion of the solution was	ESTIMATED ERRORS:
methane was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from the log of the concentration	Solubility: $\pm 6 \%$ std. dev. Temperature: $\pm 0.05$ K.
versus time plot. The experimental data values are averages of two or more replicates.	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Technol. <u>1979</u> , 13, 333.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Milang water	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$20$ $1.6 \times 10^{-3}$	2.099 2.353
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was	<ol> <li>Source and purity not given.</li> <li>pH = 8.4, 700 mg salt/dm<sup>3</sup>, 140 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 13 mg organic carbon/dm<sup>3</sup>, 110 mg suspended solid/dm<sup>3</sup>.</li> </ol>
added to the stripping vessel and the desired flow was maintained. Samples of the solution being stripped	ESTIMATED ERRORS:
ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Myponga water	Envinon. Sci. Technol, <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EVDEDIMENTAL VALUES.	
t/°C Henry's low constant	100 yu 103 y
$H/m^3$ atm mol <sup>-1</sup>	(compiler) (compiler)
$20   1.5 \times 10^{-3}$	2.239 2.515
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	<ol> <li>Source and purity not given.</li> <li>pH = 7.6, 370 mg salt/dm<sup>3</sup>, 60 mg CaCO<sub>2</sub>/dm<sup>3</sup>, 10 mg organic carbon/dm<sup>3</sup>.</li> </ol>
diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate	ESTIMATED ERRORS:
maintained. Samples of the solution being stripped were taken for analysis. After samples were extract-	Solubility: Not specified.
ed, they were analyzed using a gas chromatograph equipped with a capillary column.	Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromodichloromethane; CHBrCl <sub>2</sub> ; [75-27-4]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Myponga water + humic acid	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EXPERIMENTAL VALUES	
t/°C. Henry's law constant	100 w. 10 <sup>3</sup> r.
H/m <sup>3</sup> atm mol <sup>-1</sup>	(compiler) (compiler)
20 $1.5 \times 10^{-3}$	2.239 2.515
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was	<ol> <li>Source and purity not given.</li> <li>pH = 7.9, 20 mg humic acid/dm<sup>3</sup>, 370 mg salt/dm<sup>3</sup>, 68 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 17 mg organic carbon/dm<sup>3</sup>.</li> </ol>
added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped	ESTIMATED ERRORS:
were taken for analysis. After samples were extract- ed, they were analyzed using a gas chromatograph equipped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.

,

COMPONENTS:	EVALUATOR:
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	January 1993.

### CRITICAL EVALUATION:

76

All the available solubility data for dibromochloromethane (1) in water (2) are summarized in Table 1.

The data are in very poor agreement. Considerable discrepancies exists even between the two solubility values reported by the same workers, compare (ref. 2) and (ref. 3). In the absence of further independent studies, it is not possible to determine which value is the more reliable. Additional investigations will be necessary before even tentative values can be assigned. Even so, it may be noted that the solubility trend for halogenated hydrocarbon - water systems, as presented in the Preface, provided some support for the mass per cent (1) value, 100  $w_1$ , of 0.8124 at 298.15 K (ref. 1).

Table 1. Reported Solubility of Dibromochloromethane (1) in Water (2)

Temperature		perature	Solubility	
	°C	К	100 w <sub>i</sub>	$10^4 x_1$
	10	283.15	0.3040 (ref. 6)	2.637 (ref. 6)
	20	293.15	0.6320 (ref. 4),	5.498 (ref. 4),
			0.2375 (ref. 6)	2.059 (ref. 6)
	24.85	298.00	1.423 (ref. 5)	12.470 (ref. 5)
	25	298.15	0.8124 (ref. 1)	7.079 (ref. 1)
	30	303.15	0.1049 (ref. 2),	0.9082 (ref. 2),
			0.2514 (ref. 3)	2.179 (ref. 3)
			0.3132 (ref. 6)	2.7168 (ref. 6)
			· · ·	· · ·

### **REFERENCES:**

- 1. Symons, J. M. et al. Treatment Techniques for Controlling Trihalomethanes in Drinking Water, U. S. EPA Report 600/2-81-156, Cincinnati, OH., <u>1981</u>.
- 2. McNally, M. E.; Grob, R. L. J. Chromatogr. 1983, 260, 23.
- 3. Ibid., <u>1984</u>, 284, 105.
- 4. Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518.
- 5. Warner, H. P.; Cohen, J. M.; Ireland, J. C. Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB87-212684, Cincinnati, OH., July 1987.
- Howe, G. B.; Mullins, M. E.; Rogers, T. N. AFESC Tyndall Air Force Base, Report ESL-TR-86-66, Vol. 1, Florida, <u>Sept. 1987</u>, 86 pp. (AD-A188 571).

COMPONENTS	ODIGINAL MEASUDEMENTS:
(1) Difference the set of the child of the	Summer I. M. et al.
(1) Dibromochloromethane; CHBr <sub>2</sub> CI; $[124-48-1]$	Symons, J. M. <i>et al.</i>
(2) Water; $H_2O$ ; [7732-18-5]	Treatment Techniques for Controlling Trihalo- methanes in Drinking Water, U. S. EPA Report 600/2-81-156, Cincinatti, OH., <u>1981</u> .
VARIABLES:	PREPARED BY:
<i>T/</i> K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$25  3.5 \times 10^{-2}$	0.8124 7.080
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are given elsewhere (ref. 1).	(1) Source and purity not given.
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm 0.5$ K.
	REFERENCES:
	<ol> <li>Werner, U. S. Environmental Protection Agency, Cincinatti, OH., Personal communication, <u>1980</u>.</li> </ol>
COMPONENTS	ODIGINAL MEASUDEMENTS.
(1) Dibromochloromethane: CHBr Cl. [124.48.1]	MoNally M. E. Grab P. I
(1) Water H $_{0}$ [7722 18 5]	I Chromotoge 1092 260 22 22
(2) water, 11 <sub>2</sub> 0, [7732-16-3]	5. Chromatogr. <u>1785</u> , 200, 25-52.
VARIABLES:	PREPARED BY:
T/K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	1
t/°C 10 <sup>6</sup> w <sub>i</sub>	$100 w_{1}$ $10^{5} x_{1}$
	(compiler) (compiler)
30 1049.9	0.1049 9.081
AUXILIARY II	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Standard solutions were prepared in 100 cm <sup>3</sup>	(1) Chemical Service, West
nasks. They were shaken for 5 minutes and then al- lowed to equilibrate in a constant-temperature bath held at 30°C for 24 hours. Samples were pipetted into headspace vials, sealed, thermostated, and then ex- amined by the gas chromatographic head space ana-	<ul> <li>Cnester, PA., USA., purest grade available used as received.</li> <li>(2) Distilled and run through two Barnstead purification cartridges.</li> </ul>
lysis technique. An F. & M. Model 402 gas chro- matograph equipped with dual columns and dual	ESTIMATED ERRORS:
flame ionization detectors was used.	Solubility: $\pm$ 36.4 ppm std. dev. Temperature: $\pm$ 0.5 K (compiler).

78		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	McNally, M. E.; Grob, R. L.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chromatogr. <u>1984</u> , 284, 105-16.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/g m^{-3}$	$\begin{array}{ccc} 100 \ w_{i} & 10^{4} \ x_{i} \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
30 2509	0.2514 2.179	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Standard dibromochloromethane solutions of increasing concentration were prepared in volumetric flasks as described elsewhere (ref. 1). The aqueous organic liquid phase was 25 ml and the vapor phase above the mixture was 35 ml. The system was al-	<ol> <li>Chemical Service, West Chester, PA., USA, purest grade available.</li> <li>Distilled and run through two Barnstead purification cartridges.</li> </ol>	
based to reach equilibrium in a constant temperature bath and then a sample was injected into the gas chromatograph. Values reported are an average of at least 5 measurements.	ESTIMATED ERRORS: Solubility: ± 3.47 % std. dev. Temperature: ± 0.5 K (compiler).	
	REFERENCES:	
	(1) McNally, M. E.; Grob, R. L. J. Chromatogr. <u>1983</u> , 260, 23.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
20 8.7 $\times$ 10 <sup>-4</sup>	0.6320 5.498	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	<ol> <li>Source and purity not given.</li> <li>Distilled and deionized.</li> </ol>	
diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate	ESTIMATED ERRORS:	
were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equipped with a capillary column.	Solubility: $\pm 0.2 \times 10^{-4}$ std. dev. Temperature: $\pm 0.1$ K.	
	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.	

•

	79
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	Warner, H. P.; Cohen, J. M.; Ireland, J. C.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants, EPA Tech. Report PB87-212684, Cincinatti, OH., July 1987.
VARIABLES:	PREPARED BY:
T/K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	•
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$
24.85 $7.83 \times 10^{-4}$	1.423 1.538
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The original method and apparatus for the determination of Henry's law constants, described by Mackay <i>et al.</i> (ref. 1), were used. The general procedure was to add an excess quantity of dibromo-	<ol> <li>Purest quality available, &gt; 99 % pure, used as received.</li> <li>Distilled and deionized.</li> </ol>
the system overnight. A portion of the solution was returned to the stripping vessel. Dibromochloro- methane was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from the log of the concentration	ESTIMATED ERRORS: Solubility: ± 6 % std. dev. Temperature: ± 0.05 K.
versus time plot. The experimental data values are averages of two or more replicates.	REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Tehnol. <u>1979</u> , 13, 333.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	Howe, G. B.; Mullins, M. E.; Rogers, T. N.
(2) Water; H <sub>2</sub> O; [7732-18-5]	AFESC Tyndall Air Force Base, Report ESL-TR- 86-66. Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp. (AD-A188 571).
VARIABLES:	PREPARED BY:
T/K = 283 - 303	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10 <sup>6</sup> ₩,	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
10 3040 20 2375 30 3132	0.30402.63680.23752.05870.31322.7168
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
250 cm <sup>3</sup> bottles were filled with distilled deionized water and sealed. A measured volume of dibromochloromethane was injected into the bottles through each bottle septum using a microliter syringe. The solute was in excess of the anticipated solubility	<ol> <li>Probably a commercial reagent, at least 99 % purity, used as received.</li> <li>Distil led and deionized.</li> </ol>
limit. The bottles were shaken for one hour with a wrist-action shaker and allowed to equilibrate for about three weeks. Samples were then injected into a gas chromatograph equipped with a Carbopack column and a FID detector. The GC response was compared with the calibration plot.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).

COMPONENTS:       ORIGINAL MEASUREMENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]       ORIGINAL MEASUREMENTS:         (2) Milang water       Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. $ZNRIABLES:$ PREPARED BY: $TK = 293$ A. L. Horvath         EXPERIMENTAL VALUES: $I^{PC}$ $I^{PC}$ Henry's law constant, $I^{PC}$ Hen	80	
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Milang waterNicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1924, 18, 518-21.VARIABLES: T/K = 293PREPARED BY: A. L. HorvathIVImm at monit Imm at monit 208.5 × 10 <sup>4</sup> IVMenoy's law constant. Imm at monit 20100 w, (compiler) 0.6469METHOD/APPARATUS/PROCEDURE: The are netred through the top of the vessal and analitatined. Samples of the solution being stripping were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip 20Source and purity not given. (2) Source and purity not given. (2) ORIGINAL MEASUREMENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Wyonga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1292, 17, 533.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Wyonga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1292, 17, 533.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) 8.0 × 10 <sup>4</sup> ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1292, 17, 533.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) 8.0 × 10 <sup>4</sup> Source and purity not given. (compiler) 0.66873VARIABLES: T/K = 293Na to source and purity not gatifam, Sci. Technol. 1292, 17, 533.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) 8.0 × 10 <sup>4</sup> Source and purity not given. (compiler) 0.66873METHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel <td>COMPONENTS:</td> <td>ORIGINAL MEASUREMENTS:</td>	COMPONENTS:	ORIGINAL MEASUREMENTS:
(2) Milang water       Environ. Sci. Technol. 1924, 18, 518-21.         VARIABLES: T/K = 293       PREPARED BY: A. L. Horvath         LEXPERIMENTAL VALUES: I'C Henry's law constant, 	(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
VARIABLES: $7K = 293$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: $I^{PC}$ Henry's law constant, $H'm' atm mol'100 w,(compiler)10' x,(compiler)208.5 × 1040.64695.632AUXILIARY INFORMATIONSOURCE AND PURITY OF MATERIALS:(1) Source and purity not given,D'H = 8,47,000 mg sail/draf, 40 mgD'H = 1,47,000 mg sail/draf, 40 mgD'H = 1,67,000 mg sail/draf, $	(2) Milang water	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
$TK = 293$ A. L. Horvath         EXPERIMENTAL VALUES: $t^{PC}$ Henry's law constant, $100 \text{ w}_{1}$ (compiler)         20 $8.5 \times 10^4$ $0.6469$ AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel and the distred through the top of the vessel and the distred through th	VARIABLES:	PREPARED BY:
EXPERIMENTAL VALUES: $l'''C$ Henry's law constant, H/m' atm mol'       100 w, (compiler)       10' x, (compiler)         20 $8.5 \times 10^4$ 0.6469       5.632         AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE: The are netred through the top of the vessel and the described by Mackay et al. (cf. 1). The air entered through the top of the vessel and the desired to the stripping vessel and the desired flow rate added to the stripping vessel and the desired flow rate even taken for analysis. After samples were tracteder through the top of the vessel samples of the solution samples of the solutio	T/K = 293	A. L. Horvath
$h''C$ Henry's law constant, H'm' atm mol'100 w, (compiler)10' z, (compiler)20 $8.5 \times 10^4$ $0.6469$ $5.632$ AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel antained. Science the bottom. The solution was added to the stripping vessel and the desired flow raise chromatograph equip- ped with a capillary column.SOURCE AND PURITY OF MATERIALS: () Source and purity not given. () pH = 8.4 / 700 mg stifued. Temperature: $\pm 0.1$ K.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myonga waterORIGINAL MEASUREMENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myonga waterORIGINAL MEASUREMENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myonga waterORIGINAL MEASUREMENTS: Nicholson, B. C; Maguire, B. P; Bursill, D. B. Environ. Sci. Technol. 1924, 18, 518-21.VARIABLES: $TK = 293$ AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: $L''C$ Henry's law constant, $L''C$ Henry's law constant, $L'''$ Mackay <i>et al.</i> (ref. 1), $L''' Martin mol'(compiler)(compiler)(compiler)OURCE AND PURITY OF MATERIALS:(1) Source and purity not given.(compiler), mag salfded, et al. (P. EXTRAPLE)L'''' = 7.6, 370 mg salf/drif, 60 mgCaCO,d'm', 10 mg salf/drif, 7, 60 mgCaCO,d'm', 10 mg salf/drif, 60 mgCaCO,d'm', 10 mg salf/drif, 60 mgCaCO,d'm', 10 mg salf/drif, 60 mgCaCO,d'm', 10 $	EXPERIMENTAL VALUES:	
20 $8.5 \times 10^4$ 0.64695.632AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel and the described by Mackay et al. (fef. 1). The air entered through the top of the vessel and the diffuser was consubution being stripped were taken for analysis. After samples were extraction the dynamical extra stripping vessel and the desired flow rays added to the stripping vessel and the desired flow rays added to the stripping vessel and the desired flow rays added to the stripping vessel and the desired flow rays added to the stripping vessel and the desired flow rays the dynamical extra stripping vessel and the desired flow rays the dynamical extra stripping vessel and the desired flow rays the dynamical extra stripping vessel and the desired flow rays the dynamical extra stripping vessel and the desired flow rays temperature: $\pm 0.1$ K.Source and purity not given. (Source rays extra stripping vessel) Solubility: Not specified. Temperature: $\pm 0.1$ K.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Mynonga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1924. 18, 518-21.VARIABLES: TK = 293VARIABLES: I''C Henry's law constant, I''' Mir'' aim mol'' (compiler) 20 $8.0 \times 10^4$ OURCE AND PURITY OF MATEINALS: (IO w, (IO' $\chi$ , (compiler) (compiler) 0.6873METHOD/APPARATUS/PROCEDURE: Similar to that described by Mackay et al. (fef. 1). The air entered through the domain and desired flow ray anantained. Samples of the stripping vessel and the desired flow ray the stripping sest chromatograph equip- ped with a capillar	t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1), the air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.       SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) PH = 8.4, 700 mg salt/dm <sup>-</sup> , 140 mg CaCO_Adm <sup>-</sup> , 10 mg organic carbon/dm <sup>+</sup> , 110 mg suspended solid/dm <sup>-3</sup> .         COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water         ORIGINAL MEASUREMENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water         VARIABLES: <i>IPC</i> Henry's law constant, <i>H/m</i> <sup>-</sup> atm mol <sup>+</sup> 20 <i>ORIGINAL MEASUREMENTS:</i> Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. <i>Environ. Sci. Technol.</i> <u>1984</u> , <i>18</i> , 518-21.         MEEPARED BY: A. L. Horvath         SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) 8.0 × 10 <sup>4</sup> SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) Source and purity not given. (2) Source and purity not given. (2) (2) Markay, D.; Shiu, W. Y.; Sutherland, R. P. <i>Environ, Sci. Technol.</i> <u>1920</u> , <i>13</i> , 333.	20 8.5 × 10 <sup>-4</sup>	0.6469 5.632
METHOD/APPARATUS/PROCEDURE:       SOURCE AND PURITY OF MATERIALS:         The apparatus consisted of a stripping vessel       SOURCE AND PURITY OF MATERIALS:         imilar to that described by Mackay et al. (ref. 1).       Source and purity not given.         (2) Surge and year by gas chromatograph equipped with a capillary column.       Source and purity not given.         (2) Mynong water       Source and purity not given.         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]       REFERENCES:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]       ORIGINAL MEASUREMENTS:         (2) Mynonga water       Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. <i>TK</i> = 293       A. L. Horvath         EXPERIMENTAL VALUES:       IOO w, 10 <sup>4</sup> x, (compiler) (compiler)         20       8.0 × 10 <sup>4</sup> OSR73         20       8.0 × 10 <sup>4</sup> SOURCE AND PURITY OF MATERIALS:         METHOD/APPARATUS/PROCEDURE:       SOURCE AND PURITY OF MATERIALS:         The apparatus consisted of a stripping vessel       SOURCE AND PURITY OF MATERIALS:         (1) Surce and purity not given.       (2) Mynong water         METHOD/APPARATUS/PROCEDURE:       SOURCE AND PURITY OF MATERIALS:         (1) Surce and purity not given.       (2) PI = 7.6, 370 mg sal/dm <sup>2</sup> , 60 mg         (2) Mynong water       SOURCE AND PURITY OF MATERIALS:         (2) Mynong water <td>AUXILIARY I</td> <td>NFORMATION</td>	AUXILIARY I	NFORMATION
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1).       (1) Source and purity not given.         The air entered through the top of the vessel and the disired flow rate added to the stripping vessel and the desired flow rate added to the stripping vessel and the desired flow rate diffuser was enalyzed by gas chromatograph equiped with a capillary column.       (2) HP = 8.4, 700 mg osth/dm².         Maintained. Samples of the solution being stripped with a capillary column.       (1) Source and purity not given.         COMPONENTS:       (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.         COMPONENTS:       (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]         (2) Myponga water       ORIGINAL MEASUREMENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]       ORIGINAL MEASUREMENTS:         (2) Myponga water       PREPARED BY:         7/K = 293       A. L. Horvath         EXPERIMENTAL VALUES:       (20) 8.0 × 10 <sup>4</sup> 100 w, (compiler)       (compiler)         20       8.0 × 10 <sup>4</sup> METHOD/APPARATUS/PROCEDURE:       SOURCE AND PURITY OF MATERIALS:         The apparatus consisted of a stripping vessel and the desired flow rate and yard by gas chromatograph equiped with a capillary column.       SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given, 60 mg 24/2 Gr. 1).       Source and purity not given, 60 mg 26/2 Gr. 10 mg sali/dm², 60 mg 26/2 Gr. 10 mg sali/dm², 60 m	METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
added to the Shifting tester and the desired now rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.ESTIMATED ERRORS: Solubility: Not specified. Temperature: $\pm 0.1$ K.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1928, 18, 518-21.VARIABLES: T/K = 293ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1928, 18, 518-21.VARIABLES: T/K = 293A L. HorvathEXPERIMENTAL VALUES: $U'^C Henry's law constant,H/m' atm mol'120 8.0 × 104SOURCE AND PURITY OF MATERIALS:(compiler)0.6873 5.984SOURCE AND PURITY OF MATERIALS:(1) Source and purity not given.(2) STIMATED ERRORS:Sould do the desired flow vratemaintained. Samples of the solution being strippedwere taken for analysis. After samples were extract-ed, they were analyzed by gas chromatograph equip-ped with a capillary column.STIMATED ERRORS:Subolitity: Not specified.Technol. 1929, 13, 333.ETRMATED ERRORS:Subolity: Not specified.(they were analyzed by gas chromatograph equip-ped with a capillary column.ESTIMATED ERRORS:Subolity: Not specified.(they were analyzed by gas chromatograph equip-ped with a capillary column.$	The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired four ste	<ol> <li>Source and purity not given.</li> <li>pH = 8.4, 700 mg salt/dm<sup>3</sup>, 140 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 10 mg organic carbon/dm<sup>3</sup>, 110 mg suspended solid/dm<sup>3</sup>.</li> </ol>
dia term were analyzed by gas chromatograph equipped with a capillary column.Not specified. Temperature: $\pm 0.1$ K.REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1929, 13, 333.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> CI; [124-48-1] (2) Myponga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1928, 18, 518-21.VARIABLES: T/K = 293ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1928, 18, 518-21.VARIABLES: T/K = 293A. L. HorvathEXPERIMENTAL VALUES: $l^PC$ Henry's law constant, H/m <sup>3</sup> atm mol <sup>14</sup> 20SOURCE AND PURITY OF MATERIALS: (compiler) (compiler) 0.6873SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) PH = 7.6, 570 mg sall/dm², 60 mg CaCO/dm², 10 mg organic carbon/dm³.ESTIMATED ERRORS: Subdet ot the solution was added to the stripping vessel added to the stripping vessel added to the stripping vessel added to the stripping vessel added to the stripping vessel anitationed. Samples of the solution was added to the stripping vessel and the desired flow wrate maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Emviron. Sci. Technol. 1979, 13, 333.	maintained. Samples of the solution being stripped	ESTIMATED ERRORS:
REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518-21.VARIABLES: T/K = 293PREPARED BY: A. L. HorvathVARIABLES: $I/*^{C}$ Henry's law constant, $H/m'$ atm mol't 208.0 × 10 <sup>4</sup> 00 $W_{1}$ (compiler) (compiler) (compiler)OU WALLIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1), The air entered through the top of the vessel and the desired flow resel and the distribution was and resultion was and re	ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.
(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P.         COMPONENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]         (2) Myponga water         (2) Myponga water         VARIABLES:         7/K = 293         Korrent Merry's law constant, Hm' atm mol'         20       8.0 × 10 <sup>4</sup> 0.6873       5.984         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         The apparatus consisted of a stripping vessel and the desired flow rate maintained. Samples of the solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped with a capillary column.       Source and purity not given.         (2) Mackay, D.; Shiu, W. Y.; Sutherland, R. P.       ESTIMATED ERRORS:         Solubility:       Not specified.         catter:       ± 0.1 K.		REFERENCES:
COMPONENTS: (1) Dibromochloromethane; CHBr2CI; [124-48-1] (2) Myponga waterORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. <i>Environ. Sci. Technol.</i> 1984, 18, 518-21.VARIABLES: $T/K = 293$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: 		(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. <i>Environ. Sci. Technol.</i> 1984, 18, 518-21.(2) Myponga waterPREPARED BY: A. L. Horvath $T/K = 293$ A. L. HorvathEXPERIMENTAL VALUES: $t'^{\circ}$ CHenry's law constant, 		
(2) Myponga waterEnviron. Sci. Technol. 1984, 18, 518-21.VARIABLES: $T/K = 293$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: $l^{0}C$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> $100 w_1$ 10 <sup>4</sup> $x_1$ (compiler) 0.6873 5.984AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCED URE: similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desiref flow rate added to the stripping vessel and the desiref flow rate added to the stripping vessel and the desiref flow rate added to the stripping vessel and the desire flow rate added to the stripping vessel and the desire flow rate added to the stripping vessel and the desire flow rate exert taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) PH = 7.6, 370 mg salt/dm <sup>3</sup> , 60 mg (2) CaCO <sub>3</sub> /dm <sup>3</sup> , 10 mg organic carbon/dm <sup>3</sup> .ESTIMATED ERRORS: Solubility: Not specified. Temperature: $\pm$ 0.1 K.REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS:	ORIGINAL MEASUREMENTS:
VARIABLES: $T/K = 293$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: $t^{PC}$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 100 w, (compiler) $10^4 x_1$ (compiler)20 $8.0 \times 10^4$ $0.6873$ $5.984$ AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analyzed by gas chromatograph equip- ped with a capillary column.Solution was rest and the desired flow rate termerature: $\pm 0.1$ K.REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.PREPARED BY: A. L. Horvath	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
$T/K = 293$ A. L. HorvathEXPERIMENTAL VALUES: $l^{9}C$ Henry's law constant, $H/m^{3}$ atm mol'1 $20$ $8.0 \times 10^{4}$ $100 w$ , (compiler) $0.6873$ $10^{4} x$ , (compiler)20 $8.0 \times 10^{4}$ $0.6873$ $5.984$ METHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). 	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
EXPERIMENTAL VALUES: $t^{PC}$ Henry's law constant, $H/m^3$ atm mol'100 w, (compiler)104 x, (compiler)20 $8.0 \times 10^4$ 0.68735.984AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.SOURCE AND PURITY OF MATERIALS: 	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES:	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. <u>1984</u> , 18, 518-21. PREPARED BY:
$t^{0}$ CHenry's law constant, H/m³ atm mol <sup>-1</sup> 100 w <sub>1</sub> (compiler)10 <sup>4</sup> x <sub>1</sub> (compiler)20 $8.0 \times 10^{-4}$ $0.6873$ $5.984$ AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.SOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) pH = 7.6, 370 mg salt/dm³, 60 mg CaCO <sub>3</sub> /dm³, 10 mg organic carbon/dm³.ESTIMATED ERRORS: Solubility: Not specified. Temperature: $\pm 0.1$ K.REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. 	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. <u>1984</u> , 18, 518-21. PREPARED BY: A. L. Horvath
20 $8.0 \times 10^4$ $0.6873$ $5.984$ AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) pH = 7.6, 370 mg salt/dm <sup>3</sup> , 60 mg CaCO <sub>3</sub> /dm <sup>3</sup> , 10 mg organic carbon/dm <sup>3</sup> .         ESTIMATED ERRORS:         added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extracted, they were analyzed by gas chromatograph equipped with a capillary column.       Solubility: Not specified. Temperature: ± 0.1 K.         REFERENCES:         (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: <i>T</i> /K = 293 EXPERIMENTAL VALUES:	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. <u>1984</u> , 18, 518-21. PREPARED BY: A. L. Horvath
AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         Source and purity of MATERIALS:         (1) Source and purity not given.         (2) pH = 7.6, 370 mg salt/dm <sup>3</sup> , 60 mg CaCO <sub>3</sub> /dm <sup>3</sup> , 10 mg organic carbon/dm <sup>3</sup> .         ESTIMATED ERRORS:         maintained. Samples of the solution being stripped were taken for analysis. After samples were extracted, they were analyzed by gas chromatograph equipped with a capillary column.         REFERENCES:         (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's law constant, $H/m^{3}$ atm mol <sup>-1</sup>	ORIGINAL MEASUREMENTS:Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY:A. L. Horvath $100 w_i$ $(compiler)$ $10^4 x_i$ $(compiler)$
AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel         similar to that described by Mackay et al. (ref. 1).         The air entered through the top of the vessel and the         diffuser was close to the bottom. The solution was         added to the stripping vessel and the desired flow rate         maintained. Samples of the solution being stripped         were taken for analysis. After samples were extract-         ed, they were analyzed by gas chromatograph equip-         ped with a capillary column.         REFERENCES:         (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P.         Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t^{\circ}C$ Henry's law constant, $H/m^3$ atm mol' <sup>1</sup> 20 8.0 × 10 <sup>4</sup>	ORIGINAL MEASUREMENTS:Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY:A. L. Horvath $100 w_i$ $(compiler)$ $0.6873$ $5.984$
<ul> <li>METHOD/APPARATUS/PROCEDURE:</li> <li>The apparatus consisted of a stripping vessel similar to that described by Mackay et al. (ref. 1). The air entered through the top of the vessel and the desired flow rate added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extracted, they were analyzed by gas chromatograph equipped with a capillary column.</li> <li>SOURCE AND PURITY OF MATERIALS:</li> <li>Source and purity not given.</li> <li>(2) pH = 7.6, 370 mg sall/dm<sup>3</sup>, 60 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 10 mg organic carbon/dm<sup>3</sup>.</li> <li>ESTIMATED ERRORS:</li> <li>Solubility: Not specified.</li> <li>Temperature: ± 0.1 K.</li> <li>REFERENCES:</li> <li>(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.</li> </ul>	COMPONENTS:(1) Dibromochloromethane; CHBr2Cl; [124-48-1](2) Myponga waterVARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 20 $8.0 \times 10^{-4}$	ORIGINAL MEASUREMENTS:Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY:A. L. Horvath $100 w_i$ $(compiler)$ $0.6873$ $5.984$
The apparatus consisted of a stripping vessel similar to that described by Mackay et al. (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.(1) Source and purity not given. (2) $pH = 7.6, 370 \text{ mg salt/dm}^3, 60 \text{ mg}$ CaCO <sub>3</sub> /dm <sup>3</sup> , 10 mg organic carbon/dm <sup>3</sup> .ESTIMATED ERRORS: solubility: Not specified. Temperature: $\pm 0.1 \text{ K}.$ REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's law constant, $H/m^{3}$ atm mol <sup>-1</sup> 20 8.0 × 10 <sup>-4</sup> AUXILIARY II	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 0.6873 $0.6873$ S.984
added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extracted, they were analyzed by gas chromatograph equipped with a capillary column.       ESTIMATED ERRORS:         Solubility:       Not specified.         REFERENCES:       (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. 1979, 13, 333.	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t^{\circ}C$ Henry's law constant, $H/m^3$ atm mol' <sup>1</sup> 20 8.0 × 10 <sup>4</sup> AUXILIARY II METHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS:         Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.         Environ. Sci. Technol. 1984, 18, 518-21.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         0.6873         5.984
Imamander. Solution of the soluticance of the soluticance of the solution of th	COMPONENTS: (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1] (2) Myponga water VARIABLES: T/K = 293 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's law constant, $H/m^{3}$ atm mol <sup>-1</sup> 20 8.0 × 10 <sup>-4</sup> AUXILIARY II METHOD/APPARATUS/PROCEDURE: The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	ORIGINAL MEASUREMENTS:Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY:A. L. Horvath $100 w_i$ $(compiler)$ $0.6873$ 5.984NFORMATIONSOURCE AND PURITY OF MATERIALS:(1) Source and purity not given.(2) $pH = 7.6, 370 mg salt/dm³, 60 mg CaCO_3/dm³, 10 mg organic carbon/dm³.$
REFERENCES: (1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.	COMPONENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]         (2) Myponga water         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}C$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 20 $8.0 \times 10^{-4}$ METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel         similar to that described by Mackay <i>et al.</i> (ref. 1).         The air entered through the top of the vessel and the         diffuser was close to the bottom. The solution was         added to the stripping vessel and the desired flow rate         maintained_Samples of the sculution heing stripping	ORIGINAL MEASUREMENTS:Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY:A. L. Horvath $100 w_i$ $(compiler)$ $0.6873$ $5.984$ NFORMATIONSOURCE AND PURITY OF MATERIALS:(1) Source and purity not given.(2) pH = 7.6, 370 mg salt/dm³, 60 mg CaCO <sub>3</sub> /dm³, 10 mg organic carbon/dm³.ESTIMATED ERRORS:
(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.	COMPONENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]         (2) Myponga water         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}C$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 20 $8.0 \times 10^{-4}$ METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1).         The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped were taken for analysis. After samples were extracted, they were analyzed by gas chromatograph equipped with a capillary column.	ORIGINAL MEASUREMENTS:         Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.         Environ. Sci. Technol. 1984, 18, 518-21.         PREPARED BY:         A. L. Horvath $100 w_i$ $(compiler)$ $0.6873$ 5.984         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source and purity not given.         (2) pH = 7.6, 370 mg salt/dm², 60 mg CaCO <sub>3</sub> /dm³, 10 mg organic carbon/dm³.         ESTIMATED ERRORS:         Solubility:       Not specified.         Temperature: $\pm$ 0.1 K.
	COMPONENTS:         (1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]         (2) Myponga water         VARIABLES: $T/K = 293$ EXPERIMENTAL VALUES: $t'^{\circ}C$ Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 20 $8.0 \times 10^{-4}$ METHOD/APPARATUS/PROCEDURE:         The apparatus consisted of a stripping vessel         similar to that described by Mackay et al. (ref. 1).         The air entered through the top of the vessel and the         diffuser was close to the bottom. The solution was         added to the stripping vessel and the desired flow rate         maintained. Samples of the solution being stripped         were taken for analysis. After samples were extractied, they were analyzed by gas chromatograph equipped with a capillary column.	ORIGINAL MEASUREMENTS: Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518-21.PREPARED BY: A. L. Horvath104 $x_1$ (compiler) 0.6873104 $x_2$ (compiler) 0.6873104 $x_2$ (compiler) 0.6873NFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Source and purity not given. (2) $pH = 7.6, 370 \text{ mg salt/dm}^3, 60 \text{ mg CaCO_3/dm}^3, 10 \text{ mg organic carbon/dm}^3.ESTIMATED ERRORS:Solubility:Temperature:Not specified.\pm 0.1 \text{ K}.REFERENCES:$

	8
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Dibromochloromethane; CHBr <sub>2</sub> Cl; [124-48-1]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Myponga water + humic acid	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$
20 $7.7 \times 10^{-4}$	0.7141 6.2191
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate mainting a complex of the solution being stripped.	<ol> <li>Source and purity not given.</li> <li>pH = 7.9, 20 mg humic acid/dm<sup>3</sup>, 370 mg salt/dm<sup>3</sup>, 68 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 17 mg organic carbon/dm<sup>3</sup>.</li> </ol>
were taken for analysis. After samples were extract-	ESTIMATED ERRORS:
ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.
	-

COMPONENTS:	EVALUATOR:
<ol> <li>Tribromomethane (bromoform); CHBr<sub>3</sub>; [75-25-2]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; $H_2O$ ; [7732-18-5]	February 1993.

## CRITICAL EVALUATION:

82

The tribromomethane (1) and water (2) binary system is discussed in two parts; part 1 is tribromomethane (1) in water (2) and part 2 is water (2) in tribromomethane (1).

Part 1. All the available data for the solubility of tribromomethane (1) in water (2) were considered acceptable for use in the smoothing equation except the following reported values. The datum of Squire and Caines (ref. 1) is rejected because of the lack of details on the purity of the sample, method of the experimental work, and significant deviation from other measurements. The approximate value of Booth and Everson (ref. 2) is also rejected because it is significantly lower than later studies. The datum of McNally and Grob (ref. 3) is rejected because the reported solubility is rather higher than other determinations. The data of Wright *et al.* (ref. 4) are contrary to the general trend of the temperature dependence of the solubilities in water (formation of maximum) and are rejected.

The remaining data, mainly at 298 K, are in reasonable agreement although further studies are required before any values can be recommended without qualification.

The tentative solubility values in mass per cent for tribromomethane (1) in water (2) over the 283 to 303 K temperature range were used to obtain the following equation:

Solubility  $[100 w_1] = 2.371 - 0.01517 (T/K) + 2.7808 \times 10^{-5} (T/K)^2$ 

This regression equation was established using the combined data from (refs. 5 - 12), which yielded a standard deviation of  $1.6 \times 10^{-2}$ .

The tentative solubility values in mass per cent at 5 K intervals for tribromomethane (1) in water (2) are presented in the Table 1.

The curve obtained from the smoothing equation is shown in Figure 1. It should be noted that the solubility minimum, which is discussed in the Preface, appearing in the 283 to 303 K temperature interval is not pronounced.



Figure 1. Solubility of Tribromomethane (1) in Water (2).

(continued next page)

COMPONENTS:			EVALUATOR:	
(1) Tribromomethane (brom CHBr <sub>3</sub> ; [75-25-2]	oform);		A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.	
(2) Water; H <sub>2</sub> O; [7732-18-5]			February 1993.	
CRITICAL EVALUATION	(continued)	olubility of 1	Frihramamathana (1) in Watar (2)	
Table	. Tentative S	olubility of	Thoromomethane (1) in water (2)	
Ter	nperature	5	Solubility	
°C	K	100 w <sub>1</sub>	$10^4 x_1$	
10	283.15	0.305	2.18	
15	288.15	0.309	2.21	
20	293.15	0.314	2.24	
25	298.15	0.320	2.29	

**Part 2.** The only datum available for the solubility of water (2) in tribromomethane (1) is that of Mackay *et al.* (ref. 9); therefore, no Critical Evaluation is possible. The interested reader is referred to the relevant Compilation Sheet for the experimental solubility.

2.34

0.328

### **REFERENCES:**

1. Squire, P. W.; Caines, C. M. Pharm. J. 1905, 74, 784.

30

2. Booth, H. S.; Everson, H. E. Ind. Eng. Chem. 1948, 40, 1491.

303.15

- 3. McNally, M. E.; Grob, R. L. J. Chromatogr. 1984, 284, 105.
- 4. Wright, D. A.; Sandler, S. I.; DeVoll, D. Environ. Sci. Technol. 1992, 26, 1828.
- 5. Gross, P. M.; Saylor, J. H. J. Am. Chem. Soc. 1931, 53, 1744.
- 6. van Arkel, A. E.; Vles, S. E. Recl. Trav. Chim. Pays-Bas 1936, 55, 407.
- 7. O'Connell, W. L. Trans. Am. Inst. Mech. Eng. 1963, 226, 126.
- 8. Symons, J. M. et al. Treatment Techniques for Controlling Trihalomethanes in Drinking Water, U. S. EPA Report 600/2-81-156, Cincinnati, OH., 1981.
- 9. Mackay, D. et al. Volatilization of Organic Pollutants from Water, U. S. EPA Report 600/2-82-019, Athens, Georgia, <u>1982</u>, (PB 82-230939).
- 10. Nicholson, B. C.; Maguire, B. P.; Bursill, D. B. Environ. Sci. Technol. 1984, 18, 518.
- 11. Munz, C. D. Ph. D. Thesis, Stamford University, Stamford, CA., 1985, 306 pp.
- 12. Warner, H. P.; Cohen, J. M.; Ireland, J. C. Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB87-212684, Cincinnati, OH., July 1987.

COMPONENTS:			ORIO	GINAL MEAS	UREMENTS:
(1) Tribromomethane (bromoform);			Squi	re, P. W.; Ca	ines, C. M.
$CHBr_3; [73-23-2]$	2] 722 19 51		Pha	rm. J. <u>1905</u> , 2	74, 784-6.
(2) water; $H_2O$ ; [7	/32-18-3]				
VARIABLES:			PRE	PARED BY:	
<i>T</i> /K = 293			A. I	. Horvath	
EXPERIMENTAL	VALUES:				
	t/°C	V,/V2	(	100 w <sub>1</sub> compiler)	10 <sup>4</sup> x, (compiler)
	20	$1.25 \times 10^{-3}$		0.361	2.582
		AUXILIARY I	NFOR	MATION	
METHOD/APPARA	TUS/PROCF	DURE:	sou	RCE AND PI	IRITY OF MATERIALS:
Details are no	ot available.		(1)	Source and 1	purity not given.
			(2)	Purity not gi	ven.
			ESTIMATED ERRORS:		
			Solu	bility:	Not specified.
			Tem	perature:	Not specified.
COMPONENTS:			ORIC	GINAL MEAS	UREMENTS:
(1) Tribromometha	ne (bromoforn	n);	Gros	ss, P. M.; Say	lor, J. H.
CHBr <sub>3</sub> ; [75-25-2]			J. A	m. Chem. Soc	. <u>1931</u> , <i>53</i> , 1744-51.
(2) Water; $H_2O$ ; [7	732-18-5]				
VARIABLES:			PRE	PARED BY:	
T/K = 288 - 303			A. I	Horvath	
EXPERIMENTAL	VALUES:				
	t/°C	1000 g1/g		100 w.	$10^4 x_1$
		51 61 62	(	compiler)	(compiler)
	15 30	3.01 3.19		0.301 0.319	2.15 2.28
		AUXILIARY I	NFORI	MATION	
METHOD/APPARA	TUS/PROCE	DURE:	sou	RCE AND PU	RITY OF MATERIALS:
An excess of	tribromometh	ane in 500 g water	(1)	Eastman Koo	lak Co., further purified by
was shaken for 12 hours in a thermostat bath. Sam- ples were then withdrawn and read against water in the interfacementar made by Zairs (ref. 1)			(2)	Distilled.	ezing defore use.
A detailed description of the complete procedure is given in a Ph. D. thesis (ref. 2).		ESTI	MATED ERR	ORS:	
		Solu	bility:	± 1.5 %.	
			Tem	perature:	<b>±</b> 0.02 K.
			REFI	ERENCES:	
			(1)	Gross, P. M	. J. Am. Chem. Soc. <u>1929</u> , 51,
			(2)	Saylor, J. H.	Ph. D. Thesis, Duke Univer-
				July Pullidi	·) <u>ACKY</u> '

•

	85
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tribromomethane (bromoform);	van Arkel, A. E.; Vles, S. E.
(2) Water; $H_{2}O$ ; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.
VARIABLES:	PREPARED BY:
T/K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 100 n <sub>1</sub> /g <sub>2</sub>	$\begin{array}{ccc} 100 \ w_1 & 10^3 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$30.0  1.26 \times 10^{-3}$	0.318 0.227
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified.
	Temperature: $\pm 0.5 \text{ K}$ (compiler).
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tribromomethane (bromoform);	Booth, H. S.; Everson, H. E.
$(2) \text{ Water: } H_{-}O^{-} [7732-18-5]$	Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.
VARIABLES:	PREPARED BY:
T/K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $V_{1}/V_{2}$	$\begin{array}{ccc} 100 \ w_1 & 10^5 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
25.0 < 0.02	< 0.058 < 4.1
AUXILIARY I	NFORMATION
	SOUDCE AND DUDITY OF MATERIALS
The equilibrium was ensured by repeated	(1) Source not given Commercial reagent C. P.
shaking and centrifuging of a tribromomethane and water mixture in a stoppered Goetz tube and return- ing it to a constant temperature water bath. The	<ul> <li>(2) Distilled.</li> </ul>
difference between the total amount of tribromo- methane which was added and the amount remaining	ESTIMATED ERRORS:
methane dissolved in the known volume of water. The determination of the excess amount of tri-	Solubility: < 100 %. Temperature: ± 1 K (compiler).
promomethane added is described by Hamslick (ref. 1).	
	(1) Hanslick R S <i>Ph D Thesis</i> Columbia
	University, <u>1935</u> .

00		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform);	O'Connell, W. L.	
(2) Water: $H_{2}O$ : [7732-18-5]	Trans. Am. Inst. Mech. Eng. <u>1963</u> , 226, 126-32.	
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 100 $g_1/g_2$	$\begin{array}{ccc} 100 \ w_{I} & 10^{4} \ x_{J} \\ (compiler) & (compiler) \end{array}$	
20 0.311	0.310 2.22	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	(1) Dow Chemicals Co., used as received.	
	(2) Distilled (compiler).	
	ESTIMATED ERRORS:	
	Solubility: Not specified.	
	Temperature. <u>T</u> 0.5 K (complet).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform);	Symons, J. M. et al.	
$CHBr_{3}; [75-25-2]$	Treatment Techniques for Controlling Trihalo-	
(2) water, $n_20$ , $[7752-10-5]$	600/2-81-156, Cincinnati, OH., <u>1981</u> .	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 $2.4 \times 10^{-2}$	0.3107 2.221	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are given elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Cincinnati tap water.</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified.	
	$1 \text{ cmperature:} \pm 0.5 \text{ K},$	
	REFERENCES:	
	<ol> <li>Werner, U. S. Environmental Protection Agency, Cincinnati, OH., Personal Communication, <u>1980</u>.</li> </ol>	

		1
COMPONENTS:		ORIGINAL MEASUREMENTS:
(1) Tribromomethane (bromoform);		Mackay, D. et al.
(2) Water; H <sub>2</sub> O; [7732-18-5]		Volatilization of Organic Pollutants from Water, U. S. EPA Report 600/2-82-019, Athens, Georgia, <u>1982</u> , (PB 82-230939).
VARIABLES:		PREPARED BY:
<i>T/</i> K = 298		A. L. Horvath
EXPERIMENTAL VALUES:		·
t/°C c <sub>1</sub> /mol m <sup>-3</sup>	100 w <sub>1</sub> (compiler)	$\begin{array}{cccc} 10^4 x_1 & 100 w_2 & 10^3 x_2 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
25 12.0	0.3032	2.168 $7.14 \times 10^{-3}$ 1.0
	AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE	:	SOURCE AND PURITY OF MATERIALS:
An aqueous solution with an exce bromomethane was stirred for 1 day bef introduced into a 1 liter glass vessel. Ni entered at the bottom of the vessel throu	ess of tri- ore being trogen ugh a sintered	<ol> <li>Source and purity not given.</li> <li>Doubly distilled.</li> </ol>
soap bubble flow meter. The concentration become the properties of the soap bubble flow meter was determined	ion of tri-	ESTIMATED ERRORS:
matography. The GC was equipped with flame ionization detector and an electron detector.	n both dual n capture	Solubility: Not specified. Temperature: $\pm$ 1.0 K.
COMPONENTS:		ORIGINAL MEASUREMENTS:
CHBr <sub>3</sub> ; [75-25-2]		I Chromotogr 1984 284 105-16
(2) Water; H <sub>2</sub> ; [7732-18-5]		5. Chromatogr. <u>1204</u> , 204, 105-10.
VARIABLES:		PREPARED BY:
T/K = 303		A. L. Horvath
EXPERIMENTAL VALUES:		
t/°C g₁V	/2 <sup>-1</sup> /g m <sup>-3</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
30 39	931	0.3933 2.814
AUXILIARY I		NFORMATION
METHOD/APPARATUS/PROCEDURE	:	SOURCE AND PURITY OF MATERIALS:
Standard tribromomethane solution creasing concentration were prepared in flasks as described elsewhere (ref. 1). T organic liquid phase was 25 ml and the	ns of in- volumetric he aqueous vapor phase	<ol> <li>Chemical Service, West Chester, PA., U. S. A., purest grade available.</li> <li>Distilled and run through two Barnstead purification cartridges.</li> </ol>
above the mixture was 35 ml. The syste ed to reach equilibrium in a constant ten	m was allow- nperature bath	ESTIMATED ERRORS:
and then a sample was injected into the gas chromato- graph. Values reported are an average of at least 5 measurements.		Solubility: $\pm$ 5.92 % std. dev. Temperature: $\pm$ 0.5 K (compiler).
		REFERENCES:
		(1) McNally, M. E.; Grob, R. L. J. Cromatogr.

88	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tribromomethane (bromoform); CHBr.: (75-25-2)	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Water; $H_2O$ ; [7732-18-5]	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	1
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_1 & 10^4 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$20$ $4.3 \times 10^{-4}$	0.3093 2.211
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	<ol> <li>Source and purity not given.</li> <li>Distilled and deionized.</li> </ol>
diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate	ESTIMATED ERRORS:
were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	Solubility: $\pm 0.3 \times 10^{-4}$ std. dev. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.
COMPONENTS	OPIGINAL MEASUREMENTS
(1) Tribromomethane (bromoform);	Munz, C. D.
CHBr <sub>3</sub> ; [75-25-2]	Ph. D. Thesis, Stamford University, Stamford,
(2) water; $H_2O$ ; [7/32-18-5]	СА., <u>1985</u> , 306 рр.
VARIABLES:	PREPARED BY:
T/K = 283 - 303	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
$\begin{array}{cccc} 10 & 9.95 \times 10^{-3} \\ 20 & 1.74 \times 10^2 \\ 30 & 2.85 \times 10^{-2} \end{array}$	0.3086 2.206 0.3228 2.308 0.3437 2.458
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess amount of tribromomethane with water was introduced into 30 ml centrifuge tubes that were closed with a screw cap. The tubes were rotated for $2 + 10^{-10}$ for $2 + 10^{-10}$ km in the screw cap.	<ol> <li>Source and purity not given.</li> <li>Distilled and deionized.</li> </ol>
for 5 days inside a temperature controlled chamber at 10, 20, and $30^{\circ}$ C, respectively. Samples were taken from the tubes and analyzed by gas chemotography	ESTIMATED ERRORS:
after liquid-liquid extraction. The experiments were carried out in duplicates.	Solubility: $\pm$ 3.9 %.Temperature: $\pm$ 0.5 K.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform);	Warner, H. P.; Cohen, J. M.; Ireland, J. C.	
(2) Water; $H_2O$ ; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB-87-212684, Cincinnati, OH., July 1987.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	100	
$H/m^3$ atm mol <sup>-1</sup>	$\begin{array}{c} 100 \ w_i \\ \text{(compiler)} \\ \end{array} \begin{array}{c} 10 \ x_i \\ \text{(compiler)} \end{array}$	
24.85 $5.32 \times 10^{-4}$	0.3512 2.511	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The original method and apparatus for the determination of Henry's law constants, described by Mackay <i>et al.</i> (ref. 1), were used. The general procedure was to add an excess quantity of tribromomethane to distilled deionized water and mix	<ol> <li>Source not given. Purest quality available and was used without further purification, stated purity of &gt; 99 %.</li> <li>Distilled and deionized.</li> </ol>	
the system overnight. A portion of the solution was returned to the stripping vessel. Tribromomethane	ESTIMATED ERRORS:	
was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from the log of the concentration versus time plot. The experimental data values are averages	Solubility: $\pm 6 \%$ std. dev. Temperature: $\pm 0.05$ K.	
of two or more replicates.	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Technol. <u>1979</u> , 13, 333.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform);	Wright, D. A.; Sandler, S. I.; DeVoll, D.	
CHBr <sub>3</sub> ; [75-25-2]	Environ. Sci. Technol. <u>1992</u> , 26, 1828-31.	
(2) Water; $H_2O$ ; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 293 - 323	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C γ <sup>∞</sup>	$\begin{array}{ccc} 100 \ w_i, & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.3962.830.4543.250.3452.47	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A differential static cell equilibrium apparatus was used to measure the infinite dilution activity coefficient of tribromomethane ( $\gamma^{\infty}$ ) in water. Cells containing degased water ware submerged in a ther-	<ol> <li>Source and purity not given.</li> <li>Distilled, filtered and deionized.</li> </ol>	
mostated water bath. Tribromomethane was injected	ESTIMATED ERRORS:	
on, and the cells were allowed to equilibrate. The stirrer was then turned off and the differential pressure was recorded. The experiment was repeated at least three times at each temperature.	Solubility: See above. Temperature: $\pm$ 0.05 K.	
90		
---	---	--
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform);	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.	
(2) Milang water	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.	
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 $4.6 \times 10^4$	0.289 2.068	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was	<ol> <li>Source and purity not given.</li> <li>pH = 8.4, 700 mg salt/dm<sup>3</sup>, 140 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 13 mg organic carbon/dm<sup>3</sup>, 110 mg suspended solid/dm<sup>3</sup>.</li> </ol>	
added to the stripping vessel and the desired flow rate maintained. Samples of the solution being stripped	ESTIMATED ERRORS:	
ed, they were analyzed by gas chromatograph equipped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.	
	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ Sci Technol 1979 13 223	
	Environ. Sci. Technol. <u>1979</u> , 13, 555.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Tribromomethane (bromoform); CHBr <sub>3</sub> ; [75-25-2]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.	
(2) Myponga water	Environ. Sci. Technol. <u>1984</u> , 18(7), 518-521.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES		
$t/^{\circ}C$ Henry's law constant.	$100 w_{i}$ $10^{4} x_{i}$	
<i>H</i> /m <sup>3</sup> atm mol <sup>-1</sup>	(compilér) (compilér)	
20 4.1 × 10 <sup>-4</sup>	0.324 2.32	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	<ol> <li>Source and purity not given.</li> <li>pH = 7.6, 370 mg salt/dm<sup>3</sup>, 60 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 10 mg organic carbon/dm<sup>3</sup>.</li> </ol>	
diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate	ESTIMATED ERRORS:	
were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.	
	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.	

.

	91
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Tribromomethane (bromoform); CHBr.: [75-25-2]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Myponga water + humic acid	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$
20 $4.3 \times 10^{-4}$	0.3093 2.173
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow rate	<ol> <li>Source and purity not given.</li> <li>pH = 7.9, 20 mg humic acid/dm<sup>3</sup>, 370 mg salt/dm<sup>3</sup>, 68 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 77 mg organic carbon/dm<sup>3</sup>.</li> </ol>
maintained. Samples of the solution being stripped were taken for analysis. After samples were extract- ed, they were analyzed by gas chromatograph equip- ped with a capillary column.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.1 K.
	<ul> <li>REFERENCES:</li> <li>(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. <i>Environ. Sci. Technol.</i> <u>1979</u>, 13, 333.</li> </ul>

COMPONENTS:	EVALUATOR:
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; $H_2O$ ; [7732-18-5]	March 1993.

## **CRITICAL EVALUATION:**

92

The trichloromethane (1) and water (2) binary system is discussed in two parts; part 1 is trichloromethane (1) in water (2) and part 2 is water (2) in trichloromethane (1).

Part 1. The solubility of trichloromomethane (1) in water (2) has been studied by 54 workers with reasonable agreement of reported values from a majority of the investigators.

The experimental data of most investigators in the 273 to 333 K temperature interval are sufficiently reliable to use in the smoothing equation, see Figure 1. (The solid curve in the figure was obtained from the regression of all values shown in the figure.) The experimental work of several workers was not used for the smoothing equation for a variety of reasons. The solubility data of Herz (ref. 1), Salkowskii (ref. 2), Ababi et al. (ref. 3), Svetlanov et al. (ref. 4), Sato and Nakijima (ref. 5), Veith et al. (ref. 6), McNally and Grob (ref. 7), and Howe et al. (ref. 8) are markedly lower than other results and are therefore rejected. The data of McCollum (ref. 9), Booth and Everson (ref. 10), Aref'eva et al. (ref. 11), Coca et al. (ref. 12), and Warner et al. (ref. 13) are several per cent higher than the smoothed solubility values and are also rejected. Although, the measured solubility data of Okuda (ref. 14) above 310 K are in reasonable agreement, the low temperature data, that is, below 303 K are very uncertain. Consequently, the low temperature data were not included in the regression analysis. Similar inconsistency was found in the data of Hunter-Smith et al. (ref. 15) and were excluded from further considerations. Lincoff and Gossett (ref. 16) presented two methods for the determination of the solubility. The data determined by the Batch Air Stripping method shows more compatible results and are in good agreement, whereas it appears the EPICS method gave values too low. Therefore, only the solubility data measured by the Batch Air Stripping method were used for fitting the regression equation.

The combined data of 40 laboratories were correlated to obtain the following mass per cent (1) equation:

Solubility  $[100 w_l] = 12.5333 - 7.40557 \times 10^2 (T/K) + 1.16374 \times 10^4 (T/K)^2$ 

which shows a standard deviation of  $5.6 \times 10^{-2}$ . This equation, which shows a maximum deviation of 18 % (but much less for most of the data), may be considered as representing the combined solubility values in the 273 to 333 K temperature range.

The solubility minimum (see Figure 1) calculated from the regression equation is, in mass per cent (1), 0.752 at 318.18 K. The presence of a minimum in the solubility versus temperature behavior for the halogenated hydrocarbon-water systems is discussed in the Preface.

The recommended solubility values in mass per cent at 5 K intervals for trichloromomethane (1) in water (2) are presented in Table 1.

Table 1. Recommended Solubility of Trichloromethane (1) in Water (2)

Ter	nperature	Solub	ility
°C	ĸ	100 w <sub>1</sub>	$10^3 x_1$
0	273.15	0.988	1.503
5	278.15	0.938	1.427
10	283.15	0.895	1.361
15	288.15	0.857	1.303
20	293.15	0.825	1.254
25	298.15	0.798	1.212
30	303.15	0.778	1.182
35	308.15	0.763	1.159
40	313.15	0.755	1.147
45	318.15	0.752	1.142
50	323.15	0.755	1.147
55	328.15	0.763	1.159
60	333.15	0.778	1.182





*Т /* К

**Part 2.** The solubility of water (2) in trichloromomethane (1) has been studied by 25 workers as recorded in the Compilation Sheets immediately following this Critical Evaluation. The reported solubilities in the 263 to 323 K temperature interval are quite acceptable for using the smoothing equation as seen in Figure 2.

All the available data for the solubility of water (2) in trichloromomethane (1) have been included for the fitting of the correlating equation with the following exceptions. The measured data of Herz (ref. 1), Evans (ref. 17), De Minjer (ref. 18), Reinders and De Minjer (ref. 19), and Kudryavtseva and Krutikova (ref. 20) are rejected because they are significantly higher than later studies. The data of Donahue and Bartell (ref. 21), Tettamanti *et al.* (ref. 22), Christyakov and Shapurova (ref. 23), and Antropov *et al.* (ref. 24) are markedly lower than other determinations and are therefore rejected.

The remaining data of 16 laboratories are in good agreement and the correlated values above 253 K are recommended although the low temperature data (below 273 K) are probably dubious. Further studies are required to establish reliable values in the lower temperature range below 273 K.

The equation for the mass per cent solubility of water (2) in trichloromomethane (1) between 263 and 323 K is as follows:

Solubility  $[100 w_l] = 6.1195 \times 10^2 - 2.0623 \times 10^3 (T/K) + 7.20879 \times 10^6 (T/K)^2$ 

This equation represents the combined data with a standard deviation of 2.2 x  $10^{-2}$  and a correlation coefficient of 0.916.

The recommended mass per cent solubility values at 5 K intervals for water (2) in trichloromomethane (1) are presented in the Table 2.

COMPONENTS:	EVALUATOR:
<ol> <li>Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	March 1993.

**CRITICAL EVALUATION: (continued)** 

The curve obtained from the smoothing equation is shown in Figure 2. It may be noted that all the studies show a general increase in the solubility with temperature, which is consistent with the discussion of water solubility versus temperature in the Preface. Often the temperature dependence of the solubility is plotted on a Cox chart, that is,  $\log_{10} x_2$  versus 1/T, where a straight line represents the solubility behavior. This procedure is very useful for the illustration of data covering a limited temperature range. However, for the water (2) trichloromethane (2) system, this method of correlation was found to be less satisfactory than the regression with the normal polynomial equation.

Table 2. Recommended Solubility of Water (2) in Trichloromethane (1)





94

COMPONENTS:	EVALUATOR:
<ol> <li>Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; $H_2O$ ; [7732-18-5]	March 1993.

## **CRITICAL EVALUATION: (continued)**

Figure 3 shows the  $\log_{10} x_2$  versus the reciprocal of absolute temperature behavior for comparison with the behavior shown in Figure 2.



Figure 3. Solubility of Water (2) in Trichloromethane (1); log<sub>10</sub> x<sub>2</sub> versus 1000/T.

## **REFERENCES**:

 $\begin{pmatrix} c \\ c \end{pmatrix}$ 

-1

<u>С</u>,

- 1. Herz, W. Chem. Ber. 1898, 31, 2669-2672.
- 2. Salkowski, E. Biochem. Z. 1920, 107, 191-201.
- Ababi, V.; Popa, A.; Mihaila, Gh. Analele Stiint. Univ. Al. I. Cuza Iasi. Sect. IC. Chem. <u>1964</u>, 10(1), 71-84.
- 4. Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. I.; Treger, Yu. A.; Flid, R. M. Russ. J. Phys. Chem. 1971, 45(4), 488-490.
- 5. Sato, A.; Nakijima, T. Arch. Environ. Health 1979, 34(2), 69-75.
- Veith, G. D.; Macek, K. J.; Petrucelli, S. R.; Carroll, J. Proc. 3<sup>rd</sup>. Ann. Symp. on Aquatic Toxicology, ASTM Publ. 707, Philadelphia, <u>1980</u>, p. 116-129.
- (26) 7. McNally, M. E.; Grob, R. L. J. Chromatogr. <u>1984</u>, 284, 105-116.
  - 8. Howe, G. B.; Mullins, M. E.; Rogers, T. N. AFESC Tyndall Air Force Base, Report ESL-TR-86-66, Vol. 1, Florida, Sept. 1987, 86 pp. (AD-A188 571).

95

96		
CO	MPONENTS:	EVALUATOR:
(1)	Trichloromethane (chloroform); CHC1 <sub>3</sub> ; [67-66-3]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2)	Water; H <sub>2</sub> O; [7732-18-5]	March 1993.
CR	TICAL EVALUATION: (continued)	
RE	FERENCES:	
9.	McCollum, J. L. J. Pharmacol. Exper. Therap.	<u>1930,</u> <i>40</i> , 305.
10.	Booth, H. S.; Everson, H. E. Ind. Eng. Chem.	<u>1948, 40,</u> 1491.
11.	Aref'eva, R. P.; Korenman, I. M.; Gorokhov, A (CA. 91:113 256k).	. A. USSR Patent 672 548, July 5. 1979, 3 pp.
12.	Coca, J.; Diaz, R. M.; Pazos, C. Fluid Phase E	quilibr. <u>1980,</u> 4, 125.
13. Warner, H. P.; Cohen, J. M.; Ireland, J. C. Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Technical Report, PB-87-212684, Cincinnati, OH., July 1987.		
14. Okuda, Y. Arch. Jap. Chir. <u>1968</u> , 37, 700.		
15.	Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S. 7	ellus <u>1983</u> , 35B, 170.
16. Lincoff, A. H.; Gossett, J. M. in Gas Transfer at Water Surfaces by Brutsaert, W. and Jirka, G. H., Eds., D. Reidel Publ. Co., Dordrecht, <u>1984</u> , p. 17-25.		
17. Evans, T. W. Ind. Eng. Chem. Anal. Edit. 1936, 8, 206.		
18. De Minjer, C. H. Doctoral Dissertation, Technische Hoogeschool, Delf, 1939, 165 pp.		
19. Reinders, W.; De Minjer, C. H. Recl. Trav. Chim. Pays-Bas 1947, 66, 573.		
20. Kudryavtseva, G. I.; Krutikova, A. D. J. Appl. Chem. USSR. 1953, 26, 1129.		
21. Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1952, 56, 480.		
22. Tettamanti, K.; Nogradi, M.; Sawinsky, J. Period. Polytech, Chem. Eng. 1960, 4, 201.		
23. Chistyakov, V. M.; Shapurova, V. V. Izv. Vyssh. Uchebn. Zaved., Khim. Khim. Technol. <u>1964</u> , 7, 349.		
24.	Antropov, L. I.; Populyai, V. E.; Simonov, V. D 46, 311. (VINITI No. 3739-71).	D.; Shamsutdinov, T. M. Russ. J. Phys. Chem. <u>197</u>

14 . ,

?:`

٦ ۱,

t

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Chancel, G.; Parmentier, F.	
CHCl <sub>3</sub> ; [67-66-3]	Compt. Rend. <u>1885</u> , 100A, 773-6.	
(2) water; H <sub>2</sub> O; [//32-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 273 - 328	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
0.0 9.87	0.987 1.502 0.890 1.353	
17.4 7.12 29.4 7.05	0.713 1.082 0.708 1.075	
41.6 7.12	0.717 1.089	
<i>1.13 7.74</i>	0.764 1.171	
AUXILIARY I	NFORMATION	
	SOUDCE AND DUDITY OF MATERIALS.	
Details are not available	(1) Prepared from chloral and purified by	
	distillation. (2) Distilled (compiler).	
	(-),(-),	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.2$ K (compiler).	
COMPONENTS: ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Herz, W.	
(2) Water: $H_2O$ : [7732-18-5]	Chem. Ber. <u>1898</u> , 31, 2669-72.	
VARIABLES:	PREPARED BY:	
T/K = 295	A. L. Horvath	
EXPERIMENTAL VALUES:	· · · · · · · · · · · · · · · · · · ·	
t/°C 100 V <sub>1</sub> /V <sub>2</sub> 10	$\begin{array}{llllllllllllllllllllllllllllllllllll$	
22 0.420 0.0	520 0.152 0.102	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The mutual solubility between trichloro- methane and water was determined by a volumetric	(1) Source and purity not given, $d_4 = 1.4760 \text{ g/cm}^3 \text{ at } 22^{\circ}\text{C}.$	
method at constant temperature. The change in the volume upon mixing was measured in a calibrated	(2) Distilled (compiler), $d_4 = 0.9980 \text{ g/cm}^3 \text{ at } 22^{\circ}\text{C}.$	
cylinder. The specific weight of the pure components and the solutions were also determined.		
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: + 0.5 K (compiler).	

98		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Moore, B.; Roaf, H. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Proc. Roy. Soc., London <u>1904</u> , 73, 382-412.	
VARIABLES:	PREPARED BY:	
T/K = 286	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
t/°C 100 w₁	$\begin{array}{ccc} 10^{3} x_{i} & 100 \ w_{i}M_{i}^{-1}/\text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
13 0.95	1.445 $7.96 \times 10^{-3}$	
AUXILIARY	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Pure trichloromethane was dropped from a fine capillary pipette into a tared graduated flask containing distilled water. Samples in a series of	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
flasks of various sizes were prepared and placed on a rotary shaking machine. After several days shaking,	ESTIMATED ERRORS:	
an equilibrium was established. The solubility was determined from the decreased amount of trichloro- methane.	Solubility: $\pm$ 5 %.Temperature: $\pm$ 1 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Rex, A.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Z. Phys. Chem. <u>1906</u> , 55, 355-70.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
$t/^{\circ}$ C 100 $g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
0 1.062 10 0.895 20 0.822 30 0.776	1.05         1.60           0.887         1.35           0.815         1.24           0.770         1.17	
	•	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility of trichloromethane in water was determined in specially designed flasks with calibrat- ed capillary cylinders. After equilibration in a thermostatic bath, the volume and the weight of the samples were measured and the solubility calculated	<ol> <li>Commercial, dried, redistilled and washed before use. B. P. = 61.7°C.</li> <li>Distilled.</li> </ol>	
samples were measured and the solubility calculated.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	

	, 99	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Antonov, G. N.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chim. Phys. <u>1907</u> , 5, 372-85.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C 100 <i>w</i> <sub>1</sub>	$\begin{array}{ccc} 10^{3} x_{i} & 100 \ w_{i} M_{i}^{-i} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
20 0.79	1.20 $6.62 \times 10^{-3}$	
AUXILIARY I	NFORMATION	
	SOURCE AND PURITY OF MATERIALS	
Details are not available.	(1) Source and purity not given.	
	(2) Distilled (compiler).	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 1$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>1</sub> : [67-66-3]	Salkowski, E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Biochem. Z. <u>1920</u> , 107, 191-201.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/kg \text{ m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
20 7.5	0.746 1.13	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A volumetric method similar to that intro- duced by Alexejew was used. Trichloromethane was gradually added to water from a pipet under constant	<ol> <li>Kahlbaum reagent, used as received.</li> <li>Distilled (compiler).</li> </ol>	
agitation. The appearance of clouding was the indi- cation of saturation.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: ± 2 K (compiler).	

100		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Clifford, C. W.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	J. Ind. Eng. Chem. <u>1921</u> , 13, 631-2.	
VARIABLES:	PREPARED BY:	
T/K = 298 - 301	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
$t/^{\circ}$ C 100 $w_t$	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-1} / \text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
24.5         0.084           26.6         0.107           27.8         0.116	$\begin{array}{cccc} 0.554 & 4.66 \times 10^3 \\ 0.705 & 5.94 \times 10^3 \\ 0.764 & 6.44 \times 10^3 \end{array}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The calcium chloride method was used for the determination of water in trichloromethane. The water was measured as the moisture abstracted from	<ol> <li>Distilled (compiler).</li> <li>U. S. P. grade, used after dehydration with CaCl<sub>2</sub>.</li> </ol>	
the samples by air and absorbed in two calcium chloride tubes. The tubes were weighed before and	ESTIMATED ERRORS:	
after absorption. For further details, see the paper by Clifford (ref. 1).	Solubility: About 1 mg. Temperature: ± 0.5 K (compiler).	
	REFERENCES:	
	(1) Clifford, C. W. J. Ind. Eng. Chem. <u>1921</u> , 13, 628.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Nicloux, M.; Scotti-Foglieni, L.	
(2) Water: H-O: $(7732-18-5)$	Ann. physiol. physicochim. biol. <u>1929</u> , 5, 434-82.	
(2) water, 1105, [7752-16-5]		
VARIABLES:	PREPARED BY:	
T/K = 286 - 313	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Ostwald coefficient, L/dimensionless	$\begin{array}{ccc} 100 \ w_{j} & 10^{3} \ x_{j} \\ (compiler) & (compiler) \end{array}$	
13 12.9 20 7.7	0.974 1.48 0.787 1.20	
25 7.0 30 6.2	0.877 1.33 0.944 1.44	
37 3.8 40 3.5	0.753 1.14 0.772 1.17	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An all-glass apparatus which consisted of a liquid container with a mechanical stirrer and a gas storage tube was used for the measurements. Water	<ol> <li>Laboratory reagent, source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
was saturated with trichloromethane gas at a pressure of 750 mmHg. The absorbed gas volume was cal-	ESTIMATED ERRORS:	
culated from the initial dry gas and final water vapor saturated gas volume. The amount of water was de- termined by measuring the mass of mercury displace- ment.	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	McCollum, J. L.	
(2) Water: H O: $(7722, 18, 5)$	J. Pharmacol. Exper. Therap. <u>1930</u> , 40, 305-25.	
(2) Waler, H <sub>2</sub> O, [7752-16-5]		
VARIABLES:	PREPARED BY:	
T/K = 310	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 100 $g_1 V_2^{-1}/g$ c	$m^3$ 100 w, 10 <sup>3</sup> x, (compiler)	
37.5 1.0	0.997 1.517	
AUXILIA	RY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A colorimetric method was used. A 2.5 g chloromethane sample was pipetted into a centrif	tri- (1) Source and purity not given. (2) Distilled (compiler).	
orously shaken in a thermostat bath for 2 days. A shaking, the tube was centrifuged at high speed t	After ESTIMATED ERRORS:	
give a clear upper water layer. From this layer the concentration of trichloromethane was determined	$\begin{array}{c c} & & \\ \hline \\ I & \\ \hline \\ by & \\ \hline \\ Solubility: \\ \pm 1 \%. \end{array}$	
using a colorimetric method.	Temperature: $\pm 1$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Gibby, C. W.; Hall, J.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	J. Chem. Soc. <u>1931</u> , 691-3.	
VARIABLES:	PREPARED BY:	
T/K = 248 - 327	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^4 x_i & 100 \ w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
-25 0.6 × 10 <sup>-2</sup>	4.0 $3.33 \times 10^{-4}$	
-15 0.9 × 10 <sup>-2</sup> -1 1.4 × 10 <sup>-2</sup> 3 10 × 10 <sup>-2</sup>	$\begin{array}{cccc} 0.0 & 5.00 \times 10^{-1} \\ 9.3 & 7.77 \times 10^{-4} \\ 12.6 & 1.05 \times 10^{-3} \end{array}$	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccc} 1.05 & 10 \\ 28.4 & 2.39 \times 10^{-3} \\ 40.3 & 3.39 \times 10^{-3} \end{array}$	
$\begin{array}{cccc} 1' & 0.1 & 10^{-2} \\ 2' & 6.5 & \times 10^{-2} \\ 23 & 7.2 & \times 10^{-2} \end{array}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
$31$ $1.00 \times 10^{-1}$	$\begin{array}{cccc} 7,10 & 7,00 \\ 65.9 & 5.55 \times 10^{-3} \\ 77.7 & 6.55 \times 10^{-3} \end{array}$	
$\begin{array}{cccc} 48 & 1.44 \times 10^{-1} \\ 54 & 1.65 \times 10^{-1} \end{array}$	$\begin{array}{cccc}             94.7 & 8.00 \times 10^{-3} \\             108.3 & 9.16 \times 10^{-3}         \end{array}$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Known weight mixtures of trichloromethan and water were prepared in glass tubes. The tube were heated until all the water had dissolved. Up cooling, the temperature at which the first detect	e (1) Distilled (compiler). s (2) Source not given. Pure B. P., treated with on H <sub>2</sub> SO <sub>4</sub> , NaOH, and H <sub>2</sub> O. Dried with Na <sub>2</sub> CO <sub>3</sub> and refluxing with Mg <sub>3</sub> N <sub>2</sub> before use.	
appearance of turbidity was noted. Stirring was e sured by gentle rocking of the tubes.	n- ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 1$ K.	

102			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Gross, P. M; Saylor, J. H.		
(2) Water: $H_{-}O$ : [7732-18-5]	J. Am. Chem. Soc. <u>1931</u> , 53, 1744-51.		
(2) water, 1120, (7752-10-5)			
VARIABLES:	PREPARED BY:		
T/K = 288 - 303	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-1} / \text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
15 0.852 30 0.771	1.28 0.845 1.16 0.765		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
An excess of trichloromethane in 500 g water was shaken for 12 hours in a thermostat bath. Samples were then withdrawn and read against water in the interferometer made by Zeiss (ref. 1). A datailed description of the complete	<ol> <li>Eastman Kodak Co., shaken with H<sub>2</sub>SO<sub>4</sub>, Hg, and distilled water and fractionated before use.</li> <li>Distilled.</li> </ol>		
procedure is given in a Ph. D. thesis (ref. 2).	ECTIMATED EDDODS.		
	Solubility: $+ 10\%$		
	Temperature: $\pm$ 0.02 K.		
	REFERENCES:		
	<ol> <li>Gross, P. M. J. Am. Chem. Soc. <u>1929</u>, 51, 2362.</li> <li>Saylor, J. H. Ph. D. Thesis, Duke University, Durham, 1930.</li> </ol>		
COMPONENTS: ODIGINAL MEASUREMENTS.			
(1) Trichloromethane (chloroform);	Evans, T. W.		
CHCl <sub>3</sub> ; [67-66-3]	Ind. Eng. Chem. Anal. Edit. <u>1936</u> , 8, 206-8.		
(2) Water; $H_2O$ ; [7732-18-5]			
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C 100 $w_1$ 10 <sup>3</sup> $x_1$ 100 $w_2$ 10 <sup>2</sup> $x_2$ (compiler) (compiler)			
20 0.8 1.	21 0.2 1.3		
ALIXII LARY INFORMATION			
	SOUDCE AND DUDITY OF MATERIALS.		
An oil centrifuge hottle (so-called) which	(1) Source and purity not given		
permitted an accurate reading of both the total volume and the small volume of upper or lower layer was used for the measurements. The necks of the sol-	(2) Distilled (compiler).		
ubility bottles were graduated to 0.05 cm <sup>3</sup> over a 5 cm <sup>3</sup> volume range. These bottles held approximately	ESTIMATED ERRORS:		
50 cm <sup>3</sup> . Following the Hill method, weighted amounts of the two liquids were introduced into the bottles and shaken until equilibrium was reached.	Solubility: $\pm$ 0.2. Temperature: $\pm$ 0.5 K (compiler).		

ORIGINAL MEASUREMENTS:		
van Arkel, A. E.; Vles, S. E.		
Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.		
PREPARED BY:		
A. L. Horvath		
-		
$\begin{array}{ccc} 10^2 x_j & 100 \ w_j M_j^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
0.116 $6.46 \times 10^{-3}$		
NFORMATION		
SOURCE AND DURITY OF MATERIALS.		
(1) Source and purity not given		
(2) Distilled (compiler).		
ESTIMATED ERRORS:		
Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).		
ORIGINAL MEASUREMENTS:		
De Minjer, C. H.		
Doctoral Dissertation, Technische Hoogeschool, Delft, <u>1939</u> , 165 pp.		
PREPARED BY:		
A. L. Horvath		
$\begin{array}{ccc} 10^2 x_i & 100 \ w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
1.31 $1.11 \times 10^{-2}$		
SOURCE AND PURITY OF MATERIALS:		
(1) Distilled. (2) Commercial grade, washed with $H_2SO_4$ and		
before use.		
ESTIMATED ERRORS:		
ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.05 K (compiler).		
ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.05 K (compiler).		

104		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Staverman, A. J.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Recl. Trav. Chim. Pays-Bas <u>1941</u> , 60, 836-41.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_j \cdot $	
$\begin{array}{cccc} 0 & 5.29 \times 10^{-2} \\ 25 & 9.32 \times 10^{-2} \\ 30 & 1.130 \times 10^{-1} \end{array}$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Trichloromethane was shaken with water for about 12 hours in a paraffin thermostat bath. The	<ol> <li>Distilled.</li> <li>Source and purity not given.</li> </ol>	
water content of the organic phase was determined by the Karl Fischer titration method. All measurements were carried out in duplicates. The result is the	ESTIMATED ERRORS:	
average of two determinations. A full description of the method used is given in a thesis (ref. 1).	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	
	REFERENCES:	
	(1) Staverman, A. J. <i>Ph. D. Thesis</i> , Univ. of Leiden, Leiden, Belgium, <u>1938</u> .	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Hutchison, C. A.; Lyon, A. M.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Columbia University Report A-745, July 1, 1943.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 100 $w_{I}M_{I}^{-1}/\text{mol g}^{-1}$	$10^3 x_1$ , $100 w_1$ (compiler) (compiler)	
$25$ $4.88 \times 10^{-3}$	5.80 $8.79 \times 10^{-2}$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A 1 to 15 volume ratio mixture of water and trichloromethane was introduced into an equilibration flask and then lowered into a water thermostat bath.	<ol> <li>Distilled.</li> <li>Source not given. Purified and dried before use.</li> </ol>	
in the assembly was snaken mechanically for about 90 minutes at constant temperature. The amount of water in the organic phase was determined by a modified	ESTIMATED ERRORS:	
Karl Fischer titration. The measurement was done in	Solubility $\pm 0.1 \times 10^4$ and day	
triplicate. The description was taken from a secondary	Temperature: $\pm 0.05$ K.	
triplicate. The description was taken from a secondary source (ref. 1), because the original report is no longer available.	Temperature: ± 0.05 K.	

			105
COMPONENTS:		ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chlo	roform);	McGovern, E. W.	
(2) Water; $H_2O$ ; [7732-18-3	5]	Ind. Eng. Chem. <u>1943</u> , 35, 1230-9.	
VARIABLES:		PREPARED BY:	
T/K = 273 - 323		A L Horvath	
EXPERIMENTAL VALUE	S:		
t/°C	$100 w_i   10^3$ (com	$\begin{array}{ccc} x_1 & 100 \ w_2 & 10^3 \ x_1 \\ \text{piler} & (\text{compiler}) \end{array}$	
0	1.02 1 0.91 1	55 38	
20 25	0.82 1	25	
30	0.77 1	17 0.100 6.59	
40 50	0.76 1	.12 0.135 8.88 .15	
AUXILIARY INFORMATION			
METHOD/APPARATUS/P	ROCEDURE:	SOURCE AND PURITY OF MATERIA	ALS:
Details are not availa	ble.	<ol> <li>Source not given. Commercial re as received.</li> <li>Distilled (compiler).</li> </ol>	agent, used
		ESTIMATED ERRORS:	
		Solubility: Not specified.	
		Temperature: ± 1.0 K (compile	er).

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichloromethane (chloroform);	Reinders, W.; De Minjer, C. H.
(2) Water: $H_{-}O^{-}$ [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1947</u> , 66, 573-604.
	-
VARIABLES:	PREPARED BY:
T/K = 273 - 333	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 100 w <sub>1</sub> 10 <sup>3</sup> (comp	$\begin{array}{cccc} x_1 & 100 \ w_2 & 10^3 \ x_2 \\ \text{iler}) & (\text{compiler}) \end{array}$
0 0.98 1. 25 0.90 1.	49 0.1 6.59
42 0.71 1.1 60 0.75 1.	08 14 0.17 11.2
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturation between trichloromethane and water took place in a water bath. The composition of	<ol> <li>Commercial grade, washed with H<sub>2</sub>SO<sub>4</sub> and K<sub>2</sub>CO<sub>3</sub> solutions, followed by distillation</li> </ol>
both the organic and water phases were determined from their density and refractive index. The constant density and refractive index calibration curves were	(2) Distilled.
established by using several synthetic mixtures.	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm 0.05$ K (compiler).
COMPONENTS	ODIGINAL MEASUDEMENTS.
COMI ONEN IS.	ORIGINAL MEASUREMENTS.
(1) Trichloromethane (chloroform):	Booth, H. S.: Everson, H. E.
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.
<ol> <li>Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> <li>Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.
<ul> <li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> <li>VARIABLES:</li> </ul>	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3. PREPARED BY:
<ul> <li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> <li>VARIABLES: <i>T</i>/K = 298</li> </ul>	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3. PREPARED BY: A. L. Horvath
<ul> <li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> <li>VARIABLES: <i>T</i>/K = 298</li> <li>EXPERIMENTAL VALUES:</li> </ul>	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3. PREPARED BY: A. L. Horvath
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_{1}/V_{2}$	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath $10^3 x_i$ $(compiler)$ $(compiler)$
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath $10^3 x_i$ $100 w_i$ (compiler)       (compiler)         2.01       1.32
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_{I}/V_{2}$ 25.0 0.90 AUXILIARY I	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath $\begin{pmatrix} 10^3 x_i & 100 w_i \\ (compiler) & (compiler) \\ 2.01 & 1.32 \end{pmatrix}$ NFORMATION
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE:	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath         (compiler)         2.01         1.32         NFORMATION         SOURCE AND PURITY OF MATERIALS:
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return-	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3. PREPARED BY: A. L. Horvath $\begin{pmatrix} 10^3 x, & 100 w, \\ (compiler) & (compiler) \\ 2.01 & 1.32 \end{pmatrix}$ NFORMATION SOURCE AND PURITY OF MATERIALS: (1) Commercial reagent, C. P. grade, used as received. (2) Distilled.
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return- ing the mixture to a constant temperature water bath. The difference between the total amount of trichloro- methane which was added and the amount remaining	Booth, H. S.; Everson, H. E.Ind. Eng. Chem. 1948, 40, 1491-3.PREPARED BY:A. L. Horvath $\begin{pmatrix} 10^3 x, \\ (compiler) \\ 2.01 \\ 1.32 \\ \end{bmatrix}$ NFORMATIONSOURCE AND PURITY OF MATERIALS:(1) Commercial reagent, C. P. grade, used as received.(2) Distilled.ESTIMATED EPROPS:
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return- ing the mixture to a constant temperature water bath. The difference between the total amount of trichloro- methane which was added and the amount of trichloro- methane dissolved in the known volume of water.	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath         10 <sup>3</sup> x, 100 w, (compiler)         2.01         1.32         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1)         Commercial reagent, C. P. grade, used as received.         (2)         Distilled.         ESTIMATED ERRORS:         Solubility:
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return- ing the mixture to a constant temperature water bath. The difference between the total amount of trichloro- methane which was added and the amount remaining in excess was taken as the amount of trichloro- methane dissolved in the known volume of water. The determination of the excess amount of tri- chloromethane added is described by Hanslick (ref. 1)	Booth, H. S.; Everson, H. E.Ind. Eng. Chem. 1948, 40, 1491-3.PREPARED BY: A. L. Horvath $10^3 x$ , (compiler) 2.01 $2.01$ $1.32$ NFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Commercial reagent, C. P. grade, used as received. (2) Distilled.ESTIMATED ERRORS: Solubility: Temperature: $\pm 1 \ K \ (compiler).$
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_1/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return- ing the mixture to a constant temperature water bath. The difference between the total amount of trichloro- methane which was added and the amount remaining in excess was taken as the amount of trichloro- methane dissolved in the lemown volume of water. The determination of the excess amount of tri- chloromethane added is described by Hanslick (ref. 1).	Booth, H. S.; Everson, H. E.Ind. Eng. Chem. 1948, 40, 1491-3.PREPARED BY: A. L. Horvath $10^3 x$ , (compiler) 2.01 $2.01$ $1.32$ NFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Commercial reagent, C. P. grade, used as received. (2) Distilled.ESTIMATED ERRORS: Solubility: Temperature: $\pm 1$ K (compiler).REFERENCES:
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $V_t/V_2$ 25.0 0.90 AUXILIARY I METHOD/APPARATUS/PROCEDURE: Equilibrium was ensured through repeated shaking and centrifuging of the trichloromethane and water mixture in a stoppered Goetz tube and return- ing the mixture to a constant temperature water bath. The difference between the total amount of trichloro- methane which was added and the amount of trichloro- methane dissolved in the known volume of water. The determination of the excess amount of tri- chloromethane added is described by Hanslick (ref. 1).	Booth, H. S.; Everson, H. E.         Ind. Eng. Chem. 1948, 40, 1491-3.         PREPARED BY:         A. L. Horvath         (compiler)         2.01         1.32         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Commercial reagent, C. P. grade, used as received.         (2) Distilled.         ESTIMATED ERRORS:         Solubility:       Not specified.         Temperature:       ± 1 K (compiler).         REFERENCES:         (1) Hanslick, R. S. Ph. D. Thesis, Columbia

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Donahue, D. J.; Bartell, F. E.		
(2) Water: H.O: [7732-18-5]	J. Phys. Chem. <u>1952</u> , 56, 480-4.		
VARIABLES:	PREPARED BY:		
T/K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C 10 <sup>3</sup> $x_1$ 10 (cor	$\frac{10}{10} \frac{w_1}{x_2}$ $\frac{100}{100} \frac{w_2}{w_2}$ (compiler)		
25 1.2 0	.790 5.0 7.58 × 10 <sup>-2</sup>		
AUXILIARY	INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A mixture of trichloromethane and water was placed in a glass stoppered flask and was shaken intermittently for at last three days in a water bath held at constant temperature. The organic phase was	<ol> <li>Source not given. Reagent grade, purified by fractional distillation.</li> <li>Purified by distillation.</li> </ol>		
analyzed for water content by the Karl Fischer titra- tion method, whereas the aqueous phase was analyzed	ESTIMATED ERRORS:		
interferometrically.	Solubility: Not specified. Temperature: $\pm 0.1$ K.		
COMPONENTS	ORIGINAL MEASUREMENTS		
(1) Trichloromethane (chloroform);	Kudryavtseva, G. I.; Krutikova, A. D.		
CHCl <sub>3</sub> ; [67-66-3]	J. Appl. Chem. USSR. <u>1953</u> , 26, 1129-33.		
(2) Water; H <sub>2</sub> O; [7732-18-5]			
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	Z. Maczynska		
EXPERIMENTAL VALUES:			
$t/^{\circ}$ C 100 $w_1$ 10 <sup>3</sup> $x_1$ 100 $w_2$ 10 <sup>2</sup> $x_2$ (compiler) (compiler)			
20 0.84 1	28 0.99 6.21		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The mutual solubility between trichloro- methane and water was determined by titration from a microburet until turbidity appeared. The end point of the titration, when turbidity first appeared, was	<ul> <li>(1) Source and purity not given.</li> <li>(2) Distilled (compiler).</li> </ul>		
determined with an accuracy of 0.3 - 0.5 %. The ti- tration of the solution took place in a thermostat bath	ESTIMATED ERRORS:		
at 20°C. The caprolactam-water-trichloromethane ternary system was also studied.	Solubility: $0.3 - 0.5$ . Temperature: $\pm 0.5$ K (compiler).		

108			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Eberius, E.		
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Wasserbestimmung mit Karl-Fischer-Lösung, Verlag Chemie, GMBH, Weinheim, <u>1954</u> , p. 67.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^{3} x_{i} & 100 \ w_{i}M_{i}^{-1}/\text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$		
20 0.084 ± 0.003	5.54 $4.66 \times 10^{-3}$		
AUXILIARY II	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A 50 g trichloromethane sample was shaken with pure water in a 100 cm <sup>3</sup> Erlenmeyer flask. When equilibrium had been attained and the two	<ol> <li>Distilled.</li> <li>Source and purity not given.</li> </ol>		
phases separated, the organic phase was analyzed for water using the Karl-Fischer titration method. Five	ESTIMATED ERRORS:		
determinations were performed.	Solubility: See above.		
	Temperature: $\pm$ 0.5 K (compiler).		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Water; H <sub>2</sub> O; [7732-18-5]	Zielinski, A. Z.		
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Chem. Stosowana <u>1959</u> , 3, 377-84.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
<i>t/</i> °C ρ <sub>1</sub> /kg m <sup>-3</sup>	10 <sup>3</sup> x, 100 w, (compiler) (compiler)		
25 1.44	6.41 9.73 × 10 <sup>-2</sup>		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The concentration of water in trichloromethane was determined by the Karl Fischer titration method (ref. 1).	<ol> <li>Distilled (compiler).</li> <li>Gliwice Chemicals, pure grade reagent, used as received.</li> </ol>		
	ESTIMATED ERRORS:		
	Solubility: Not specified. Temperature: $\pm 1$ K (compiler).		
	REFERENCES:		
	(1) Mitchell, J.; Smith, D. M. Aquametry: A Treatise on Methods for the Determination of Water, Wiley, New York, <u>1952</u> , p. 73 and 260.		

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Desnoyer, M.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Dosage de L'Eau dand les Solvants Organiques par Absorption Infra-Rouge et Mesure des Constantes Dielectriques, Centre D'Etudes Nucleaires de Saclay, Rept. No. 1254, Saclay, June 23, 1959.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 299	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}\mathbf{C}$ 100 $w_{t}$	$\begin{array}{ccc} 10^{3} x_{i} & 100 \ w_{i} M_{i}^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
26 8.6×10 <sup>-2</sup>	5.67 $4.77 \times 10^{-3}$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The saturation of trichloromethane with water took place at room temperature under vigorous stir- ring. The concentration of water in the organic rich phase was determined by infrared spectrometry.	<ol> <li>Distilled.</li> <li>Source not given. Commercial reagent, washed and dried before use.</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified.	
	Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Tettamanti, K.; Nogradi, M.; Sawinsky, J.	
(2) Water: H O: $(7732, 18, 5)$	Periodica Polytech. <u>1960</u> , 4, 201-18.	
(2) water, n <sub>2</sub> 0, [//32-16-5]		
VARIABLES:	PREPARED BY:	
T/K = 293	Z. Maczynska	
EXPERIMENTAL VALUES:		
t/°C 100 $w_1$ 10 <sup>3</sup> $x_1$ 100 $w_2$ 10 <sup>3</sup> $x_2$ (compiler) (compiler)		
20 0.82 1.	25 $7.0 \times 10^{-2}$ 4.62	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Turbidimetric titrations were used for both the	(1) Source and purity not given.	
aqueous and the organic phases. First, trichloro- methane was titrated with water from a microburet	(2) Distilled (compiler).	
titrated with trichloromethane. The amount of tri-		
an accuracy of 0.01 g. The temperature was main-	ESTIMATED ERRORS:	
trichloromethane-water-caprolactam ternary system	Solubility: $\pm 1.0$ %. Temperature: $\pm 0.1$ K.	

110		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Conti, J. J.; Othmer, D. F.; Gilmont, R.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chem. Eng. Data <u>1960</u> , 5, 301-7.	
VARIABLES:	PREPARED BY:	
T/K = 298 - 329	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
<i>t/</i> °C 100 w <sub>1</sub> 10 (con	$y_1^{\lambda} x_1 = 100 w_2 = 10^3 x_2$ npiler) (compiler)	
25 0.75 1. 56.1 0.80 1.3	139 $8.0 \times 10^{-2}$ $5.278$ 216 $1.7 \times 10^{-1}$ $11.16$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A flask was charged with a trichloromethane and water mixture of approximately the desired com- position. The system was allowed to equilibrate in a water bath at the desired temperature. Samples were withdrawn from the two layers and concentra-tions	<ol> <li>U. S. P. grade. Washed with H<sub>2</sub>SO<sub>4</sub> and K<sub>2</sub>CO<sub>3</sub> solutions and then distilled. Only the center cut was used.</li> <li>Deionized, distilled twice before use.</li> </ol>	
were determined. The analytical techniques are de- scribed elsewhere (ref. 1).	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.1$ K.	
	REFERENCES:	
	(1) Conti, J. J.; Othmer, D. F. Unpublished data.	
COMPONENTS: ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Ababi, V.; Popa, A.; Mihaila, Gh.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Analele Stiint. Univ. Al. I. Cuza Iasi. Sect. IC. Chem. <u>1964</u> , 10, 71-84.	
VARIABLES:	PREPARED BY:	
T/K = 294	Z. Maczynska	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-i} / \text{mol } g^{\cdot 1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
20.5 0.7	1.4 $5.86 \times 10^{-3}$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The isothermic titration method described by Ababi and Popa (refs. 1, 2) was used. A trichloro- methane-water-allyl alcohol ternary system was also	<ol> <li>Source and purity not given, n<sub>n</sub> = 1.4454 at 20°.C.</li> <li>Distilled (compiler).</li> </ol>	
article.	ESTIMATED ERRORS:	
	Solubility: Not specified	
	Temperature: $\pm 0.1$ K.	
	Temperature: ± 0.1 K. REFERENCES:	

COMPONENTS:		ORIC	GINAL MEAS	SUREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]		Chis	tyakov, V. M	.; Shapurova, V. V.
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]		Izv. <u>19</u> 0	Vyssh. Ucheb. 54. 7, 349-50.	n. Zaved. Khim. Khim. Tekhnol.
VARIABLES:		PRE	PARED BY:	
<i>T</i> /K = 293		A. L	. Horvath	
EXPERIMENTAL VALUES:		•		
<i>t/</i> °C 100	w <sub>i</sub>	(co	10 <sup>3</sup> x, ompiler)	100 $w_i M_i^{-1}$ /mol g <sup>-1</sup> (compiler)
20 6.3	× 10 <sup>-2</sup>		4.16	$3.5 \times 10^{-3}$
At	JXILIARY II	NFORM	MATION	
METHOD/APPARATUS/PROCEDURE		SOUI	RCE AND PI	IRITY OF MATERIALS
Details are not available.		(1) (2)	Distilled (co Source and p	mpiler). purity not given.
		ESTI	MATED ERR	ORS:
		Solu Tem	bility: perature:	Not specified. $\pm 1 \text{ K}$ (compiler).
COMPONENTS:		ORIC	SINAL MEAS	SUREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]		Mast	erton, W. L.	; Gendrano, M. C.
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]		J. Pl	hys. Chem. <u>19</u>	9 <u>66</u> , <i>70</i> , 2895-8.
VARIABLES:		PREF	ARED BY:	
<i>T/</i> K = 298		A. L	Horvath	
EXPERIMENTAL VALUES:		1		
<i>t/°</i> C <i>c<sub>1</sub>/mc</i>	ol m <sup>-3</sup>	(0	100 w <sub>i</sub> compiler)	$10^3 x$ , (compiler)
25 73	3.8	8.	.98 × 10 <sup>-2</sup>	5.92
AUXILIARY INFORMATION				
METHOD/APPARATUS/PROCEDURE:		SOU	RCE AND PL	IRITY OF MATERIALS:
Trichloromethane was equilibrated with water in an apparatus described elsewhere (ref. 1). The equilibration was allowed to take place for 2 days with the entire apparatus immersed in a water bath. The organic phase was analyzed for water by the Karl Fischer method (dead-stop end point). The microburet used for the titration was read to $\pm$ 0.01 cm <sup>3</sup> . All		(1) (2)	Distilled. Source not g distilled wate with GC.	iven. Reagent grade, washed with er and the purity was checked
		ESTI	MATED ERR	ORS:
nitrogen.	e or ary	Solu Tem	bility: perature:	$\pm$ 0.2 × 10 <sup>-3</sup> avg. dev. $\pm$ 0.05 K.
		REFE	ERENCES:	
		(1)	Christian, S. Chem. Soc.	; Affsprung, H.; Johanson, J. <i>J</i> . <u>1963</u> , 1896.

112				
COMPONENTS:		ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);		Lowe, H. J.		
(2) Water; $H_2O$ ; [7732-18-5]		in <i>Theory and Application of Gas Chromatography</i> , H. S. Kroman and S. R. Bender, Eds., Grune and Stratton, New York, <u>1968</u> , p. 194-209.		
VARIABLES:		PREPARED BY:		
T/K = 310		A. L. Horvath		
EXPERIMENTAL VALUES:				
t/°C Ostwald L/dim	1 coefficient, ensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$		
37 3.8	0	0.753 1.14		
	AUXILIARY IN	NFORMATION		
METHOD/APPARATUS/PROCEDUR	RE:	SOURCE AND PURITY OF MATERIALS:		
The direct injection technique w investigation of the solubility of trichle water. Ostwald coefficients were deter equilibration of trichloromethane with	vas used for the promethane in mined by water in a	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>		
capped syringe. After thoroughly mixi was withdrawn and analyzed in a gas	ng, a sample chromatograph	ESTIMATED ERRORS:		
equipped with a hydrogen flame detector. The con- centration was proportional to the recorded peak height.		Solubility: Not specified. Temperature: ± 1 K (compiler).		
COMPONENTS:		ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);		Okuda, Y.		
CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5]		Arch. Jap. Chir. <u>1968</u> , 37, 700-16.		
VARIABLES:		PREPARED BY:		
T/K = 288 - 313		A. L. Horvath		
EXPERIMENTAL VALUES:				
t/°C Ostwald L/dim	l coefficient, ensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
15 7.15 20 5.95	± 2.03 + 1.35	0.5915 8.971 0.6092 9.241		
25 5.08 30 4.40	$ \frac{1}{\pm} $ 0.83 $ \pm$ 0.53	0.6378 9.677 0.6717 10.195		
35 3.85 37 3.70	± 0.40 ± 0.39	0.7091 10.766 0.7501 11.392		
40 3.40	± 0.34	0.7499 11.389		
AUXILIARY INFORMATION				
METHOD/APPARATUS/PROCEDUR	E:	SOURCE AND PURITY OF MATERIALS:		
A gas chromatographic analysis combined with a crushed ampoule technique was used. The solubility values were determined by equilibrating trichloro- methane with water in a sealed ampoule. Then, after the ampoules were broken, the samples were analyz- ed using a gas chromatograph equipped with a flame ionization detector.		<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
		ESTIMATED ERRORS:		
		Solubility: See above. Temperature: $\pm$ 0.5 K (compiler).		

-	113		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Lowe, H. J.; Hagler, K.		
(2) Water; $H_2O$ ; [7732-18-5]	in Gas Chromatography, Biology and Medicine, R. Poster, Ed., Churchill, London, <u>1969</u> , p. 86-112.		
VARIABLES:	PREPARED BY:		
T/K = 310	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Ostwald coefficient, L/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$		
37 3.91	0.788 1.197		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The direct sample injection method was used. The sample containing the equilibration mixture of trichloromethane and water was injected into a	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
chromosorb column equipped with an ionization de- tector. The response areas were compared with the	ESTIMATED ERRORS:		
gas standards analyzed under similar conditions.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).		
COMPONENTS	ORIGINAL MEASUREMENTS		
(1) Trichloromethane (chloroform):	Bachofen, H.; Farhi, L. E.		
CHCl <sub>3</sub> ; [67-66-3]	J. Appl. Physiol. <u>1971</u> , 30, 136-9.		
(2) Water; H <sub>2</sub> O; [7732-18-5]			
VARIABLES:	PREPARED BY:		
T/K = 310	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Ostwald coefficient, L/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$		
37 4.04	0.80 1.216		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The solubility was determined in a manometric apparatus consisting of a chamber and a transducer. The bottom plate of the chamber contained two wells; one was filled with mercury and the other was filled with introduced water and trickloromethane. The	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>		
mixing and equilibrium between trichloromethane and water took place in the chamber under vigorous stirring. With the pressures before and after the	ESTIMATED ERRORS:		
equilibrium measured, the Ostwald coefficient was calculated from the chamber volume and the injected liquid volume.	Temperature: $\pm 0.5$ K (compiler).		

114		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ul><li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li></ul>	Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. I.; Treger, Yu. A.; Flid, R. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Russ. J. Phys. Chem. <u>1971</u> , 45, 488-90.	
VARIABLES:	PREPARED BY:	
T/K = 288 - 333	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 1000 $g_{l}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$	
15         4.82           30         3.20           45         2.05           60         1.30	0.4807.270.3194.830.2053.100.1301.96	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The dynamic method was used for the deter- mination of the solubility. It consisted of the saturation of nitrogen with trichloromethane vapor	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
which was then bubbled through a thermostatically controlled flask filled with a definite volume of	ESTIMATED ERRORS:	
water. After 2 - 3 hours the solution became saturated and samples were taken periodically (ref. 1).	Solubility: $\pm 15\%$ .	
The concentration of trichloromethane in water was determined by GLC. On average, 3 - 5 measurements	Temperature: $\pm 0.05$ K.	
were taken at each temperature.	REFERENCES:	
	(1) Treger, Yu. A.; Flid, R. M.; Spektor, S. S. <i>Russ. J. Phys. Chem.</i> <u>1964</u> , 38, 253.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Karger, B. L; Chatterjee, A. K.; King, J. W.	
(2) Water; $H_2O$ ; [7732-18-5]	Techn. Rept. No. 3, Department of Chemistry, Northeastern University, Boston, Mass., <u>May 10,</u> <u>1971</u> .	
VARIABLES:	PREPARED BY:	
T/K = 286	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ Partition coefficient, <sup>1</sup> $K_L/dimensionless$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$	
12.5 13.8	1.02 1.55	
<sup>1</sup> Gas-liquid chromatographic parameter, from instrument calibration.		
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Gas-liquid chromatography was used for the determination of the partition of trichloromethane with thin layers of water coated on Porasil D. Helium	<ol> <li>J. T. Baker Chemical Co., reagent grade.</li> <li>Distilled (compiler).</li> </ol>	
carrier gas was presaturated with water at the column temperature. The eluents were detected with an F &	ESTIMATED ERRORS:	
$(K_{L})$ was calculated from the slope of the straight line plot of $V_{N}/A_{L}$ versus $V_{L}/A_{L}$ .	Solubility: Not specified. Temperature: $\pm$ 0.05 K (compiler).	

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Antropov, L. 1.; Populyai, V. E.; Simonov, V. D.; Shamsutdinov, T. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Russ. J. Phys. Chem. <u>1972</u> , 46, 311-2 (VINITI No. 3739-71).	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 288 - 299	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/°</i> C 100 w <sub>1</sub> 10 (com	$x_1 = 100 w_2$ $10^3 x_2$ upiler) (compiler)	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The Alexejev's synthetic method of solubility determination was used (ref. 1). A fixed weight of trichloromethane and water was sealed in a tube and	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
subjected to gradually increasing temperature with constant agitation. The appearance of opalescence or	ESTIMATED ERRORS:	
clouding was the indication of the saturation temper- ature. The temperature was then allowed to fall and,	Solubility: Not specified.	
ature of the first appearance of opalescence was re- corded. The observation was repeated several times.	$\frac{1}{2} = \frac{1}{2} \frac{1}{1} $	
	REFERENCES:	
	(1) Alexejev, W. Ann. Phys. Chem. <u>1886</u> , 28, 305.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Ödberg, L.; Löfvenberg, A.; Högfeldt, E.; Fredlund, F.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	J. Inorg. Nucl. Chem. <u>1972</u> , 34, 2605-16.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C c₁/mol m⁻³	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 73.0	$8.87 \times 10^{-2}$ 5.849	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The equilibrium between water and trichloro- methane was reached after the samples were left at least 40 hours in an equilibration vessel. The Karl	<ol> <li>Distilled.</li> <li>Source not given. Purity is &gt; 99.9 %.</li> </ol>	
Fischer titration method was used for the water analysis once equilibrium had been established. At	ESTIMATED ERRORS:	
least two samples were studied.	Solubility: Not specified. Temperature: $\pm 0.3$ K.	

116				
COMPONENTS:		,	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);		;	McConnell, G.; Ferguson, D. M.; Pearson, C. I	ર.
$CHCl_{3}; [67-66-3]$			Endeavour <u>1975</u> , 34, 13-8.	
(2) water, H <sub>2</sub> O, [7752-16-5]			1	
VARIABLES:			PREPARED BY:	
T/K = 293			A. L. Horvath	
EXPERIMENTAL	VALUES:			
	t/°C	$10^6 g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
	20	8200	0.813 1.24	
		AUXILIARY II		
METHOD/APPARA	ATUS/PROCED	URE:	SOURCE AND PURITY OF MATERIALS:	
The solubility determined by using (GLC) equipped with	y of trichlorome g a gas-liquid ch th an electron c	thane in water was aromatograph apture detector.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
linked mass-spectro	meter (MS).	confirmed by a	ESTIMATED ERRORS:	
			Solubility:Not specified.Temperature: $\pm$ 0.1 K (compiler).	
COMPONENTS:			ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);		;	Pearson, C. R.; McConnell, G.	
(2) Water; $H_2O$ ; [7732-18-5]			Proc. Roy. Soc. B. <u>1975</u> , 189, 305-32.	
VARIABLES:			PREPARED BY:	
<i>T</i> /K = 293			A. L. Horvath	
EXPERIMENTAL	VALUES:			
	t/°C	$10^6 g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
	20	8200	0.813 1.24	
AUXILIARY INFORMATION				
METHOD/APPARATUS/PROCEDURE:		URE:	SOURCE AND PURITY OF MATERIALS:	
Saturated soluter temperature thermost ed with n-pentane a	utions were prep stat. Water sam nd an aliquot of	pared in a constant ples were extract- the extract taken	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
for GLC analysis. T with a <sup>63</sup> Ni electron	The gas chromat capture detector	ograph was fitted	ESTIMATED ERRORS:	
			Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
1				

	117	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ul><li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li></ul>	Pavlovskaya, E. M.; Charykov, A. K.; Tikhomirov, V. I.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Gen. Chem. USSR. <u>1977</u> , 47, 2230-4.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}\mathbb{C}$ $c_{1}/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
20 67.5	0.807 8.02	
AUXILIARY I	NFORMATION	
	SOUDCE AND DUDITY OF MATERIALS.	
Trichloromethane was added from a micro- buret to an aqueous solution of octanoic acid containing KCl under vigorous stirring. The pH of	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
the solution at first remained constant right up to the appearance of a second phase, i. e., up to the satura-		
tion of the aqueous phase with trichloromethane, after which it began to rise. The solubility was calculated	ESTIMATED ERRORS:	
from the volume of the aqueous phase and the volume of trichloromethane used for the saturation.	Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Sato, A.; Nakijima, T.	
$CHCl_{3}$ ; [67-66-3]	Arch. Envir. Health <u>1979</u> , 34, 69-75.	
(2) water, n <sub>2</sub> O, [7732-16-5]		
VARIABLES:	PREPARED BY:	
T/K = 310	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Ostwald coefficient, L/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$	
37 3.5	0.677 1.027	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Trichloromethane vapor was equilibrated in an airtight vial between water and the overlying air. When equilibrium was reached, a portion of the e-	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
quilibrated air in the vessel was withdrawn using an airtight syringe and was injected into a gas chro-	ESTIMATED ERRORS:	
matograph and analyzed. The height of the chromat- ogram peak was used to calculate the partition coef- ficient.	Solubility: $\pm$ 0.3 std. dev. Temperature: $\pm$ 0.5 K (compiler).	

,

-

118		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Freed, V. H.; Chiou, C. T.; Schmedding, D.; Kohnert, R.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Health Perspect <u>1979</u> , 30, 75-80.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 10 <sup>6</sup> w <sub>1</sub>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 7950	0.795 1.208	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A known volume of water was mixed with tri- chloromethane in an Erlenmeyer flask. The flask was immersed in a water bath and magnetically stirred. Samples were removed for analysis at regular inter- vals. All samples were analyzed using a gas-liquid	<ol> <li>Source not given. Analytical standard grade, greater than 95 % purity.</li> <li>Distilled and run through a resin column before use.</li> </ol>	
chromatograph equipped with an electron capture detector. The solubility given is the average of five	ESTIMATED ERRORS:	
consecutive samples with less than 5 % variation.	Solubility: 5 % variation of 5 samples. Temperature: $+ 0.5$ K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Aref'eva, R. P.; Korenman, I. M.; Gorokhov, A. A.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	USSR Patent 672 548, July 5, 1979, 3 pp. (CA. 91:113 256k).	
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 12.7 ± .2	1.274 1.944	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Equal quantities of diphenylthiocarbazone were added to aqueous solutions with different amounts of trichloromethane in several flasks. The contents of the flasks were shaken for 5 minutes and left to stand	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
for 1 hour. The mixtures were filtered through filter paper before their analysis by spectrophotometry. The	ESTIMATED ERRORS:	
turning point on the graph of optical density versus		

	119	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Coca, J.; Diaz, R. M.; Pozas, C.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Fluid Phase Equilibr. <u>1980</u> , 4, 125-36.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
t/°C 100 w, 10 (con	$\begin{array}{cccc} {}^{4}x_{1} & 100 \ w_{2} & 10^{4} \ x_{2} \\ \text{npiler} & & (\text{compiler}) \end{array}$	
25 1.19 1.	814 8.0 $\times$ 10 <sup>-2</sup> 5.278	
AUXILIARY I	NFORMATION	
	SOURCE AND BUDITY OF MATERIALS.	
METHOD/APPARATUS/PROCEDURE:	(1) Ehrle reagent Eurther purified by distillation	
The inductal solubility data were determined by the method as described by Othmer <i>et al.</i> (ref. 1). Ten cm <sup>3</sup> trichloromethane was added to water from a buret and agitated until the solution became turbid.	<ol> <li>(1) Fluka reagent. Further purfied by distillation in a heli-packing column.</li> <li>(2) Distilled.</li> </ol>	
The appearance of turbidity indicated the formation of a second phase. The solubility was calculated from	ESTIMATED ERRORS:	
known densities and volumes.	Solubility: Not specified.	
	Temperature: $\pm 0.5$ K (compiler).	
	REFERENCES:	
	(1) Othmer, D. F.; White, R. E.; Trueges, E. Ind. Eng. Chem. <u>1941</u> , 33, 1240.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Balls, P. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ph. D. Thesis, Univ. of East Anglia, Norwich, U. K., <u>July 1980</u> , 375 pp.	
VARIABLES:	PREPARED BY:	
T/K = 293 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$\begin{array}{cccc} 20 & 0.112 \pm 0.038 \\ 25 & 0.166 \pm 0.100 \end{array}$	0.847 1.393 0.699 1.149	
	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The method involved repeated equilibrations of nitrogen with a water sample containing trichloro- methane. The sample was shaken vigorously at the	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
required temperature in a constant temperature bath for 30 minutes. The equilibrated gas phase was then injected into a gas chamategraph. The experiment	ESTIMATED ERRORS:	
were performed at least three times at 20 and 25°C; each experiment involved about six equilibrations.	Solubility: See above. Temperature: $\pm 0.5$ K (compiler).	

,

120		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Hutchinson, T. C.; Hellebust, J. A.; Tam, D.; et al.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Hydrocarbons and Halogenated Hydrocarbons in the Aquatic Environment, Plenum Press, New York, <u>1980</u> , p. 577-86.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	I	
$t/^{\circ}C$ $\rho_1/kg m^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 7.95	0.7973 1.21	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess quantity of trichloromethane was added to water and stirred vigorously for 24 hours. After settling at 25°C for 48 hours, the solution was extracted with evaluation and the solution was	<ol> <li>Aldrich Chemicals, highest grade, used as received.</li> <li>Double distilled.</li> </ol>	
lyzed using an Aminco-Bowman spectrophotofluoro- meter. A detailed description of the method has been	ESTIMATED ERRORS:	
reported elsewhere (ref. 1).	Solubility: Not specified. Temperature: $\pm 0.2$ K (compiler).	
	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y. J. Chem. Eng. Data <u>1977</u> , 22, 399.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Veith, G. D.; Macek, K. J.; Petrucelli, S. R.; Carroll, J.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Proc. 3 <sup>rd</sup> Ann. Symp. on Aquatic Toxicology, ASTM Publ. 707, Philadelphia, <u>1980</u> , p. 116-29.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:	-	
$t/^{\circ}C$ $n_{1}V_{2}^{-1}/\text{mol m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 61.7	0.732 1.11	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of trichloromethane was added to 10 cm <sup>3</sup> distilled water in a 50 cm <sup>3</sup> flask. The mixture was magnetically stirred in a constant temperature	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
water bath. Samples taken from the solution were analyzed using radioactive techniques. The <sup>14</sup> C	ESTIMATED ERRORS:	
liquid scintillation spectrometer by recording the number of counts per minute.	Solubility: Not specified. Temperature: $\pm 1$ K.	

.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C.	
(2) Water; $H_2O$ ; [7732-18-5]	Environ. Sci. Techn. <u>1980</u> , 14, 1227-9.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $n_{I}V_{2}^{-1}/\text{mol m}^{-3}$	$100 w_i$ $10^3 x_i$	
25 60.5	0.719 1.09	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of trichloromethane was added to a stainless steel tube containing water and the tube was sealed. It was allowed to equilibrate during shaking	<ol> <li>New England Nuclear, used as received.</li> <li>Distilled.</li> </ol>	
for a week at 25°C. After centrifugation, the sample was analyzed for radioactivity using a liquid scintil-	ESTIMATED ERRORS:	
ried out at least twice for each sample and the analy- sis was done in duplicate.	Solubility: $\pm 4.3\%$ std. dev. Temperature: $\pm 0.3$ K.	
(1) Trichloromethane (chloroform):	ORIGINAL MEASUREMENTS: Symons I M · et al	
CHCl <sub>3</sub> ; [67-66-3]	Treatment Techniques for Controlling Trihalo-	
(2) Water; $H_2O$ ; [7732-18-5]	<i>methanes in Drinking Water</i> , U. S. EPA Report 600/2-81-156, Cincinnati, OH., <u>1981</u> .	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant H/dimensionless	$\begin{array}{ccc} 100 \ w, & 10^3 \ x, \\ (compiler) & (compiler) \end{array}$	
25 0.152	0.8205 1.256	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are given elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Cincinnati tap water.</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm$ 0.5 K.	
	REFERENCES:	
	(1) Warner, U. S. Environmental Protection Agency, Cincinnati, OH., Personal communication, <u>1980</u> .	

,

122			
COMPONENTS:		ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]		Wu, X.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]		Huaxue Shiji <u>1981</u> , 221-4.	
VARIABLES:		PREPARED BY:	
T/K = 278 - 328		A. L. Horvath	
EXPERIMENTAL VALUES:			
t/°C	10 <sup>6</sup> w <sub>1</sub>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
5 10 15 20 25 30 35 40 45 50 55	417 568 606 758 872 909 1061 1099 1251 1289 1402	$\begin{array}{ccccccc} 0.0417 & 2.76 \\ 0.0568 & 3.75 \\ 0.0606 & 4.00 \\ 0.0758 & 5.00 \\ 0.0872 & 5.75 \\ 0.0909 & 5.99 \\ 0.1061 & 6.99 \\ 0.1099 & 7.24 \\ 0.1251 & 8.23 \\ 0.1289 & 8.48 \\ 0.1402 & 9.22 \end{array}$	
AUXILIARY INFORMATION		NFORMATION	
METHOD/APPARATUS/PROCEDURE: A known volume of water was added to dried trichloromethane to prepare a series of standard samples for establishing calibration curves and obtaining the saturated solutions at each temperature. The determination of the water content of trichloro- methane was made with a RM-250 NMR spectromet- er (radiation frequency = 250 MHz). The water peak amplitudes were the mean of four readings taken at a particular temperature.		<ol> <li>SOURCE AND PURITY OF MATERIALS:</li> <li>(1) Distilled.</li> <li>(2) Source not given. Grade 2 purity was washed with KOH solution, H<sub>2</sub>SO<sub>4</sub>, and distilled water. Dried on molecular sieve before use.</li> </ol>	
		ESTIMATED ERRORS: Solubility: ± 72 std. dev. Temperature: 1 - 2 K.	

·

COMPONENTS:	OMPONENTS:		ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);		Leighton, D. T.; Ca	llo, J. M.	
(2) Water; $H_2O$ ; [7732-18-5]		J. Chem. Eng. Data <u>1981</u> , 26, 382-5.		
VARIABLES:		PREPARED BY:		
T/K = 275 - 298		A. L. Horvath		
EXPERIMENTAL VALUES		1		
t/°C	Distribution coefficient <sup>1</sup> , $D_L$ /dimensionless	, 100 w <sub>i</sub> (compiler)	$\begin{array}{c} 10^3 x_1 \\ \text{(compiler)} \end{array}$	
1.9 13.5 15.7 17.1 22.0 24.9	68.9 128.7 146.7 155.0 194.5 204.8	0.8251 0.7846 0.7565 0.7676 0.7669 0.8297	1.254 1.192 1.149 1.166 1.165 1.261	
<sup>1</sup> Gas-liquid sy	vstem analysis parameter,	from calibration meas	urements.	
	AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PRO	OCEDURE:	SOURCE AND PUR	ITY OF MATERIALS:	
A 5 $\mu$ L trichloromethan into a 2.3 liter equilibration c water. After the cell was shall	ne sample was injected ell containing distilled ten vigorously for about	(1) Source and put (2) Distilled.	rity not given.	
5 minutes, the homogeneity of the liquid sample was maintained with a magnetic stirrer. Compressed air was passed through the cell and the gas flow was measured with a soap film flowmeter. The trichloro- methane content was extracted and analyzed using a dual flame ionization detector gas chromatograph.		ESTIMATED ERRO	RS:	
		Solubility: Temperature:	± 10.0 %. ± 0.5 K.	
COMPONENTS:		ORIGINAL MEASU	REMENTS:	
(1) Trichloromethane (chloroform);		Mackay, D.; et al.		
CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5]		Volatilization of Org U. S. EPA Report Georgia, <u>1982</u> (PB	canic Pollutants from Water, 600/3-82-019, Athens, 82-230 939).	
VARIABLES:	VARIABLES:		PREPARED BY:	
<i>T</i> /K = 293		A. L. Horvath		
EXPERIMENTAL VALUES:		1		
t/°C	$\rho_l/\text{kg m}^{-3}$	100 w <sub>i</sub> (compiler)	10 <sup>3</sup> x, (compiler)	
25	7.95	0.7950	1.203	
METHOD/APPARATUS/PROCEDURE: An aqueous solution with an excess amount of trichloromethane was stirred for 1 day before being introduced into a 1 liter glass vessel. Nitrogen entered at the bottom of the vessel through a sintered glass disk. The exit gas flow rate was measured by a soap bubble flow meter. The concentration of tri- chloromethane in water was determined using gas chromatography. The GC was equipped with both dual flame ionization detector and an electron capture detector.		(1) Source and pur (2) Double distille	ITY OF MATERIALS: rity not given. d.	
		ESTIMATED ERRO Solubility: Temperature:	RS: Not specified. ± 1.0 K.	
		I emperature:	± 1.0 K.	

123

124		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Ohtsuka, K.; Kazama, K.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Sen'i Seihin Shohi Kagaku Kaishi <u>1982</u> , 22, 197-201.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$	
25 1.5	0.101 6.66	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Water was added gradually to 50 cm <sup>3</sup> tri- chloromethane in a flask which was then lowered into a thermostat bath. The flash was shaken vigorously until the first cloud (turbidity) appeared. The sample	<ol> <li>Distilled (compiler).</li> <li>Commercial JTS extra pure reagent. Further purified by conventional methods.</li> </ol>	
water content was determined using the Karl Fischer titration method.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform);	Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S.	
(2) Water; $H_2O$ ; [7732-18-5]	Tellus <u>1983</u> , 35B, 170-6.	
VARIABLES:	PREPARED BY:	
T/K = 293 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 0.1044 25 0.1686	0.9845 1.498 0.7456 1.132	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The multiple equilibration technique was used to measure the Henry's law constant as described elsewhere (ref. 1). A known volume of water in a surinear was shown with a law and the state of the state	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
chloromethane. After equilibration, the headspace was separated and analyzed for trichloromethane. A	ESTIMATED ERRORS:	
gas chromatograph fitted with an electron capture detector was used for the analysis.	Solubility: $\pm$ 5.5 % std. dev. Temperature: $\pm$ 0.2 K.	
	REFERENCES:	
	(1) McAuliffe, C. D. Chem. Techn. <u>1971</u> , 1, 46.	

COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform); CHCl <sub>1</sub> : [67-66-3]	Orlandini, M.; Fermeglia, M.; Kikic, I.; Alessi, P.			
(2) Water; H <sub>2</sub> O; [7732-18-5]	Chem. Eng. J. <u>1983</u> , 16, 245-50.			
VARIABLES:	PREPARED BY:			
T/K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
$t/^{\circ}C$ 10 <sup>3</sup> $x_1$ 10 <sup>3</sup> $x_2$	$100 w_1$ $100 w_2$			
20 1 22 5 17	(compiler) (compiler)			
20 1.23 5.17	0.809 0.0784			
AUXILIARY I	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A mixture of water and trichloromethane was	(1) Fluka reagent, further purified by distillation			
agitated overnight and left at equilibrium for 24 hours. Three samples were withdrawn from each phase using a syringe and analyzed using cas	(2) Distilled (compiler).			
chromatography. The gas chromatograph (C. Erba ATc/t) was equipped with a Shimadzu Chromatopac	ESTIMATED ERRORS:			
E-1A integrator. The stainless column was packed with 100 - 200 mesh Chromosorb W DMCD coated	Solubility: Not specified.			
with heptadecane nitrile.	Temperature: $\pm 0.1$ K.			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform); CHCL: [67-66-3]	Lincoff, A. H.; Gossett, J. M.			
(2) Water; $H_2O$ ; [7732-18-5]	in Gas Transfer at Water Surfaces, W. Brutsaert and G. H. Jirka, Eds., D. Reidel Publ. Co., Dordrecht, <u>1984</u> , p. 11-25.			
VARIABLES:	PREPARED BY:			
T/K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
EPICS Method				
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$			
20 $3.33 \times 10^{-3}$	0.7442 1.130			
$\log_e H = -\frac{4180}{T(K)} + 8.$	553			
	(continued)			
AUXILIARY INFORMATION				
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
Henry's law constants were measured by	(1) Source and purity not given.			
EPICS and batch air stripping methods. The Equili- brium Partitioning in the Closed Systems technique	(2) Distilled.			
of headspace samples. In the Batch Air Stripping	ESTIMATED ERRORS:			
bottles and the equilibrated headspaces contained	Solubility: + 5 - 10 %.			
concentrations proportional to original concentrations in the aqueous samples. Full equilibrium was achieved in a few hours.	Temperature: $\pm 0.1$ K.			
126				
--	--	--	--	--
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Lincoff, A. H.; Gossett, J. M.			
(2) Water; H <sub>2</sub> O; [7732-18-5]	in Gas Transfer at Water Surfaces, W. Brutsaert and G. H. Jirka, Eds., D. Reidel Publ. Co., Dordrecht, <u>1984</u> , p. 11-25.			
VARIABLES:	PREPARED BY:			
<i>T</i> /K = 293	A. L. Horvath			
EXPERIMENTAL VALUES: (continued)				
Batch Air Str	ipping Method			
t/°C Henry's law constant, H/m' atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$			
20 $3.04 \times 10^{-3}$	0.8152 1.239			
$\log_e H = -\frac{4322}{T(K)} + 8$	. 956			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	McNally, M. E.; Grob, R. L.			
(2) Water; $H_2O$ ; [7732-18-5]	J. Chromatogr. <u>1984</u> , 284, 105-16.			
VARIABLES:	PREPARED BY:			
<i>T/</i> K = 303	A. L. Horvath			
EXPERIMENTAL VALUES:				
$t/^{\circ}$ C $g_{1}V_{2}^{-1}/\text{kg m}^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_j \\ (compiler) & (compiler) \end{array}$			
30 2.524	0.2529 3.825			
AUXILIARY I	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
Standard increasing concentration solutions of trichloromethane were prepared in volumetric flasks as described elsewhere (ref. 1). The aqueous-organic phase was 25 cm <sup>3</sup> and the vapor phase above was 35 cm <sup>3</sup> . The system was allowed to reach equilibrium in	<ol> <li>Chemical Service, West Chester, PA., U. S. A., purest grade available.</li> <li>Distilled and run through two Barnstead purification cartridges.</li> </ol>			
a constant temperature bath and then a sample was injected into the gas chromatograph. Values reported are an average of at least five measurements.	ESTIMATED ERRORS: Solubility: $\pm$ 1.85 % std. dev. Temperature: $\pm$ 0.5 K (compiler).			
	REFERENCES: (1) McNally, M. E.; Grob, R. L. J. Chromatogr. <u>1983</u> , 260, 23.			

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Techn. <u>1984</u> , 18, 518-21.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 293	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$		
20 $3.0 \times 10^{-3}$	0.8257 1.255		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the	<ol> <li>Source and purity not given.</li> <li>Distilled and deionized.</li> </ol>		
added to the stripping vessel and the desired flow was	ESTIMATED ERRORS:		
were taken for analysis. After extraction of the samples, the extracts were analyzed by gas chromat- ograph equipped with a capillary column.	Solubility: $\pm 0.1 \times 10^{-3}$ std. dev. Temperature: $\pm 0.1$ K.		
	REFERENCES:		
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Techn. <u>1979</u> , 13, 333.		
	ODICINAL MEASUDEMENTS.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
CHCl <sub>3</sub> ; [67-66-3]	Ph D Thesis Stamford University Stamford		
(2) Water; H <sub>2</sub> O; [7732-18-5]	CA., <u>1885</u> , 306 pp.		
VARIABLES:	PREPARED BY:		
T/K = 283 - 303	A. L. Horvath		
EXPERIMENTAL VALUES:	1		
t/°C Henry's Law Constant, ρ <sub>1</sub> /kg H/dimensionless	$m^{-3}$ 100 $w_i$ 10 <sup>3</sup> $x_i$ (compiler) (compiler)		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0 0.8750 1.330 0 0.8320 1.264 0 0.7880 1.197		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The multiple equilibrium technique with direct aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law constants.	<ol> <li>J. T. Baker Chemicals Co., N.J. &gt; 99.9 % pure.</li> <li>Distilled and deionized.</li> </ol>		
A known volume of liquid and gas was introduced in- to a syringe and allowed to equilibrate. The ratio of	ESTIMATED ERRORS:		
gas to inquid volume was maintained constant. While vapor samples were injected directly into a gas chro- matograph, the liquid samples were preconcentrated by liquid-liquid extraction before analysis. The ex- periments were carried out in duplicates.	Solubility: $\pm 2.22 \times 10^{-3}$ std. dev. Temperature: $\pm 0.5$ K.		

128				
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Takano, J.; Ishihara, Y.; Yasuoka, T.; Mitsuzawa, S.			
(2) Water; H <sub>2</sub> O; [7732-18-5]	Nippon Kagaku Kaishi, <u>1985</u> , 2116-9.			
VARIABLES:	PREPARED BY:			
T/K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
$t/^{\circ}C$ $\rho_{I}/kg m^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_j \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
20 7.77 9.33 6.33 8.55	0.77701.1800.93301.4190.63300.96040.85501.300			
AUXILIARY I	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A mixture of 1 cm <sup>3</sup> trichloromethane and 30 cm <sup>3</sup> water was introduced into a centrifuge tube which was then shaken for 60 minutes in a shaker at room temperature. After an additional shaking of 48 hours each use of literad through four turns of	<ol> <li>Tokyo Chemical Co., analytical grade reagent, redistilled before use.</li> <li>Double distilled.</li> </ol>			
filter paper in order to free the solution from emulsions. The concentrations of the aqueous solu- tions were determined by the TOC (Total Organic Carbon) method. Nine measurements were perform- ed for each solution.	ESTIMATED ERRORS: Solubility: 0 - 2.5 % coeff. of var. Temperature: ± 1.0 K.			
(1) Trichloromethane (chloroform):	ORIGINAL MEASUREMENTS:			
CHCl <sub>3</sub> ; [67-66-3]	Anal. Chem. <u>1986</u> , 58, 1596-7.			
(2) Water; $H_2O$ ; [7732-18-5]				
VARIABLES:	PREPARED BY:			
T/K = 298	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C 100 w <sub>1</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$			
25 0.82	1.25 $6.87 \times 10^{-3}$			
AUXILIARY I	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A 5 cm <sup>3</sup> trichloromethane sample containing labeled complex cobalt(III) pyrrolidinecarbodithioate was placed together with 1000 cm <sup>3</sup> of water into a separation funnel. After vigorous shaking for 5	<ol> <li>E. Merck reagent, used as received.</li> <li>Demineralized and subboiled.</li> </ol>			
minutes, the mixture was left for 30 minutes before the separation of the two phases. Samples were with- drawn for activity measurement using a NaI (T1) scintillation detector. Several factors were adjusted to obtain accurate measurements.	ESTIMATED ERRORS: Solubility: ± 0.02 std. dev. Temperature: ± 1.0 K (compiler).			

COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Barr, R. S.; Newsham, D. M. T.		
(2) Water: $H_2O$ : [7732-18-5]	Fluid Phase Equilibr. <u>1987</u> , 35, 189-205.		
VARIABLES:	PREPARED BY:		
T/K = 293 - 323	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}C$ 10 <sup>3</sup> $x_{i}$ (co	$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
20 1.218 35 1.121	.8016 5.06 0.0767 .7382 8.29 0.126		
50 1.099	.7238 11.6 0.177		
Mole fraction of trichloromethane $(x_i)$ dilution $(\gamma^{\infty})$ using the relation $x_i = 1/\gamma^{\infty}$ .	was calculated from the activity coefficient at infinite		
AUXILIARY	INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The water-rich mixture was studied using gas- liquid chromatography in which a mixture of methan	(1) Koch Light Laboratories Ltd., used as received.		
and trichloromethane vapor passed through the samp ing loop. The eluted trichloromethane was determine	(2) Double distilled tap water.		
with a flame ionization detector. The Organic-rich mixture was studied using ar	ESTIMATED ERRORS:		
isopiestic method in which a CaCl <sub>2</sub> solution control- led the partial pressure of water. The water concen-	Solubility: Not specified.		
Fischer titration apparatus.	Temperature: $\pm 0.05$ K.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	ORIGINAL MEASUREMENTS: Gossett, J. M.		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5]	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. <u>1987</u> , 21, 202-8.		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES:	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. <u>1987</u> , 21, 202-8. PREPARED BY:		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 283 - 308	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. 1987, 21, 202-8. PREPARED BY: A. L. Horvath		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 283 - 308 EXPERIMENTAL VALUES:	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. <u>1987</u> , 21, 202-8. PREPARED BY: A. L. Horvath		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: t/°C Henry's Law Constant, CV	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. 1987, 21, 202-8. PREPARED BY: A. L. Horvath $100 w_l$ $10^3 x_l$		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CW$ $H/m^3$ atm/mol <sup>-1</sup> %	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. <u>1987</u> , 21, 202-8. PREPARED BY: A. L. Horvath 100 w, 10 <sup>3</sup> x, (compiler) (compiler)		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> % 9.6 1.50 × 10 <sup>-3</sup> 6.1 17.5 2.46 × 10 <sup>-3</sup> 1.5 24.8 2.66 × 10 <sup>-3</sup> 1.5	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         6       1.0320         1.571         0.8980       1.365         0.8424       1.365		
COMPONENTS:         (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 283 - 308$ EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> %         9.6       1.50 × 10 <sup>-3</sup> 6.1         17.5       2.46 × 10 <sup>-3</sup> 1.5         24.8       3.67 × 10 <sup>-3</sup> 3.7         34.6       5.63 × 10 <sup>-3</sup> 3.7	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         (compiler)         (compiler)         0.8980         1.365         0.8424         0.8332		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3 atm/mol^{-1}$ % 9.6 1.50 × 10 <sup>-3</sup> 6.1 17.5 2.46 × 10 <sup>-3</sup> 1.5 24.8 3.67 × 10 <sup>-3</sup> 3.7 34.6 5.63 × 10 <sup>-3</sup> 3.7 1 CV = coefficient of variation (=	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath $100 w_i$ $10^3 x_i$ (compiler)       (compiler) $6$ $1.0320$ $1.571$ $6$ $0.8980$ $1.365$ $0.8332$ $1.2662$ $100$ S.D./mean).		
COMPONENTS:         (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 283 - 308$ EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> $9.6$ $1.50 \times 10^{-3}$ $17.5$ $2.46 \times 10^{-3}$ $24.8$ $3.67 \times 10^{-3}$ $34.6$ $5.63 \times 10^{-3}$ $1$ CV = coefficient of variation (=	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         (compiler)         (compiler)         0.8980         1.365         0.8424         1.280         6         0.8332         1.2662         100 S.D./mean).		
COMPONENTS:         (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 283 - 308$ EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> %         9.6       1.50 × 10 <sup>-3</sup> 6.1         17.5       2.46 × 10 <sup>-3</sup> 1.5         24.8       3.67 × 10 <sup>-3</sup> 3.7         34.6       5.63 × 10 <sup>-3</sup> 3.7         I CV = coefficient of variation (=         AUXILIARY         METHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath         100 w, 10 <sup>3</sup> x, (compiler)         6       1.0320         1.0320       1.571         6       0.8980         1.365         0.8424       1.280         100 S.D./mean).         INFORMATION         SOURCE AND PURITY OF MATERIALS:		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> % 9.6 1.50 × 10 <sup>-3</sup> 6.1 17.5 2.46 × 10 <sup>-3</sup> 1.5 24.8 3.67 × 10 <sup>-3</sup> 3.7 34.6 5.63 × 10 <sup>-3</sup> 3.7 <sup>1</sup> CV = coefficient of variation (= AUXILIARY METHOD/APPARATUS/PROCEDURE: A modification of the EPICS method was used to measure Henry's law constants. Precise quantities of trichloromethane were injected into serum bottles	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath         100 w, 10 <sup>3</sup> x, (compiler)         6         1.0320         1.571         0.8980         1.365         0.8424         1.280         0.8332         1.2662         100 S.D./mean).         INFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Mullinckrodt reagent, ACS, preserved with 0.75 % ethanol.         (2) Distilled.		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3 atm/mol^{-1}$ % 9.6 1.50 × 10 <sup>-3</sup> 6.1 17.5 2.46 × 10 <sup>-3</sup> 1.5 24.8 3.67 × 10 <sup>-3</sup> 3.7 34.6 5.63 × 10 <sup>-3</sup> 3.7 1 CV = coefficient of variation (= AUXILIARY METHOD/APPARATUS/PROCEDURE: A modification of the EPICS method was used to measure Henry's law constants. Precise quantities of trichloromethane were injected into serum bottles which contained distilled water. The bottles were incubated for 18 - 24 hours at four desired tempera- tures in a conteneer in bottle bath. The bottles were	ORIGINAL MEASUREMENTS:         Gossett, J. M.         Environ. Sci. Technol. 1987, 21, 202-8.         PREPARED BY:         A. L. Horvath $100 w_i$ (compiler)         6       1.0320         1.571         6       0.8980         1.365         0.8332       1.2662         100 S.D./mean).         INFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Mullinckrodt reagent, ACS, preserved with 0.75 % ethanol.         (2) Distilled.		
COMPONENTS: (1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 283 - 308 EXPERIMENTAL VALUES: $t/^{\circ}C$ Henry's Law Constant, $CV$ $H/m^3$ atm/mol <sup>-1</sup> % 9.6 1.50 × 10 <sup>-3</sup> 6.1 17.5 2.46 × 10 <sup>-3</sup> 1.5 24.8 3.67 × 10 <sup>-3</sup> 3.7 34.6 5.63 × 10 <sup>-3</sup> 3.7 <sup>1</sup> CV = coefficient of variation (= AUXILIARY METHOD/APPARATUS/PROCEDURE: A modification of the EPICS method was used to measure Henry's law constants. Precise quantities of trichloromethane were injected into serum bottles which contained distilled water. The bottles were incubated for 18 - 24 hours at four desired tempera- tures in a reciprocating shaker bath. The headspace concentrations of the EPICS bottles were measured by a gas chromatograph which was exercibed was used	ORIGINAL MEASUREMENTS: Gossett, J. M. Environ. Sci. Technol. 1987, 21, 202-8.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) (compiler) $100 w_i$ (compiler) $100 s.D./mean$ ).INFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Mullinckrodt reagent, ACS, preserved with 0.75 % ethanol. (2) Distilled.ESTIMATED ERRORS: Solubility:Solubility:See above		

130				
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Warner, H. P.; Cohen, J. M.; Ireland, J. C.			
(2) Water; $H_2O$ ; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants U. S. EPA Techn. Rept. PB87-212684, Cincinnati, OH., July 1987.			
VARIABLES:	PREPARED BY:			
<i>T</i> /K = 298	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
24.85 3.39×10 <sup>-3</sup>	0.9169 1.397			
AUXILIARY II	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
The original method and apparatus for the determination of Henry's law constants have been described elsewhere (ref. 1). The general procedure was to add an excess of trichloromethane to distilled,	<ol> <li>Purest quality available and was used without further purification, that is &gt; 99 %.</li> <li>Distilled and deionized.</li> </ol>			
overnight. A portion of the solution was returned to the stripping vessel. Trichloromethane was stripped isothermally from the solution at a known gas flow rate. The Henry's law constant was calculated from the log of the concentration versus time plot. The	ESTIMATED ERRORS: Solubility: $\pm 6 \%$ std. dev. Temperature: $\pm 0.05$ K.			
experimental data values are averages of two or more replicates.	REFERENCES:			
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Technol. <u>1979</u> , 13, 333.			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Howe, G. B.; Mullins, M. E.; Rogers, T. N.			
CHCl <sub>3</sub> ; [67-66-3] (2) Water; H <sub>2</sub> O; [7732-18-5]	AFESC Tyndall Air Force Base, Report ESL-TR- 86-66, Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp. (AD-A188 571).			
VARIABLES:	PREPARED BY:			
T/K = 283 - 303	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C 10 <sup>6</sup> w <sub>1</sub>	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
10 3471 20 2963 30 3596	0.3471 5.2535 0.2963 4.4827 0.3596 5.4432			
AUXILIARY IN	IFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
250 cm <sup>3</sup> bottles were filled with distilled deionized water and sealed. A measured volume of trichloromethane was injected into the bottles through each bottle septum using a microliter syringe. Tri- chloromethane was in excess of the anticipated solu-	<ol> <li>Probable a commercial reagent at least 99 % purity. Used as received.</li> <li>Distilled and deionized.</li> </ol>			
bility limit. The bottles were shaken for 1 hour with a wrist-action shaker and allowed to equilibrate for about 3 weeks. Samples were then injected into a gas chromatograph equipped with a Carbopack column and a FID detector. The GC response was compared with the calibration plot.	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiled).			

COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform); CHCI: [67-66-3]	Vogel, A. I., rev. by Furniss, B. S.; et al.			
(2) Water; $H_2O$ ; [7732-18-5]	Vogel's Textbook of Practical Organic Chemistry, 5 <sup>th</sup> ed., Longman, London, <u>1989</u> , p. 1442.			
VARIABLES:	PREPARED BY:			
<i>T</i> /K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C 100 w <sub>j</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-i} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$			
20.0 0.82	1.25 $6.87 \times 10^{-3}$			
	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
The solubility determination was done in small test tubes (e. g., $100 \times 12$ mm) to permit vigorous shaking of the trichloromethane and water mixture. During the experiment, 0.20 cm <sup>3</sup> trichloromethane	<ol> <li>Commercial grade, dried over anhydrous CaCl<sub>2</sub> before use.</li> <li>Distified.</li> </ol>			
was added to 3.0 cm <sup>2</sup> of water and snaken. The amount of trichloromethane in the water was analyzed by passing the sample through a chromatographic	ESTIMATED ERRORS:			
column.	Solubility: Not specified. Temperature: ± 1 K (compiler).			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Wright, D. A.; Sandler, S. I.; DeVoll, D.			
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Technol. <u>1992</u> , 26, 1828-31.			
VARIABLES:	PREPARED BY:			
T/K = 293 - 323	A. L. Horvath			
EXPERIMENTAL VALUES:	í.			
t/°C γ <sup>∞</sup>	$\begin{array}{ccc} 10^3 x_1 & 100 w_1 \\ (compiler) \end{array}$			
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	1.22 0.803 1.18 0.777 1.16 0.764			
AUXILIARY II	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A differential static cell equilibrium apparatus was used for measuring the infinite dilution activity coefficient of trichloromethane ( $\gamma^{\infty}$ ) in water. The degassed water cells were submerged in a thermostat-	<ol> <li>Source and purity not given.</li> <li>Distilled, filtered and deionized.</li> </ol>			
water pain. I richloromethane was injected into the mixture cell and a magnetic stirrer was turned on. The mixture in the cells was allowed to equilibrate	ESTIMATED ERRORS:			
The stirrer was then turned off and the differential pressure was recorded. The experiment was repeated at least three times at each temperature.	Solubility: See above. Temperature: $\pm$ 0.05 K.			

132	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Trichloromethane (chloroform); CHCl.: [67-66-3]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.
(2) Milang water	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$
20 3.3×10 <sup>-3</sup>	0.7506 1.1407
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The solution was added to the stripping vessel and the desired flow was	<ol> <li>Source and purity not given.</li> <li>pH = 8.4, 700 mg salt/dm<sup>3</sup>, 140 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 15 mg organic carbon/dm<sup>3</sup>, 110 mg suspended solid/dm<sup>3</sup>.</li> </ol>
was taken for analysis. After extraction of the sam-	ESTIMATED ERRORS:
equipped with a capillary column.	Solubility: Not specified. Temperature: $\pm 0.1$ K.
	REFERENCES:
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.
COMPONENTS:	ORIGINAL MEASUREMENTS:
CHCl <sub>3</sub> ; [67-66-3]	Ph. D. Thesis, Stamford University, Stamford,
(2) MQ-Water	CA., <u>1985</u> , 306 pp.
VARIABLES:	PREPARED BY:
T/K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$
20 0.244	0.4207 6.371
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The multiple equilibrium technique with direct aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law con-stant. A known volume of trichloromethane and MQ-water was introduced into a syringe and allowed to equili- brate. The set of each to live up up a more	<ol> <li>J. T. Baker Chemicals Co., NJ., &gt; 99.5 % purity.</li> <li>0.25 mg total organic carbon per dm<sup>3</sup>, 0.75 mg COD/dm<sup>3</sup>.</li> </ol>
tained constant. While vapor samples were injected direct into a gas chromatograph. the liquid samples	ESTIMATED ERRORS:
were preconcentrated by liquid-liquid extraction before analysis. The experiments were carried out in duplicates.	Solubility: $\pm$ 0.016 std. dev. Temperature: $\pm$ 0.5 K.

	133			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Munz, C. D.			
(2) Municipal tap water (PASE)	Ph. D. Thesis, Stamford University, Stamford, CA., <u>1985</u> , 306 pp.			
VARIABLES:	PREPARED BY:			
T/K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
20 0.324	0.3174 4.803			
AUXILIARY II	NFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
The multiple equilibrium technique with direct aqueous injection into a gas chromatograph was used to determine the dimensionless Henry's law constant. A known volume of trichloromethane and municipal tap water was introduced into a syringe and allowed	<ol> <li>J. T. Baker Chemicals Co., NJ., &gt; 99.5 % purity.</li> <li>10 mg total organic carbon per dm<sup>3</sup>, 40 mg COD/dm<sup>3</sup>.</li> </ol>			
to equilibrate. The ratio of gas to liquid volume was maintained constant. While vapor samples were in- jected direct into a gas chromatograph, the liquid samples were preconcentrated by liquid-liquid extraction before analysis. The experiments were carried out in duplicate.	ESTIMATED ERRORS: Solubility: ± 0.020 std. dev. Temperature: ± 0.5 K.			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform);	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.			
(2) Myponga water	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.			
VARIABLES:	PREPARED BY:			
T/K = 293	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$			
20 2.9×10 <sup>-3</sup>	0.8542 1.299			
AUXILIARY II	VFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
The apparatus consisted of a stripping vessel similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser was close to the bottom. The mixture was	<ol> <li>Source and purity not given.</li> <li>pH = 7.6, 370 mg salt/dm<sup>3</sup>, 60 mg CaCO<sub>3</sub>/dm<sup>3</sup>, 10 mg organic carbon/dm<sup>3</sup>.</li> </ol>			
added to the stripping vessel and the desired flow was maintained. A sample of the solution being stripped	ESTIMATED ERRORS:			
sample, the extracts were analyzed by gas chromato- graph equipped with a capillary column.	Solubility: Not specified. Temperature: $\pm$ 0.1 K.			
· ·	REFERENCES:			
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.			

134	• · · · · · · · · · · · · · · · · · · ·		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform); CHCL: [67-66-3]	Nicholson, B. C.; Maguire, B. P.; Bursill, D. B.		
(2) Myponga water + humic acid	Environ. Sci. Technol. <u>1984</u> , 18, 518-21.		
MADIADI EQ			
VARIABLES: $TV = 202$	A L Horwith		
11K = 295			
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/m' atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$		
20 $3.1 \times 10^{-3}$	0.7991 1.2145		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The apparatus consisted of a stripping vessel	(1) Source and purity not given.		
similar to that described by Mackay <i>et al.</i> (ref. 1). The air entered through the top of the vessel and the diffuser use close to the bottom. The mixture user	(2) pH = 7.9, 20 mg humic acid/cm <sup>3</sup> , 370 mg salt/dm <sup>3</sup> , 68 mg CaCO <sub>3</sub> /dm <sup>3</sup> , 17 mg organic		
added to the stripping vessel and the desired flow was maintained. A sample of the solution being stripped	ESTIMATED ERRORS:		
was taken for analysis. After extraction of the sample, the extracts were analyzed by gas chromato-	Solubility: Not specified.		
graphy equipped with a capillary column.	Temperature: $\pm 0.1$ K.		
	REFERENCES:		
	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. P. Environ. Sci. Technol. <u>1979</u> , 13, 333.		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);	Balls, P. W.		
(2) Seawater	Ph. D. Thesis, Univ. of East Anglia, Norwich,		
	5. K., Part 1200, 500 pp.		
VARIABLES:	PREPARED BY:		
T/K = 283 - 298	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ (compiler) & (compiler) \end{array}$		
$10  0.112 \pm 0.025$	0.569 9.33		
$\begin{array}{cccc} 13 & 0.129 \pm 0.039 \\ 20 & 0.161 \pm 0.027 \\ 25 & 0.197 \pm 0.057 \end{array}$	0.604 10.07 0.605 9.92 0.603 9.89		
	0.005		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
The method involved repeated equilibrations of nitrogen with seawater sample containing trichloro- methane. The sample was shaken vigorously at the	<ol> <li>Source and purity not given.</li> <li>Salinity = 20 ‰.</li> </ol>		
required temperatures in a constant temperature bath for 30 minutes. The equilibrated gas phase was then	ESTIMATED ERRORS:		
injected into a gas chromatograph. The experiments			

COMPONENTS:		ORIGINAL MEASUREMENTS:		
(1) Trichloromethane (chloroform);		Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S.		
CHCl <sub>3</sub> ; [67-66-3]		Tellus <u>1983</u> , 35B, 170-6.		
VARIABLES:			PREPARED BY:	
T/K = 283 - 298			A. L. Horvath	
EXPERIMENTAL	VALUES	:	I	
	t/℃	Henry's law constant, H/dimensionless	100 w, (compiler)	$10^4 x_i$ (compiler)
	10 15 20 25	0.1046 0.1309 0.1624 0.2001	0.6500 0.6470 0.6428 0.6377	9.864 9.818 9.753 9.675
The Henry's law	v constants	s (dimensionless) were de	rived from the best-	fit lines of van't Hoff plots:
	log	$T_e H = -\frac{3649}{T(K)} + 10$	). 63	
		AUXILIARY II	NFORMATION	
METHOD/APPARA	ATUS/PR	OCEDURE:	SOURCE AND P	URITY OF MATERIALS:
The multiple measure the Henry' elsewhere (ref. 1). syringe was shaken	equilibriu s law cons A known with a kn	m technique was used to stant as described volume of seawater in a own volume of tri-	<ol> <li>Source and</li> <li>Samples fro late 1981.</li> </ol>	purity not given. m 48° N to 65° S in the Atlantic in
chloromethane. After equilibration, the headspace was separated and analyzed for trichloromethane. A		ESTIMATED ERI	RORS:	
detector was used f	gas chromatograph fitted with an electron capture detector was used for the analysis.		Solubility: Temperature:	± 5.5 % std. dev. ± 0.2 K.
			<b>REFERENCES:</b>	
			(1) McAuliffe,	C. D. Chem. Techn. <u>1971</u> , 1, 46.

136							
COMPONENTS:				ORIG	INAL MEASU	IREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]			Pavlovskaya, E. M.; Charykov, A. K.; Tikhomirov, V. I.				
(2) Cesium chloride; CsCl; [7647-17-8]			J. Ge	en. Chem. USS	R, <u>1977</u> , <i>4</i> 7, 2230-4.		
(3) Water; $H_2O$ ;	[7732-18-5]						
VARIABLES:				PREP	ARED BY:		
T/K = 293 Concentration				A. L	. Horvath		
EXPERIMENTA	L VALUES:						
	t/°C	<i>c</i> ₂/mol dm <sup>-3</sup>	<i>c</i> ,/m	ol m <sup>.3</sup>	100 w <sub>i</sub> (compiler)	$10^3 x_1$ (compiler)	
	20	0.5	64 4	5	0 6828	1 221	
		1.5 2.0	58.5	5	0.5830	1.143	
		2.5 3.0	56.0 52.5	5	0.5081 0.4560	1.237 1.263	
		AUXIL	JARY IN	IFORM	ATION		
METHOD/APPA	RATUS/PRO	CEDURE:	T	SOURCE AND PURITY OF MATERIALS:			
Trichlorom microburet into a chloride which co indicator. The mi pH of the solution to the appearance saturation of the a methane, after wh solubility was cal aqueous phase an	ethane was i n aqueous so ontained octai xture was vig 1 at first rem of a second aqueous phass nich the pH b culated from d the volume	ntroduced from a lution of cesium noic acid as an gorously stirred. ained constant rip phase, i. e., up e with trichloro- egan to rise. The the volume of th of trichlorometh	The ght up to the e he hane	(1) (2) (3) ESTIM Solub Temp	Source and pu Source and pu Distilled (com ATED ERRO ility: erature:	rity not given. rity not given. piler). RS: ± 10 %. ± 0.5 K (compiler).	
used for the satur	ation.						

						13
COMPONENTS:			ORIGINAL MEASUREMENTS:			
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]			Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. I.; Treger, Yu. A.; Flid, R. M.			
(2) Hydrogen chloride; HCl; [7647-01-0]			Russ	s. J. Phys. Ch	em. <u>1971</u> , 45, 488-90.	
(3) Water; H <sub>2</sub> O; [7732-18-5]						
VARIABLES:			PRE	PARED BY:		
T/K = 288 - 333	3		A. I	. Horvath		
EXPERIMENTAI	L VALUES:		-			
Solubility of CHC	Cl <sub>3</sub> in 10 % HCl s	solution:				
	t/℃	$10^3 w_I$	(	100 <i>w</i> , compiler)	$10^4 x_i$ (compiler)	
	15	6.70		0.670	10.71	
	30 45 60	4.44 2.36		0.444	7.086 3.759 2.132	
	00	1.54		0.154	2.152	
Solubility of CHC	Cl <sub>3</sub> in 20 % HCl s	olution:				
	t/°C	10 <sup>3</sup> w <sub>1</sub>	(	100 <i>w</i> , compiler)	$10^4 x_i$	
	15	9.20	(	0.920	15.58	
	30 45	5.16 2.84		0.516 0.284	8.706 4.781	
	60	1.50		0.150	2.522	
· <u>·····</u> _···						
METHOD/APPA	RATUS/PROCED	OURE:		RCE AND PU	URITY OF MATERIALS:	
solubility determin nitrogen with the bubbled through a	nation consisted o solute vapor which thermostatically	of the saturation of the was then controlled flask	(1) (2) (3)	Source and p Source and p Distilled (con	burity not given. mpiler).	
filled with a defin After 2 - 3 hours,	the solution beca	the saturated and	ESTI	MATED ERR	ORS:	
samples were taken periodically (ref. 1). The tri- chloromethane concentration in the aqueous solution was determined by GLC. On average, 3 - 5 measure- ments were taken at each temperature		Solu Tem	bility: perature:	± 15 %. ± 0.05 K.		
	•		REFI	ERENCES:		
			(1)	Treger, Yu. Russ. J. Phy	A.; Flid, R. M.; Spektor, S. S. s. Chem. <u>1964</u> , 38, 253.	
			I			

138		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ul><li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li></ul>	Pavlovskaya, E. M.; Charykov, A. K.; Tikhomirov, V. I.	
(2) Lithium chloride; LiCl; [7447-41-8]	J. Gen. Chem. USSR, <u>1977</u> , 47, 2230-4.	
(3) Water; H <sub>2</sub> O; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 293 Concentration	A. L. Horvath	
EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_2/\text{mol dm}^{-3}$ $c_1/\text{mol m}^{-3}$ 100 $w_1$ 10 <sup>3</sup> $x_1$ (compiler) (compiler)		
20 0.5 66. 1.0 64. 1.5 60. 2.0 58. 2.5 56. 3.0 52.	8         0.7880         1.212           5         0.7503         1.168           0         0.6921         1.091           5         0.6675         1.065           0         0.6320         1.022           0         0.5805         1.003	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Trichloromethane was introduced from a microburet into an aqueous solution of lithium chloride which contained octanoic acid as indicator. The mixture was vigorously stirred. The pH of the solution at first remained constant right up to the	<ol> <li>Source and purity not given.</li> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
appearance of a second phase, i. e., up to the saturation of the aqueous phase with trichloro- methane, after which the pH began to rise. The solubility was calculated from the volume of the aqueous phase and the volume of trichloromethane used for the saturation.	ESTIMATED ERRORS: Solubility: ± 10 %. Temperature: ± 0.5 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Pavlovskaya, E. M.; Charykov, A. K.; Tikhomirov, V. I.	
(2) Potassium chloride; KCl; [7447-40-7]	J. Gen. Chem. USSR, <u>1977</u> , 47, 2230-4.	
(3) Water; H <sub>2</sub> O; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 293 Concentration	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C $c_2/\text{mol dm}^{-3}$ $c_1/\text{m}$	ol m <sup>-3</sup> 100 w, $10^3 x_i$ (compiler) (compiler)	
20 0.5 56.0	0 0.6535 1.021 (continued)	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Trichloromethane was introduced from a microburet into an aqueous solution of potassium chloride which contained octanoic acid as indicator. The mixture was vigorously stirred. The pH of the solution at first remained constant right up to the appearance of a second phase, i. e., up to the	<ol> <li>Source and purity not given.</li> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol> ESTIMATED ERRORS:	
saturation of the aqueous phase with trichloro- methane, after which the pH began to rise. The solubility was calculated from the volume of the aqueous phase and the volume of trichloromethane used for the saturation.	Solubility: $\pm$ 10 %. Temperature: $\pm$ 0.5 K (compiler).	

							139
COMPONENTS	:			ORI	GINAL MEASU	IREMENTS:	
(1) Trichlorome CHCl <sub>3</sub> ; [67-	thane (chlo 66-3]	roform);		Pav Ti	ovskaya, E. M khomirov, V. I	; Charykov, A. K.;	
(2) Potassium c	hloride; KC	1; [7447-40-7]		J. C	ien. Chem. USS	R, <u>1977</u> , <i>4</i> 7, 2230-4.	
(3) Water; H <sub>2</sub> O	; [7732-18-5	]					
VARIABLES:				PRE	PARED BY:		
T/K = 293 Concentration				A. I	. Horvath		
EXPERIMENTA	L VALUE	S: (continued)					
	t/℃	<i>c</i> ₂/mol dm <sup>-3</sup>	<i>c</i> ,/m	ol m <sup>.3</sup>	100 w <sub>i</sub> (compiler)	$\begin{array}{c} 10^3 x_1 \\ \text{(compiler)} \end{array}$	
	20	1.0 1.5	50. 41.	D D	0.5692 0.4591	0.9160 0.7610	
		2.0 2.5 3.0	29. 24.	2	0.3131 0.2546	0.5542 0.4639	
COMPONENTS	:			ORIC	SINAL MEASU	REMENTS:	
(1) Trichlorome CHCl <sub>3</sub> ; [67-4	thane (chlor 66-3]	roform);		Pavl Ti	ovskaya, E. M. chomirov, V. I.	; Charykov, A. K.;	
(2) Rubidium cl	hloride; Rb(	Cl; [7791-11-9]		J. G	en. Chem. USS	R, <u>1977</u> , 47, 2230-4.	
(3) Water; $H_2O$	; [7732-18-5	]					
VARIABLES:				PRE	PARED BY:		
T/K = 293 Concentration				A. I	Horvath		
EXPERIMENTA	L VALUE	S:					
	t/°C	<i>c</i> ₂/mol dm <sup>-3</sup>	<i>c</i> 1/m	ol m <sup>-3</sup>	100 w <sub>ر</sub> (compiler)	$\begin{array}{c} 10^3 \ x_i \\ \text{(compiler)} \end{array}$	
	20	0.5 1.0	58.5	5	0.6454	1.0920	
		1.5 2.0 2.5	46.( 43.	)	0.4738 0.4635	0.9049 0.8751	
		3.0	39.	1	0.3852	0.8449	
		AUXIL	JARY II	NFORM	MATION		
METHOD/APPA	ARATUS/PI	OCEDURE:		SOU	RCE AND PUR	ITY OF MATERIALS:	
Trichloron microburet into a chloride which c The mixture was	nethane was an aqueous ontained oc vigorously	s introduced from solution of rubidiu anoic acid as indic stirred. The pH o	a m cator. f the	(1) (2) (3)	Source and pu Source and pu Distilled (com	rity not given. rity not given. piler).	
solution at first r appearance of a	remained co second phas	nstant right up to the	the	ESTI	MATED ERRO	RS:	
saturation of the methane, after w solubility was ca aqueous phase a used for the satu	aqueous ph which the pH lculated fro nd the volut ration.	ase with trichloro- began to rise. Th m the volume of t ne of trichloromet	he he hane	Solu Tem	bility: perature:	$\pm$ 10 %. $\pm$ 0.5 K (compiler).	

140		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
<ul><li>(1) Trichloromethane (chloroform); CHCl<sub>3</sub>; [67-66-3]</li></ul>	Pavlovskaya, E. M.; Charykov, A. K.; Tikhomirov, V. I.	
(2) Sodium chloride; NaCl; [7647-14-5]	J. Gen. Chem. USSR, 47, 2230-4.	
(3) Water; H <sub>2</sub> O; [7732-18-5]		
VARIABLES:	PREPARED BY:	
T/K = 293 Concentration	A. L. Horvath	
EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_2/\text{mol dm}^{-3}$ $c_1/\text{mol m}^{-3}$ 100 $w_1$ 10 <sup>3</sup> $x_1$ (compiler) (compiler)		
20 0.5 57. 1.0 51. 1.5 42. 2.0 37. 2.5 30. 3.0 25.	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Trichloromethane was introduced from a microburet into an aqueous solution of sodium chloride which contained octanoic acid as indicator. The mixture was vigorously stirred. The pH of the solution at first remained constant right up to the	<ol> <li>Source and purity not given.</li> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
appearance of a second phase, i. e., up to the saturation of the aqueous phase with trichloro- methane, after which the pH began rise. The solubility was calculated from the volume of the aqueous phase and the volume of trichloromethane used for the saturation.	ESTIMATED ERRORS: Solubility: ± 10 %. Temperature: ± 0.5 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water-d <sub>2</sub> ; D <sub>2</sub> O; [7789-20-0]	Hutchison, C. A.; Lyon, A. M.	
(2) Trichloromethane (chloroform); CHCl <sub>3</sub> ; [67-66-3]	Columbia University Report A-745, July 1, 1943.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES: t/°C 100 w <sub>t</sub> M <sub>t</sub> -t/mol g <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$25$ $4.15 \times 10^{-3}$	$8.31 \times 10^{-2}$ 4.93	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A 1 to 15 volume ratio mixture of heavy water and trichloromethane was introduced into an equilibration flask and then lowered into a water thermeetter. The generative walks and the machanically	<ol> <li>Source and purity not given.</li> <li>Source not given. Purified and dried before use.</li> </ol>	
for about 90 minutes at constant temperature. The	ESTIMATED ERRORS:	
determined by a modified Karl Fischer titration method. The determination was done in triplicate. The description was taken from a secondary	Solubility: $\pm 0.6 \times 10^{-4}$ avg. dev. Temperature: $\pm 0.05$ K.	
source (ref. 1). The original report is no longer available.	REFERENCES:	
	<ol> <li>Eidinoff, M. L.; Joris, G. G.; Taylor, H. S.; Urey, H. C., Eds., Production of Heavy Wa- ter, McGraw-Hill, New York, <u>1955</u>, p. 129.</li> </ol>	

	141
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Triiodomethane (iodoform);	Dehn, W. M.
(2) Water; $H_2O$ ; [7732-18-5]	J. Am. Chem. Soc. <u>1917</u> , 39, 1399-404.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 293 - 298	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}$ C 100 $g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^6 \ x_i \\ (compiler) & (compiler) \end{array}$
$20 - 25$ $1.0 \times 10^{-1}$	$1.0 \times 10^{-1}$ 4.57
	YINFORMATION
AUXILIAR	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Excess truodomethane in 5 cm <sup>2</sup> water sample were enclosed in small vials which, when stoppere were staken until equilibria were established. The	d, (2) Distilled (compiler).
solutions were filtered into weighed crucibles and reweighed. After drying in vaccum desiccators, the	ESTIMATED ERRORS:
crucibles were again weighed and the loss of water calculated. No great accuracy was claimed for the	Solubility: Not specified.
experiments.	Temperature: ± 5 K.
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Triiodomethane (iodoform);	van Arkel, A. E.; Vles, S. E.
(2) Water; $H_2O$ ; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 303	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C m₁/mol kg <sup>·1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^6 \ x_i \\ (compiler) & (compiler) \end{array}$
$30 3 \times 10^{-4}$	$1.18 \times 10^{-2}$ 5.41
AUXILIAR	Y INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified.
	T = T + C (complete).

142		-	
COMPONENTS:		ORIGINAL MEASUR	REMENTS:
<ul> <li>(1) Triiodomethane (iodoform); CHI<sub>3</sub>; [75-47-8]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ul>		Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.	
VARIABLES:		PREPARED BY:	
T/K = 298		A. L. Horvath	
EXPERIMENTAL VALUES:			
t/°C	$g_1 V_2^{-1} / \text{kg m}^{-3}$	100 w <sub>i</sub> (compiler)	10 <sup>s</sup> x, (compiler)
25.0	< 0.2	< 0.02	< 4.6
		NFORMATION	
	AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCE	AUXILIARY I	NFORMATION SOURCE AND PURI	TY OF MATERIALS:
METHOD/APPARATUS/PROCE Equilibrium was ensured the shaking and centrifuging of a triin water mixture in a stoppered Goe turning it to a constant temperatur difference between the temperature	AUXILIARY I BDURE: hrough repeated odomethane and tz tube and re- ire water bath. The ut of triedo	NFORMATION SOURCE AND PURI (1) Commercial rea received. (2) Distilled.	TY OF MATERIALS: agent, C. P. grade, used as
METHOD/APPARATUS/PROCE Equilibrium was ensured ti shaking and centrifuging of a trii water mixture in a stoppered Goe turning it to a constant temperatu difference between the total amou methane which was added and the	AUXILIARY I BDURE: hrough repeated odomethane and tz tube and re- ire water bath. The int of triiodo- e amount remaining t of triiodo-	NFORMATION SOURCE AND PURI (1) Commercial rea received. (2) Distilled. ESTIMATED ERROR	TY OF MATERIALS: agent, C. P. grade, used as
METHOD/APPARATUS/PROCE Equilibrium was ensured the shaking and centrifuging of a triis water mixture in a stoppered Goe turning it to a constant temperatu difference between the total amoun methane which was added and the in excess was taken as the amoun dissolved in the known volume o termination of the excess amount added is described by Hanslick (r	AUXILIARY I BDURE: hrough repeated odomethane and etz tube and re- re water bath. The int of triiodo- e amount remaining it of triiodomethane f water. The de- of triiodomethane ref. 1).	NFORMATION SOURCE AND PURI (1) Commercial rea received. (2) Distilled. ESTIMATED ERROR Solubility: Temperature:	TY OF MATERIALS: agent, C. P. grade, used as SS: < 100 %. ± 1 K (compiler).
METHOD/APPARATUS/PROCE Equilibrium was ensured ti shaking and centrifuging of a trii water mixture in a stoppered Goe turning it to a constant temperatu difference between the total amou methane which was added and th in excess was taken as the amoun dissolved in the known volume o termination of the excess amount added is described by Hanslick (r	AUXILIARY I BDURE: hrough repeated odomethane and etz tube and re- ire water bath. The unt of triiodo- e amount remaining it of triiodomethane f water. The de- of triiodomethane ref. 1).	NFORMATION SOURCE AND PURI (1) Commercial rea received. (2) Distilled. ESTIMATED ERROR Solubility: Temperature: REFERENCES:	TY OF MATERIALS: agent, C. P. grade, used as S: < 100 %. ± 1 K (compiler).

COMPONENTS:	EVALUATOR:
<ul><li>(1) Bromochloromethane; CH₂BrCl;</li><li>[74-97-5]</li></ul>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.
(2) Water; H <sub>2</sub> O; [7732-18-5]	March 1993.

### CRITICAL EVALUATION:

The bromochloromethane (1) and water (2) binary system is discussed in two parts; part 1 is bromochloromethane (1) in water (2) and part 2 is water (2) in bromochloromethane (1).

The solubility of bromochloromethane (1) in water (2) has been studied by three workers, whereas the solubility of water (2) in bromochloromethane (1) has been reported in one paper only.

**Part 1.** The solubility of bromochloromethane (1) in water (2) has been studied by three workers. The experimental data of Tewari *et al.* (ref. 1) and Miller *et al.* (ref. 2) are in excellent agreement and the mass per cent (1) value of 1.646 at 298.15 K is recommended. In both cases the modified generator column method was used, but the analytical method was more advanced in the latter case. (For a comparison, see the two compilation sheets.) The single mass per cent (1) data point of 1.5 g(1)/100 g soln. at 293.15 K, reported by O'Connell (ref. 3) is considered as tentative. Whether the solubilities at 293.15 and 298.15 K are consistent with the general trend of temperature dependence remains to be investigated.

**Part 2.** The solubility of water (2) in bromochloromethane (1) has been reported in only one paper, that of O'Connell (ref. 3) and there are no other measurements for comparison. Therefore, it is not possible to perform a critical evaluation. The readers are referred to the compilation sheet for further details.

### **REFERENCES:**

- <sup>7</sup><sup>3</sup> 1. Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E. J. Chem. Eng. Data <u>1982</u>, 27, 451.
- ⊃ 3 4 2. Miller, M. M.; Wasik, S. P.; Huang, T.-C.; Shiu, W.-Y.; Mackay, D. Environ. Sci. Technol. <u>1985</u>, 19, 522.

3. O'Connell, W. L. Trans. Am. Ins. Mech. Eng. 1963, 226, 126.

144	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromochloromethane; $CH_2BrCl$ ; [74-97-5]	O'Connell, W. L.
(2) Water; $H_2O$ ; [7732-18-5]	Trans. Am. Inst. Mech. Eng. <u>1963</u> , 226, 126-32.
VARIABLES:	PREPARED BY:
<i>T/</i> K = 293	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 100 g <sub>1</sub> /g <sub>2</sub> 10 (con	$x_1 = \frac{100 g_2/g_1}{(compiler)}$
20 1.5 2	$.08    9.0 \times 10^{-2}    6.43$
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Details are not available.	<ol> <li>Dow Chemicals Co., used as received.</li> <li>Distilled (compiler).</li> </ol>
	ESTIMATED ERRORS:
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).
COMPONENTS:	ORIGINAL MEASUREMENTS:
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5]	ORIGINAL MEASUREMENTS: Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5]	<ul> <li>ORIGINAL MEASUREMENTS:</li> <li>Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.</li> <li>J. Chem. Eng. Data <u>1982</u>, 27, 451-4.</li> </ul>
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES:	ORIGINAL MEASUREMENTS: Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E. J. Chem. Eng. Data <u>1982</u> , 27, 451-4. PREPARED BY:
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 298	ORIGINAL MEASUREMENTS: Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E. J. Chem. Eng. Data 1982, 27, 451-4. PREPARED BY: A. L. Horvath
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: <i>T</i> /K = 298 EXPERIMENTAL VALUES:	ORIGINAL MEASUREMENTS: Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E. J. Chem. Eng. Data 1982, 27, 451-4. PREPARED BY: A. L. Horvath
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/\text{mol m}^{-3}$	ORIGINAL MEASUREMENTS:Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.J. Chem. Eng. Data 1982, 27, 451-4.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) $10^3 x_i$ (compiler)
COMPONENTS:         (1) Bromochloromethane; $CH_2BrCl;$ [74-97-5]         (2) Water; $H_2O;$ [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ 25.0       125	ORIGINAL MEASUREMENTS:         Tewari, Y. B.; Miller, M. M.; Wasik, S. P.;         Martire, D. E.         J. Chem. Eng. Data 1982, 27, 451-4.         PREPARED BY:         A. L. Horvath $100 w_i$ $10^3 x_i$ (compiler)       1.646         2.325
COMPONENTS:         (1) Bromochloromethane; CH <sub>2</sub> BrCl;         [74-97-5]         (2) Water; H <sub>2</sub> O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t'^{\circ}$ C $t''^{\circ}$ C $t'''^{\circ}$ C      <	ORIGINAL MEASUREMENTS:Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.J. Chem. Eng. Data 1982, 27, 451-4.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 1.646 $2.325$
COMPONENTS:         (1) Bromochloromethane; CH2BrCl;         [74-97-5]         (2) Water; H2O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}$ C $c_1$ /mol m <sup>-3</sup> 25.0       125         AUXILIARY II	ORIGINAL MEASUREMENTS:Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.J. Chem. Eng. Data 1982, 27, 451-4.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 1.646 $1046$ 2.325NFORMATION
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ 25.0 125 AUXILIARY II METHOD/APPARATUS/PROCEDURE:	ORIGINAL MEASUREMENTS:Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.J. Chem. Eng. Data 1982, 27, 451-4.PREPARED BY: A. L. Horvath $\begin{pmatrix} 100 & w_i \\ (compiler) \\ 1.646 \end{pmatrix}$ 1.6462.325NFORMATIONSOURCE AND PURITY OF MATERIALS:
COMPONENTS:         (1) Bromochloromethane; CH2BrCl; $[74-97-5]$ (2) Water; H2O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t'^{\circ}C$ $c_1/mol m^{-3}$ 25.0         125         METHOD/APPARATUS/PROCEDURE:         The modified generator column method was used. A chromatographic column was coated with bromochloromethane. An aqueous phase was generated by pumping water through the coated generator	ORIGINAL MEASUREMENTS:         Tewari, Y. B.; Miller, M. M.; Wasik, S. P.;         Martire, D. E.         J. Chem. Eng. Data 1982, 27, 451-4.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         1.646         2.325         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source not known, at least 99 % pure by GC analysis.         (2) Baker-analyzed HPLC grade.
COMPONENTS:         (1) Bromochloromethane; CH2BrCl; $[74-97-5]$ (2) Water; H2O; [7732-18-5]         VARIABLES: $T/K = 298$ EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ 25.0         125         METHOD/APPARATUS/PROCEDURE:         The modified generator column method was used. A chromatographic column was coated with bromochloromethane. An aqueous phase was generated by pumping water through the coated generator column. The aqueous solution was analyzed by a gas chromatograph. The gas chromatograph was equipated by a gas chromatograph.	ORIGINAL MEASUREMENTS:         Tewari, Y. B.; Miller, M. M.; Wasik, S. P.;         Martire, D. E.         J. Chem. Eng. Data 1982, 27, 451-4.         PREPARED BY:         A. L. Horvath         100 w, (compiler)         1.646         2.325         NFORMATION         SOURCE AND PURITY OF MATERIALS:         (1) Source not known, at least 99 % pure by GC analysis.         (2) Baker-analyzed HPLC grade.         ESTIMATED ERRORS:
COMPONENTS: (1) Bromochloromethane; CH <sub>2</sub> BrCl; [74-97-5] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES: T/K = 298 EXPERIMENTAL VALUES: $t/^{\circ}C$ $c_1/mol m^{-3}$ 25.0 125 AUXILIARY I METHOD/APPARATUS/PROCEDURE: The modified generator column method was used. A chromatographic column was coated with bromochloromethane. An aqueous phase was generat- ed by pumping water through the coated generator column. The aqueous solution was analyzed by a gas chromatography. The gas chromatograph was equip- ped with a flame ionization detector. On average, at least three measurements were taken.	ORIGINAL MEASUREMENTS:Tewari, Y. B.; Miller, M. M.; Wasik, S. P.; Martire, D. E.J. Chem. Eng. Data 1982, 27, 451-4.PREPARED BY: A. L. Horvath $100 w_i$ (compiler) 1.646 $100 w_i$ (compiler) 1.646 $100 w_i$ (compiler) 2.325NFORMATIONSOURCE AND PURITY OF MATERIALS: (1) Source not known, at least 99 % pure by GC analysis. (2) Baker-analyzed HPLC grade.ESTIMATED ERRORS: Solubility: Temperature: $\pm 1.0 %$ . $\pm 0.1 K.$

			14
COMPONENTS:		ORIGINAL MEASUR	EMENTS:
<ol> <li>Bromochloromethane; CH<sub>2</sub>BrCl; [74-97-5]</li> <li>Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>		Miller, M. M.; Wasik, S. P.; Huang, GL.; Shiu, WY.; Mackay, D. <i>Environ. Sci. Technol.</i> <u>1985</u> , 19, 522-9.	
T/K = 298		A. L. Horvath	
EXPERIMENTAL VALUES:			
t/°C	$n_1 V_2^{-1} / \text{mol m}^{-3}$	100 w <sub>i</sub> (compiler)	$10^3 x$ , (compiler)
25	129.0	1.646	2.325
	AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCH	EDURE:	SOURCE AND PURIT	Y OF MATERIALS:
A saturated aqueous soluti passing water through a column with glass beads coated with brow The saturated solution was then p	on was prepared by which was packed mochloromethane. pumped through an	<ol> <li>Eastman Kodak used as received</li> <li>Double distilled.</li> </ol>	Co., Commercial quality,
extractor column and bromochlor tracted into an analytical column	omethane was ex- The samples were	ESTIMATED ERROR	S:
analyzed with a liquid chromatog both an UV absorbance and a flu with excitation filters. The peak	analyzed with a liquid chromatograph equipped with both an UV absorbance and a fluorescent detector with excitation filters. The peak area was established		Not specified. 1 K (compiler).
using a newlett-rackard integrat	UI.		

COMPONENTS:	EVALUATOR:
<ol> <li>(1) Dibromomethane; CH<sub>2</sub>Br<sub>2</sub>; [74-95-3]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K. March 1993.

#### **CRITICAL EVALUATION:**

146

The dibromomethane (1) and water (2) binary system is discussed in two parts; part 1 is dibromomethane (1) in water (2) and part 2 is water (2) in dibromomethane (1).

**Part 1.** The solubility of dibromomethane (1) in water (2) has been studied by 6 workers. The Compilation Sheets immediately follow this Critical Evaluation.

The experimental data of all investigators in the 273 to 323 K temperature range are sufficiently reliable for use in the smoothing equation (see Figure 1) with the exception of the experimental work of Booth and Everson (ref. 1) which is markedly higher than other results and is therefore rejected. The combined data of 5 laboratories were correlated to give the following equation for mass per cent (1) solubility:

Solubility  $[100 w_l] = 10.1890 - 0.06336789 (T/K) + 1.10906 \times 10^4 (T/K)^2$ 

which produced a standard deviation of  $4.4 \times 10^{-2}$ . This equation represents the combined solubility data with a maximum deviation of 6.0 % (usually less) and may be considered recommended to represent the data in the 273 to 323 K temperature interval.

The recommended mass per cent solubility values at 5 K intervals for dibromomethane (1) in water (2) are recorded in Table 1.

Table 1. Recommended Solubility of Dibromomethane (1) in Water (2)

Solubility
$10^3 x_1$
1.209
1.198
1.191
1.191
1.198
1.209
1.226
1.251
1.279
1.313
1.356

**Part 2.** The solubility of water (2) in dibromomethane (1) has been reported in one work (ref. 2) and, therefore, no Critical Evaluation has not been performed. The readers are referred to the relevant Compilation Sheet.

(continued next page)



# **REFERENCES:**

- 1. Booth, H. S.; Everson, H. E. Ind. Eng. Chem. 1948, 40, 1491.
  - 2. O'Connell, W. L. Trans. Am. Inst. Mech. Eng. 1963, 226, 126.

2

t

; 1

148		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromomethane; CH <sub>2</sub> Br <sub>2</sub> ; [74-95-3]	Rex, A.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Z. Phys. Chem. <u>1906</u> , 55, 355-70.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 100 $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
0 1.173	1.15 1.20	
20 1.148 30 1.76	1.13 1.16 1.18 1.18	
	I	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility of dibromomethane in water was determined in specially designed flasks with calibrat- ed capillary cylinders. After the samples reached	<ol> <li>Merck reagent, redistilled and washed before use, b. p. = 97.8°C.</li> <li>Distilled.</li> </ol>	
equilibrium in a thermostatic bath, their volumes and weights were used to determine the solubility values.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
	-	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromomethane; $CH_2Br_2$ ; [74-95-3]	Gross, P. M.; Saylor, J. H.	
(2) Water; $H_2O$ ; [7732-18-5]	J. Am. Chem. Soc. <u>1931</u> , 53, 1744-51.	
VARIABLES:	PREPARED BY:	
T/K = 288 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:	· · · · · · · · · · · · · · · · · · ·	
$t/^{\circ}C$ 1000 $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
15 11.70 30 11.93	1.16 1.21 1.18 1.24	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of dibromomethane in 50 g water was shaken for 12 hours in a thermostat bath. Samples were then withdrawn and read against water in an interferometer made by Zeiss (ref. 1).	<ol> <li>Eastman Kodak Co., purified by fractionation through a Hempel column before use.</li> <li>Distilled.</li> </ol>	
A detailed description of the complete procedure is given in a Ph. D. thesis (ref. 2).	ESTIMATED ERRORS:	
	Solubility: $\pm 0.5$ %. Temperature: $\pm 0.02$ K.	
	REFERENCES:	
	(1) Gross, P. M. J. Am. Chem. Soc. 1929. 51.	
	<ul> <li>2362.</li> <li>(2) Saylor, J. H. Ph. D. Thesis, Duke University, Durham, <u>1930</u>.</li> </ul>	

	149	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromomethane; CH <sub>2</sub> Br <sub>2</sub> ; [74-95-3]	van Arkel, A. E.; Vles, S. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 303	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
$t/^{\circ}C$ $m_{i}/mol kg^{-1}$	$10^2 x_i$ $100 w_i$ (compiler)	
$30.0  ext{ } 6.86  imes 10^2$	0.123 1.174	
AUXILIARY	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified.	
	Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dibromomethane; CH <sub>2</sub> Br <sub>2</sub> ; [74-95-3]	Botth, H. S.; Everson, H. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
$t/^{\circ}C$ 100 $V_{1}/V_{2}$	$100 w_i$ $10^3 x_i$	
25.0 0.70	1.71 1.80	
	SOURCE AND DUDITY OF MATERIALS	
The equilibrium was ensured through repeated	(1) Commercial reagent C P grade used as	
shaking and centrifuging of a dibromomethane and water mixture in a stoppered Goetz tube and return- ing the sample to a constant temperature water bath.	(2) Distilled.	
The difference between the total amount of dibromo- methane which was added and the amount remaining	ESTIMATED ERRORS:	
ane dissolved in the known volume of water. The determination of the excess amount of dibromo- methane added is described by Hanslick (ref. 1).	Solubility: Not specified. Temperature: ± 1 K (compiler).	
	REFERENCES:	
	(1) Hanslick, R. S. Ph. D. Thesis, Columbia University, <u>1935</u> .	
	Oniversity, <u>1222</u> .	

-

COMPONENTS:		ORIGINAL MEASUREMENTS:		
(1) Dibromomethane; CH <sub>2</sub> Br <sub>2</sub> ; [74-95-3]		O'Connell, W. L.		
(2) Water; H <sub>2</sub> O; [7732-18-5]		Trans. Am.	Inst. Mech.	Eng. <u>1963</u> , 226, 126-32.
VARIABLES:		PREPARED	BY:	
<i>T</i> /K = 293 - 298		A. L. Horva	ath	
EXPERIMENTAL VALUES:				
t/°C 100 w₁	10 (com	'x, l ppiler)	100 w <sub>2</sub>	$10^3 x_2$ (compiler)
20 - 25 1.1	- 1.	14 .	0.07 -	6.71
AUX	ILIARY II	NFORMATION	N	
METHOD/APPARATUS/PROCEDURE:		SOURCE AN	ND PURITY	OF MATERIALS:
Details are not available.		<ol> <li>Dow Chemicals Co., used as received.</li> <li>Distilled (compiler).</li> </ol>		
		ESTIMATEI	O ERRORS:	
		Solubility: Temperature	: ±	Not specified. 0.5 K (compiler).
COMPONENTS		ODIGINAL	MEASUDEN	IENTS.
(1) Dibromomethane: CH Br $\cdot$ [74-95-3]		Wright D	A · Sandler	S I · DeVoll D
(1) $M_{2}$ (1)		Environ Sci	$T_{ach}$ 100 <sup>o</sup>	3. 1., Devon, D.
(2) water, 1120, [7752-10-5]		Linni on. Sci	. 1een. <u>1994</u>	<u>2</u> , 20, 1020-51.
VARIABLES:		PREPARED	BY:	
<i>T</i> /K = 293 - 323		A. L. Horva	ath	
EXPERIMENTAL VALUES:				
t/°C γ <sup>∞</sup>		$10^3 x_1$		100 w, (compiler)
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		1.15		1.10
50 740 ± 32		1.35		1.29
AUX	ILIARY IN	NFORMATION	Ν	
METHOD/APPARATUS/PROCEDURE:		SOURCE AN	ND PURITY	OF MATERIALS:
A differential static cell equilibrium apparatus was used to measure the infinite dilution activity coefficient for dibromomethane ( $\gamma^{\infty}$ ) in water. Degas- sed water cells were submerged in a thermostated water bath. Dibromomethane was injected into a mix- ture cell, a magnetic stirrer was turned on, and the cells were allowed to equilibrate. The stirrer was then turned off and the differential pressure was recorded. The experiment was repeated at least three times at each temperature.		(1) Source (2) Distille	and purity and filtered a	not given. nd deionized.
		ESTIMATED	ERRORS:	
		Solubility: Temperature	: ±(	See above. D.05 K.

COMPONENTS:	EVALUATOR:
<ol> <li>(1) Dichloromethane; CH<sub>2</sub>Cl<sub>2</sub>; [75-09-2]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K. March 1993.

### **CRITICAL EVALUATION:**

The dichloromethane (1) and water (2) binary system is discussed in two parts; part 1 is dichloromethane (1) in water (2) and part 2 is water (2) in dichloromethane (1).

**Part 1.** The solubility of dichloromethane (1) in water (2) has been studied by at least 32 groups of workers. Most of the data reported have been considered for inclusion in the smoothing equation with the following exclusions. The experimental data of Salkowski (ref. 1) are markedly lower than the other results and are therefore rejected. The solubility data of Booth and Everson (ref. 2), Alexandrova *et al.* (ref. 3), and Sadovnikova *et al.* (refs. 4 and 5) are several per cent higher than the smoothed solubility values and are also rejected. The data of Sabinin *et al.* (ref. 6) between 293.15 and 303.15 K show a temperature dependence of solubility much too large and are therefore rejected. The data of Svetlanov *et al.* (ref. 7) also show a change in solubility far too large between 288.15 and 333.15 K and are rejected; however, the value at 303.15 K is retained. The solubilities calculated from the distribution coefficients of Prosyanov *et al.* (ref. 8) are in very poor agreement, providing no confidence in their values which are regarded as dubious. The solubility data of Howe *et al.* (ref. 9) between 283.15 and 303.15 K show a maximum which is inconsistent with the trend of temperature dependence of solubility and are rejected; however, the value at 293.15 K is retained.

The remaining data of 25 laboratories by Rex (ref. 10), van Arkel and Vles (ref. 11), Bakowski and Treszczanovicz (ref. 12), Niini (ref. 13), McGovern (ref. 14), Donahue and Bartell (ref. 15), Kudryavtseva and Krutikova (ref. 16), Maretic and Sirocic (ref. 17), du Pont (ref. 18), Svetlanov *et al.* (ref. 7), Karger *et al.* (ref. 19), Antropov *et al.* (ref. 20), McConnell *et al.* (ref. 21), Pearson and McConnell (ref. 22), Archor and Stevens (ref. 23), Sato and Nakijima (ref. 24), Coca *et al.* (ref. 25), Hutchinson *et al.* (ref. 26), Leighton and Calo (ref. 27), Lincoff and Gossett (ref. 28), Gossett (ref. 29), Warner *et al.* (ref. 30), Howe *et al.* (ref. 9), Vogel (ref. 31), and Wright *et al.* (ref. 32), were used to obtain the following smoothing equation for solubility in mass per cent (1):

Solubility  $[100 w_l] = 58.838 - 0.38224 (T/K) + 6.3928 \times 10^4 (T/K)^2$ 

This equation represents the combined data with a 0.37 standard deviation in the 273 to 310 K temperature range.

The curve obtained from the smoothing equation shows a distinct minimum at 299 K as seen in Figure 1. The appearance of the minimum for the solubility versus temperature behavior is discussed in the Preface.

The recommended mass per cent solubility values at 5 K intervals for dichloromethane (1) in water (2) are presented in the Table 1.

Table 1. Recommended Solubility of Dichloromethane (1) in Water (2)

Tem	perature	Solubil	ity
°C	ĸ	100 w <sub>1</sub>	$10^{3} x_{1}$
0	273.15	2.126	4.587
5	278.15	1.977	4.260
10	283.15	1.860	4.004
15	288.15	1.766	3.799
20	293.15	1.722	3.703
25	298.15	1.701	3.657
30	303.15	1.712	3.681
35	308.15	1.754	3.773

(continued next page)

# COMPONENTS:

- (1) Dichloromethane;  $CH_2Cl_2$ ; [75-09-2]
- (2) Water;  $H_2O$ ; [7732-18-5]

EVALUATOR:

A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.

March 1993.

# CRITICAL EVALUATION: (continued)





**Part 2.** The solubility of water (2) in dichloromethane (1) has been reported from 19 laboratories. The solubility measurements cover the 253 to 313 K temperature range as represented in Figure 2.

The data reported have been considered for inclusion in the smoothing equation with the following exclusions. The data of Bakowski and Terszczanowicz (ref. 12), Niini (ref. 13), Donahue and Bartell (ref. 15), and Kudryavtseva and Krutikova (ref. 16) are markedly higher than other results and are therefore rejected. The measured solubility of Alexandrova et al. (ref. 3), and Sadovnikova *et al.* (refs. 4 and 5) are several per cent lower than the smoothed solubility and are also rejected. The solubilities calculated from the distribution coefficients of Prosyanov *et al.* (ref. 33), are in very poor agreement, providing no confidence in their values which are regarded as dubious.

The remaining data of 11 laboratories by Staverman (Ref. 34), McGovern (ref. 14), Davies *et al.* (ref. 35), Maretic and Sirocic (ref. 17), Sabinin *et al.* (ref. 6). Lees and Sarram (ref. 36), Antropov *et al.* (ref. 20), Archer and Stevens (ref. 23), Coca *et al.* (ref. 25), Wu (ref. 37), and Ohtsuka and Kazama (ref. 38) were compiled or used for the smoothing equation. The fitting equation used was:

$$\log_{10} x_2 = 1.8214 - 1164.63/(T/K)$$

The equation represents the combined data with a standard deviation of  $5.8 \times 10^{-2}$  in the 253 to 313 K temperature range.

The recommended mass per cent solubility values at 5 K intervals for water (2) in dichloromethane (1) are presented in the Table 2.

The linear relation between the solubility expressed as  $\log_{10} x_2$  versus the reciprocal of absolute temperature behavior is illustrated in Figure 2. The straight line behavior of such plots is characteristic of the solubility of water in halogenated hydrocarbons, as discussed in the Preface.

(continued next page)

### COMPONENTS:

(1) Dichloromethane; CH<sub>2</sub>Cl<sub>2</sub>; [75-09-2]

**EVALUATOR:** 

A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K.

(2) Water;  $H_2O$ ; [7732-18-5]

March 1993.

Table 2. Recommended Solubility of Water (2) in Dichloromethane (1)

# CRITICAL EVALUATION: (continued)

Ter	nperature	Solub	ility
°C	ĸ	100 w <sub>2</sub>	$10^3 x_2$
-20	253.15	0.0353	1.663
-15	258.15	0.0434	2.042
-10	263.15	0.0522	2.487
-5	268.15	0.0640	3.008
0	273.15	0.0768	3.612
5	278.15	0.0917	4.308
10	283.15	0.109	5.109
15	288.15	0.128	6.021
20	293.15	0.151	7.057
25	298.15	0.176	8.226
30	303.15	0.204	9.541
35	308.15	0.236	11.015
40	313.15	0.271	12.656



Figure 2. Solubility of Water (2) in Dichloromethane (1).

(continued next page)

154	4		
C	OMPONENTS:	EVALUATOR:	
(1)	) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	A. L. Horvath, Imperial Chemical Industries	
(2	) Water; H <sub>2</sub> O; [7732-18-5]	March 1993.	
CI	RITICAL EVALUATION: (continued)		
RI	EFERENCES:		
1. I.	. Salkowski, E. Biochem. Z. <u>1920</u> , 107, 191.		
2	. Booth, H. S.; Everson, H. E. Ind. Eng. Chem.	1 <u>948,</u> <i>40</i> , 1491.	
3.	. Alexandrova, M. V.; Sadovnikova, L. V.; Martir <u>1972</u> , 14, 146.	nov, W. W. Sb. Nauch. Tr., Ivanov. Energ. Inst.	
4.	<ul> <li>Sadovnikova, L. V.; Komarova, V. P.; Alexandre <u>1972</u>, 14, 205.</li> </ul>	ova, M. V. Sb. Nauch. Tr., Ivanov. Energ. Inst.	
5.	5. Sadovnikova, L. V.; Komarova, V. P.; Alexandrova, M. V.; Serafimov, L. A. Izv. Vyssh. Ucheb. Zaved. Khim. Khim. Tekhnol. <u>1972</u> , 15, 1891.		
6.	6. Sabinin, V. E.; Kiya-Oglu, N. V.; Gorichnina, V. P. J. Appl. Chem. USSR. <u>1970</u> , 43, 1788.		
7.	7. Svetlanov, E. B.; Velichko, S. M.; Levinskii, M. I.; Treger, Yu. A.; Flid, R. M. Russ. J. Phys. Chem. <u>1971</u> , 45, 488.		
8.	8. Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. KhimTekhnol. Inst. 1973, 183.		
9.	<ol> <li>Howe, G. B.; Mullins, M. E.; Rogers, T. N. AFESC Tyndall Air Force Base, Report ESL-TR-86-66, Vol. I, Florida, Sept. <u>1987</u>, 86 pp. (AD-A188 571).</li> </ol>		
10	10. Rex, A. Z. Phys. Chem. <u>1906</u> , 55, 355.		
11	11. van Arkel, A. E.; Vles, S. E. Recl. Trav. Chim. Pays-Bas 1936, 55, 407.		
12	12. Bakowski, S.; Treszczanowicz, E. Przemysl Chemiczny 1937, 21, 204.		
13	13. Niini, A. Suomen Kemistilehti 1938, 11A, 19.		
14	14. McGovern, E. W. Ind. Eng. Chem. <u>1943</u> , 35, 1230.		
15.	15. Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1952, 56, 480.		
16.	16. Kudryavtseva, G. I.; Krutikova, A. D. J. Appl. Chem. USSR. <u>1953</u> , 26, 1129.		
17	17. Maretic, M.; Sirocic, V. Nafta (Zagreb) 1962, 13, 126.		
18.	<ol> <li>du Pont de Nemours &amp; Company, Solubility Relationship of the Freon Fluorocarbon Compounds, Techn. Bull. <u>B-7</u>, Wilmington, Del., <u>1966</u>, 16 pp.</li> </ol>		
19.	19. Karger, B. L.; Chatterjee, A. K.; King, J. W. Techn. Rept. No. 3, Dept. of Chem., Norheast Univ., Boston, Mass., <u>May 10</u> , <u>1971</u> .		
20.	. Antropov, L. I.; Populyai, V. E.; Simonov, V. D 46(2), 311-312. (VINITI No. 3739-71).	.; Shamsutdinov, T. M. Russ. J. Phys. Chem. <u>1972</u> ,	
		(continued next page)	

		155	
	COMPONENTS: (1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2] (2) Water; H <sub>2</sub> O; [7732-18-5]	EVALUATOR: A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K. March 1993.	
	CRITICAL EVALUATION: (continued)		
	REFERENCES:		
)	21. McConnell, G.; Ferguson, D. M.; Pearson, C. R	. Endeavoir <u>1975</u> , 34, 13.	
- /	22. Pearson, C. R.; McConnell, G. Proc. Roy. Soc.	B, <u>1975</u> , <i>189</i> , 305.	
2	23. Archer, W. L.; Stevens, V. L. Ind. Eng. Chem.	Prod. Res. Dev. <u>1977</u> , 16, 319.	
J'.	24. Sato, A.; Nakijima, T. Arch. Environ. Health 19	<u>79,</u> <i>34</i> , 69.	
15	25. Coca, J.; Diaz, R. M.; Pazos, C. Fluid Phase Ed	quilibr. <u>1980,</u> 4, 125.	
27'	26. Hutchinson, T. C.; Hellebust, J. A.; Tam, D. et al. Hydrocarbons and Halogenated Hydrocarbons in the Aquatic Environment, Plenum Press, New York, <u>1980</u> , p. 577-586.		
o15	27. Leighton, D. T.; Calo, J. M. J. Chem. Eng. Data 1981, 26, 382.		
515	<ol> <li>Lincoff, A. H.; Gossett, J. M. in Gas Transfer at Water Surfaces by W. Brutsaert and G. H. Jirka, Eds., D. Reidel Publ. Co., Dordrecht, <u>1984</u>, p. 17-25.</li> </ol>		
۶ <b>۱</b> ۵	29. Gossett, J. M. Environ. Sci. Techn. <u>1987</u> , 21, 202.		
コフフ	<ol> <li>Warner, H. P.; Cohen, J. M.; Ireland, J. C. Determination of Henry's Law Constants of Selected Priority Pollutants, U. S. EPA Techn. Rept., PB87-212684, Cincinnati, OH., July 1987.</li> </ol>		
379	<ol> <li>Vogel, A. I., rev. by Furniss, B. S. et al. Vogel's Textbook of Practical Organic Chemistry, 5<sup>th</sup> ed., Longman, London, <u>1989</u>, p. 1442.</li> </ol>		
510	32. Wright, D. A.; Sandler, S. I.; DeVoll, D. Environ. Sci. Technol. 1992, 26, 1828.		
212	33. Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. KhimTekhnol. Inst. 1974, 55.		
	34. Staverman, A. J. Recl. Trav. Chim. Pays-Bas 1941, 60, 836.		
່	35. Davies, W.; Jagger, J. B.; Whalley, H. K. J. Soc. Chem. Ind. (London) 1949, 68, 26.		
<i>``</i>	36. Lees, F. P.; Sarram, P. J. Chem. Eng. Data <u>1971</u> , 16, 41.		
,,,)	37. Wu, X. Huaxue Shiji <u>1981</u> , 221.		
>7"	38. Ohtsuka, K.; Kazama, K. Sen'i Seihin Shohi Kag	aku Kaishi <u>1982,</u> 22, 197.	

156		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Rex, A.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Z. Phys. Chem. <u>1906</u> , 55, 355-70.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 100 $g_{1}/g_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (\text{compiler}) & (\text{compiler}) \end{array}$	
0 2.363 10 2.122 20 2.000 30 1.969	2.304.972.084.481.964.221.934.16	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The dichloromethane in water solubility was determined in specially designed flasks with cali- brated capillary cylinders. After samples were equi- librated in a thermostatic bath, their volumes and weights were determined in order to calculate the	<ol> <li>Kahlbaum, redistilled and washed before use, b.p. = 41°C.</li> <li>Distilled.</li> </ol>	
solubilities.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Salkowski, E.	
(2) Water; $H_2O$ ; [7732-18-5]	Biochem. Z. <u>1920</u> , 107, 191-201.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/kg m^{-3}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
20 12.24	1.211 2.59	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A volumetric method similar to that intro- duced by Alexejew was used. Dichloromethane was gradually added to water from a pipet under constant oritotion. The appearance of clouding was the indi-	<ol> <li>Kahlbaum, used as received.</li> <li>Distilled (compiler).</li> </ol>	
cation of saturation.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: ± 2 K (compiler).	

	157	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	van Arkel, A. E.; Vles, S. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 10 <sup>2</sup> $x_{i}$	$m_i/\text{mol kg}^{-1}$ 100 $w_i$	
30.0 0.416	0.232 1.931	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; $CH_2Cl_2$ ; [75-09-2]	Bakowski, S.; Treszczanowicz, E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Przemysł Chemiczny <u>1937</u> , 21, 204-6.	
VARIABLES:	PREPARED BY:	
T/K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 100 $w_{I}$ 10	$x_1 = 100 w_2 = 10^3 x_2$	
20 2.0 4.32 0.5 17.5		
AUXILIARY IN	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The water content of a liquid mixture was determined in a specially constructed apparatus which consisted of a 250 cm <sup>3</sup> flask, a Vigreux type column, two water condensers, a U-shaped measuring tube, an	<ol> <li>Merck reagent, dried by means of CaCl<sub>2</sub> before use.</li> <li>Distilled (compiler).</li> </ol>	
electric heater, and a thermometer. The volume of water was determined in a calibrated tube with an	ESTIMATED ERRORS:	
accuracy of 0.01 cm <sup>2</sup> .		

158		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Niini, A.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Suomen Kemistilehti <u>1938</u> , 11A, 19-20.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
<i>t/°</i> C 100 <i>w<sub>1</sub></i> 10 (con	$x_1^3 x_1 = 100 w_2 = 10^2 x_2$ npiler) (compiler)	
20 1.629 3.	50 0.239 1.12	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The changes resulting from the addition of dichloromethane to water were determined by the measurement of both the density and refractive index of the saturated solutions. A conventional refractom-	<ol> <li>Kahlbaum, dried over P<sub>2</sub>O<sub>5</sub> and redistilled before use.</li> <li>Distilled.</li> </ol>	
eter and a dilatometer were used for the measure- ments.	ESTIMATED ERRORS:	
	Solubility: $\pm 12$ %.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Staverman, A. J.	
(2) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Recl. Trav. Chim. Pays-Bas <u>1941</u> , 60, 836-41.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 273 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}\mathbf{C}$ 100 $w_{i}$	$\begin{array}{ccc} 10^{5} x_{i} & 100 \ w_{i}M_{i}^{-i}/\text{mol g}^{-i} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
$\begin{array}{cccc} 0 & 8.49 \times 10^{-2} \\ 25 & 1.67 \times 10^{-1} \\ 30 & 1.96 \times 10^{-1} \end{array}$	$\begin{array}{cccc} 400 & 4.71 \times 10^{-3} \\ 788 & 9.27 \times 10^{-3} \\ 925 & 10.90 \times 10^{-3} \end{array}$	
MEINOD/APPAKATUS/PKOCEDURE:	SOURCE AND FURITY OF MATERIALS:	
about 12 hours in a paraffin thermostat bath. The water content of the organic phase was determined using the Karl Fischer titration method. All	(2) Source and purity not specified.	
measurements were carried out in duplicate and the average of the two measurements was reported. A	ESTIMATED ERRORS:	
full description of the method is given in a thesis (ref. 1).	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
	REFERENCES:	
	(1) Staverman, A. J. <i>Ph. D. Thesis</i> , Univ. of Leiden, Leiden, Belgium, <u>1938</u> .	
25 1.67 × 10 <sup>-1</sup> 30 1.96 × 10 <sup>-1</sup> AUXILIARY I METHOD/APPARATUS/PROCEDURE: Dichloromethane was shaken with water for about 12 hours in a paraffin thermostat bath. The water content of the organic phase was determined using the Karl Fischer titration method. All measurements were carried out in duplicate and the average of the two measurements was reported. A full description of the method is given in a thesis (ref. 1).	<ul> <li>925 927 × 10<sup>-3</sup> 925 10.90 × 10<sup>-3</sup></li> <li>NFORMATION</li> <li>SOURCE AND PURITY OF MATERIALS: <ul> <li>(1) Distilled.</li> <li>(2) Source and purity not specified.</li> </ul> </li> <li>ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).</li> <li>REFERENCES: <ul> <li>(1) Staverman, A. J. Ph. D. Thesis, Univ. of Leiden, Leiden, Belgium, <u>1938</u>.</li> </ul> </li> </ul>	

.

	159	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; $CH_2Cl_2$ ; [75-09-2]	McGovern, E. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ind. Eng. Chem. <u>1943</u> , 35, 1230-9.	
VARIABLES:	PREPARED BY:	
T/K = 263 - 308	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub> 10 (con	$x_1 = 100 w_2 = 10^3 x_2$ npiler) (compiler)	
-10 0 2.00 4. 10 1.62 3. 20 1.40 3. 25 1.32 2. 30 1.29 2. 35 1.27 2. Solubility data as a function of temperative tabulated data point at 25°C.	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ol> <li>Source not given, commercial grade.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 1$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Booth, H. S.; Everson, H. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ind. Eng. Chem. <u>1948</u> , 40, 1491-3.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 100 $V_{1}/V_{2}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
25.0 2.60	3.32 7.23	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Equilibrium was ensured through repeated shaking and centrifuging of a dichloromethane and water mixture in a stoppered Goetz tube and re- turning the complete temperature under	<ol> <li>Commercial reagent, C. P. grade, used as received.</li> <li>Distilled.</li> </ol>	
bath. The difference between the total amount of dichloromethane which was added and the amount re-	ESTIMATED ERRORS:	
maining in excess was taken as the amount of di- chloromethane dissolved in the known volume of water. The determination of the excess amount of dichloromethane added is described by Handlick (act	Solubility: Not specified. Temperature: ± 1 K (compiler).	
1).	REFERENCES:	
	(1) Hanslick, R. S. Ph. D. Thesis, Columbia University, <u>1935</u> .	

160			
COMPONENTS:		ORIGINAL MEASU	JREMENTS:
(1) Water; H <sub>2</sub> O; [7732-18-5]		Davies, W.; Jagger, J. B.; Whalley, H. K.	
(2) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]		J. Soc. Chem. Ind.	(London) <u>1949</u> , 68, 26-31.
VARIABLES:		PREPARED BY:	
T/K = 253 - 313		A. L. Horvath	
EXPERIMENTAL VALUES:		-	
t/°C 10	00 w <sub>i</sub>	$10^3 x_i$ (compiler)	$\begin{array}{c} 100 \ w_{I}M_{I}^{-1}/\text{mol g}^{-1} \\ (\text{compiler}) \end{array}$
-20 0. -10 0. 0 0. 10 1. 20 1. 30 1. 40 2.	$\begin{array}{r} 3 \times 10^{-1} \\ 4 \times 10^{-1} \\ 7 \times 10^{-1} \\ 0 \times 10^{-1} \\ 4 \times 10^{-1} \\ 9 \times 10^{-1} \\ 6 \times 10^{-1} \end{array}$	1.4 1.9 3.3 4.7 6.6 8.9 12.1	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
	AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDUR	RE:	SOURCE AND PUR	RITY OF MATERIALS:
A mixture of water and dichloro agitated in a flask until equilibrium wa The apparatus was assembled in a water bath The equilibrium was established	omethane was as established. er thermostat after 30	<ol> <li>Distilled.</li> <li>Commercial q given.</li> </ol>	uality, source and purity not
minutes. The pipette used for withdraw	ving samples	ESTIMATED ERRO	DRS:
at the tip to prevent ice particles from being with- drawn at low temperatures. The organic layer was titrated with Karl Fischer reagent.		Solubility: Temperature:	$\pm$ 5 %. $\pm$ 0.5 K (compiler).
COMPONENTS:		ORIGINAL MEASU	IREMENTS:
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]		Donahue, D. J.; Ba	rtell, F. E.
(2) Water; H <sub>2</sub> O; [7732-18-5]		J. Phys. Chem. <u>195</u>	<u>2</u> , <i>56</i> , 480-4.
VARIABLES:		PREPARED BY:	
T/K = 298		A. L. Horvath	
EXPERIMENTAL VALUES:			
t/°C 10 <sup>3</sup> x	(com	$0 w_1   10^2 x_2$ piler)	100 w <sub>2</sub> (compiler)
25 4.2	1.9	950 1.13	0.242
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDUR	E:	SOURCE AND PURITY OF MATERIALS:	
A mixture of dichloromethane and water was placed in a glass stoppered flask and was shaken intermittently for at least three days in a water bath held at constant temperature. The organic phase was analyzed for water content using the Karl Fischer titration methods. The concentration of dichloro- methane in the aqueous phase was analyzed interfer- ometrically.		<ol> <li>Reagent grade distillation bef</li> <li>Purified by dis</li> </ol>	, purified by fractional ore use. stillation.
		ESTIMATED ERRORS:	
		Solubility: Not specified. Temperature: $\pm 0.1$ K.	

	161	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Kudryavtseva, G. I.; Krutikova, A. D.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Appl. Chem. USSR. <u>1953</u> , 26, 1129-33.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 293	Z. Maczynska	
EXPERIMENTAL VALUES:	,	
t/°C 100 w <sub>1</sub> 10 (con	$x_1 x_2$ 100 $w_2$ 10 <sup>2</sup> $x_2$ (compiler) (compiler)	
20 1.89 4.	07 1.92 8.45	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The mutual solubility between dichloromethane and water was determined by titration from a micro- buret until a turbidity appeared. The end point of the	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
ed with an accuracy of 0.3 - 0.5 %. The titration of the solutions was done in a thermostat bath at $20^{\circ}$	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Maretic, M.; Sirocic, V.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Nafta (Zagreb) <u>1962</u> , 13, 126-31.	
VARIABLES:	PREPARED BY:	
T/K = 298 - 346 P/mmHg = 750	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C 100 <i>w<sub>1</sub></i> 10 (corr	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
25 1.4 3.	00 0.2 9.36	
t/°C P/mmHg	$10^2 x_i$ 100 $w_i$ (compiler)	
38.3 750	0.39 1.81	
44.0 750 58.0 750	0.19 0.89 0.15 0.70	
73.0 750	0.10 0.47	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility was determined in a 50 cm <sup>3</sup> flask containing a weighed quantity of water. Di- chloromethane was added dropwise to the water until the first excess drop created turbidity. A slight	<ol> <li>Laboratory sample dried with CaCl<sub>2</sub> and distilled. The middle fraction was used.</li> <li>Distilled.</li> </ol>	
turoidity immediately indicated the saturation point.	ESTIMATED ERRORS:	
	Solubility: $\pm 0.2$ % by mass. Temperature: $\pm 2.0$ K.	
162		
---	--	--
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	du Pont de Nemours & Company,	
(2) Water; $H_2O$ ; [7732-18-5]	Solubility Relationship of the Freon Fluorocarbon Compounds, Tech. Bull. B-7, Wilmington, Del., 1966, 16 pp.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	- ·	
<i>t/°</i> C 100 <i>w<sub>1</sub></i>	$\begin{array}{ccc} 10^{3} x_{i} & 100 \ w_{i} M_{i}^{-1} / \text{mol } g^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
25 1.32	2.83 $1.55 \times 10^{-2}$	
AUXILIAR	Y INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility data were taken from a second ary source (ref. 1). Details and the original report a not available.	- (1) Source and purity not given. re (2) Distilled (compiler).	
	ESTIMATED ERRORS:	
	Solubility:Not specified.Temperature: $\pm 1 \text{ K}$ (compiler).	
	REFERENCES:	
	<ol> <li>Sanders, P. A. Handbook of Aerosol Technology, 2<sup>nd</sup> ed., Van Nostrand Reinhold Co., New York, <u>1979</u>, p. 42.</li> </ol>	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Sabinin, V. E.; Kiya-Oglu, N. V.;	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Gorichnina, V. P.	
	J. Appl. Chem. USSR. <u>1970</u> , 43, 1788-90.	
VARIABLES:	PREPARED BY:	
T/K = 293 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 100 w <sub>1</sub> (c	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
20 0.6 30 2.6	1.28         0.15         7.03           6.63         0.19         8.89	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The saturation of dichloromethane and water mixtures occurred in a flask fitted with a thermostat jacket and a powerful stirrer. After a stirring time of 10 - 20 minutes, samples were withdrawn from both the organic and aqueous layers and the compositions were determined from their refractivity indices.	<ul> <li>(1) Source not given. Purified by repeated distillation; density = 1.3252 g/cm<sup>3</sup> and refractive index = 1.4233 at 20 °C.</li> <li>(2) Distilled.</li> </ul>	
	Solubility Not specified	
	Temperature: $\pm 0.1$ K.	

•

			·	163
COMPONENTS:			ORIGINAL MEAS	UREMENTS:
(1) Water; $H_2O$ ; [7732-18-5]		Lees, F. P.; Sarran	m, P.	
(2) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]		J. Chem. Eng. Dai	na <u>1971</u> , <i>16</i> , 41-4.	
VARIABLES:		PREPARED BY:		
<i>T/</i> K = 298			A. L. Horvath	
EXPERIMENTAL V	ALUES:		l	
	t/℃	100 w <sub>i</sub>	$10^3 x_i$ (compiler)	$\frac{100 \ w_i M_i \ '/mol \ g^1}{(compiler)}$
	25	0.13	6.10	$7.2 \times 10^{3}$
		AUXILIARY I	NFORMATION	
METHOD/APPARAT	US/PROCE	DURE:	SOURCE AND PU	RITY OF MATERIALS:
An apparatus c chloromethane and wa bath. The stirring of t by using a glass rod r	ontaining the ater was plac the mixture v otated by a l	e mixture of di- ed in a thermostat vas accomplished horseshoe magnet	<ol> <li>Distilled.</li> <li>Source not g before use.</li> </ol>	iven. Analytical grade, degassed
for about 48 hours. T organic phase was de	the concentratermined by t	tion of water in the the Karl Fischer	ESTIMATED ERR	ORS:
titration method.			Solubility: Temperature:	± 2 %. ± 0.1 K.
COMPONENTS:			ORIGINAL MEAS	UREMENTS:
(1) Dichloromethane;	$CH_2Cl_2; [75]$	09-2 <b>]</b>	I.; Treger, Yu. A	A.; Flid, R. M.
(2) water; $H_2O$ ; [77]	02-10-JJ		Russ. J. Phys. Chem. <u>1971</u> , 45, 488-90.	
VARIABLES:			PREPARED BY:	
T/K = 288 - 333			A. L. Horvath	
EXPERIMENTAL V	ALUES:		L	
	t/°C	1000 g <sub>1</sub> /g <sub>2</sub>	100 <i>w<sub>i</sub></i> (compiler)	$\begin{array}{c} 10^3 x_{i} \\ \text{(compiler)} \end{array}$
	15	25.00 15.60	2.44	5.28
	45 60	8.80 5.30	0.872 0.527	1.86 1.12
•				
		AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PU	RITY OF MATERIALS:	
A dynamic method was used for the solubility determination. It consisted of the saturation of nitrogen with dichloromethane vapor which was then bubbled through a thermostatically controlled flask filled with a definite volume of water. After 2 - 3 hours, the solution became saturated and samples were taken periodically (ref. 1). The concentration of dichloromethane in water was determined by gas- liquid chromatography. On average, 3 - 5 measure- ments were taken at each temperature.		<ol> <li>Source and p</li> <li>Distilled (corr</li> </ol>	ourity not given. mpiler).	
		ESTIMATED ERR	ORS:	
		Solubility: Temperature:	± 15 %. ± 0.05 K.	
			_	
		<b>REFERENCES:</b>		
		(1) Treger, Yu. Russ. J. Phy.	A.; Flid, R. M.; Spektor, S. S. s. Chem. <u>1964</u> , 38, 253.	

164		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; $CH_2Cl_2$ ; [75-09-2]	Karger, B. L.; Chatterjee, A. K.; King, J. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Tech. Rept. No: 3, Dept. of Chemistry, North- eastern Univ., Boston, Mass., May 10, 1971.	
VARIABLES:	PREPARED BY:	
T/K = 286	A. L. Horvath	
EXPERIMENTAL VALUES: t/°C Partition coefficient <sup>1</sup> , K <sub>L</sub> /dimensionless 12.5 19.2 <sup>1</sup> Gas-liquid chromatographic parameter	$\begin{array}{ccc} 100 & w_i & 10^3 & x_i \\ (compiler) & (compiler) \\ 2.31 & 4.99 \\ er, from instrument calibration. \end{array}$	
AUXILIARY	INFORMATION	
	SOURCE AND PURITY OF MATERIALS	
Gas-liquid chromatography was used for the determination of the partition of dichloromethane wit thin layers of water coated on Porasil D. Helium carrier gas was presaturated with water at the column temperature. The eluents were detected with an F &	<ul> <li>(1) J. T. Baker Chemical Co., reagent grade, used as received.</li> <li>(2) Distilled (compiler).</li> </ul>	
M than ionization detector. The partition coefficient was obtained from the slope of the straight line plot of $V_N/A_L$ versus $V_L/A_L$ .	ESTIMATED ERRORS: Solubility: Not specified. Temperature: ± 0.5 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Alexandrova, M. V.; Sadovnikova, L. V.;	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Martinov, w. w. Sb. Nauchn. Tr., Ivanov. Energ. Inst. <u>1972</u> , 14, 146-54.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	Z. Maczynska	
EXPERIMENTAL VALUES:		
<i>t/</i> °C 100 w <sub>1</sub> 1 (co	$\begin{array}{ccccc} 0^3 x_1 & 100 w_2 & 10^3 x_2 \\ mpiler) & (compiler) \end{array}$	
20 2.76	i.98 0.10 4.70	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The titration method was used. Solubility data were also reported for the ternary dichloromethane-water-acetic acid system.	<ol> <li>Source not given. C. P. grade, used as received; n<sub>D</sub> = 1.4242 at 20 °C.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.1$ K.	

	165	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Antropov, L. I.; Populyai, V. E.; Simonov, V. D.; Shamsutdinov, T. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Russ. J. Phys. Chem. <u>1972</u> , 46, 311-2 (VINITI No: 3739-71).	
VARIABLES:	PREPARED BY:	
T/K = 291 - 299	A. L. Horvath	
EXPERIMENTAL VALUES:	1	
t/°C 100 w <sub>1</sub> 10 (cor	$x_1 = 100 w_2$ $10^3 x_2$ npiler) (compiler)	
18 - 20 2.0 4	0.14 6.57	
22 - 26 -	- 0.16 7.50 - 0.18 8.43	
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Alexejev's synthetic solubility determination method was used (ref. 1). A fixed weight of di-	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
chloromethane and water was sealed in a tube and the mixture was subjected to gradually increasing	ESTIMATED ERRORS:	
temperature with constant agitation. The appearance of opalescence or clouding was the indication of the saturation temperature. The temperature was then allowed to fall and an observation made, while the	Solubility: Not specified. Temperature: ± 0.2 K (compiler).	
tube was constantly agitated, of the temperature of the first appearance of opalescence. The observation	REFERENCES:	
was repeated several times.	(1) Alexejew, W. Ann. Phys. Chem. <u>1886</u> , 28, 305.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Sadovnikova, L. V.; Komarova, V. P.;	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Alexandrova, M. V. Sb. Nauchn. Tr., Ivanov. Energ. Inst. <u>1972</u> , 14,	
	205-9.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 311	Z. Maczynska	
EXPERIMENTAL VALUES:		
<i>t/</i> °C 100 <i>w<sub>1</sub></i> 10 (cor	$x_1 = 100 w_2 = 10^3 x_2$ npiler) (compiler)	
38 5.26 11	$1.0 \times 10^{-2}$ 0.47	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The titration method was used. The solubility was determined at the normal boiling point tempera- ture. An ebuliometer was used to check the boiling	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
methane-water-acetic acid system were also reported.	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.2$ K (compiler).	

166		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Sadovnikova, L. V.; Komarova, V. P.; Alexandrova, M. V.; Serafimov, L. A	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Izv. Vyssh. Ucheb. Zaved., Khim. Khim. Tekhnol.	
	<u>1972</u> , <i>15</i> , 1891-4.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 293	Z. Maczynska	
EXPERIMENTAL VALUES:		
<i>t/</i> °C 100 w <sub>1</sub> 10 (con	$x_1 = 100 w_2 = 10^3 x_2$ npiler) (compiler)	
19.85 2.76 5.	98 0.1 5.0	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available. Solubility data for the ternary dichloromethane-water-isobutyric acid system were also reported.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm$ 0.2 K (compiler).	
COMPONENTS	ODICINAL MEASUDEMENTS.	
(1) Dichloromethane: CH-Cl.: 175-09-21	Prosvanov N N · Shalvgin V A · Zel'venskij	
(2) Water; $H_2O$ ; [7732-18-5]	Ya. D.	
	Tr. Mosk. KhimTekhnol. Inst. <u>1973</u> , 183-6.	
VARIABLES:	PREPARED BY:	
T/K = 298 - 373	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Distribution coefficient <sup>1</sup> , D <sub>L</sub> /dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^4 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25.1 3920 50.0 2480	0.0693 1.471 0.2696 5.731	
70.0 1650 90.0 1240	0.7419 15.83 1.6652 35.79	
96.0 1130 100 1080	2.1071 45.45 2.4173 52.27	
<sup>1</sup> Gas-liquid system analysis parameter,	from calibration measurements.	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The distribution coefficient was determined by distillation. An inert gas was used to take samples from the vapor phase at equilibrium. The concentra-	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
tion of dichloromethane in both liquid and vapor phases was determined by using <sup>36</sup> Cl labeled com-	ESTIMATED ERRORS:	
pound.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	

	167	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii,	
(2) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Tr. Mosk. KhimTekhnol. Inst. <u>1974</u> , 55-6.	
VARIABLES:	PREPARED BY:	
T/K = 280 - 310	A. L. Horvath	
EXPERIMENTAL VALUES:	I	
$\log_{10} \alpha = \frac{547.33}{T(K)} - 1.$	0416	
where $\alpha$ = Distribution Coe	efficient	
At the normal boiling point of CH <sub>2</sub> Cl <sub>2</sub> , 313.15	K, $\alpha = 50.0$ , the activity coefficient, $\gamma_1 = 67.0$ ,	
and the mole fraction of (1) at saturation, $x_i =$	$3.953 \times 10^{-3}$ (compiler).	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
After equilibration of a dichloromethane and water mixture, the water content of the organic phase	<ol> <li>Distilled (compiler).</li> <li>Source and purity not given.</li> </ol>	
water. The experimental method is described in great- er detail elsewhere (ref 1)	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	
	REFERENCES:	
	<ol> <li>Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. Khim. Tekhnol. Inst. <u>1973</u>, 100.</li> </ol>	
COMPONENTS	ORIGINAL MEASUREMENTS	
(1) Dichloromethane: CH <sub>2</sub> Cl <sub>2</sub> : [75-09-2]	McConnell, G.: Ferguson, D. M.: Pearson, C. R.	
(2) Water; $H_2O$ ; [7732-18-5]	Endeavour <u>1975</u> , 34, 13-8.	
VARIABLES:	PREPARED BY:	
<i>T/</i> K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 10 <sup>6</sup> $g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 13200	1.30 2.79	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility of dichloromethane in water was determined using gas-liquid chromatography (GLC). The instrument was equipped with an electron capture detector. Where possible, identification was confirm- ed by using a linked mass-spectrometer (MS).	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: ± 0.2 K (compiler).	
l		

168		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Pearson, C. R.; McConnell, G.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Proc. Roy. Soc. B. <u>1975</u> , 189, 305-32.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	·	
t/°C 10 <sup>6</sup> w <sub>i</sub>	$100 w_i$ $10^3 x_i$ (compiler) (compiler)	
25 13200	1.320 2.84	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Saturated solutions were prepared and placed in a constant temperature thermostat bath. Water samples were extracted with n-pentane and an aliguot	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
of the extract taken for analysis using a gas-liquid chromatograph. The gas chromatograph was fitted	ESTIMATED ERRORS:	
with a "Ni electron capture detector.	Solubility: Not specified.	
	Temperature: $\pm 0.5$ K (complet).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Archer, W. L.; Stevens, V. L.	
(2) WAter; H <sub>2</sub> O; [7732-18-5]	1&EC Prod. Res. Dev. <u>1977</u> , 16, 319-25.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	·	
t/°C 100 w <sub>1</sub> 10 (com	$x_1 = 100 w_2$ $10^3 x_2$	
25 2.0 4.	22 0.20 9.34	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available. Data were reported elsewhere (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERKORS:	
	Temperature: $\pm 1$ K (compiler).	
	REFERENCES:	
	<ol> <li>News Release from Dow Chemicals Co., U. S. A., dated <u>October 23</u>, <u>1975</u>.</li> </ol>	

	169
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Sato, A.; Nakijima, T.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Arch. Envir. Health <u>1979</u> , 34, 69-75.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 310	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C Partition coefficient <sup>1</sup> , K <sub>1</sub> /dimensionless	$\begin{array}{ccc} 100 \ w_{i} & 10^{3} \ x_{i} \\ (compiler) & (compiler) \end{array}$
37 7.2	2.2 4.75
<sup>1</sup> Gas-liquid chromatographic parameter,	, from instrument calibration.
AUXILIARY I	NFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Dichloromethane vapor was equilibrated in an airtight vial between water and the overlying air. When equilibrium was established, a portion of the	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>
equilibrated air in the vessel was withdrawn using an airtight syringe and was injected into a gas chro-	ESTIMATED ERRORS:
matogram was used to calculate the partition coef- ficient.	Solubility: $\pm$ 0.6 std. dev. Temperature: $\pm$ 0.5 K (compiler).
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Coca, J.; Diaz, R. M.; Pazos, C.
(2) Water; H <sub>2</sub> O; [7732-18-5]	Fluid Phase Eqilibr. <u>1980</u> , 4, 125-36.
VARIABLES:	PREPARED BY:
<i>T</i> /K = 298	A. L. Horvath
EXPERIMENTAL VALUES:	
<i>t/°</i> C 100 <i>w<sub>i</sub></i> 10 (com	$x_1 = 100 w_2 = 10^3 x_2$ piler) (compiler)
25 1.37 2.5	04 0.18 8.43
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The mutual solubility data were determined by the method described by Othmer <i>et al.</i> (ref. 1). A 10 $cm^3$ dichloromethane sample was added to water from a burst and particular duratil the solution became turbid	<ol> <li>Probus reagent, further purified by distillation in a heli-packing column.</li> <li>Distilled.</li> </ol>
The appearance of the turbidity indicated the forma- tion of a second phase. The solubilities were calcu-	ESTIMATED ERRORS:
lated by means of known densities and volumes.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).
	REFERENCES:
	(1) Othmer, D. F.; White, R. E.; Trueges, E. Ind. Eng. Chem. <u>1941</u> , 33, 1240.

170		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Hutchinson, T. C.; Hellebust, J. A.;	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Hydrocarbons and Halogenated Hydrocarbons in the Aquatic Environment, Plenum Press, New York, <u>1980</u> , p. 577-86.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
<i>t/</i> °C ρ <sub>1</sub> /kg m <sup>-3</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_j \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25 19.4	1.946 4.19	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
An excess of dichloromethane was added to water and stirred vigorously for 24 hours. After settling at 25°C for 48 hours, the solution was	<ol> <li>Aldrich Chemicals, highest grade, used as received.</li> <li>Double distilled.</li> </ol>	
extracted with cyclohexane. The samples were ana- lyzed using an Aminco-Bowman spectrophoto-	ESTIMATED ERRORS:	
fluorometer. A detailed description of the method is reported elsewhere (ref. 1).	Solubility: Not specified. Temperature: ± 0.2 K (compiler).	
	REFERENCES:	
	(1) Mackay, D.; Shiu, W. Y. J. Chem. Eng. Data <u>1977</u> , 22, 399.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Wu, X.	
(2) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Huaxue, Shiji <u>1981</u> , 221-4.	
VARIABLES:	PREPARED BY:	
T/K = 278 - 308	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ 10 <sup>6</sup> $w_{i}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
5 1313 10 1440	0.1313 6.16 0.1440 6.75	
15 1525 20 1609	0.1525 7.15 0.1609 7.54	
25 1779 30 1948	0.1779 8.33 0.1948 9.12	
35 2075	0.2075 9.71	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A known volume of water was added to dried dichloromethane to prepare a series of standard solutions for establishing calibration curves and obtaining the acturated colutions of each terms	<ol> <li>Distilled.</li> <li>Source not given, chromatographically pure, used as received.</li> </ol>	
The determination of the water content of dichloro- methone was made with an PM 250 NMP spectrom	ESTIMATED ERRORS:	
eter (radiation frequency of 250 MHz). The reported water peak amplitude was the mean of four readings taken at a particular temperature.	Solubility: ± 24 std. dev. Temperature: 1 - 2 K.	

•

	171	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Leighton, D. T.; Calo, J. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Chem. Eng. Data <u>1981</u> , 26, 382-5.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 275 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:	•	
$t/^{\circ}C$ Partition coefficient <sup>1</sup> , $K_L/dimensionless$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
$\begin{array}{ccccc} 1.9 & 61.4 \\ 13.5 & 111.5 \\ 15.7 & 121.5 \\ 17.1 & 141.6 \\ 22.0 & 157.1 \\ 24.9 & 161.9 \end{array}$	1.58713.4091.48863.1951.49043.1991.36512.9271.51443.2511.65693.561	
<sup>1</sup> Gas-liquid chromatographic parameter, from i	nstrument calibration.	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A 5 $\mu$ L dichloromethane sample was injected into a 2.3 liter equilibration cell containing distilled water. After the cell was shaken vigorously for about 5 minutes, the homogeneity of the liguid sample was	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
maintained with a magnetic stirrer. Compressed air was passed through the cell and the gas flow was measured with a soap film flowmeter. The dichloro- methane content was extracted and analyzed using a dual flame ionization detector gas chromatograph.	ESTIMATED ERRORS: Solubility: ± 4.8 %. Temperature: ± 0.5 K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]	Ohtsuka, K.; Kazama, K.	
(2) Dichloromethane; $CH_2Cl_2$ ; [75-09-2]	Sen'i Seihin Shohi Kagaku Kaishi <u>1982</u> , 22, 197-201.	
VARIABLES:	PREPARED BY:	
T/K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $g_{1}V_{2}^{-1}/kg m^{-3}$	$\begin{array}{ccc} 100 & w_i & & 10^3 & x_i \\ \text{(compiler)} & & \text{(compiler)} \end{array}$	
25 2.00	0.151 7.08	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Water was added gradually to 50 cm <sup>3</sup> dichloro- methane in a flask and then lowered into a thermostat bath. The flask was then shaken vigorously until the first cloud (turbidity) appeared. The work control of	<ol> <li>Distilled (compiler).</li> <li>Commercial JIS extra pure reagent, further purified by conventional methods before use.</li> </ol>	
the sample was determined by the Karl Fischer titra- tion method.	ESTIMATED ERRORS:	
	Solubility:Not specified.Temperature: $\pm$ 0.5 K (compiler).	

COMPONENTS: (1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2] (2) Water; H <sub>2</sub> O; [7732-18-5] VARIABLES:		ORIGINAL MEASU	REMENTS:					
		Lincoff, A. H.; Gossett, J. M. in <i>Gas Transfer at Water Surface</i> , by W. Brutsaert and G. H. Jirka, Eds., D. Reidel Publ. Co., Dordrecht, <u>1984</u> , p. 17-25. PREPARED BY:						
				<i>T</i> /K = 293			A. L. Horvath	
				EXPERIMENTAL	. VALUES	5:	1	
EPICS Method:								
	t/°C	Henry's law constant, <i>H</i> /m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{c} 100 \ w_{I} \\ (compiler) \end{array}$	$10^3 x_1$ (compiler)				
	20	2.25×10 <sup>-3</sup>	1.7786	3.826				
	loį	$g_e H = -\frac{4191}{T(K)} + 8$	. 200					
Batch Air Strippin	g Method: t/°C	Henry's law constant,	100 w,	$10^3 x_i$				
Batch Air Strippin	g Method: <i>t/</i> °C 20	Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup>	100 <i>w</i> , (compiler) 2.0314	10 <sup>3</sup> x, (compiler) 4.379				
Batch Air Strippin	g Method: t/°C 20 log	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$	100 <i>w</i> , (compiler) 2.0314 . 035	$10^{3} x_{i}$ (compiler) 4.379				
Batch Air Strippin	g Method: t/°C 20 log	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$	100 <i>w</i> , (compiler) 2.0314 . 035	10 <sup>3</sup> x, (compiler) 4.379				
Batch Air Strippin	g Method: t/°C 20 log	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$	100 w, (compiler) 2.0314 . 035	10 <sup>3</sup> x, (compiler) 4.379				
Batch Air Strippin	g Method: t/°C 20 log	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$ AUXILIARY II	100 w, (compiler) 2.0314 . 035 NFORMATION	$10^{3} x_{i}$ (compiler) 4.379				
Batch Air Strippin	g Method: t/°C 20 log	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$ AUXILIARY IN COCEDURE:	100 w, (compiler) 2.0314 . 035 . 035 NFORMATION SOURCE AND PURI	10 <sup>3</sup> x, (compiler) 4.379 ITY OF MATERIALS:				
Batch Air Strippin Batch Air Strippin METHOD/APPAF Henry's law EPICS and Batch librium Partitionin technique compare injection of headsp	g Method: t/°C 20 log RATUS/PR / constants Air Strippi g in Close id the GL pace sampl	Henry's law constant, $H/m^3$ atm mol <sup>-1</sup> 1.97×10 <sup>-3</sup> $g_e H = -\frac{4472}{T(K)} + 9$ AUXILIARY II COCEDURE: were determined by ng methods. The Equi- d Systems (EPICS) peak heights upon direct es.	100 w, (compiler) 2.0314 . 035 . 035 NFORMATION SOURCE AND PURI (1) Source and pur (2) Distilled.	10 <sup>3</sup> x, (compiler) 4.379 4.379				

•

	173	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Gossett, J. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Environ. Sci. Technol. <u>1987</u> , 21, 202-8.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 308	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's Law Constant, CV <sup>1</sup> H/m <sup>3</sup> atm mol <sup>-1</sup> %	$\begin{array}{ccc} 100 \ w_1 & 10^3 \ x_1 \\ (\text{compiler}) & (\text{compiler}) \end{array}$	
$\begin{array}{ccccccc} 9.6 & 1.15 \times 10^{-3} & 19.2 \\ 17.5 & 1.31 \times 10^{-3} & 19.9 \\ 24.8 & 2.19 \times 10^{-3} & 17.4 \\ 34.6 & 3.26 \times 10^{-3} & 2.37 \end{array}$	2.2513       4.862         2.7492       5.960         2.1422       4.842         2.2150       4.782	
<sup>1</sup> CV = coefficient of variation (= 1	00 S.D./mean).	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A modification of the EPICS (Equilibrium Partitioning in Closed Systems) procedure was used for measuring Henry's law constants. A precise quantity of dichloromethane was injected into serum bottles which contained distilled water. The bottles	<ol> <li>Burdick &amp; Jackson reagent, pesticide grade. Distilled in a glass set-up before use.</li> <li>Distilled.</li> </ol>	
were incubated for 18 - 24 hours at four desired temperatures in a reciprocating shaker bath. The headspace concentrations of the EPICS bottles were measured using a gas chromatograph which was equipped with a flame ionization detector. The mean of the coefficient of variation values was approxi- mately 4.3 %.	ESTIMATED ERRORS: Solubility: See above. Temperature: ± 0.1 K.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Warner, H. P.; Cohen, J. M.; Ireland, J. C.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Determination of Henry's Law Constants of Selected Priority Pollutants Cincinnati, OH., July 1987.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	·	
t/°C Henry's law constant, H/m <sup>3</sup> atm mol <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
24.85 $3.19 \times 10^{-3}$	1.599 3.436	
AUXILIARY II	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The original method and apparatus for the determination of Henry's law constant, as described by Mackay <i>et al.</i> (ref. 1), was used. The general	<ol> <li>Source not given. Purest quality available, used as received. Stated purity &gt; 99 %.</li> <li>Distilled and deionized.</li> </ol>	
procedure was to add an excess quantity of dichloro- methane to distilled deionized water, place the sample	ESTIMATED ERRORS:	
A portion of this solution was returned to the strip- ping vessel. Dichloromethane was stripped isother- mally from the solution at a known gas flow rate.	Solubility: $\pm 6\%$ std. dev. Temperature: $\pm 0.05$ K.	
The Henry's law constant was calculated from the log of the concentration versus time plot. The experi-	REFERENCES:	
replicates.	(1) Mackay, D.; Shiu, W. Y.; Sutherland, R. D. Environ. Sci. Technol. <u>1979</u> , 13, 333.	

174		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]	Howe, G. B.; Mullins, M. E.; Rogers, T. N.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	AFESC Tyndall Air Force Base, Report ESL-TR- 86-66, Vol. 1, Florida, <u>Sept. 1987</u> , 86 pp. (AD-A188 571).	
VARIABLES:	PREPARED BY:	
T/K = 283 - 303	A. L. Horvath	
EXPERIMENTAL VALUES:	·	
t/°C 10 <sup>6</sup> w,	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
10 11092 20 13508 30 11211	1.1092       2.3736         1.3508       2.8961         1.1211       2.3993	
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
250 cm <sup>3</sup> bottles were filled with distilled deionized water and sealed. Measured volumes of di- chloromethane were injected into the bottles through each bottle septum using a microliter syringe. Di-	<ol> <li>Probably a commercial reagent at least 99 % pure. Used as received.</li> <li>Distilled and deionized.</li> </ol>	
bility limit. The bottles were shaken for one hour	ESTIMATED ERRORS:	
with wrist-action shaker and allowed to equilibrate for about 3 weeks. Samples were than injected into a gas chromatograph equipped with a Carbopack col- umn and a FID detector. The GL response was com- pared with the calibration plot.	Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Dichloromethane; $CH_2Cl_2$ ; [75-09-2]	Vogel, A. I., rev. by Furniss, B. S.; et al.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Vogen's Textbook of Practical Organic Chemistry, 5 <sup>th</sup> ed., Longman, London, <u>1989</u> , p. 1442.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 298	A. L. Horvath	
EXPERIMENTAL VALUES:	-	
t/°C 100 w <sub>t</sub>	$\begin{array}{ccc} 10^3 x_i & 100 \ w_i M_i^{-1} / \text{mol g}^{-1} \\ \text{(compiler)} & (\text{compiler}) \end{array}$	
25.0 1.30	2.79 $1.53 \times 10^{-2}$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The solubility determination was carried out in small test tubes (e. g., $100 \times 12$ mm) to permit vigorous shaking of dichloromethane and water mixtures. During the experiment 0.20 cm <sup>3</sup> dichloromethane upp added is 2.0 cm <sup>3</sup> of upstress and shelled	<ol> <li>Commercial reagent, purified by washing with concentrated sulfuric acid before use.</li> <li>Distilled.</li> </ol>	
The amount of dichloromethane dissolved in water	ESTIMATED ERRORS:	
chromatographic column.	Solubility: Not specified. Temperature: ± 1 K (compiler).	

					175
COMPONENTS:			ORIGINAL MEASU	JREMENTS:	
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]		Wright, D. A.; Sandler, S. I.; DeVoll, D.			
(2) Water; H <sub>2</sub> O; [773	32-18-5]		Environ. Sci. Techr	nol. <u>1992</u> , 26, 1828-31.	
VARIABLES:			PREPARED BY:		
T/K = 283 - 303			A. L. Horvath		
EXPERIMENTAL V	ALUES:				
	t/⁰C	γ"	100 $w_i$ (compiler)	$10^3 x_1$ (compiler)	
	10 22 20 22 30 23	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$	1.977 1.849 1.858	4.26 3.98 4.00	
		AUXILIARY IN	NFORMATION		
METHOD/APPARAT	US/PROCED	URE:	SOURCE AND PUR	TTY OF MATERIALS:	
A differential st was used to measure t coefficient of dichloror containing degassed w	atic cell equil he infinite dil methane $(\gamma^{\infty})$	ibrium apparatus ution activity in water. Cells merged in a ther	(1) Source and pu (2) Distilled, filter	rity not given. red and deionized.	
containing degassed water were submerged in a ther- mostated water bath. Dichloromethane was injected into the mixture cells and a magnetic stirrer was turned on. The cells were allowed to equilibrate. The stirrer was then turned off and the differential pres- sure was recorded. The experiment was repeated at least three times at each temperature		ESTIMATED ERRO Solubility: Temperature:	DRS: See above. ± 0.05 K.		

176					
COMPONENTS:		ORIGINAL MEASUREMENTS:			
(1) Dichloromethane; CH <sub>2</sub> Cl <sub>2</sub> ; [75-09-2]		Svet	lanov, E. B.;	Velichko, S. M.; Levinskii, M.	
(2) Hydrogen chlor	ide; HCI; [7647-	01-0]	Rus	I Phys Ch	em 1971 AS A88-90
(3) Water; H <sub>2</sub> O; [7732-18-5]		Λμ5.	s. J. Phys. Ch	<i>eni</i> . <u>1771</u> , 43, 400-90.	
VARIABLES:			PRE	PARED BY:	
T/K = 288 - 333			A. 1	Horvath	
HCI concentration					
EXPERIMENTAL	VALUES:				
Solubility of CH <sub>2</sub> Cl <sub>2</sub>	2 in 10 % HCl so	olution:			
	t/°C	10 <sup>3</sup> w.		100 w.	$10^3 x_1$
			(	compiler)	(compiler)
	15 30	29.40 18.50		2.940 1.850	6.733 4.198
	45 60	12.50 6.00		1.250 0.600	2.822 1.347
Solubility of CH <sub>2</sub> Cl <sub>2</sub>	2 in 20 % HCl so	olution:			
	t/℃	$10^3 w_1$		100 w <sub>1</sub>	$10^3 x_1$
			(	compiler)	(compiler)
	30 45	24.50 12.00		2.450 1.200	5.909 2.862
	60	6.50		0.650	1.543
AUXILIARY INFORMATION					
METHOD/APPARATUS/PROCEDURE:		SOU	RCE AND PU	RITY OF MATERIALS:	
The dynamic method was used for the de- termination of the solubility. It consisted of the saturation of nitrogen with dichloromethane vapor which was then bubbled through a thermostatically controlled flask filled with a definite volume of solution. After 2 - 3 hours the solution became saturated and samples were taken periodically (ref. 1). The concentration of dichloromethane in the		(1)	Source and p	urity not given.	
		(3)	Distilled (con	mpiler).	
		FCTI		ORS	
		Solu		+ 15 %.	
solution was determ measurements were	ined by GLC. Of taken at each ter	n average, 3 - 5 nperature.	Tem	perature:	± 0.05 κ.
		-	REF	ERENCES:	
			(1)	Treger, Yu.	A.; Flid, R. M.; Spektor, S. S.
				Russ. J. Phy.	s. Chem. <u>1964</u> , 38, 253.

COMPONENTS:	EVALUATOR:
<ol> <li>(1) Diiodomethane; CH<sub>2</sub>I<sub>2</sub>; [75-11-6]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K. March 1993.

177

#### CRITICAL EVALUATION:

The diiodomethane (1) and water (2) binary system is discussed in two parts; part 1 is diiodomethane (1) in water (2) and part 2 is water (2) in diiodomethane (1).

**Part 1.** The solubility of diiodomethane (1) in water (2) has been studied by four groups of workers. The experimental data of Gross and Saylor (ref. 1) and Andrews and Keefer (ref. 2) are supported by the descriptions of the methods used for determining the reported data. Details are not available for methods, apparatus, and procedures used by van Arkel and Vles (ref. 3) and O'Connell (ref. 4) for obtaining the solubility data.

The three solubility values, see (refs. 1, 3, and 4), between 293.15 and 303.15 K compare favorably with the selected value, whereas the datum of Andrews and Keefer (ref. 2) at 298.15 K is low and classified as doubtful. More accurate solubility determinations are required, meanwhile, the tentative mas per cent solubility of diiodomethane (1) in water (2) is 0.124 at 303.15 K.

**Part 2.** The only datum available for the solubility of water (2) in diiodomethane (1) is that of Hutchison and Lyon (ref. 5) and so no Critical Evaluation is possible. The interested reader is referred to the relevant Compilation Sheet for the experimental solubility.

#### **REFERENCES:**

- 1. Gross, P. M; Saylor, J. H. J. Am. Chem. Soc. 1931, 53, 1744.
- 2. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. 1951, 73, 5733.
- 3. van Arkel, A. E.; Vles, S. E. Recl. Trav. Chim. Pays-Bas 1936, 55, 407.
- 4. O'Connell, W. L. Trans. Am. Inst. Mech. Eng. 1963, 226, 126.
- 5. Hutchison, C. A.; Lyon, A. M. Columbia University Report A-745, July 1, 1943.

) ) ) ) ) )

25 X

178			
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Diiodomethane; $CH_2I_2$ ; [75-11-6]	Gross, P. M.; Saylor, J. H.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Am. Chem. Soc. <u>1931</u> , 53, 1744-51.		
VARIABLES:	PREPARED BY:		
<i>T/</i> K = 303	A. L. Horvath		
EXPERIMENTAL VALUES:	•		
$t/^{\circ}$ C 1000 $g_1/g_2$	$\begin{array}{ccc} 100 \ w_i & 10^5 \ x_i \\ (compiler) & (compiler) \end{array}$		
30 1.24	0.124 8.35		
AUXILIARY I	NFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
An excess of diiodomethane in 500 g water was shaken for 12 hours in a thermostat bath. Samples were then withdrawn and read against water	<ol> <li>Eastman Kodak Co., shaken with Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> solution. washed with distilled water, dried, and fractionally frozen twice before use.</li> </ol>		
in an interferometer made by Zeiss (ref. I). A detailed description of the complete method and procedure is given in 2 Ph. D. thesis (ref. 2)	(2) Distilled.		
procedure is given in a r n. D. mesis (iei. 2).	Solubility: $\pm 2.0$ %.		
	Temperáture: $\pm$ 0.02 K.		
	REFERENCES:		
	(1) Gross, P. M. J. Am. Chem. Soc. <u>1929</u> , 51, 2362.		
	(2) Saylor, P. M. <i>Ph. D. Thesis</i> , Duke University, Durham, <u>1930</u> .		
COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Diiodomethane; CH <sub>2</sub> I <sub>2</sub> ; [75-11-6]	van Arkel, A. E.; Vles, S. E.		
(2) Water; H <sub>2</sub> O; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.		
VARIABLES:	PREPARED BY:		
<i>T</i> /K = 303	A. L. Horvath		
EXPERIMENTAL VALUES:			
t/°C m <sub>i</sub> /mol kg <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$		
$30    4.6 \times 10^{-3}$	0.123 8.28		
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>		
	ESTIMATED ERRORS:		
	Solubility: Not specified.		
	$\mathbf{T} = \mathbf{T} \mathbf{U} \cdot \mathbf{T} \mathbf{V} \cdot \mathbf{T} \mathbf{V} \cdot \mathbf{T} \mathbf{U} \cdot \mathbf{U} \mathbf{U} \mathbf{U} \mathbf{U} \mathbf{U} \mathbf{U} \mathbf{U} \mathbf{U}$		

COMPONENTS:		ORIGINAL MEASUREMENTS:	
(1) Water; H <sub>2</sub> O; [7732-18-5]		Hutchison, C. A.; Lyon, A. M.	
(2) Diiodomethane; $CH_2I_2$ ; [75-11-6]		Columbia University Report A-745, July 1, 1943.	
VARIABLES:		PREPARED BY:	
<i>T</i> /K = 298		A. L. Horvath	
EXPERIMENTAL VALUES:	<u></u>		
t/°C	$100 \ w_1 M_1^{-1} / \text{mol g}^{-1}$	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
25	$1.207 \times 10^{-3}$	$2.174 \times 10^{-2}$ 3.222	
	AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROC	EDURE:	SOURCE AND PURITY OF MATERIALS:	
A 1 to 15 volume ratio n diiodomethane was introduced i flask and then the flask was low	nixture of water and nto an equilibration vered into a water	<ol> <li>Distilled.</li> <li>Source not given, purified and dried before use.</li> </ol>	
thermostat bath. The assembly cally for about 90 minutes at co	was shaken mechani- nstant temperature.	ESTIMATED ERRORS:	
termined by a modified Karl Fit The determination was done in The description was take	scher titration method. triplicate. n from a secondary	Solubility: $\pm 1.5 \times 10^{-5}$ avg. dev. Temperature: $\pm 0.05$ K.	
source (ref. 1). The original report is no longer available.		REFERENCES:	
		<ol> <li>Eidinoff, M. L.; Jorris, G. G.; Taylor, H. S.; Urey, H. C., eds. "Production of Heavy Wa- ter," McGraw-Hill, New York, <u>1955</u>, p. 129.</li> </ol>	
COMPONENTS		ORIGINAL MEASUREMENTS	
(1) Diiodomethane; CH <sub>2</sub> I <sub>2</sub> ; [75-	-11-6]	Andrews, L. J.; Keefer, R. M.	
(2) Water; H <sub>2</sub> O; [7732-18-5]		J. Am. Chem. Soc. <u>1951</u> , 73, 5733-6.	
VARIABLES:		PREPARED BY:	
<i>T</i> /K = 298		A. L. Horvath	
EXPERIMENTAL VALUES:		·	
t/°C	$c_i/\text{mol m}^{\cdot 3}$	$\begin{array}{ccc} 100 \ w_i & 10^s \ x_i \\ (compiler) & (compiler) \end{array}$	
25.0	3.11	$8.35 \times 10^2$ 5.62	
AUXILIARY INFORMATION			
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:	
Water was mixed with diiodomethanme in a glass-stoppered Erlenneyer flask which was then placed in a constant temperature bath and rotated for 30 hours. The organic halide content of the aqueous solution was extracted with n-hexane. The optical density of the extract was measured against a n- hexane blank using a Beckman spectrophotometer (ref. 1).		<ol> <li>Prepared from iodoform at the University of California; b. p. = 71.0 - 71.5 °C at 18 mmHg.</li> <li>Distillad (compiled)</li> </ol>	
		ESTIMATED ERRORS	
		Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
		REFERENCES:	
		(1) Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1949</u> , 71, 3644.	

ORIGINAL MEASUREMENTS:
O'Connell, W. L.
Trans. Am. Inst. Mech. Eng. <u>1963</u> , 226, 126-32.
PREPARED BY:
A. L. Horvath
I
$100 w_i$ $10^5 x_i$
0.124 8.34
NFORMATION
SOURCE AND PURITY OF MATERIALS:
<ol> <li>Dow Chemicals Co., used as received.</li> <li>Distilled (compiler).</li> </ol>
ESTIMATED ERRORS:
Solubility: Not specified. Temperature: $\pm$ 0.5 K (compiler).
ORIGINAL MEASUREMENTS:
Hutchison, C. A.; Lyon, A. M.
Columbia University Report A-745, July 1, 1943.
PREPARED BY:
A. L. Horvath
$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$
$2.04 \times 10^{-2}$ 2.721
NFORMATION
SOURCE AND PURITY OF MATERIALS:
<ol> <li>Source and purity not given.</li> <li>Source not given, purified and dried before use.</li> </ol>
ESTIMATED ERRORS:
Solubility: $\pm 0.3 \times 10^{-5}$ avg. dev. Temperature: $\pm 0.05$ K.
REFERENCES:
<ol> <li>Eidinoff, M. L.; Joris, G. G.; Taylor, H. S.; Urey, H. C., Eds., "Production of Heavy Wa- ter," McGraw-Hill, New York, <u>1955</u>, p. 129.</li> </ol>

COMPONENTS:	EVALUATOR:
<ol> <li>(1) Iodomethane; CH<sub>3</sub>I; [74-88-4]</li> <li>(2) Water; H<sub>2</sub>O; [7732-18-5]</li> </ol>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, U.K. March 1993.

#### CRITICAL EVALUATION:

The solubility of iodomethane (1) in water (2) has been studied by 8 groups of workers; whereas, the solubility of water (2) in iodomethane (1) has not been investigated yet.

The experimental data of Glew (ref. 1) covering the 273 to 323 K temperature range are significantly higher than the likely solubility. Furthermore, the continuous decrease in solubility with temperature between 273 and 323 is not consistent with the formation of a minimum shown by most similar binary systems. Therefore, these data are classed as doubtful and are rejected. The presence of a minimum in the solubility versus temperature behavior for the halogenated hydrocarbon-water systems is discussed in the Preface.

The solubility data of Balls (ref. 2) between 283 and 298 K show a continuous rise with temperature which is contrary to the trends shown by all other investigators and are therefore rejected. The single data point of Liss and Slater (ref. 3) at 293 K is lower than other results and is likewise rejected.

The remaining data of Rex (ref. 4), Fühner (ref. 5), van Arkel and Vles (ref. 6), Swain and Thornton (ref. 7), and Hunter-Smith *et al.* (ref. 8) are in reasonable agreement, although further studies are required before any values can be recommended. The tentative mass per cent solubilities of iodomethane (1) in water (2) over the 273 to 313 K temperature range were used to obtain the following equation:

Solubility  $[100 w_l] = 19.0036 - 0.11723 (T/K) + 1.95356 \times 10^4 (T/K)^2$ 

This regression equation was established using the combined data from (refs. 4 - 8), which yielded a standard deviation of  $4.3 \times 10^2$ .

The tentative mass per cent solubility values at 5 K intervals for iodomethane (1) in water (2) are presented in Table 1. The curve obtained from the smoothing equation is shown in Figure 1. It shows a distinct minimum at 300.0 K, as discussed in the Preface.

Table 1. Tentative Solubility of Iodomethane (1) in Water (2)

Tem	perature,	Solubil	ity,
°C	K	100 w <sub>1</sub>	$10^{3} x_{1}$
0	273.15	1.557	2.003
5	278.15	1.510	1.942
10	283.15	1.472	1.893
15	288.15	1.444	1.856
20	293.15	1.425	1.831
25	298.15	1.417	1.821
30	303.15	1.418	1.822
35	308.15	1.430	1.838
40	313.15	1.450	1.864

(continued next page)

181



- 29.5 2. Balls, P. W. Ph. D. Thesis, University of East Anglia, Norwich, U. K. July 1980, 375 pp.
- 591 3. Liss, P. S.; Slater, P. G. Nature 1974, 247, 181.
  - Rex, A. Z. Phys. Chem. 1906, 55, 355. 4.
  - 5. Fühner, H. Ber. <u>1924</u>, 57, 510.

2<sup>9°</sup>

, 4

ۍ. ۲

- van Arkel, A. E.; Vles, S. E. Recl. Trav. Chim. Pays-Bas 1936, 55, 407. 6.
- 7. Swain, C. G.; Thornton, E. R. J. Am. Chem. Soc. 1962, 84, 822.
- 8. Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S. Tellus, 1983, 35B, 170.

COMPONENTS:ORIGINAL MEASUREMENTS:(1) lodomethane; CH_3I; [74-88-4]Rex, A.(2) Water; H_2O; [7732-18-5]Z. Phys. Chem. 1906, 55, 355-70.VARIABLES:PREPARED BY: $T/K = 273 - 303$ A. L. HorvathEXPERIMENTAL VALUES: $100 g_1/g_2$ $t^{PC}$ $100 g_1/g_2$ $10 \cdot 1.565$ $1.540$ $10 \cdot 1.446$ $1.425$ $10 \cdot 1.446$ $1.425$ $10 \cdot 1.446$ $1.425$ $10 \cdot 1.429$ $1.400$ $1.30 \cdot 1.429$ $1.409$ $1.811$ METHOD/APPARATUS/PROCEDURE:The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their yolumes andSOURCE AND PURITY OF MATERIALS: (2) Distilled.(2) Distilled.		
(1) Iodomethane; CH_3I; [74-88-4] (2) Water; H_2O; [7732-18-5]Rex, A. Z. Phys. Chem. 1906, 55, 355-70.VARIABLES: $T/K = 273 - 303$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: $100 g_1/g_2$ (compiler) $100 w_1$ (compiler)01.565 1.540 1.425 $1.981$ 1.431 20 3001.565 1.419 30 $1.425$ 1.400 1.793 1.409AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their yournes andSOURCE AND PURITY OF MATERIALS: (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.		
(2) Water; H2O; [7732-18-5]Z. Phys. Chem. 1906, 55, 355-70.VARIABLES: $T/K = 273 - 303$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES:A. L. Horvath $t^{\circ}C$ 100 $g_1/g_2$ 100 $w_1$ (compiler)01.5651.540101.4461.4191.400201.419301.4291.4091.811		
VARIABLES: $T/K = 273 - 303$ PREPARED BY: A. L. HorvathEXPERIMENTAL VALUES: $t/^{\circ}C$ 100 $g_1/g_2$ 100 $w_1$ (compiler) $10^3 x_1$ (compiler)01.5651.5401.981 1.831101.4461.4251.831 2.0201.4191.4001.793 3.0301.4291.4091.811METHOD/APPARATUS/PROCEDURE: The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes andSOURCE AND PURITY OF MATERIALS: (2) Distilled.		
$T/K = 273 - 303$ A. L. HorvathEXPERIMENTAL VALUES: $t/^{\circ}C$ $100 g_1/g_2$ $100 w_1$ $10^3 x_1$ 0 $1.565$ $1.540$ $1.981$ 10 $1.446$ $1.425$ $1.831$ 20 $1.419$ $1.400$ $1.793$ 30 $1.429$ $1.409$ $1.811$ MUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE:SOURCE AND PURITY OF MATERIALS:(1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.(2) Distilled.		
EXPERIMENTAL VALUES: $t^{0}$ C100 $g_1/g_2$ 100 $w_1$ 10^3 $x_1$ 01.5651.5401.981101.4461.4251.831201.4191.4001.793301.4291.4091.811MUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes andSOURCE AND PURITY OF MATERIALS: (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.(2)Distilled.		
$t/^{\circ}C$ 100 $g_1/g_2$ 100 $w_1$ (compiler)10^3 $x_1$ (compiler)01.5651.5401.981101.4461.4251.831201.4191.4001.793301.4291.4091.811AUXILIARY INFORMATIONMETHOD/APPARATUS/PROCEDURE: The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes andSOURCE AND PURITY OF MATERIALS: (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.(2)Distilled.		
0       1.565       1.540       1.981         10       1.446       1.425       1.831         20       1.419       1.400       1.793         30       1.429       1.409       1.811         AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes and       SOURCE AND PURITY OF MATERIALS:         (1)       Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.       (2)         (2)       Distilled.		
AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:       SOURCE AND PURITY OF MATERIALS:         The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes and       (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.         (2) Distilled.       (2) Distilled.		
AUXILIARY INFORMATION         METHOD/APPARATUS/PROCEDURE:         The solubility of iodomethane in water was         determined using specially designed flasks with         calibrated capillary cylinders. After samples were         equilibrated in a thermostatic bath, their volumes and    SOURCE AND PURITY OF MATERIALS:     (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C.		
METHOD/APPARATUS/PROCEDURE:SOURCE AND PURITY OF MATERIALS:The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes and(1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C. (2) Distilled.		
The solubility of iodomethane in water was determined using specially designed flasks with calibrated capillary cylinders. After samples were equilibrated in a thermostatic bath, their volumes and (1) Kahlbaum reagent, redistilled and washed before use, b.p. = 42.8 °C. (2) Distilled.		
calculated. ESTIMATED ERRORS:		
Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).		
COMPONENTS: ORIGINAL MEASUREMENTS:		
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4] Fühner, H.		
(2) Water; H <sub>2</sub> O; [7732-18-5] Ber. <u>1924</u> , 57, 510-5.		
VARIABLES: PREPARED BY:		
7/K = 295 A. L. Horvath		
EXPERIMENTAL VALUES:		
$t/^{\circ}$ C 100 $w_i$ 10 <sup>3</sup> $x_i$ 100 $w_i M_i^{-1}$ /mol g <sup>-1</sup> (compiler) (compiler)		
22 1.362 1.75 $9.59 \times 10^{-3}$		
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS:		
Iodomethane was added gradually to 100 cm <sup>3</sup> water from a pipette under continuous stirring and shaking in an oil bath. The appearance of the first cloud (turbidity) indicated the saturation of the solubility under conducted from the		
volume of iodomethane added to a known quantity of ESTIMATED ERRORS:		
Solubility: Not specified. Temperature: ± 0.5 K (compiler).		

184		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	van Arkel, A. E.; Vles, S. E.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Recl. Trav. Chim. Pays-Bas <u>1936</u> , 55, 407-11.	
VARIABLES:	PREPARED BY:	
T/K = 303	A. L. Horvath	
EXPERIMENTAL VALUES: <i>t</i> /°C <i>m<sub>i</sub></i> /mol kg <sup>-1</sup>	$\begin{array}{ccc} 100 \ w_1 & 10^3 \ x_1 \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
30 0.101	1.413 1.816	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available.	<ol> <li>Source and purity not given.</li> <li>Distilled (compiler).</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 0.5$ K (compiler).	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Glew, D. N.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ph. D. Thesis, University of Cambridge, Cambridge, U. K. <u>1952</u> , 245 pp.	
VARIABLES:	PREPARED BY:	
T/K = 273 - 323 A. L. Horvath		
EXPERIMENTAL VALUES: t/°C P <sub>1</sub> /c <sub>1</sub> Henry's law /dm <sup>3</sup> mmHg mol <sup>-1</sup> H/m <sup>3</sup> atm mol <sup>-1</sup>	constant, $100 w_i$ , $10^3 x_i$ (compiler) (compiler) (compiler)	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The experiment was performed in an Ostwald type solubility apparatus consisting of a gas burette and a solubility pipette. The gas burette containing iodomethane and the solubility pipette filled with water ware concentrate and allowed the consideration in a	<ol> <li>British Drug House, washed with KOH solution, dried, redistilled and dried again with Ca metal t rnings.</li> <li>Gas free, deionizes by ion exchange resin.</li> </ol>	
thermostat bath. The volume, pressure, and tempera- ture were monitored and Henry's law constants were	ESTIMATED ERRORS:	
calculated. A summary of the experiment is given elsewhere (ref. 1).	Solubility: $\pm 0.5 \%$ . Temperature: $\pm 0.2 $ K.	
	REFERENCES:	
	(1) Glew, D. N.; Moelwyn-Hughes, E. A. Faraday Discussion Chem. Soc. <u>1953</u> , 150.	

	185	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Swain, C. G.; Thornton, E. R.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	J. Am. Chem. Soc. <u>1962</u> , 84, 822-6.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 303 - 313 Pressure	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C P <sub>1</sub> /c <sub>1</sub> Henry's law /dm <sup>3</sup> mmHg mol <sup>-1</sup> H/m <sup>3</sup> atm mol <sup>-1</sup>	constant, $100 w_1$ , $10^3 x_1$ (compiler) (compiler) (compiler)	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The measuring apparatus was described by Brown <i>et al.</i> (ref. 1). Iodomethane gas was admitted to the measurement apparatus from a storage bulb by	<ol> <li>Eastman Kodak Co., redistilled before use.</li> <li>(2) Double distilled.</li> </ol>	
distillation. The gas volume absorbed by water was determined using a gas buret. The temperature was controlled by a water bath and the equilibrium was aided by a magnetic stirrer in the equilibration bulb. The concentration in the solution was calculated from the pressure and volume measured	Solubility:       ± 3 %.         Temperature:       ± 0.01 K.         Pressure:       ± 0.05 mmHg.	
	<ul> <li>REFERENCES:</li> <li>(1) Brown, H. C.; Taylor, M. D.; Gerstein, M. J. Am. Chem. Soc. <u>1944</u>, 66, 431.</li> </ul>	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Balls, P. W.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	Ph. D. Thesis, University of East Anglia, Norwich, U. K., July 1980, 375 pp.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_i & 10^3 \ x_i \\ (compiler) & (compiler) \end{array}$	
$\begin{array}{ccccc} 10 & 0.12 \pm 0.04 \\ 15 & 0.15 \pm 0.05 \\ 20 & 0.17 \pm 0.03 \\ 25 & 0.20 \pm 0.04 \end{array}$	1.4641.8821.8291.8291.5141.9471.5391.979	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The measurements involved repeated equili- bration of nitrogen with a water sample containing iodomethane. The sample was shaken vigorously at	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
the required temperature in a constant temperature bath for 30 minutes. The equilibrated gas phase was	ESTIMATED ERRORS:	
then injected into a gas chromatograph. The experi- ments were performed at least three times at 10, 15, 20, and 25 °C. Each experiments involved about six equilibrations.	Solubility: See above. Temperature: ± 0.5 K (compiler).	

186		
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Hunter-Smith, R. J.; Balls, P. W.; Liss, P. S.	
(2) Water; H <sub>2</sub> O; [7732-18-5]	<i>Tellus <u>1983</u>, 35B</i> , 170-6.	
VARIABLES:	PREPARED BY:	
T/K = 283 - 298	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, H/dimensionless	$\begin{array}{ccc} 100 \ w_I & 10^3 \ x_I \\ (compiler) & (compiler) \end{array}$	
10 0.1147 15 0.1425	1.5317 1.971 1.4982 1.927	
20 0.1757 25 0.2151	1.4645 1.883 1.4308 1.839	
The Henry's law constant (H) was deriv	ed from the best-fit lines of van't Hoff plots:	
$\log_e H = -\frac{3541}{T(K)} + 10$	0.34	
AUXILIARY I	NFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The multiple equilibration technique as de- scribed elsewhere (ref. 1) was used to measure the	<ol> <li>Source and purity not given.</li> <li>Distilled.</li> </ol>	
Henry's law constants. A known volume of water in a syringe was shaken with a known volume of pure	ESTIMATED ERRORS:	
iodomethane. After equilibration, the headspace was separated and analyzed for the compound of interest.	Solubility: ± 5.5 % std. dev.	
A gas chromatograph fitted with an electron capture detector was used for the analysis.	Temperature: $\pm 0.2$ K.	
	REFERENCES:	
	(1) McAuliffe, C. D. Chem. Techn. <u>1971</u> , 1, 46.	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Liss, P. S.; Slater, P. G.	
(2) Seawater	Nature <u>1974</u> , 247, 181-4.	
VARIABLES:	PREPARED BY:	
<i>T</i> /K = 293	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C Henry's law constant, <i>H</i> /dimensionless	$\begin{array}{ccc} 100 \text{ w}_1 & 100 \text{ w}_l M_l^{-l}/\text{mol g}^{-1} \\ \text{(compiler)} & \text{(compiler)} \end{array}$	
20 0.24	1.093 7.7× 10 <sup>-3</sup>	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Details are not available except that the data originate from Lovelock (ref. 1).	<ol> <li>Source and purity not given.</li> <li>Samples from Atlantic ocean.</li> </ol>	
	ESTIMATED ERRORS:	
	Solubility: Not specified. Temperature: $\pm 2 K$ (compiler).	
	REFERENCES:	
	(1) Lovelok, J. E. Personal communication.	

· · · · · · · · · · · · · · · · · · ·	187			
COMPONENTS:	ORIGINAL MEASUREMENTS:			
(1) Iodomethane; CH <sub>3</sub> I; [74-88-4]	Swain, C. G.; Thornton, E. R.			
(2) Water-d <sub>2</sub> ; D <sub>2</sub> O; [7789-20-0]	J. Am. Chem. Soc. <u>1962</u> , 84, 822-6.			
VARIABLES:	PREPARED BY:			
T/K = 303 - 313	A. L. Horvath			
EXPERIMENTAL VALUES:				
t/°C P <sub>1</sub> /c <sub>1</sub> Henry's law /dm <sup>3</sup> mmHg mol <sup>-1</sup> H/m <sup>3</sup> atm mol	constant, $100 w_1$ , $10^3 x_1$ , $(compiler)$ (compiler)			
29.43 5450 7.171 × 29.45 5340 7.026 ×	$10^3$ 1.131 1.611 $10^3$ 1.154 1.645			
40.34 7420 9.763 × 40.35 7450 9.803 ×	$10^{-3}$ 1.231 1.755 $10^{-3}$ 1.226 1.748			
	1.220 1.740			
AUXILIARY INFORMATION				
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
The measuring apparatus was described by	(1) Eastman Kodak Co., redistilled before use.			
Brown et al. (ref. 1). Iodomethane gas was admitted	(2) Liquid Carbonic Co., 99.5 % pure, used as			
distillation. The gas volume absorbed by heavy water				
was determined in the gas buret. The temperature was controlled by a water bath and the equilibrium	ESTIMATED ERRORS:			
was aided by a magnetic stirrer in the equilibration hulb. The concentration of the solution was calculated	Solubility: ± 3 %			
from the pressures and volumes measured.	Temperature: $\pm 0.01$ K.			
٠	Pressure: $\pm 0.05$ mmHg.			
	REFERENCES:			
	(1) Brown H C Towlor M D Garstain M			
	J. Am. Chem. Soc. <u>1944</u> , 66, 822.			
······				

# SYSTEM INDEX

Page numbers preceded by E refer to evaluation texts whereas those not preceded by E refer to compiled tables. Substances are listed as in the Chemical Abstracts.

Bromoform	
see methane, tribromo-	
Carbon tetrabromide	
see methane, tetrabromo-	
Carbon tetrachloride	
see methane, tetrachloro-	
Chloroform	
see methane, trichloro-	
Iodoform	
see methane, triiodo-	
Magnesium sulfate	
see sulfuric acid, magnesium salt	
Methane, bromochloro-	
+ water	E143, 144-145
Methane, bromodichloro-	,,,,,,,
+ water	E71, 72-74
+ water, Milang	74
+ water. Myponga	75
+ water, Myponga	
+ humic acid	75
Methane dibromo-	
+ water	E146-E147 148-150
Methane dibromochloro-	E140-E147, 140-130
	E76 77-70
+ water Milane	E70, 77-79
+ water, Milliang	6U 9A
+ water, Myponga	80
+ water, Myponga	01
+ numic acid	81
Methane, dichloro-	
+ water	E151-E155, 156-175
+ hydrogen chloride (aqueous)	
+ water	176
Methane, diiodo-	
+ water	E177, 178-180
Methane, iodo-	
+ water	E181-E182, 183-186
+ water- $d_2$	187
+ water, sea-	186
Methane, tetrabromo-	
+ water	2
Methane, tetrachloro-	
+ water	E17-E21, 22-26, 28-30, 32, 34, 36, 39-44, 46-49, 51-57
+ water, MQ-	58
+ water, Municipal tap, (PASE)	58
+ water. sea-	59-60
+ calcium chloride (aqueous)	
+ water	61
+ hydrogen chloride (aqueous)	
+ water	62
+ lithium chloride (aqueous)	
+ water	63
+ potassium chloride (aqueous)	65
+ water	65
r waici L potassium hydroxida (agusous)	05
+ polassium nyuroxide (aqueous)	66
+ water	00
+ sodium chioride (aqueous)	(7)
+ water	0/

# SYSTEM INDEX

Methane, tetrachloro-	
+ sulfuric acid, dipotassium salt (aqueous)	
+ water	66-67
+ sulfuric acid, disodium salt (aqueous)	
+ water	68
+ sulfuric acid, magnesium salt (aqueous)	
+ water	64
Methane tribromo-	
+ water	E82-E83 84-80
+ water Milang	90
+ water, Munonga	90
+ water, Myponga	50
+ humic acid	01
Methane tribromofluoro	91
wethane, information	1
+ walci Mathana triahlara	1
Methane, tricinoro-	
+ water	E92-E96, 97-103, 105-107, 109-110, 112-121, 123-131
+ water, Milang	132
+ water, MQ-	132
+ water, Municipal tap, (PASE)	133
+ water, Myponga	133
+ water, Myponga	
+ humic acid	134
+ water, sea	134-135
+ cesium chloride (aqueous)	
+ water	136
+ hydrogen chloride (aqueous)	
+ water	137
+ lithium chloride (aqueous)	
+ water	138
+ potassium chloride (aqueous)	
+ water	138-139
+ rubidium chloride (aqueous)	
+ water	139
+ sodium chloride (aqueous)	
+ water	140
Methane, trichlorofluoro-	
+ water	E3-E5, 6-13
+ water, sea-	13-16
Methane, triiodo-	
+ water	141-142
Potassium sulfate	
see sulfuric acid, dipotassium salt	
Sodium sulfate	
see sulfuric acid, disodium salt	
Water	
+ methane, bromotrichloro-	1
+ methane, dichloro-	- 158 160 163 167 170-171
+ methane, dijodo-	170
+ methane, tetrachloro-	177 17 17 21 22 25 40 42 42 45 49 50 52 55
+ methane, trichloro-	22-27, 31-33, 33-40, 42-43, 43, 40, 30, 32, 33 100-101 103-104 108-100 111 115 122 124
Water-d	100-101, 103-104, 106-107, 111, 113, 122, 124
t methane dijodo.	190
T methane tetrachlara	100 40 70
T methane, terachloro	140
- memane, memoro-	140

# **REGISTRY NUMBER INDEX**

Page numbers preceded by E refer to evaluation texts whereas those not preceded by E refer to compiled tables.

56-23-5	E17-E21, 22-70
67-66-3	E92-E96, 97-140
74-88-4	E181-E182, 183-187
74-95-3	E146-E147, 148-150
74-97-5	E143, 144-145
75-09-2	E151-E155, 156-176
75-11-6	E177, 178-180
75-25-2	E82-E83, 84-91
75-27-4	E71, 72-75
75-47-8	141-142
75-62-7	1
75-69-4	E3-E5, 6-16
124-48-1	E76, 77-81
353-54-8	1
558-13-4	2
1310-58-3	66
7447-40-7	65, 138-139
7447-41-8	63, 138
7487-88-9	64
7647-01-0	62, 137, 176
7647-14-5	67, 149
7647-17-8	136
7732-18-5	1-2, E3-E5, 6-13, E17-E21, 22-57, 61-68, E71, 72-74, E76, 77-79, E82-E83, 84-89, E92-E96, 97-131, 136-142, E143, 144-145, E146-E147, 148-150, E151-155, 156-176, E177, 178-180, E181-E182, 183-186
7757-82-6	68
7778-80-5	66-67
7789-20-0	69-70, 140, 180, 187
7791-11-9	139
10043-52-4	61

Pages preceded by an E refer to evaluation texts while those not preceded by E refer to compiled tables.

Ababi, V. 110 35, 37 Affsprung, H. E. 53, 125 Alessi, P. Alexandrova, M. V. 164-166 179 Andrews, L. J. Antonov, G. N. 99 Antropov, L. I. 41, 115, 165 Archer, W. L. 168 47, 118 Aref'eva, R. P. Bachofen, H. 113 Bakowski, S. 157 Balls, P. W. 9, 13, 48, 52, 59, 60, 119, 124, 134-135, 185, 186 49, 121 Banerjee, S. 129 Barr, R. S. Bartell, F. E. 30, 107, 160 9 Battino, R. 25 Bell, R. P. Booth, H. S. 29, 85, 106, 142, 159 Botth, H. S. 149 Buchowski, H. 48 Bursill, D. B. 73-75, 78, 80-81, 88, 90-91, 127, 132-134 84 Caines, C. M. 51, 123, 171 Calo, J. M. Carroll, J. 49, 120 Cave, G. C. B. 45 97 Chancel, G. 117, 136, 138-140 Charykov, A. K. 40, 114, 164 Chatterjee, A. K. Chiou, C. T. 46, 118 111 Chistyakov, V. M. 30 Chitwood, B. G. 35, 37 Christian, S. D. Clifford, C. W. 22, 100 10, 14-15 Cline, J. D. 47, 119, 169 Coca, J. Cohen, J. M. 11, 56, 74, 79, 89, 130, 173 Conti, J. J. 110 35 Cristian, S. D. 160 Davies, W. De Minjer, C. H. 103, 106 141 Dehn, W. M. 33, 109 Desnover, M. DeVoll, D. 57, 89, 131, 150, 175 Diaz, R. M. 47, 119, 169 Donahue, D. J. 30, 107, 160 du Pont de Nemours & Co. 6, 162 31, 108 Eberius, E. 69 Epherre, P. 102 Evans, T. W. Everson, H. E. 29, 85, 107, 142, 149, 159 113 Farhi, L. E. 7, 44, 116, 167 Ferguson, D. M.

Fermeglia, M. Fitzgerald, M. E. Flid, R. M. Fox, J. J. Fraga, D. W. Fredlund, F. Freed, V. H. Fühner, H. Furniss, B. S. Gendrano, M. C. Gibby, C. W. Gillam, W. S. Gilmont, R. Glasoe, P. K. Glew, D. N. Goldman, S. Gorichnina, V. P. Gorokhov, A. A. Gossett, J. M. Greinacher, E. Griffing, V. Grigsby, R. D. Grob, R. L. Gross, P. M. Hagler, K. Hall, J. Hellebust, J. A. Herz, W. Hildebrand, J. H. Högfeldt, E. Horvath, A. L. Howe, G. B. Huang, T.-C. Hunter, W. J. A. Hunter-Smith, R. J. Hutchinson, T. C. Hutchison, C. A. Ireland, J. C. Ishihara, Y. Jagger, J. B. Johnson, J. R. Karger, B. L. Kazama, K. Keefer, R. M. Kikic, I. King, J. W. Kirchnerova, J. Kiya-Oglu, N. V. Kleeberg, H. Klein, D. Köhl, A. Kohnert, R.

53, 125 32 39, 62, 114, 137, 163, 176 26 31 37, 115 46, 118 183 57, 131, 174 111 101 37 110 40, 70 184 43 162 47, 118 55, 125-126, 129, 172, 173 32 32 36 72, 73, 77-78, 87, 126 2, 23-24, 64-65, 84, 102, 148, 178 113 101 120, 170 97 31 37, 115 E3-E5, E17-E21, E71, E76, E82-E83, E92-E96, E143, E146-E147, E151-E155, E177, E181-E182 12, 56, 79, 130, 174 34, 61, 63-68, 145 37 52, 60, 124, 135, 186 120, 170 27, 69, 104, 140, 179, 180 11, 56, 74, 79, 89, 130, 173 128 160 35-36 40, 114, 164 52, 124, 171 179 53, 125 40, 114, 164 45 162 55 55 66 46, 118

Komarova, V. P. Korenman, I. M. Krutikova, A. D. Ksiazczak, A. Kudryavtseva, G. I. Lees, F. P. Leighton, D. T. Levinskii, M. I. Lincoff, A. H. Liss, P. S. Liu, J.-L. Lo, J. M. Löfvenberg, A. Lowe, H. J. Luck, W. A. P. Lüttke, W. Lyon, A. M. Macek, K. J. Mackay, D. Maguire, B. P. Maretic, M. Martin, A. K. Martinov, W. W. Martire, D. E. Masterton, W. L. McCollum, J. L. McConnell, G. McCoy, W. H. McGovern, E. W. McNally, M. E. Mecke, R. Mihaila, Gh. Miller, M. M. Mitsuzawa, S. Moore, B. Mullins, M. E. Munz, C. D. Nakijima, T. Newsham, D. M. T. Nicholson, B. C. Nicloux, M. Niini, A. Nogradi, M. O'Connell, W. L. Ödberg, L. Ohtsuka, K. Okuda, Y. Olling, M. Orlandini, M. Ose, Y. Othmer, D. F. Park, T. Parmentier, F.

165-166 47, 118 107, 161 48 107, 161 163 51, 123, 171 39, 62, 114, 137, 163, 176 125-126, 172 52, 59-60, 124, 135, 186 34, 61, 63-68 54, 128 115 112-113 55 32 27, 69, 104, 140, 179-180 49, 120 51, 88, 123, 144 73-75, 78, 80-81, 88, 90-91, 127, 132-134 161 26 164 144 111 101 7-8, 44, 116, 167-168 37 28, 159, 105 72-73, 77-78, 87, 126 32 110 144-145 128 98 12, 56, 79, 130, 174 53, 58, 88, 127, 132-133 46, 117, 169 129 73-75, 78, 80-81, 88, 90-91, 127, 132-134 100 26, 158 34, 109 1, 86, 144, 150, 180 115 52, 124, 171 112 7 53, 125 54 110 9 97

193

Priors

Pavlovskaya, E. M. Pazos, C. Pearson, C. R. Peterson, D. Petrucelli, S. R. Pogulyai, V. E.K Popa, A. Same Popova, L. N. Populyai, V. E. Powell, J. F. Pozas, C. Prosyanov, N. N. Rauws, A. G. Reinders, W. Rettich, T. R. Rex, A. Roaf, H. E. Rogers, T. N. Rosenbaum, C. K. Rotariu, G. J. Roth, E. Sabinin, V. E. Sadovnikova, L. V. Salkowski, E. Sandler, S. I. Sarram, P. Sato, A. Sato, T. Sawinsky, J. Saylor, J. H. Schmedding, D. Schultz, S. D. Scotti-Foglieni, L. Serafimov, L. A. Shalygin, V. A. Shamsutdinov, T. M. Shapurova, V. V. Shiu, W.-Y. Simonov, V. D. Sirocic, V. Slater, P. G. Squire, P. W. Staverman, A. J. Stevens, V. L. Sukomick. B. Sullivan, J. Svetlanov, E. B. Swain, C. G. Symons, J. M. Takano, J. Tam, D. Tettamanti, K. Tewari, Y. B. Thornton, E. R. Tikhomirov, V. I. Treadwell, W. D.

117, 136, 138-140 47, 169 7-8, 44, 116, 167-168 9 49, 120 38, 41, 43 110 43 115, 165 28 119 41-42, 166-167 7 106 9 22, 98, 148, 156, 183 98 12, 56, 79, 130, 174 23-24 31 69 162 164-166 99, 156 57, 89, 131, 150, 175 163 46, 117, 169 54 34, 109 2, 24, 84, 102, 148, 178 46, 118 40, 70 100 166 41-42, 166, 167 38, 41, 43, 115, 165 111 145 38-39, 41, 43, 115, 165 161 59, 186 84 27, 104, 158 168 12 32 39, 62, 114, 137, 163, 176 185, 187 72, 77, 86, 121 128 120, 170 34, 109 144 185, 187 117, 136, 138-140 66

194

. ۲<sup>41</sup>

Treger, Yu. A. Treszczanowicz, E. Tseng, C. L. 54, Valvani, S. C. van Arkel, A. E. Veith, G. D. Velichko, S. M. Vles, S. E. Vogel, A. I. Walton, J. H. Warner, M. J. Warner, H. P. Wasik, S. P. Weiss, R. F. Whalley, H. B. Wibowo, A. E. Wilhelm, E. Wisegarver, D. P. Wright, D. A. Wu, X. Yalkowsky, S. H. Yang, J. Y. Yasuoka, T. Yoshioka, Y. Zeininger, H. Zel'venskii, Ya. D.

Zielinski, A. Z.

39, 62, 114, 137, 163, 176 157 128 49, 121 2, 25, 85, 103, 141, 149, 157, 178, 184 49, 120 39, 62, 114, 137, 163, 176 2, 25, 85, 103, 141, 149, 157, 178, 184 57, 131, 174 23-24 10-11, 15-16 11, 56, 74, 79, 89, 130, 173 144, 145 10-11, 15-16 160 7 9 10, 14-15 57, 89, 131, 150, 175 50, 122, 170 49, 121 54, 128 128 54 8,13

41-42, 166-167 33, 108

# SOLUBILITY DATA SERIES

Volume 1	H. L. Clever, Helium and Neon
Volume 2	H. L. Clever, Krypton, Xenon and Radon
Volume 3	M. Salomon, Silver Azide, Cyanide, Cyanamides, Cyanate, Selenocyanate
	and Thyocyanate
Volume 4	H. L. Clever, Argon
Volume 5/6	C. L. Young, Hydrogen and Deuterium
Volume 7	R. Battino, Oxygen and Ozone
Volume 8	C. L. Young, Oxides of Nitrogen
Volume 9	W. Hayduk, Ethane
Volume 10	R. Battino, Nitrogen and Air
Volume 11	B. Scrosati and C. A. Vincent, Alkali Metal, Alkaline Earth Metal and
	Ammonium Halides, Amide Solvents
Volume 12	C. L. Young, Sulfur Dioxide, Chlorine, Fluorine and Chlorine Oxides
Volume 13	S. Siekierski, T. Mioduski and M. Salomon, Scandium, Yttrium,
	Lanthanum and Lanthanide Nitrates
Volume 14	H. Miyamoto, M. Salomon and H. L. Clever, Alkaline Earth Metal
	Halates
Volume 15	A. F. M. Barton, Alcohols with Water
Volume 16/17	E. Tomlinson and A. Regosz, Antibiotics: 1,β-Lactam Antiobiotics
Volume 18	O. Popovych, Tetraphenylborates
Volume 19	C. L. Young, Cumulative Index: Volumes 1–18
Volume 20	A. L. Horvath and F. W. Getzen, Halogenated Benzenes, Toluenes and
	Phenols with Water
Volume 21	C. L. Young and P. G. T. Fogg, Ammonia, Amines, Phosphine, Arsine,
	Stibine, Silane, Germane and Stannane in Organic Solvents
Volume 22	T. Mioduski and M. Salomon, Scandium, Yttrium, Lanthanum and
	Lanthanide Halides in Nonaqueous Solvents
Volume 23	T. P. Dirkse, Copper, Silver, Gold and Zinc, Cadmium, Mercury Oxides
	and Hydroxides
Volume 24	W. Hayduk, Propane, Butane and 2-Methylpropane
Volume 25	C. Hirayama, Z. Galus and C. Guminski, Metals in Mercury
Volume 26	M. R. Masson, H. D. Lutz and B. Engelen, Sulfites, Selenites and
	Tellurites
Volume 27/28	H. L. Clever and C. L. Young, <i>Methane</i>
Volume 29	H. L. Clever, Mercury in Liquids, Compressed Gases, Molten Salts and
	Other Elements
Volume 30	H. Miyamoto and M. Salomon, Alkali Metal Halates, Ammonium lodate
	and Iodic Acid
Volume 31	J. Eysseltová and T. P. Dirske, Alkali Metal Orthophosphates
Volume 32	P. G. T. Fogg and C. L. Young, Hydrogen Sulfide, Deuterium Sulfide and
	Hydrogen Selenide
Volume 33	P. Franzosini, Molten Alkali Metal Alkanoates
Volume 34	A. N. Paruta and R. Piekos, 4-Aminobenzenesulfonamides.
	Part I: Non-cyclic Substitutents
Volume 35	A. N. Paruta and R. Piekos, 4-Aminobenzenesulfonamides.
	Part 11: 5-Membered Heterocyclic Substituents

Volume 36	A. N. Paruta and R. Piekos, 4-Aminobenzenesulfonamides. Part III: 6-Membered Heterocyclic Substituents and Miscellaneous
Volume 37	D. G. Shaw, Hydrocarbons with Water and Seawater.
Volume 38	D. G. Shaw, Hydrocarbons with Water and Seawater.
Volume 39	C I Young Cumulative Inder: Volumes 20-38
Volume 40	J. Hala, Halides, Oxyhalides and Salts of Halogen Complexes of Titanium,
	Zirconium, Hafnium, Vanadium, Niobium and Tantalum
Volume 41	CY. Chan, I. N. Lepeshkov and K. H. Khoo, Alkaline Earth Metal Perchlorates
Volume 42	P. G. T. Fogg and W. Gerrard, Hydrogen Halides in Non-aqueous
Valuma 42	B. W. Cargill Carbon Monorida
Volume 45	K. W. Cargin, Carbon Monorale H. Miyamoto, F. M. Woolloy and M. Solomon, Conner and Silver Halates
Volume 44	R P T Tomkins and N P Bansal Gases in Molten Salts
Volume 47	R Cohen-Adad and I W I orimer Alkali Metal and Ammonium
Volume 47	Chlorides in Water and Heavy Water (Rinary Systems)
Volume 48	F. Getzen, G. Hefter and A. Maczynski, <i>Esters with Water</i> .
Volume 49	F. Getzen, G. Hefter and A. Maczynski, Esters with Water. Part 11: Esters 7-C to 32-C
Volume 50	P. G. T. Fogg, Carbon Dioxide in Non-aqueous Solvents at Pressures Less Than 200 kPa
Volume 51	J. G. Osteryoung, M. M. Schreiner, C. Guminski and Z. Galus, Intermetallic Compounds in Mercury
Volume 52	I. Lambert and H. L. Clever, Alkaline Earth Hydroxides in Water and Aqueous Solutions
Volume 53	C. L. Young, Cumulative Index: Volumes 40–52
Volume 54	W. E. Acree, Jr, Polycyclic Aromatic Hydrocarbons in Pure and Binary Solvents
Volume 55	S. Siekierski and S. L. Phillips, Actinide Nitrates
Volume 56	D. Shaw, A. Skrzecz, J. W. Lorimer and A. Maczynski, <i>Alcohols with Hydrocarbons</i>
Volume 57	W. Hayduk, Ethene
Volume 58	W. E. Acree, Jr, Polycyclic Aromatic Hydrocarbons: Binary Non-aqueous
	Systems. Part I: Solutes A–E
Volume 59	W. E. Acree, Jr, Polycyclic Aromatic Hydrocarbons: Binary Non-aqueous Systems. Part II: Solutes F-Z