SOLUBILITY DATA SERIES

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Editor-in-Chief A. S. KERTES

Volume 20

HALOGENATED BENZENES, TOLUENES ``` AND PHENOLS WITH WATER

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If the knowledge is undigested or simply wrong, more is not better

How to communicate and disseminate numerical data effectively in chemical science and technology has been a problem of serious and growing concern to IUPAC, the International Union of Pure and Applied Chemistry, for the last two decades. The steadily expanding volume of numerical information, the formulation of new interdisciplinary areas in which chemistry is a partner, and the links between these and existing traditional subdisciplines in chemistry, along with an increasing number of users, have been considered as urgent aspects of the information problem in general, and of the numerical data problem in particular.

Among the several numerical data projects initiated and operated by various IUPAC commissions, the *Solubility Data Project* is probably one of the most ambitious ones. It is concerned with preparing a comprehensive critical compilation of data on solubilities in all physical systems, of gases, liquids and solids. Both the basic and applied branches of almost all scientific disciplines require a knowledge of solubilities as a function of solvent, temperature and pressure. Solubility data are basic to the fundamental understanding of processes relevant to agronomy, biology, chemistry, geology and oceanography, medicine and pharmacology, and metallurgy and materials science. Knowledge of solubility is very frequently of great importance to such diverse practical applications as drug dosage and drug solubility in biological fluids, anesthesiology, corrosion by dissolution of metals, properties of glasses, ceramics, concretes and coatings, phase relations in the formation of minerals and alloys, the deposits of minerals and radioactive fission products from ocean waters, the composition of ground waters, and the requirements of oxygen and other gases in life support systems.

The widespread relevance of solubility data to many branches and disciplines of science, medicine, technology and engineering, and the difficulty of recovering solubility data from the literature, lead to the proliferation of published data in an ever increasing number of scientific and technical primary sources. The sheer volume of data has overcome the capacity of the classical secondary and tertiary services to respond effectively.

While the proportion of secondary services of the review article type is generally increasing due to the rapid growth of all forms of primary literature, the review articles become more limited in scope, more specialized. The disturbing phenomenon is that in some disciplines, certainly in chemistry, authors are reluctant to treat even those limited-in-scope reviews exhaustively. There is a trend to preselect the literature, sometimes under the pretext of reducing it to manageable size. The crucial problem with such preselection - as far as numerical data are concerned - is that there is no indication as to whether the material was excluded by design or by a less than thorough literature search. We are equally concerned that most current secondary sources, critical in character as they may be, give scant attention to numerical data.

On the other hand, tertiary sources - handbooks, reference books and other tabulated and graphical compilations - as they exist today are comprehensive but, as a rule, uncritical. They usually attempt to cover whole disciplines, and thus obviously are superficial in treatment. Since they command a wide market, we believe that their service to the advancement of science is at least questionable. Additionally, the change which is taking place in the generation of new and diversified numerical data, and the rate at which this is done, is not reflected in an increased third-level service. The emergence of new tertiary literature sources does not parallel the shift that has occurred in the primary literature. With the status of current secondary and tertiary services being as briefly stated above, the innovative approach of the Solubility Data Project is that its compilation and critical evaluation work involve consolidation and reprocessing services when both activities are based on intellectual and scholarly reworking of information from primary sources. It comprises compact compilation, rationalization and simplification, and the fitting of isolated numerical data into a critically evaluated general framework.

The Solubility Data Project has developed a mechanism which involves a number of innovations in exploiting the literature fully, and which contains new elements of a more imaginative approach for transfer of reliable information from primary to secondary/tertiary sources. The fundamental trend of the Solubility Data Project is toward integration of secondary and tertiary services with the objective of producing in-depth critical analysis and evaluation which are characteristic to secondary services, in a scope as broad as conventional tertiary services.

Fundamental to the philosophy of the project is the recognition that the basic element of strength is the active participation of career scientists in it. Consoludating primary data, producing a truly crutically-evaluated set of numerical data, and synthesizing data in a meaningful relationship are demands considered worthy of the efforts of top scientists. Career scientists, who themselves contribute to science by their involvement in active scientific research, are the backbone of the project. The scholarly work is commissioned to recognized authorities, involving a process of careful selection in the best tradition of IUPAC. This selection in turn is the key to the quality of the output. These top experts are expected to view their specific topics dispassionately, paying equal attention to their own contributions and to those of their peers. They digest literature data into a coherent story by weeding out what is wrong from what is believed to be right. To fulfill this task, the evaluator must cover all relevant open literature. No reference is excluded by design and every effort is made to detect every bit of relevant primary source. Poor quality or wrong data are mentioned and explicitly disqualified as such. In fact, it is only when the reliable data are presented alongside the unreliable data that proper justice can be done. The user is bound to have incomparably more confidence in a succinct evaluative commentary and a comprehensive review with a complete bibliography to both good and poor data.

It is the standard practice that the treatment of any given solute-solvent system consists of two essential parts: I. Critical Evaluation and Recommended Values, and II. Compiled Data Sheets.

The Critical Evaluation part gives the following information:

- (i) a verbal text of evaluation which discusses the numerical solubility information appearing in the primary sources located in the literature. The evaluation text concerns primarily the quality of data after consideration of the purity of the materials and their characterization, the experimental method employed and the uncertainties in control of physical parameters, the reproducibility of the data, the agreement of the worker's results on accepted test systems with standard values, and finally, the fitting of data, with suitable statistical tests, to mathematical functions;
- (ii) a set of recommended numerical data. Whenever possible, the set of recommended data includes weighted average and standard deviations, and a set of smoothing equations derived from the experimental data endorsed by the evaluator;
- (iii) a graphical plot of recommended data.

The Compilation part consists of data sheets of the best experimental data in the primary literature. Generally speaking, such independent data sheets are given only to the best and endorsed data covering the known range of experimental parameters. Data sheets based on primary sources where the data are of a lower precision are given only when no better data are available. Experimental data with a precision poorer than considered acceptable are reproduced in the form of data sheets when they are the only known data for a particular system. Such data are considered to be still suitable for some applications, and their presence in the compilation should alert researchers to areas that need more work.

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The typical data sheet carries the following information:

- (1) components - definition of the system - their names, formulas and Chemical Abstracts registry numbers;
- (11) reference to the primary source where the numerical information is reported. In cases when the primary source is a less common periodical or a report document, published though of limited availability, abstract references are also given;
- (1ii) experimental variables;
- (iv) identification of the compiler;
 - (v) experimental values as they appear in the primary source. Whenever available, the data may be given both in tabular and graphical form. If auxiliary information is available, the experimental data are converted also to SI units by the compiler.

Under the general heading of Auxiliary Information, the essential experimental details are summarized:

- (vi) experimental method used for the generation of data;
- (vii) type of apparatus and procedure employed;
- (viii) source and purity of materials; (ix) estimated error; (x) references relevant to the generation of experimental data as cited in the primary source.

This new approach to numerical data presentation, formulated at the initiation of the project and perfected as experience has accumulated, has been strongly influenced by the diversity of background of those whom we are supposed to serve. We thus deemed it right to preface the evaluation/compilation sheets in each volume with a detailed discussion of the principles of the accurate determination of relevant solubility data and related thermodynamic information.

Finally, the role of education is more than corollary to the efforts we are seeking. The scientific standards advocated here are necessary to strengthen science and technology, and should be regarded as a major effort in the training and formation of the next generation of scientists and engineers. Specifically, we believe that there is going to be an impact of our project on scientific-communication practices. The quality of consolidation adopted by this program offers down-to-earth guidelines, concrete examples which are bound to make primary publication services more responsive than ever before to the needs of users. The self-regulatory message to scientists of the early 1970s to refrain from unnecessary publication has not achieved much. A good fraction of the literature is still cluttered with poor-quality articles. The Weinberg report (in 'Reader in Science Information', ed. J. Sherrod and A. Hodina, Microcard Editions Books, Indian Head, Inc., 1973, p. 292) states that 'admonition to authors to restrain themselves from premature, unnecessary publication can have little effect unless the climate of the entire technical and scholarly community encourages restraint...' We think that projects of this kind translate the climate into operational terms by exerting pressure on authors to avoid submitting low-grade material. The type of our output, we hope, will encourage attention to quality as authors will increasingly realize that their work will not be suited for permanent retrievability unless it meets the standards adopted in this project. It should help to dispel confusion in the minds of many authors of what represents a permanently useful bit of information of an archival value, and what does not.

If we succeed in that aim, even partially, we have then done our share in protecting the scientific community from unwanted and irrelevant, wrong numerical information.

A. S. Kertes

PREFACE

This volume contains a comprehensive collection and critical evaluation of published solubility data for halogenated benzenes, halogenated toluenes, and halogenated phenois in water prior to 1983. In addition, the solubilities of water in some of the mentioned compounds are also included as well as the mutual solubilities between heavy water (D₂O) and some of the compounds. It should be readily apparent to the reader that there is a great scarcity of data available from the published literature on these systems. This is due primarily to the limited use of the organic compounds in commercial quantities.

All halogenated benzenes, toluenes, and phenols are liquids or solids at room temperature. Consequently, this volume is concerned entirely with the solubility behavior for condensed systems only (solid or liquid solutes in liquid solvents). For several of the compounds considered, only one single measured solubility value is available. It is particularly noteworthy, for example, that only one source (1) has provided the only reported data for solubilities in water of almost the entire series of halogenated, substituted phenols. As the critical evaluators often observe, further experimental investigations are needed for an improvement of the reported solubilities and for an extension of the solubilities over wider temperature ranges. Of course, higher temperature (above the normal boiling point of water) and higher pressure (above atmospheric pressure) solubility data are extremely scarce and available for only a very few systems.

The evaluators for this volume have often been faced with the difficult task of examining and evaluating the reliability of a single solubility with only a limited description of the experimental procedures used for the measurement. In such cases, the evaluator's experience in the techniques of measurement has proved invaluable. Also, some of the reported solubility data are several decades old. However, despite the dated experimental techniques, these published measurements have been found to be no less reliable than those of more recent investigations. Consequently, a recently reported result of a solubility measurement has not automatically become the most reliable or recommended value. In every case, efforts have been made to provide the most reliable and realistic solubility value regardless of the period of origin.

Because of their nature, mixtures of water and halogenated benzenes, halogenated toluenes, and halogenated phenols at room temperature over a wide range of proportions form two distinct phases, one aqueous rich and the other organic rich. However, at higher temperatures and at high pressures, the mutual solubilities usually increase significantly as shown, for example, in Figure 1 for 1,4-difluorobenzene in water system (from the reported work of Jockers and Schneider (2)). With changing temperature and pressure, the two liquid phases generally become identical at a critical, or consolute, point. With changing pressure, the critical solution temperature will change, forming a critical solution line. However, applied pressure exerts only a small effect on the critical temperature.

Systems containing halogen derivatives of benzenes, toluenes, and phenols with water are classified as non-regular type solutions. Consequently, those theories and relationships which were developed by J. H. Hildebrand and his co-workers over several decades for explaining solubility behavior are not generally useful for such systems. However, for the organic rich phase, where the less-well-behaved and more non-ideal water molecules are in very low concentrations, the solubility can be correlated against Absolute temperature by means of the equation:

$$\log x(2) = A - B/T$$

where: x(2) = mole fraction solubility of water in the organic phase T = Absolute temperature A,B = Adjustable constants

That is, for water as the solute, the logarithm of its mole fraction composition is a linear (straight line) function of the reciprocal of the Absolute temperature (3-6). This linearity of the solubility behavior indicates that the organic-rich phase may approach the random distribution of molecules in solution which is one of the basic assumptions in the regular solution theory as detailed by Hildebrand and Scott (7). However, the strictly straight line correlation given above is not always followed for a wide temperature interval, e.g., from the triple point to the critical point. Even so, for the solubility of water in the halogenated benzenes, toluenes, and phenols, a good fit can be anticipated in the temperature range between the triple point of the solvent and the normal boiling point of water (373.15 K).

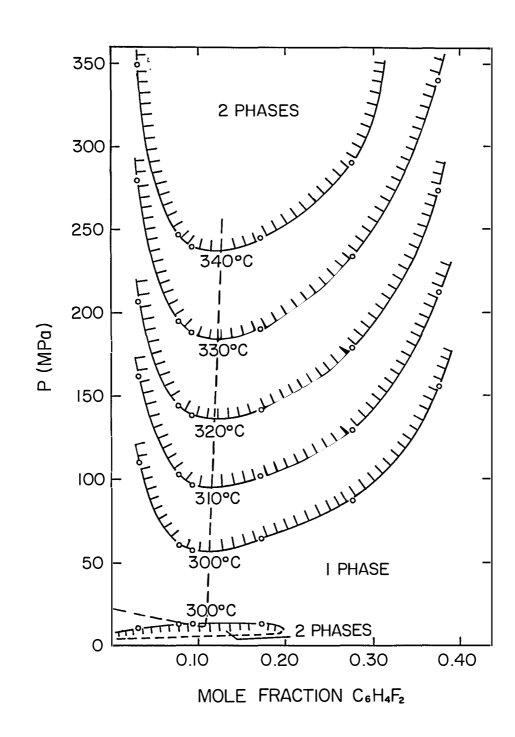


Figure 1. Solubility behavior of 1,4-difluorobenzene and water versus pressure and temperature (2).

Preface

Since each system has been treated separately in the evaluation procedure, it is appropriate here in the Introduction to present the general behavior of the evaluated solubilities of water in the various organic constituents as a function of temperature. These relationships are shown in Figure 2 as the logarithm of mole fraction solubility versus the reciprocal of Absolute temperature. It can be seen that, despite the similarity of the organic solvents, the solubility curves do not show any sort of strict regularity. However, according to Horvath (8), some relatively simple pattern of behavior such as parallel straight lines should be expected. It is clear from Figure 2 that, for example, the solubility curve for iodobenzene shows a slope which differs from the remaining system. Indeed, further studies are needed as additional solubility data become available.

One very simple and practical relationship has been found to be quite useful for solubility data correlation. A plot of the logarithm of solubility in water expressed as molarity against the solute molar volume at 25°C follows a simple linear behavior and such plots may be used to check reported solubility data (9-11). The great advantage of this procedure is that the only information required is the density of the solute at 25°C which is relatively easy to measure when not available in standard references. Such a relation, which was reported by Horvath (8), has been used in this volume and is shown in Figure 3 for eight halogenated benzenes. However, the values shown in the figure do not include the recently reported measurements by Yalkowsky et al. (12).

The utility of the logarithm of molar saturation versus molar volume correlation was recognized in its application to the evaluation of the solubility of 1,3-dibromobenzene in water where the two independently determined measurements were in conflict. The solubility data for eighteen of the substances reported by Yalkowsky et al., including the 1,3-dibromobenzene, were correlated as shown in Figure 4. This graph suggests that the errors for the measured solubilities may, in fact, not agree well with the \pm 10 percent which was reported by Yalkowsky (13) for the measurements. The graph suggests also that the reported solubility of 1,3-dibromobenzene in water may be too high in relation to the other solubilities reported by Yalkowsky et al.

It is, of course, relevant to consider the existence of trends and regularities observed in the behavior of physical parameters and properties, such as their thermodynamic constants and hydrophobic interactions, as determined from solubility measurements involving groups of compounds. In the case of solution enthalpies, for example, Gill et al. (14) have investigated their positive linear dependence with temperature for slightly soluble aromatic compounds. The calorimetric enthalpy of solution determinations for slightly soluble aromatic compounds over a temperature interval provide a very useful relationship for the temperature dependence of the solubility in water. The heat capacity change derived from the enthalpy of solution varies slightly with temperature (over limited, but reasonable, ranges of temperature). Consequently, thermodynamic expressions can provide useful descriptions of the temperature dependence of solubilities.

The curve for solubility versus temperature for all liquid aromatic hydrocarbons shows a minimum for zero heat of solution. This temperature minimum is calculated from the equation:

 $\Delta H_{soln}^{\infty} (T) = \Delta H_{soln}^{\infty} (T_{o}) + \Delta C_{p,soln}^{\infty} (T - T_{o})$ by use of the condition: $\Delta H_{soln}^{\infty} (T) = 0 \text{ at } T = T_{min}$ so that: $T_{min} = T_{o} - \Delta H_{soln}^{\infty} (T_{o}) / \Delta C_{p,soln}^{\infty}$

In other words, the minimum solubility temperatures, T_{min} , for liquid aromatic hydrocarbons in water are calculated from the measured solution enthalpies at T_o = 298.15 K, $\Delta H^{\infty}_{\rm soln}$ (T_o) and the heat capacity change at T_o = 298.15 K, $\Delta C^{\infty}_{\rm p,soln}$.

The minimum solubility phenomenon is a characteristic of the solubility versus temperature curves in the 285 through 320 K range for liquid aromatic hydrocarbon in water systems. This minimum is illustrated in the cases of 1,2-dichlorobenzene, 1,3-dichlorobenzene, and chlorobenzene in water where solubilities have been reported over wide ranges of temperature.

The solubility in water behavior for 1,4-dichlorobenzene has been evaluated in two parts, one below and the other above the normal melting point (326.25 K) of the compound. This is necessary because of the distinct break in the solubility versus temperature behavior at the compound melting point. Here it is important to note that the solubility versus

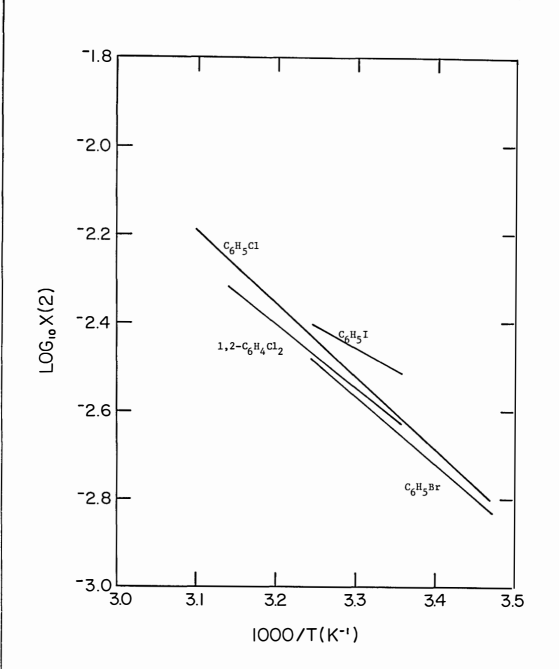
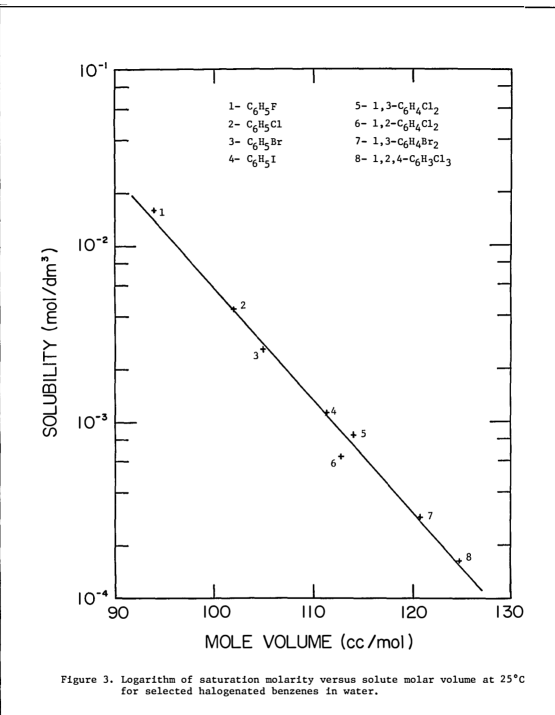


Figure 2. Logarithm of mole fraction water versus the reciprocal of Absolute temperature for the solubility of water in selected halogenated aromatic solvents.



Correlation equation: $\log_{10}S_1 (mole/dm^3) = 4.17442 - 6.40668 \times 10^{-2} \bar{v}_{25°C} (cc/mol)$

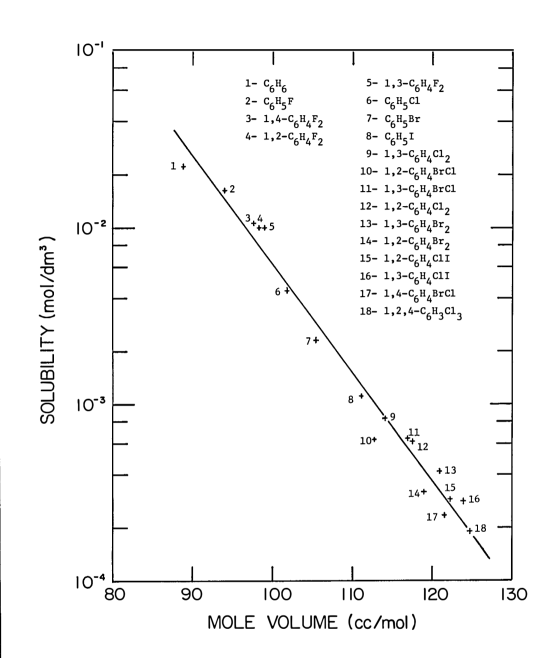


Figure 4. Logarithm of saturation molarity versus solute molar volume at 25°C for selected aromatics in water as reported by Yalkowsky et al. (13).

Correlation equation:

 $\log_{10}S_1(\text{mol/dm}^3) = 3.94385 - 6.15067 \times 10^{-2} \bar{v}_{25°C}(\text{cc/mol})$

To	Bunsen coeff. ^B 2	Ostwald coeff.	Technical coeff. ^T 2	Kuenen coeff. ^K 2	Weight Percent ^W 2	Mole Fraction
Bunsen coefficient, B ₂	1	$\frac{273.15 \text{ 0}_2}{\text{T}}$	d T ₂ 0.9678	d K ₂	$\frac{22415 \text{ z d } \text{W}_2}{(100-\text{W}_2) \text{ M P}}$	$\frac{22415 \text{ z d } X_2}{(1 - X_2) M_L^P}$
Ostwald coefficient, O ₂	$\frac{T B_2}{273.15}$	1	d T T ₂ 264.35	d TK ₂ 273.15	22415 zdTW ₂ 273.15(100-W ₂)MP	$\frac{22415 \text{ z d T X}_2}{273.15(1 - X_2) \text{ M}_L^P}$
Technical coefficient, T ₂	$\frac{0.9678 \text{ B}_2}{\text{d}}$	$\frac{264.35 \text{ O}_2}{\text{d T}}$	1	0.9678 к ₂	0.9678 · 22415 z W ₂ (100-W ₂) M P	$\frac{0.9678 \cdot 22415 \text{ z X}_2}{(1 - \text{X}_2) \text{ M}_L^P}$
Kuenen coefficient, K ₂	$\frac{B_2}{d}$	273.15 0 ₂ d T	$\frac{T_2}{0.9678}$	1	22415 zW ₂ (100-W ₂)MP	$\frac{0.9678 \cdot 22415 \text{ z } \text{X}_2}{(1 - \text{X}_2) \text{ M}_L^P}$
Weight percent, W ₂	$\frac{100}{1 + \frac{22415 \text{ z d}}{\text{M P B}_2}}$	$\frac{\frac{100}{1 + \frac{22415 \text{ z d T}}{273.15 \text{ MPO}_2}}$	$\frac{100}{1 + \frac{22415 \text{ z } 0.9678}{\text{MPT}_2}}$	$\frac{100}{1 + \frac{22415 \text{ z}}{\text{MPK}_2}}$	1	$\frac{\frac{100}{1 + \frac{(1 - X_2)M_L}{MX_2}}}$
Mole fraction, X 2	$\frac{1}{1+\frac{22415 \text{ z d}}{\text{M}_{L}\text{ P B}_{2}}}$	$\frac{1}{1 + \frac{22415 \text{ z d T}}{273.15 \text{ M}_{\text{L}}^{\text{PO}}2}}$	$\frac{1}{1 + \frac{22415 \ z \ 0.9678}{M_L P \ T_2}}$	$\frac{1}{1 + \frac{22415 \text{ z}}{\text{M}_{L} \text{ P K}_{2}}}$	$\frac{\frac{W_2/M}{M}}{\frac{W_2}{M} + \frac{100 - W_2}{M_L}}$	1

Table I. Conversion Factors between Solubility Units

d = density of liquid solvent [g/ml], z = compressibility factor of gaseous solute, T = absolute temperature [K], P = partial pressure of solute [atm], M = molecular weight of solute, M_L = molecular weight of solvent.

temperature curves for the solid aromatic hydrocarbons such as 1,4-dichlorobenzene do not pass through minima around ambient temperature. Thus, the heats of solution do not become zero in this temperature range.

While the theory of isotope effects upon physical properties of compounds has developed considerably during the last three decades, some areas of understanding still involve qualitative descriptions despite a large number of investigations. In this connection, a large amount of work on the physicochemical properties of heavy water and its effect upon other properties has been done (15). The objective of various studies has been the establishment of the effects of nuclear masses upon intermolecular bond energies and upon the physical properties of liquids relative to their molecular structure, temperature, pressure, and chemical behavior (16).

The solubilities of ordinary water and heavy water have been determined in 29 organic liquids and in all cases the heavy water was found less soluble than the ordinary water (17). This can be explained simply by the difference in the total molecular surface areas. With increasing temperature, the solubility differences decrease markedly.

In the case of the solubility of liquid organics in ordinary water and in heavy water, there is good indication that the dissolving power of heavy water is less than that of ordinary water under the same conditions (18). As they do for solubilities of ordinary water and heavy water in organics, the isotopic effects decrease in extent with increasing temperature for organic liquid solubilities in the two solvents. With respect to the mutual solubilities of halogenated benzenes, toluenes, and phenols with ordinary water and heavy water, the available data do not show exceptions or unusual irregularities. In all cases, the usual behavior, as described above, is observed.

The solubility data found in the literature have been reported in various units. While the users of solubility data very often prefer mole fraction or weight percentage concentration units, others have been employed. As a convenience, the appropriate conversion factors between the various concentration units are presented in Table I from Horvath (19). Also, it should be pointed out that some original measurement conversions have been made in order to present the reported data in currently used concentration units. However, no assumed parameters have been involved in these conversions. In addition, the actual reported values have been used to calculate concentrations in two other concentration units. These values are reported routinely to one more significant figure than the measured value to aid the user in further calculations. One should not assume that these calculated concentration values have any greater precision than the originally measured values. Care has been taken to identify the reported concentration values throughout the volume.

Finally, it should be indicated that the halogenated compounds have been organized according to the Hill System. The heavy water follows the ordinary water.

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Preface

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		Ari L. Horvath Imperial Chemical Industries PLC Runcorn, England

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March 1984

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LIQUID-LIQUID SOLUBILITY: INTRODUCTORY INFORMATION

Allan F.M. Barton

The Solubility Data Series is made up of volumes of comprehensive and critically evaluated solubility data on chemical systems in clearly defined areas. Data of suitable precision are presented on data sheets in a uniform format, preceded for each system by a critical evaluation if more than one set of data are available. In those systems where data from different sources agree sufficiently, recommended values are proposed. In other cases, values may be described as "tentative" or "rejected".

This volume is primarily concerned with liquid-liquid systems, but a limited number or related solid-liquid and multicomponent (organic-water-salt) systems have been included where it was 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 artifical 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. Such data have not been found for systems in this volume. Generally, however, data on reported miscibility gap regions and upper and lower critical solution temperatures are included where appropriate and when available.

TERMINOLOGY

In this volume, a mixture (1,2) or a solution (1,2) refers to a single liquid phase containing components 1 and 2. In a mixture, no distinction is made between solvent and solute.

The solubility of a substance is the relative proportion of 1 in a mixture which is saturated with respect to component 1 at a specified temperature and pressure. (The term "saturated" implies the existence of equilibrium with respect to the processes of mass transfer between phases.)

QUANTITIES USED AS MEASURES OF SOLUBILITY

<u>Mole fraction</u> of component 1, x_1 or x(1):

$$x_1 = n_1 / \sum_i n_i = \frac{m_1 / M_1}{\sum (m_i / M_i)}$$

where n_i is the amount of substance (number of moles) of component i, m_i is the mass of substance i and M_i is its molar mass.

<u>Mole per cent</u> of component 1 is $100x_1$

<u>Mass fraction</u> of component 1, w_{1}

$$w_1 = m_1 / \sum_{i}^{\Sigma m_i} i$$

where m_i is the mass of component i.

<u>Mass per cent</u> of component 1 is $100\omega_1$, and may be described as g(1)/100g in which the 100g without a subscript means that it is mass of solute relative to solution and not to solvent. The equivalent terms "weight fraction" and "weight per cent" are not used. The mole fraction solubility is related to the mass fraction solubility in a binary system by

$$x_{1} = \frac{\omega_{1}/M_{1}}{\omega_{1}/M_{1} + (1 - \omega_{1})/M_{2}}$$

<u>Amount</u> of substance concentration of component i in a solution of volume V,

 $c_i = n_i/V$

is expressed in units of mol L^{-1} . The terms "molarity" and "molar" and the unit symbol *M* are not used. However, the unit mol(1)/dm³ is used frequently rather than "molarity" (where the dm³ without a subscript means solution and not solvent volume). Within the range of experimental errors usually encountered in reported solubility measurements, this unit may be taken as equal to "molarity".

<u>Mass ratio</u> is frequently used for a two-component solution in the form g(1)/g(2), mg(1)/g(2), etc. The term "parts per million" (ppm) is not used, but may be expressed as mg(1)/kg or $g(1)/10^6 g$, etc.

<u>Molality</u> of component 1 in component 2 is often used in solid-liquid systems defined as $\overline{m_1} = n_1/n_2M_2$, but is not used in liquid-liquid systems where the distinction between "solute" 1 and "solvent" 2 is inappropriate. The term molality alone in inadequate, and the units (mol kg(2)⁻¹, mmol kg(2)⁻¹) must be stated.

Mole fractions 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.

ORDERING OF SYSTEMS

It is necessary to establish a method of ordering chemical compounds, to be used for the lists of saturating components which define each chemical system. This order is also used for ordering systems within volumes.

The systems are ordered first on the basis of empirical formula according to the Hill system (ref. 2). The organic compounds, within each Hill formula are ordered as follows:

- (i) by degree of unsaturation, then
- (ii) by order of increasing chain length in the parent hydrocarbon, then
- (iii) by order of increasing chain length of hydrocarbon branches, then
- (iv) numerically by position of unsaturation, then
- (v) numerically by position by substitution, then

(vi) alphabetcially by IUPAC name.

C₅H₈

For example,

cyclopentane
2-methy1-1,3-butadiene
1,4-pentadiene
1-pentyne

^C 5 ^H 10	cyclopentane 3-methyl-l-butene 2-methyl-2-butene 1-pentene 2-pentene
C ₅ H ₁₂	2,2-dimethylpropane 2-methylbutane pentane
c5H120	2,2-dimethyl-1-propanol 2-methyl-1-butanol 2-methyl-2-butanol 3-methyl-1-butanol 3-methyl-2-butanol 1-pentanol 2-pentanol 3-pentanol
C ₆ H ₁₂ O	cyclohexanol 4-methyl-l-penten-3-ol 1-hexen-3-ol 4-hexen-3-ol

Deuterated (²H) compounds immediately follow the corresponding ¹H compounds.

GUIDE TO THE COMPILATIONS AND EVALUATIONS

The format used for the compilations and evaluations has been discussed in the Forward. Additional information on the individual sections of each sheet is now provided in the folowing.

"Components". Each component is listed by its IUPAC name (ref. 1), chemical formula according to the Hill system, and Chemical Abstracts Registry Number. Also included are the "Chemical Abstracts" name if this differs from the IUPAC name, and trivial name or names if appropriate. IUPAC and common names are cross-referenced to "Chemical Abstracts" names in the System Index.

"Original Measurements". References are expressed in "Chemical Abstracts" style, journal names being abbreviated, and if necessary transliterated, in the form given by the "Chemical Abstracts Service Source Index" (CASSI). In the case of multiple entries (for example translations) an asterisk indicates the publication used for the data compilation.

"Variables". Ranges of variations of temperature, pressures, etc. are indicated here.

"Prepared by". The compiler is name here.

"Experimental Values". Components are described as (1) and (2), as defined in "Components". The experimental data are presented in the units in the original paper. Thus the temperature is expressed $t/^{\circ}C$ or $t/^{\circ}F$ as in the original, and conversion to T/K is made only in the critical evaluation. However, the author's units are expressed according to IUPAC recommendations (ref. 3,4) as far as possible.

In addition, compiler-calculated values of mole fractions and/or mass per cent are included if the original data do not use these units. 1975 or 1977 atomic weights (ref. 5) are used in such calculations. If densities are reported in the original paper, conversions from concentrations to mole fractions are included in the compilation sheets, 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.

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.

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.

"Methods". An outline of the method is presented, reference being made to sources of further detail if these are cited in the original paper. "Chemical Abstracts" abbreviations are often used in this text.

"Source and Purity of Materials". For each component, referred to as (1) and (2), the following information (in this order and in abbreviated form) is provided if it is available in the original papers:

source and specification

method of preparation

properties

degree of purity

"Estimated Error". If this information was omitted by the authors, and if the necessary data are available in the paper, 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, the type of apparatus, and other relevant information. Methods used by the compilers for reporting estimated errors are based on the papers of Ku and Eisenhart (ref. 6).

"References". These are the references (usually cited in the original paper) which the compiler considers particularly useful in discussing the method and material.

"Evaluator". The information provided here is the name of the evaluator, the evaluator's affiliation, and the date of the evaluation.

"Critical Evaluation". The evaluator aims, to the best of his or her ability, to check that the compiled data are correct, to assess their reliability and quality, to estimate errors where necessary, and to recommend numerical values. The summary and critical review of all the data supplied by the compiler include the following information:

(a) Critical text. The evaluator produces a text evaluating all the published data for the particular system being discussed, reviewing their merits or shortcomings. Only published data (including theses and reports) are considered, and even some of the published data may only be referred to in this text if it is considered that inclusion of a data compilation sheet is unjustified.

(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 in the compilation sheets, stating the limits within which it should be used.

(c) Graphical summary. This may be provided in addition to tables and/or fitting equations.

(d) *Recommended values*. Data are *recommended* if the results of at least two independent experimental groups are available and are in good agreement, and if the evaluator has no doubt as to the adequacy and reliability of the experimental and computational procedures used.

Data are reported as *tentative* if only one set of measurements is available, or if the evaluator is uncertain of the reliability of some aspect of the experimental or computational method but judges that it should cause only minor error, or if the evaluator considers some aspect of the computational or experimental method undesirable but believes the data to have some value in those instances when an approximate value of the solubility is needed.

Data determined by an inadequate method or under ill-defined conditions are *rejected*, the reference being included in the evaluation together with a reason for its rejection by the evaluator.

(e) *References*. All pertinent references are listed here, including all those publications appearing in the accompanying compilation sheets and also those which have been rejected and not compiled.

(f) Units. The final recommended values are reported in SI units (ref. 3). It should be noted that in most cases the rounded absolute temperature values (e.g. 298 K) actually refer to 298.15 K, etc., although very few solubilities are known with such precision that the differences are significant.

Continuation Sheets. These are used for both compilations and evaluations, and include sections listing the "Components" and also the "Original measurement" or "Evaluator" as well as the word "continued". Compilation continuation sheets may include a section headed "Comments and/or Additional Data".

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COMPONENTS :	EVALUATOR:
(1) Hexachlorobenzene; C ₆ Cl ₆ ; [118-74-1]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku
(2) Water; H ₂ O; [7732-18-5]	September 1982

CRITICAL EVALUATION:

Little is known about the solubility of hexachlorobenzene in water at normal temperatures despite the fact that large quantities of this compound are used commercially. In recent times, however, toxicologists and environmental chemists have begun to study the aqueous solubilities of chlorinated aromatic hydrocarbons in connection with pollution analyses. Currently, solubility data for this compound at ordinary temperatures exist in two published papers. A third work reports the solubility of hexachlorobenzene at higher temperatures.

The work of Weil, Dure, and Quentin (1) discusses the determination of extremely small solubilities. Their saturation process was carried out by a chromatographic technique to avoid contamination on glass surfaces and formation of possible colloidal aggregates of the solute. In their procedure, Mg silica gel, overcoated with the solute, was eluted with water. However, the analytical procedure was not reported, a factor which makes it difficult to estimate the precision of the measured values. It is known that certain problems and complications arise in the course of the determination of solubilities of this magnitude. First of all, the effects of a third component (glass or silica gel) may be significant. Also, the purity of the solvent (water) can exert a dominant influence on results in cases of highly diluted, saturated systems. In addition to distillation and deionization, the solvent should also be degassed, a procedure not done for the reported measurements. For such reasons, the values of Weil et al. may involve large absolute errors.

Hollifield (2) applied a nephelometric procedure for the estimation of the solubility of hexachlorobenzene in water. His analytical method involved the preparation of a standard curve by successive dilution of an acetone solution of the test substance with water. The turbidities of the solutions were measured nephelometrically. Extrapolation of this curve to an intensity of the reagent blank gave an estimate of the solubility in water. The solubility value reported for 297 ± 2 K was 110 µg(1)/kg. The method evidently gives rough estimates of the solubilities and it has the advantage of being rapid. The lack of precision in the procedure can be appreciated from a review of the reported literature values. Consequently, these solubility values are not considered further in this evaluation.

Sharov (3) has reported the solubility of hexachlorobenzene in water at higher temperatures. Two variables (temperature and pressure) make a comparison with other solubility data difficult. Only the value referred to 353 K was probably measured at normal pressure. Even so, its value of 10 mg(1)/kg(2) cannot be used to support the solubility value of Weil et al.

The following solubility value for hexachlorobenzene in water is based upon the value reported by Weil et al. However, for the reasons stated, the value should be considered doubtful.

T/K	10^8 mol(1)/dm ³	10 ⁶ g(1)/kg	$10^{10}x(1)$
298.15	1.8	5.0	3.2

REFERENCES

1. Weil, L.; Dure, G.; Quentin, K-E. Wasser Abwasser Forsch. <u>1974</u>, 7, 169-75.

2. Hollifield, H.C. Bull. Environ. Contam. Toxicol. 1979, 23(4/5), 579-86.

3. Sharov, V. G. V. Sb. Dostizheniya Nauk-Neftekhim-Vam. 1975, 138-41.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Hexachlorobenzene; C ₆ Cl ₆ ; [118-74-1]	Weil, L.; Dure, G.; Quentin, K-E. Wasser Abwasser Forsch. 1974, 7, 169-75.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{6}g(1)/dm^{3}$ a $10^{8}mol(1)$	$/dm^{3 b}$ $10^{10}x(1)^{b}$
25 5.0 1.76	3.17
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturation process involved a chromato- graphic technique. A 600 mg Florisil powder	C ₆ Cl ₆ : Source and purity not specified.
sample (15.5% MgO, 84.0% SiO, 0.5% Na ₂ SO ₄), 60/100 mesh, was overcoated with 10 mg solute	H ₂ 0: Twice distilled.
in 10 ml acetone solution. After the acetone was removed by evaporation, the solid mix-	
ture was transferred to a column (5 mm dia- meter, 7 cm length) and the solute was eluted	
from the Mg silica gel coating with water. The organic solute was then extracted from the water solution. The procedure for deter-	
mining the composition of the extracted sam-	
ple was not specified.	ESTIMATED ERROR: Solubility: >20% (This evaluation is based on the table I in the original paper. In the table the effect of the ratio of the organic
	table the effect of the ratio of the organic compound and Florisil was tested with five determinations in each case.) Temperature: ±0.2 K (evaluator).
	Temperature: ±0.2 K (evaluator). REFERENCES:

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	Vesala, Department of Chemistry and chemistry, University of Turku.	
(2) Water; H ₂ ⁰ ; [7732-18-5] Sep	chemistry, University of lurku.	
	September 1982.	

CRITICAL EVALUATION:

The solubility of pentachlorobenzene in water has been measured by Yalkowsky, Orr, and Valvani (1) and by Banerjee, Yalkowsky, and Valvani (2). The measurements differ primarily in the analytical procedures involved. The principal motivation for the studies was to obtain certain correlations for solubilities of halogenated benzenes in water.

In the former measurements (1), a conventional experimental procedure was employed. Commercial reagents without further purification were used for the measurements which were done at room temperature ($25 \pm 1^{\circ}C$). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (3), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The latter investigation (2) was based upon a radiochemical analysis. Although temperature regulation and time for equilibration seemed to have received sufficient attention, the result deviates far more than 10 percent from the value reported in (1). The accuracy of the latter measurements also depends upon the radiochemical purity of the substrate, a property which was not tested. The uncertainty of radiochemical purity may account in part for the deviation from the spectrophotometrically determined value.

The solubility of pentachlorobenzene in water is reported here as a tentative value based upon the spectrophotometric determinations (1):

T/K	10^{6} mol(1)/dm ³	10 ⁴ g(1)/kg	$10^{8}x(1)$
298.15	2.2	5.52	3.95

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environ. Sci. Technol. <u>1980</u>, 14(10), 1227-9.
- 3. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :				
	ORIGINAL MEASUREMENTS:			
(1) Pentachlorobenzene; C ₆ HCl ₅ ; [608-93-5]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.			
(2) Water; H ₂ O; [7732-18-5]				
-				
VARIABLES:	PREPARED BY:			
One temperature	A. Vesala			
EXPERIMENTAL VALUES:				
EXPERIMENTAL VALUES:				
4 3 9 6	3 h 8 a			
$t/^{\circ}C$ $10^{4}g(1)/dm^{3}a$ $10^{6}mol(1)$	$/dm^{5}$ 10 ⁵ x(1) ⁴			
25 5.51 2.2	3.95			
a. Calculated by F. W. Getzen.				
b. Reported.				
AUXILIARY	INFORMATION			
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:			
A small excess of solute in water was agi-	C ₆ HCl ₅ : Commercial reagent (Aldrich or			
tated for a period of 4-48 hours (the exact	Eastman), used as received.			
time for equilibration was not reported in single cases) and then filtered. The satu-	H ₂ 0: Source and purity not specified.			
rated solution was extracted with a small	2			
volume of methylene chloride which was then				
assayed spectrophotometrically. At least two independent determinations were carried				
out.				
	ESTIMATED ERROR:			
	Solubility: ±10% (authors).			
	Temperature: ±1 K (authors).			
	REFERENCES :			

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Pentachlorobenzene; C ₆ HCl ₅ ; [608-93-5]	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. <i>Environ. Sci. Techn.</i> 1980, <i>14(10)</i> , 1227-9.
(2) Water; H ₂ 0; [7732-18-5]	1227-9.
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁶ mol(1)	$/dm^{3 b} 10^{8} x(1)^{a}$
25 1.332 5.32	7.015
a. Calculated by F. W. Getzen b. Reported.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The equilibrium was performed in sealed	SOURCE AND PURITY OF MATERIALS: C ₂ HCl ₅ : Commercial reagent, the ¹⁴ C-
stainless steel centrifuge tubes with con- stant or intermittent shaking. The equili-	labeled compound was purchased by NEN, the nonlabeled one by Aldrich.
brium was generally complete within 1 week. The mixture was then centrifuged for 60	H ₂ 0: Distilled water.
minutes after which aliquots of the solution were removed for analysis either by a pipet	
or syringe. Liquid scintillation counting with ¹⁴ C-labelled solute was employed in the	
solubility determinations. The entire pro-	
cedure was carried out at least twice and each analysis was also conducted in dupli-	
cate.	ESTIMATED ERROR: Solubility: ±4.9% (std. deviation estimated
	by authors). Temperature: ±0.2 K (equilibration) ±0.3 K (centrifugation).
	REFERENCES:

COMPO	DNENTS:	EVALUATOR:
(1)	Pentachlorophenol; C ₆ HCl ₅ 0; [87-86-5]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

CRITICAL EVALUATION:

The solubility of pentachlorophenol in water was first reported by Carswell and Nason (1) in 1938 in connection with their study of properties and uses of this compound. The determinations were done very rapidly and the method, the classical "synthetic" one, allowed the solubility measurements to be made at various temperatures. When the reported values are fit by linear regression to the logarithm of solubility versus the reciprocal of Absolute temperature, the following equation results:

$$\log_{10}(S_1(g(1))/kg) = (3.517 \pm 0.239) - (1589 \pm 74)/T$$
 [1]

The coefficient of correlation for equation [1] is 0.997. The calculated value for the solubility of pentachlorophenol from equation [1] is 0.0154 g(1)/kg at 298.15 K. As is evident, the magnitude of the standard error estimate is in the order of 6-7 percent.

Two other reports on the solubility of pentachlorophenol in water take into account the variation of the solubility with hydrogen ion activity. The earlier work published in 1955 by Blackman, Parke, and Garton (2) reported the pH value at which the solubility was measured. The later study by Toyota and Kuwahara (3) in 1967 provided the variation of solubility with pH in water, in aqueous KCl solutions, and in aqueous NH₄Cl solutions. Also provided in their work was an empirical equation for the solubility in water with respect to pH.

The ability of pentachlorophenol to protolyze in water makes the question of its solubility somewhat complicated. In order to establish the solubility behavior generally, one must measure solubilities with pH as one of the variables. The information provided in (2) and (3) is, however, too limited to allow further calculations in this respect. Thus, in reality, it is practical to evaluate and report only the total solubility of pentachlorophenol/phenoxide at the pH prevailing in the saturated solution. This total solubility can be compared with the value obtained from equation [1].

Some critical observations can be made concerning the measurements of Toyota and Kuwahara. First of all, no description of the sources and purities of reagents was provided. Also, the method of analysis was gravimetric based upon the assumption that the pentachlorophenol was insoluble at pH values lower than 3. That is, the saturation value was established by weighing the pentachlorophenol precipitated from a saturated solution with its pH suppressed below 3. Thus, the method rejected the solubility of pentachlorophenol at pH values lower than 3. Such a method permits the determination of total concentration of saturated solutions only at elevated pH values (from pH 6 and up; pH 5 seems too low). The accuracy of the method is not sufficient to determine the cosolute effects of urea, KCl, and NH₂Cl either. This can be seen clearly from the plots of $log(s/s_0)$ versus the concentration of the cosolute. The complicated behavior is understandable in view of the fact that the cosolutes may have dissimilar effects on the phenol and the phenoxide ion. Therefore, the values must be discarded and the solubilities in water must be checked against other values available. Blackman et al. reported the total solubility of pentachlorophenol at pH 5.1. This hydrogen ion activity was established by the use of a phosphate buffer in amounts which may have been insufficient to affect the solubility. Therefore, this value can be taken as comparable with the value of Carswell and Nason provided the pH of the solution is taken to be in the range 4.5-5.5.

On the basis of the three solubilities available, 1.4×10^{-2} g(1)/kg from (1), a 9.6 x 10^{-3} g(1)/kg from (2), and 1.4×10^{-2} g(1)/kg from (3), the following is a tentative solubility value. (The last of these three values was calculated from the equation provided in (3) and it refers to 300 K.) Also, it must be emphasized that, as for any protolyzing solute, the solubility of pentachlorophenol in water is affected to a degree by changes in pH.

т/к	10^{5} mol(1)/dm ³	10 ² g(1)/kg	$10^{7}x(1)$	
298.15	3.7	1.0	6.8	

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COMPONENTS :	EVALUATOR:		
<pre>(1) Pentachlorophenol; C₆HCl₅0; [87-86-5]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.		
(2) Water; H ₂ 0; [7732-18-5]	November 1979.		
CRITICAL EVALUATION: (Continued)			

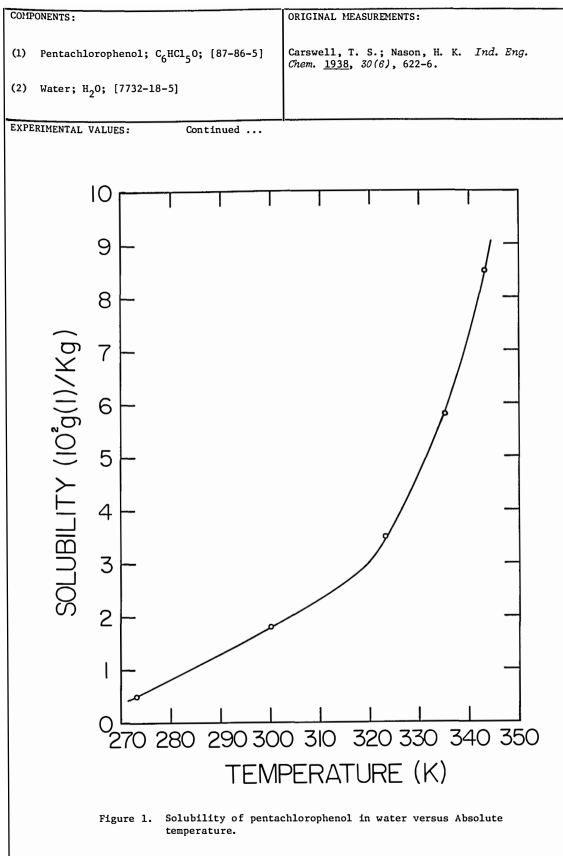
REFERENCES

1. Carswell, T. S.; Nason, H. K. Ind. Eng. Chem. 1938, 30(6), 622-6.

2. Blackman, G. E.; Park, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

3. Toyota, H.; Kuwahara, M. Nippon Dojohiryogaku Zasshi 1967, 38(2,3), 93-97.

COMPONENTS :	ORIGINAL MEASUREMENTS:	
 Pentachlorophenol; C₆HCl₅0; [87-86-5] Water; H₂0; [7732-18-5] 	Carswell, T. S.; Nason, H. K. Ind. Eng. Chem. <u>1938</u> , 30(6), 622-6.	
VARIABLES:	PREPARED BY:	
Temperature: 0 - 70°C	A. Vesala	
EXPERIMENTAL VALUES:		
t/°C $10^2 g(1)/kg^{a}$ $10^5 mol(1)$		
0 0.5 1.9	3.4	
27 1.8 6.76	12.2	
50 3.5 13.1	23.7	
62 5.8 21.8	39.2	
70 8.5 31.9	57.5	
b. Calculated by F. W. Getzen. Measurements are shown graphically in	Figure 1. Continued	
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A known amount of pure pentachlorophenol was added to cold water. Then, the tempera- ture was raised slowly with agitation until dissolution was complete.	C ₅ HCl ₅ O: Source and purity not specified. H ₂ O: No specifications given.	
	ESTIMATED ERROR: Solubility: <6% (estimated by the evaluator on the basis of the smoothed curve of log c vs. 1/T, where c denotes solubility).	
	REFERENCES:	



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Pentachlorophenol

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Pentachlorophenol; C ₆ HCl ₅ O; [87-86-5]	Blackman, G. E.; Parke, M. H.; Garton, G.
	Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	
One pH: 5.1	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)	$/dm^{3} b = 10^{7} x(1)^{a}$
25 9.59 3.6	6.51
a. Calculated by F. W. Getzen.	
a. Calculated by F. W. Getzen.b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS;
The samples were equilibrated in a thermostat	C ₆ HCl ₅ O: Probably a commercial reagent.
bath with intermittent shaking over periods	6 5 . riobably a commercial reagenc.
of 3-4 weeks. During the equilibration time,	H ₂ 0: Distilled water.
the pH values of the solutions were control-	-
led by dropwise addition of phosphate buffer solution. The analysis of the solute con-	
centration in the saturated samples was done	
spectrophotometrically either directly or	
by using proper colorizing agents.	
	ESTIMATED ERROR:
	Solubility: <5% (evaluated on the basis of
	the reported results of the two
	techniques of analysis).
	REFERENCES:

COMPONENTS :		<u> </u>	OPICINAL	MEASUREMENTS :	
					Nippon Dojohiryogaku
 Pentachlorophenol; C₆HCl₅0; [87-86-5] Potassium chloride; KCl; [7447-40-7] 		Zasshi 1	<u>967</u> , <i>38(2,3)</i> , 93	-97 .	
		4/-40-/]			
(3) Water; H ₂ C	; [//32-18-5]				
VARIABLES:	······································		PREPARED	BY:	
pH, concentrati	ons of KCl at 27°C		H. Ohtak		
EXPERIMENTAL VAL		All Measure	ments at 2	27°C	
. 3	Initial pH = 5.0		3 h	6 a.c	d
mol(KCl)/dm	$10^2 g(1)/dm^3 a$			$10^{6}x(1)^{a,c}$	pH ^d
1.0	1.68	6.3		1.13	5.32
0.50	2.32	8.7		1.56	5.45
0.20	1.65	6.2		1.12	5.25
0.10	2.08	7.8	l	1.41	5.34
0.05	3.20	12		2.17	5.28
0.02	2.16	8.1		1.46	5.20
0.0	3.20	12		2.17	5.35
	Initial pH = 6.	0			
mol(KCl)/dm ³	10g(1)/dm ³ a)/dm ^{3 b}	10 ⁶ x(1) ^{a,c}	pH d
1.0	1.01	3.8	l	6.81	5.96
0.50	0.879	3.3	l	5.93	5.89
0.20	0.985	3. 7	,	6.67	6.07
	<u></u>			C	ontinued
		AUXILIARY			
METHOD/APPARATUS	S/PROCEDURE: nol and various amo	unts of	SOURCE AN	D PURITY OF MATE	RIALS:
NaOH and KC1 we	ere dissolved in wa lutions were left	ter at	Nothing specified.		
with occasional	stirring (once a	day for 1			
	the undissolved pen ered off and the p				
	es were measured wi each aliquot of th				
was added hydro	ochloric acid to ma The precipitates	intain their			
sulted were age		which ie-			
			ESTIMATED) ERROR: No esti	mation was given for
			errors in ever, the	n the solubility e reported values	measurements. How- s of S/S° (S the
			solubili tions and	ty of pentachlor d S its solubil	ophenol in KCl solu- ity in pure water) do oth curve, but rather
			spread.	Therefore, the of the larger than a	errors may be esti-
			REFERENCE	-	
			 Nose, K.; Fukunaga, K. Noyaku Seisan Gijutsu 1952, 6, 30. 		
			<i>u</i> 1	<u> </u>	

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Pentachlorophenol

COMPONENTS:		(DRIGINAL MEASUREMENTS:			
(2) Potassium	cophenol; C ₆ HCl ₅ O; chloride; KCl; [744 D; [7732-18-5]		Toyota, H.; Kuwahara, M. Z <i>asshi</i> <u>1967</u> , <i>38(2,3)</i> , 93	Nippon Dojohiryogaku -97.		
EXPERIMENTAL VA	ALUES: Continue	I_	**************************************	·····		
	Initial pH = 6					
3	-		3 6 9 0	đ		
mol(KCl)/dm ³	$10g(1)/dm^{3}$ a	10 [°] mol(1)	$/dm^{3 b}$ $10^{6}x(1)^{a,c}$	pH d		
0.10	1.17	4.4	7.94	6.00		
0.05	1.39	5.2	9.38	5.95		
0.02	1.54	5.8	10.5	5.98		
0.0	0.613	2.3	4.15	5.70		
	Taitial -11 - 7	0				
	Initial pH = 7.		$/dm^{3 b} 10^{5} x(1)^{a,c}$	d		
mol(KCl)/dm ³	g(1)/dm ^{- 1}	10 ⁻ mol(1)	$/dm^{-1}$ $10^{-}x(1)^{-1}$	pH d		
1.0	1.01	3.8	6.81	6.81		
0.50	1.07	4.0	7.20	6.79		
0.20	1.41	5.3	9.56	6.82		
0.10	1.25	4.7	8.49	6.77		
0.05	1.46	5.5	9.94	6.86		
0.02	1.33	5.0	9.03	8.82		
0.0	2.05	7.7	13.9	6.92		
	Initial pH = 8.	0				
mol(KCl)/dm ³	$10^{-1}g(1)/dm^{3}a$	10 ² mol(1)	$/dm^{3 b} 10^{3} x(1)^{a,c}$	pH d		
1.0	1.68	6.3	1.15	7.78		
0.50	1.57	5.9	1.08	7.67		
0.20	2.18	8.2	1.51	7.93		
0.10	1.68	6.3	1.15	7.70		
0.05	2.45		1.70	8.35		
		9.2				
0.02	1.92	7.2	1.32	7.67		
0.0 1.94 7.3 1.34 7.80 a. Calculated by F. W. Getzen. b. Reported. c. Calculated mole fraction based upon complete dissociation of KC1. d. pH value of solution equilibrated with precipitate.						
COMMENTS AND/OR	ADDITIONAL DATA:					
Solubilities of pentachlorophenol in water versus hydrogen ion activity, in the absence of KCl, at 27°C were measured and calculated. The stoichiometric amounts of dissolved pentachlorophenol were calculated from mass balance based upon incomplete dissociation. The dissociation constant was given as $K_p = [C_6C_{15}O^-][H^+]/[C_6C_{15}OH] = 3.2 \times 10^{-5} \text{ mol/dm}^3$. This value together with the activity of the hydrogen ion, $\{H^+\}$, as determined from pH measurements was incorporated in an equation for calculating the solubility: $S = \frac{1.796 \times 10^{-6}}{\{H^+\} 0.170} \left[1 + \frac{3.2 \times 10^{-5}}{\{H^+\}} \right]$ Observed and calculated values reported:						
рН	5.0	6.0				
	10^{5} mol(1)/dm ³					
Observed value: Calculate value	4.05 : 5.34	6.0				
		5.2				

Pentachiorophenoi						
COMPONENTS :	ENTS: ORIGINAL MEASUREMENTS:					
(1) Pentachloro	Pentachlorophenol; C ₆ HC1 ₅ 0; [87-86-5]			Toyota, H.; Kuwahara, M. <i>Nippon</i> Dojohiryogaku Zasshi <u>1967</u> , 38(2,3), 93-97.		
(2) Ammonium ch	loride; NH ₄ Cl; [121	25-02-9]				
(3) Water; H ₂ O;	[7732-18-5]					
VARIABLES:						
			PREPARED : H. Ohtak			
ph, concentratio	ons of NH ₄ Cl at 27°C	i	n. Untak	1		
EXPERIMENTAL VALU		Measurem	ents at 27	°C		
2	Initial pH = 5.0	5	3 6	7 ас	đ	
	$10^{2}g(1)/dm^{3}a$	10 [°] mo1(1	L)/dm ⁵		pH ^d	
1.0	0.8390	3.3	15	5.690	5.12	
0.50	1.377			9.335	5.20	
0.05	3.3161	12.	45	22.472		
0.0	3.0657	11.9	51	20.774	5.35	
	Initial pH = 5.5					
$mol(NH,C1)/dm^3$	$\frac{10^2 g(1)}{dm^3} a$	4 10 [°] mol(1	L)/dm ^{3 b}	$10^{6}x(1)^{a,c}$	pH d	
0.05	6.153	2.	31	4.170	5.73	
$mol(NH cl)/dm^3$	Initial pH = 6.0 $10g(1)/dm^3 a$	10 ⁴ mo1(1)/dm ³ b	10^{5} (1) a,c	pH d	
1.0	10g(1)/dm 1.489		59	1.010		
0.50	1.345)5	0.9120	6.05	
0.30	2.9645	11.1		2.0099	6.19	
0.20	2.650	9.9		1. 797	6.14	
0.0		2.2		0.4061		
					Continued	
MEMILOS		AUXILIARI	INFORMATIO			
METHOD/APPARATUS/	PROCEDURE: ol and various amoun	ts of	SOURCE AN	D PURITY OF MAT	ERIALS:	
NaOH and NH,C1 w	vere dissolved in wa	ter at	Nothing specified.			
2/°C and the sol with occasional	utions were left fo sitrring (once a da	r 3 days v for 1				
hour). Then, th	e undissolved penta	chloro-				
of the filtrates	red off and the pH were measured with	values a glass				
electrode. To e	ach aliquot of the	solutions				
Was added hydroc	hloric acid to main an 3. The precipit	tain ates				
which resulted w	vere aged and weighe	d.				
			ESTIMATED	ERROR: No est	imation was given f	or
			errors in	n the solubilit	y measurements. Ho	
			solubili	ty of pentachlo	es of S/S° (S the rophenol in KCl sol	u-
			do not fa	all on a single	lity in pure water) smooth curve, but	
	rati		rather spread. Therefore, the errors may b estimated to be larger than 2-3%.		be	
			REFERENCE			
			I			

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Pentachlorophenol

COMPONENTS:		ORIGINAL MEASUREMENTS:			
(1) Pentachlorophenol; C ₆ HC1 ₅ 0; [87-86-5]			Toyota, H.; Kuwahara, M. <i>Nippon</i> <i>Dojohiryogaku Zasshi</i> <u>1967</u> , <i>38(2,3)</i> , 93-97.		
(2) Ammonium chloride; NH ₄ Cl; [12125-02-9]					
(2) Water; H ₂ O	; [7732-18-5]				
EXPERIMENTAL VALU	ES: Continued	••••			
	Initial pH = 6.5		_		
mol(NH ₄ Cl)/dm ³	10g(1)/dm ^{3 a}	10 ³ mo1(1)	$/dm^{3 b}$ $10^{5}x(1)^{a,c}$	pH d	
1.0	2.352	0.883	1.595	6.46	
0.50	3.9819	1.495	2.7005	6.50	
0.10	5.966	2.24	4.045	6.40	
0.0	6.233	2.34	4.226	6.52	
	Initial pH = 7.0				
mol(NH ₄ Cl)/dm ³	10g(1)/dm ^{3 a}	10 ³ mol(1)	$/dm^{3 b}$ $10^{5}x(1)^{a,c}$	pH d	
1.0	2.9432	1.105		6.97	
0.50	5.114	1.92	3.469	6.98	
0.30	7.777	2.92	5.275	7.03	
0.20	8.736	3.28	5.926	7.10	
0.10	11.13	4.18	7.553	6.86	
0.05	14.76	5.54	10.01	6.92	
0.0	20.40	7.66	13.85	6.92	
	Initial pH = 7.5				
mol(NH ₄ Cl)/dm ³	$g(1)/dm^{3}a$	10^{3} mo1(1)	$/dm^{3 b}$ 10 ⁴ $x(1)^{a,c}$	pH d	
0.10	1.675	6.29	1.137	7.50	
mol(NH ₄ Cl)/dm ³	Initial pH = 8.0 $g(1)/dm^3 a$	10 ³ mol(1)	$/dm^{3 b}$ $10^{5}x(1)^{a,c}$	pH d	
1.0	0.3276	1.23		8.02	
0.50	0.5780	2.17	3.920	8.00	
0.30	0.6339	2.38	4.299	8.05	
0.20	1.247	4.68	8.458	8.01	
0.10	1.851	6.95	12.57	8.01	
0.05	2.903	10.9	19.73	7.98	
0.0	19.39	72.8	133.8	7.80	
 a. Calculated by F. W. Getzen. b. Reported. c. Calculated mole fraction based upon complete dissociation of NH₄Cl. d. pH value of solution equilibrated with precipitate. 					
d. pH value of COMMENTS AND/OR	-	.ca wren pre	cipitate.		
Solubilities of pentachlorophenol in water versus hydrogen ion activity, in the absence of NH_4Cl , at 27°C were measured and calculated. The stoichiometric amounts of dissolved pentachlorophenol were calculated from mass balance based upon incomplete dissociation. The dissociation constant was given as $K_p = [C_6Cl_5O^-][H^+]/[C_6Cl_5OH] = 3.2 \times 10^{-5} \text{ mol/dm}^3$. This value together with the activity of the hydrogen ion, $\{H^+\}$, as determined from pH measurements was incorporated in an equation for calculating the solubility:					
$S = \frac{1.796 \times 10^{-6}}{(H^+)^{0.170}} \left[1 + \frac{3.2 \times 10^{-5}}{(H^+)} \right]$					
Observed and cal pH	culated values repo 5.0	6.0	7.0		
	10 ⁵ mol(1)/dm ³ 1	.0 ⁴ mo1(1)/dm	1^{3} 10 ³ mo1(1)/dm ³		
Observed value:	4.05	6.05	8.85		
1 Coloulate welver	5 3/	6 20	8 90		

8.90

5.34

Calculate value:

6.20

COMPONENTS:		EVALUATOR:
(1)	1,2,4,5-Tetrabromobenzene; C ₆ H ₂ Br ₄ ; [636-28-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,2,4,5-tetrabromobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of ± 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2,4,5-tetrabromobenzene in water is reported here as a tentative value:

т/к	$10^{7} \text{mol}(1)/\text{dm}^{3}$	10 ⁵ g(1)/kg	$10^{9}x(1)$
298.15	1.1	4.34	1.99

REFERENCES

 Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2,4,5-Tetrabromobenzene; C ₆ H ₂ Br ₄ ; [636-28-2]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	
One temperature	PREPARED BY: A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁵ g(1)/dm ³ a 10 ⁷ mol(1)/d	10^{3} b $10^{9}x(1)^{a}$
25 4.33 1.1	1.99
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi- tated for a period of 4-48 hours (the exact time for equilibration was not reported in	C ₆ H ₂ Br ₄ : Commercial product (Aldrich or Eastman), used as received.
single cases) and then filtered. The satu- rated solution was extracted with a small volume of methylene chloride which was then assayed spectrophotometrically. At least two independent determinations were carried out.	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR: Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES :

COMPONENTS:		EVALUATOR:	
(1)	1,2,3,4-Tetrachlorobenzene; C ₆ H ₂ Cl ₄ ; [634-66-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.	
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.	

A single solubility measurement of 1,2,3,4-tetrachlorobenzene in water has been reported by Yalkowsky, Orr and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2,3,4-tetrachlorobenzene in water is reported here as a tentative value:

T/K	10^{5} mol(1)/dm ³	10 ³ g(1)/kg	$10^{7}x(1)$
298.15	2.0	4.33	3.61

REFERENCES

1. Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. 1979, 18(4),

2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 1,2,3,4-Tetrachlorobenzene; C₆H₂Cl₄; [634-66-2]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)/dr	$10^{3} b 10^{7} x(1)^{a}$
25 4.32 2.0	3.61
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi-	⁶ ⁴ ² ⁴
tated for a period of 4-48 hours (the exact	Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The satu-	H ₂ 0: Source and purity not reported.
rated solution was extracted with a small	2
volume of methylene chloride which was then assayed spectrophotometrically. At least	
two independent determinations were carried out.	
out.	
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES :

COMPONENTS:	EVALUATOR:	
<pre>(1) 1,2,3,5-Tetrachlorobenzene; C₆H₂Cl₄; [634-90-2]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.	
(2) Water; H ₂ O; [7732-18-5]	September 1982.	

The solubility of 1,2,3,5-tetrachlorobenzene in water has been measured by Yalkowsky, Orr and Valvani (1) and by Banerjee, Yalkowsky, and Valvani (2). The measurements differ primarily in the analytical procedures involved. The principal motivation for the studies was to obtain certain correlations for solubilities of halogenated benzenes in water.

In the former measurements (1), a conventional experimental procedure was employed. Commercial reagents without further purification were used for the measurements which were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (3), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The latter investigation (2) was based upon a radiochemical analysis. Although temperature regulation and time for equilibration seemed to have received sufficient attention, the result deviates far more than 10 percent from the value reported in (1). The accuracy of the latter measurements also depends upon the radiochemical purity of the substrate, a property which was not tested. The uncertainty of radiochemical purity may account in part for the deviation from the spectrophotometrically determined value.

The solubility of 1,2,3,5-tetrachlorobenzene in water is reported here as a tentative value based upon the spectrophotometric determinations (1):

т/к	10^{5} mol(1)/dm ³	10 ³ g(1)/kg	$10^{7}x(1)$
298.15	1.6	3.46	2.89

REFERENCES

- 1. Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environ. Sci. Technol. <u>1980</u>, 14(10), 1227-9.
- 3. Yalkowsky, S. H., Personal Communication, 1979.

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COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 1,2,3,5-Tetrachlorobenzene; C₆H₂Cl₄; [634-90-2]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
3 3 2 5	3 b 7 c
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)/d	10'x(1)
25 3.45 1.6	2.89
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi-	C ₆ H ₂ Cl ₄ : Commercial product (Aldrich or
tated for a period of 4-48 hours (the exact time for equilibration was not reported in	Eastman), used as received.
single cases) and then filtered. The satu-	H ₂ 0: Source and purity not specified.
rated solution was extracted with a small volume of methylene chloride which was then	-
assayed spectrophotometrically. At least	
two independent determinations were carried out.	
	ESTIMATED ERROR: Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES :

ORIGINAL MEASUREMENTS:
Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. <i>Environ. Sci. Techn.</i> <u>1980</u> , <i>14(10)</i> , 1227-9.
PREPARED BY:
A. Vesala
, , , , , , , , , , , , , , , , , , ,
dm^{3} b $10^{7}x(1)$ a
3.361
INFORMATION
SOURCE AND PURITY OF MATERIALS: 14 C ₆ H ₂ Cl ₄ : Commercial reagent, the ¹⁴ C- labeled compound was purchased by NEN, the nonlabeled by Aldrich. H ₂ O: Distilled water. ESTIMATED ERROR: Solubility: ±4.0% (std. deviation estimated by authors). Temperature: ±0.2 K (equilibration), ±0.3 K (centrifugation). REFERENCES:

COMP	ONENTS:	EVALUATOR:
(1)	1,2,4,5-Tetrachlorobenzene; C ₆ H ₂ Cl ₄ ; [95-94-3]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,2,4,5-tetrachlorobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2,4,5-tetrachlorobenzene in water is reported here as a tentative value:

т/к	10 ⁶ mo1(1)/dm ³	10 ⁴ g(1)/kg	$10^8 x(1)$
298.15	2.8	6.06	5.06

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :	
COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,2,4,5-Tetrachlorobenzene; C₆H₂Cl₄; [95-94-3]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁴ g(1)/dm ^{3 a} 10 ⁶ mol(1)/d	$m^{3 b} 10^8 x(1)^a$
25 6.05 2.8	5.06
a. Calculated by F. W. Getzen. b. Reported.	
b. Reported.	
	······
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi- tated for a period of 4-48 hours (the exact	C ₆ H ₂ Cl ₄ : Commercial reagent (Aldrich or Eastman), used as received.
time for equilibration was not reported in	
single cases) and then filtered. The satu- rated solution was extracted with a small	H ₂ 0: Source and purity not specified.
volume of methylene chloride which was then	
assayed spectrophotometrically. At least two independent determinations were carried	
out.	
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES:
l	

COMP	ONENTS:	EVALUATOR:
(1)	2,3,4,6-Tetrachlorophenol; C ₆ H ₂ Cl ₄ O; [58-90-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

One study has been reported in the literature concerning the solubility of 2,3,4,6tetrachlorophenol in water by Blackman, Parke, and Garton (1). Their concern was the relation of physical properties of certain substituted phenols to their physiological activities. In order to standardize the measuring conditions, they regulated the pH of all their saturated solutions to a value of 5.1 by the slight addition of a phosphate buffer. The electrolytic effects of this buffer addition on the equilibrium solubility can be neglected. However, the effects of pH changes upon the solubility are more complicated and should not be overlooked. In order to establish the solubility behavior generally, one must measure solubilities with pH as one of the variables. Such data, however, do not exist for 2,3,4,6-tetrachlorophenol. The only alternative here is to consider the total solubility of the compound in water, considering total solubility as the sum of the concentrations of the phenol and the phenolate ion. In the final analysis, the overall influence of pH upon the solubility is controlled by the acidity constant of the 2,3,4,6-tetrachlorophenol which was reported to be 5.3 by Blackman et al. Evidently the regulation of pH to a value of 5.1 in this case provides a higher concentration of the phenolate ion relative to the undissociated phenol as compared to that in a solution simply saturated with the pure phenol. Thus, the solubility value at the pH of 5.1 should have a somewhat higher value than that for the saturated solution of the pure phenol.

The information concerning the purity and further treatment of the reagents was also incomplete. The experimental procedures seem quite satisfactory however. The following solubility value based entirely upon the work reported by Blackman et al., must be considered as doubtful.

т/к	10^4 mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
298.15	7.9	1.8	1.4

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

2,3,4,6- I etrac	niorophenol 25
COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,3,4,6-Tetrachlorophenol; C₆H₂Cl₄0; [58-90-2]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	
One temperature One pH: 5.1	PREPARED BY: A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm	$3 b 10^5 x(1)^a$
25 1.83 7.9	1.43
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The samples were equilibrated in a thermostat bath with intermittent shaking over periods of 3-4 weeks. During the equilibration time, the pH values of the solutions were control- led by dropwise addition of phosphate buffer solution. The analysis of the solute con- centration in the saturated samples was done spectrophotometrically either directly or by using proper colorizing agents.	SOURCE AND PURITY OF MATERIALS: C ₆ H ₂ Cl ₄ O: Not specified, probably a commercial product.

COMPO	DNENTS:	EVALUATOR:
(1)	1,2,4-Tribromobenzene; C ₆ H ₃ Br ₃ ; [615-54-3]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ O; [7732-18-5]	November 1979.

A single solubility measurement of 1,2,4-tribromobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of ± 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2,4-tribromobenzene in water is reported here as a tentative value:

т/к	10^{5} mol(1)/dm ³	10 ² g(1)/kg	$10^{7}x(1)$
298.15	3.2	1.01	5.78

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,2,4-Tribromobenzene; C ₆ H ₃ Br ₃ ; [615-54-3]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
-	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{5}mol(1)/dt$	$m^{3 b} 10^{7} x(1)^{a}$
25 1.01 3.2	5.78
a. Calculated by F. W. Getzen.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi-	$C_6H_3Br_3$: Commercial reagent (Aldrich or 66)
tated for a period of 4-48 hours (the exact	Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The satu-	$H_{2}0$: Source and purity not specified.
rated solution was extracted with a small	² ² ² ²
volume of methylene chloride which was then assayed spectrophotometrically. At least	
two independent determinations were carried	
out.	
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES :

COMP	ONENTS:	EVALUATOR:
(1)	1,3,5-Tribromobenzene; C ₆ H ₃ Br ₃ ; [626-39-1]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,3,5-tribromobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,3,5-tribromobenzene in water is reported here as a tentative value:

т/к	10^{6} mol(1)/dm ³	10 ⁴ g(1)/kg	$10^8 x(1)$
298.15	2.5	7.89	4.52

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,3,5-Tribromobenzene; C₆H₃Br₃; [626-39-1]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
2	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁴ g(1)/dm ^{3 a} 10 ⁶ mol(1)/dm	$m^{3 b} 10^8 x(1)^a$
25 7.87 2.5	4.52
a. Calculated by F. W. Getzen.	
b. Reported.	
·	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi-	$C_{63}H_{3}Br_{3}$: Commercial product (Aldrich or
tated for a period of 4-48 hours (the exact	Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The satu-	H ₂ 0: Source and purity not specified.
rated solution was extracted with a small	2
volume of methylene chloride which was then assayed spectrophotometrically. At least	
two independent determinations were carried out.	
out.	
	ESTIMATED ERROR:
	Solubility: ±10% (authors)
	Temperature: ±1 K (authors).
	REFERENCES :

	esala, Department of Chemistry and hemistry, University of Turku.
(2) Water; H ₂ O; [7732-18-5] Novem	mber 1979.

The only two available published works concerning the solubility of 2,4,6-tribromophenol in water are old and information provided in these works concerning the methods of analysis is incomplete. Also, the purity of the reagents used in the measurements is open to question. However, the solubility value of 70 mg(1)/kg(2) at 288 K determined by Werner (1) in 1884 agrees quite well with the value of 76 mg(1)/kg(2) at 17-18°C measured by Ogston (2). (These concentrations have been established in the same units by assuming the density of the saturated solution to be 1.0 g/cm³.)

It should be noted that the solubility of 2,4,6-tribromophenol in water is considerably dependent upon the acidity of the solution. Here, it is assumed that the solubility established refers to that of a solution having a pH value prevailing for saturated 2,4,6-tribromophenol in water.

The solubility of 2,4,6-tribromophenol in water is reported here as a doubtful value:

T/K	10^4 mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
283.15	2.1	7.0	3.8

REFERENCES

1. Werner, E. Ann. Chim. Phys. Ser. G 1884, 572-3.

2. Ogston, A. G. J. Chem. Soc. <u>1936</u>, Part II, 1713.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 2,4,6-Tribromophenol; C ₆ H ₃ Br ₃ O; [118-79-6]	Werner, E. Ann. Chim. Phys. Ser. G <u>1884</u> , 572-3.
(2) Water; H ₂ 0; [7732-18-5]	
:	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3}a = 10^{4}mol(1)/dm^{3}a$	$m^{3} b 10^{6} x(1)^{b}$
15 7 2.1	3.8
	510
a Departed	
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Information concerning the equilibration	C ₆ H ₃ Br ₃ O: Synthesized product, reported
procedure was incomplete. The analysis was	melting point 92°C.
done by titration of the saturated solutions with a standard barium hydroxide solution.	H _a O: Source and purity not specified.
with a standard barium hydroxide solution.	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR:
	REFERENCES:

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4,6-Tribromophenol; C₆^H3^{Br}3^O; [118-79-6]</pre>	Ogston, A. G. J. Chem. Soc. <u>1936</u> , Part II, 1713.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ² g(1)/dm ³ a 10 ⁴ mol(1)/dm	$n^{3 b} 10^{6} x(1)^{a}$
17-18 7.61 2.3	4.15
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
There were no specifications in the original paper about the experimental methods. Also,	C ₆ H ₃ Br ₃ O: Source and purity not specified.
the temperature at which the measurements were made was not well defined.	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR:
	REFERENCES :

COMPO	DNENTS:	EVALUATOR:
(1)	1,2,3-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [87-61-6]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,2,3-trichlorobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principla motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature $(25 \pm 1^{\circ}C)$. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2,3-trichlorobenzene in water is reported here as a tentative value:

T/K	10^4 mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
298.15	1.7	3.09	3.07

REFERENCES

- 1. Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2,3-Trichlorobenzene; C ₆ H ₃ Cl ₃ ;	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.
[87-61-6]	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{4}mol(1)/dm^{3}$	$m^{3 b} 10^{6} x(1)^{a}$
25 3.08 1.7	3.07
a Calculated by F. U. Cataoa	
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi-	C ₆ H ₃ Cl ₃ : Commercial product (Aldrich or
tated for a period of 4-48 hours (the exact time for equilibration was not reported in	Eastman), used as received.
single cases) and then filtered. The satu- rated solution was extracted with a small	H ₂ 0: Source and purity not specified.
volume of methylene chloride which was then	
assayed spectrophotometrically. At least two independent determinations were carried	
out.	
]	
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES :

COMP	ONENTS:	EVALUATOR:
(1)	l,2,4-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [120-82-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) !	Water; H ₂ 0; [7732-18-5]	January 1983.

The solubility of 1,2,4-trichlorobenzene in water has been measured by Meleshchenko (1), Yalkowsky et al. (2), and Chiou et al. (3). However, there appears to be a serious dis-Crepancy between the three reported solubility values. According to the authors, the error estimates in their measurements were \pm 8 percent for Meleshchenko, \pm 10 percent for Yalkowsky et al., and \pm 5 percent for Chiou et al. Consequently, in view of these uncertainties, the recommended solubility value must be taken as the arithmetical mean for the three reported measured values.

With reference to Figures 3 and 4 in the Introduction, the recommended solubility of 1,2,4-trichlorobenzene in water given below for 298.15 K is within an acceptable margin of expectation.

The single data point available for the solubility of water in 1,2,4-trichlorobenzene at 298.15 K by Eidinoff (4) is accepted as a tentative value.

т/к	10^{3} mol(1)/dm ³	10 ² g(1)/kg	$10^4 x(1)$
298.15	3.05	3.79	3.82
т/к	10^2 mol(2)/dm ³	10g(2)/kg	$10^{3}x(2)$
298.15	1.63	2.02	2.03

REFERENCES

1. Maleshchenko, K. F. Gigiena i Sanit. 1960, 25(5), 54-57.

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 3. Chiou, C. T.; Schmedding, D. W.; Maines, M. Environ. Sci. Technol. <u>1982</u>, 16(1), 4-10.
- 4. Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M., Urey, H. C., Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u>; Part II, Chapter 7, pp 129-44.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) Water; H₂0; [7732-18-5] (2) 1,2,4-Trichlorobenzene; C₆H₃Cl₃; [120-82-1]</pre>	Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M., Urey, H. C., Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II, Chapter 7, pp 129- 44.
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ² mol(1)/kg	b $10^3 x(1)^{a}$
25 2.0232 1.123	2.0340
a. Calculated by compiler. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Addition of 2 ml H ₂ O to 100 ml C ₆ H ₃ Cl ₃ in an equilibration flask was performed in a dry box. The closed flask was then placed in a water thermostat bath and the assembly was shaken for 90 min. The samples for analysis were taken with a sampling pipet while the equilibrium temperature was maintained. The water content in the sample was determined by a modified Karl Fischer titration.	SOURCE AND PURITY OF MATERIALS: C ₆ H ₃ Cl ₃ : Purified and dried before use. H ₂ O: Distilled. ESTIMATED ERROR: Solubility: Average dev. ±0.00006 mol(1)/ kg.
	Temperature: ±0.05 K. REFERENCES:

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2,4-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [120-82-1]	Maleshchenko, K. F. <i>Gigiena i Sanit</i> . <u>1960</u> , 25(5), 54-57.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{4}mol(1)/d$	$m^{3 b} 10^{6} x(1)^{c}$
19 3.0 1.65	2.98
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
A 100 mg trichlorobenzene in 1 liter water	C ₆ H ₃ Cl ₃ : Not specified.
mixture was shaken periodically for 5 days at 18-20°C. After filtration, a determined quantity of ether was added and extracted three times. The ether was evaporated and Oxidation of the residue was accomplished with chlorosulphonic acid. The chlorine	H ₂ O: Twice distilled before use.
content was determined by iodometric titra- tion.	
	ESTIMATED ERROR:
	Solubility: ±8%.
	Temperature: ±1 K.
	REFERENCES:

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2,4-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [120-82-1]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3}a = 10^{4}mol(1)/dm^{3}$	10^{3} b $10^{6}x(1)$ a
25 3.466 1.91	3.451
a. Calculated by compiler. b. Reported.	
l	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,2,4-trichlorobenzene in water was agitated at room temperature for about 24 hours and then filtered. The fil-	C ₆ H ₃ Cl ₃ : Aldrich commercial grade, used as received.
trate was diluted and assayed spectrophoto- metrically. The determination was carried	H ₂ 0: Deionized.
out in duplicate.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES :

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,2,4-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [120-82-1]	Chiou, C. T.; Schmedding, D. V.; Manes, M. Environ. Sci. Technol. <u>1982</u> , 16(1), 4-10.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES :	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{4}mol(1)/d$	$m^{3 b} 10^{6} x(1)^{a}$
25 4.881 2.69	4.861
Note: The solubility was also reported "unpublished results".	in (1), as
a. Calculated by compiler. b. Reported.	
	INFORMATION
METHOD /APPARATUS/PROCEDURE: An excess of 1,2,4-trichlorobenzene was	SOURCE AND PURITY OF MATERIALS:
equilibrated with water in screwcapped bot- tles in a reciprocal shaker for 24 hours.	C ₆ H ₃ Cl ₃ : Not specified. H ₂ O: Distilled.
After two days settling, samples were taken from the solution for analysis by gas chroma-	2
tography. The chromatograph was equipped with a Ni ⁶³ EC detector. Analyses were con-	
tinued until a constant concentration was observed.	
observeu.	
	ESTIMATED ERROR: Solubility: ±5% (compiler).
	Temperature: ±0.5 K.
	REFERENCES: 1. Chiou, C. T. Hazard Assess. Chem. Current Dev. <u>1981</u> , 1, 117-53.
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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Water-d ₂ ; D ₂ 0; [7789-20-0]	Hutchison, C. A.; Lyon, A. M. Columbia
(2) 1,2,4-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [120-82-1]	University Report A-745, July 1, <u>1943</u> .
VARIABLES:	
One temperature	PREPARED BY: G. Jancso
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ³ mol(1)/kg ¹	$10^{3}x(1)^{a}$
25.0 1.999 9.98	1.808
a. Calculated by F. W. Getzen. b. Reported (average of two experimenta)	l measurements).
	INFORMATION
METHOD/APPARATUS/PROCEDURE: Between 25 and 100 ml 1,2,4-trichlorobenzene and 1 to 2 ml D_20 were placed in a flask and shaken for about 90 min. The thermostat water bath temperature was maintained within ± 0.05 °C. Then, a sample was removed and the amount of D_20 dissolved was determined by a modified Karl Fischer titration (1). The original report was unavailable; however, the method and the results were described in sufficient detail in (1). The solubility of H ₂ 0 in 1,2,4-trichlorobenzene was also determined and found to be 0.01123 mol(1)/kg. The average deviation for two experiments was ± 0.00006 mol(1)/kg.	SOURCE AND PURITY OF MATERIALS: C ₆ H ₃ Cl ₃ : Carefully purified and dried before use. Source and method not given. D ₂ O: Source not specified.
	ESTIMATED ERROR: Solubility: av. dev. $\pm 2.1 \times 10^{-5}$ mol D ₂ 0/100 g solution. Temperature: ± 0.05 K.
	REFERENCES: 1. Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M., Urey, H. C., Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II, Chapter 7, pp 129-44.

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COMPONENTS:	EVALUATOR:
<pre>(1) 1,3,5-Trichlorobenzene; C₆H₃Cl₃; [108-70-3]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2) Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,3,5-trichlorobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature $(25 \pm 1^{\circ}C)$. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubility correlations.

The solubility of 1,3,5-trichlorobenzene in water is reported here as a tentative value:

т/к	10^{5} mol(1)/dm ³	10 ³ g(1)/kg	$10^{7}x(1)$
298.15	3.6	6.55	6.51

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

HBTPW-C

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,3,5-Trichlorobenzene; C ₆ H ₃ Cl ₃ ; [108-70-3]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)/dm	$n^{3 b} 10^{7} x(1)^{a}$
25 6.53 3.6	6.51
a. Calculated by F. W. Getzen. b. Reported.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
tated for a period of 4-46 hours (the exact	C ₆ H ₃ Cl ₃ : Commercial reagent (Aldrich or Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The satu-	H ₂ O: Source and purity not specified.
rated solution was extracted with a small volume of methylene chloride which was then	-
assayed spectrophotometrically. At least two independent determinations were carried	
out.	
	ESTIMATED ERROR: Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	-
	REFERENCES :

COMP	DNENTS:	EVALUATOR:
(1)	2,4,5-Trichlorophenol; C ₆ H ₃ Cl ₃ O; [95-95-4]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

The solubility 2,4,5-trichlorophenol in aqueous buffer solution has been reported by Blackman, Parke, and Garton (1) in 1954. The principal motivation of this study was to observe the relationships between physical properties and physiological activities of certain substituted phenols. The experimental conditions were standardized with respect to pH by the slight addition of a phosphate buffer to the test solutions. The final pH value of the saturated solutions was regulated to 5.1. Under saturation equilibrium conditions, however, the concentration of the buffer was relatively dilute having a concentration of about 6 mmol/dm³ relative to the added phosphate, KH_2PO_4 .

The salt effect of the added buffer can be neglected, but the effect of the adjusted pH upon the acid dissociation equilibrium of the phenol must be given some consideration. The acidity constant for the 2,4,5-trichlorophenol was reported by Blackman et al. to have a magnitude in the order of 10^{-7} . Consequently, in the absence of the added buffer, the pH of the saturated solution should fall below the 5.1 value. Therefore, the adjustment of the pH to a value of 5.1 with the buffer should result in an increase in the concentration of the phenoxide ion relative to the undissociated phenol. This situation would result in a higher total solubility compared to the solution without the added buffer.

The authors did undertake tests of their method of analysis. They established the difference between the values measured by what they called "colorimetric" and "spectrophotometric" methods to be approximately 4 percent. However, the information concerning the purity of the reagents used was inadequate. Considering all such factors, the solubility of 2,4,5-trichlorophenol in water must be reported as a doubtful value.

The solubility of 2,4,5-trichlorophenol in water at a pH of 5.1 is reported here as a doubtful value:

T/K	10^{3} mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
298.15	5.0	9.9	9.0

REFERENCES

 Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u>, 54(1), 55-71.

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4-5-Trichlorophenol; C₆H₃Cl₃O; [95-95-4]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	A. Vesala
one ph: 5.1	
EXPERIMENTAL VALUES:	
t/°C 10g(1)/cm ^{3 a} 10 ³ mo1(1)/dm	$^{3 b}$ $10^{5}x(1)^{a}$
25 9.48 4.8	8.68
a Calculated by F. W. Cataoa	
a. Calculated by F. W. Getzen. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The samples were equilibrated in a thermostat bath with intermittent shaking over periods	C ₆ H ₃ Cl ₃ O: Source and purity not specified, probably a commercial reagent.
of 3-4 weeks. During the equilibration time, the pH values of the solutions were control-	
led by dropwise addition of phosphate buffer solution. The analysis of the solute con-	
centration in the saturated samples was done	
spectrophotometrically either directly or by using proper colorizing agents.	
	ESTIMATED ERROR:
	Solubility: <5% (evaluated here on the basis of the reported results of the two techniques of analysis).
	REFERENCES:

COMP	DNENTS:	EVALUATOR:
(1)	2,4,6-Trichlorophenol; C ₆ H ₃ Cl ₃ O; [88-06-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

Daccomo (1) has provided the first reported measurement of the solubility of 2,4,6-trichlorophenol in water. These measurements were made at least as early as 1885. They can be judged as somewhat unreliable for several reasons. First of all, the melting point of the substrate was 2-3 degrees below that reported later (69.5°C) in the "Handbook of Chemistry and Physics" (2). The measurements themselves were not described. However, it is porbable that the so-called "classical synthetic method" was applied. In this P^{roc} edure, known amounts of solute and solvent are mixed and their miscibilities tested with temperature. The reported value of 0.85 g(1)/kg(2) at 298 K from three experimental Points does not allow one to establish a reliable estimate of experimental errors.

Another value for the solubility of 2,4,6-trichlorophenol in water has been reported by Blackman, Parke, and Garton (3). Their determination was made at a solution pH of 5.1 through the slight addition of a phosphate buffer solution. Because the phenol in question has an acid pK value of 10.9 according to Blackman et al., the buffer solution pH evidently suppresses the ionization. This results in a decrease in concentration of the phenolate ion relative to the undissociated phenol. The net result is that the solubility should be suppressed somewhat for the buffered system relative to that for a system containing only the phenol itself in water. Converting the solubility value to the same units as those of Daccomo (using a solution density of 1.0 g/cm³), one obtains a value of 0.443 g(1)/kg(2) at the pH value of 5.1. This solubility value is quite possible, relative to the value provided by Daccomo, for the reasons indicated.

Since the variation of solubility with pH is not known in this case, only a doubtful value for the solubility of 2,4,6-trichlorophenol in water can be reported here. The value of Daccomo refers to a solution saturated with only the phenol at the pH which prevails for a saturated solution. Therefore, this value has been selected as the most appropriate saturation value at 298.15 K.

 $^{\mathrm{The}}$ solubility of 2,4,6-trichlorophenol in water is reported here as a doubtful value:

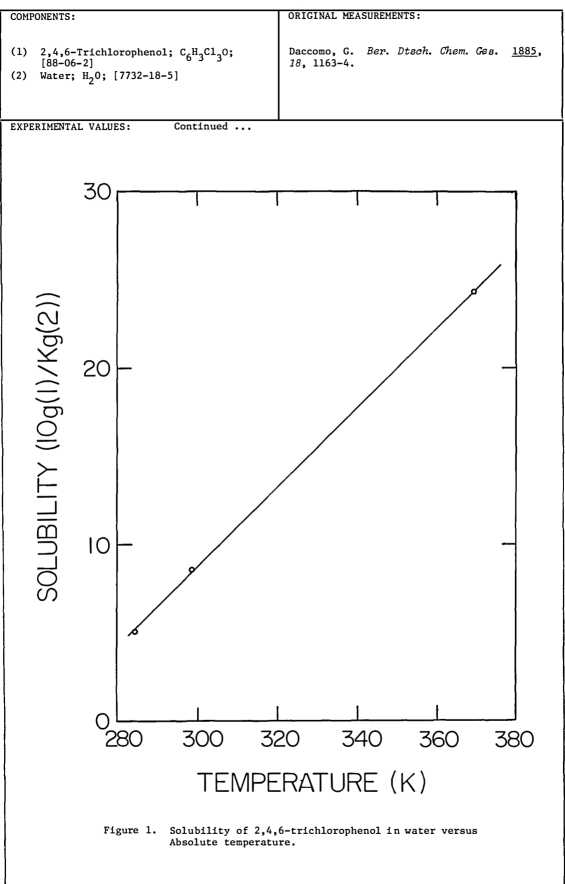
т/к	10^3 mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
298.15	2.2	4.3	4.0

REFERENCES

- 1. Daccomo, G. Ber. Dtsch. Chem. Ges. 1885, 18, 1163-4.
- "Handbook of Chemistry and Physics", 50th ed.; Weast, R. C., Ed.; CRC Press: Cleveland, Ohio, <u>1969</u>; p C-426.
- 3. Blackman, G. E.; Park, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> 54(1), 55-71.

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COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) 2,4,6-Tri [88-06-2]	chlorophenol; C ₆ H ₃ C	21 ₃ 0;	Daccomo, G. Ber. Dtsch. Chem. Ges. <u>1885</u> , 18, 1163-4.
(2) Water; H ₂	0; [7732-18-5]		
VARIABLES:			PREPARED BY:
Temperature			A. Vesala
EXPERIMENTAL VA	LUES:	<u></u>	
t/°C	10g(1)/kg(2) ^a	10 ³ mol(1)	$)/kg^{b}$ 10 ⁵ x(1) ^b
11.2	5.10	2.581	4.653
25.4	8.58	4.341	7.828
96	24.3	12.28	22.17
	rted (parts(1) per ulated by F. W. Get		2) in original work).
Maggurom	onto ano chorm area	hiaallu in T	
Measurem	ents are shown grap	nically in f	rigure 1.
			Continued
		AUXILIARY	INFORMATION
METHOD / APPARATU	S/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:
No specifications so-called synt	ons given. Probably hetic method.	y the	C ₆ H ₃ Cl ₃ O: Synthesized both from 2-chloro- phenol and 4-chlorophenol, melting point 67°C.
			H ₂ 0: Source and purity not specified.
			ESTIMATED ERROR:
			REFERENCES :



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COMPONENTS:	EVALUATOR:
(1) 1-Bromo-2-chlorobenzene; C ₆ H ₄ BrC1; [694-80-4]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ O; [7732-18-5]	Мау 1979

The experimental results consists of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions was assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated ben-Zenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1-bromo-2-chlorobenzene in water is tentative:

T/K	10^{4} mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
298.15	6.46	1.24	1.17

REFERENCES

- Yalkowsky, S.H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

HBTPW-C*

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 1-Bromo-2-chlorobenzene; C₆H₄BrCl; [694-80-4]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}$ C $10g(1)/dm^{3}$ a $10^{4}mol(1)/dm^{3}$ b $10^{5}x(1)$ a	
25 1.237 6.46	1.167
a. Calculated by compiler. b. Reported.	
AUXILIARY INFORMATION	
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
A small excess of 1-brom-2-chlorobenzene in water was agitated at room temperature for a period of about 24 hours and then fil- tered. The saturated filtrate solution was diluted and assayed spectrophotometrically. Two independent determinations were carried out.	C ₆ H ₄ BrCl: Aldrich commercial grade, used as received. H ₂ O: Deionized.
	ESTIMATED ERROR: Solubility: ±10%.
	Temperature: ±1°K.
	REFERENCES :

COMPONENTS:	EVALUATOR:
<pre>(1) 1-Bromo-3-chlorobenzene; C₆H₄BrC1; [108-37-2]</pre>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ 0; [7732-18-5]	May 1979

The experimental results consist of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The sautrated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1-bromo-3-chlorobenzene in water is tentative:

т/к	$10^4 mol(1) / dm^3$	10g(1)/kg	$10^{5}x(1)$
298.15	6.14	1.18	1.11

REFERENCES

 Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1-Bromo-3-chlorobenzene; C₆H₄BrCl; [108-37-2]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm	3^{5} 10 ⁵ x(1) ^a
25 1.181 6.17	1.115
a. Calculated by compiler.	
b. Reported.	
AUXILIARY INFORMATION METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS:	
A small excess of 1-bromo-3-chlorobenzene in	SOURCE AND PURITY OF MATERIALS: C _c H _c BrCl: Aldrich commercial grade, used
water was agitated at room temperature for a period of about 24 hours and then fil-	as received.
tered. The saturated filtrate solution was diluted and assayed spectrophotometrically.	H ₂ 0: Deionized.
Two independent determinations were carried out.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES:

COMPONENTS: -	EVALUATOR:
<pre>(1) 1-Bromo-4-chlorobenzene; C₆H₄BrCl; [106-39-8]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku
(2) Water; H ₂ O; [7732-18-5]	November 1979

A single solubility measurement of 1-bromo-4-chlorobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. All measurements were done at room temperature $(25 \pm 1^{\circ}C)$. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a Private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of ± 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubility correlations.

The solubility of 1-bromo-4-chlorobenzene in water is reported here as a tentative value:

т/к	$10^4 mol(1)/dm^3$	10 ² g(1)/kg	$10^{6}x(1)$
298.15	2.3	4.42	4.16

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1-Bromo-4-chlorobenzene; C₆H₄BrC1; [106-39-8]</pre>	Yalkowsky, S. H.; Ore, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{4}mol(1)/dm^{3}$	$m^{3 b} 10^{6} x(1)^{a}$
25 4.40 2.3	4.16
a. Calculated by F. W. Getzen. b. Reported.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	
A small excess of solute in water was agi-	SOURCE AND PURITY OF MATERIALS: C ₂ H ₂ BrCl: Commercial product (Eastman or
tated for a period of 4-48 hours (the exact	6 4 Aldrich), used as received.
time for equilibration was not reported in single cases) and then filtered. The saturated solution was diluted and assayed spectrophotometrically. At least two inde- pendent determinations were carried out.	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR: Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES:

COMPONENTS :	EVALUATOR:
<pre>(1) 1-Bromo-4-iodobenzene; C₆H₄BrI; [589-87-7]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku
(2) Water; H ₂ O; [7732-18-5]	November 1979

A single solubility measurement of 1-bromo-4-iodobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. All measurements were done at room temperature $(25 \pm 1^{\circ}C)$. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the Precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of ± 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so that total 10 percent error seems reasonable. The possible systematic errors should have no effect on the Correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1-bromo-4-iodobenzene in water is reported here as a tentative value:

т/к	10^5 mol(1)/dm ³	10 ³ g(1)/kg	$10^{7}x(1)$
298.15	2.8	7.94	5.06

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1-Bromo-4-iodobenzene; C₆H₄BrI; [589-87-7]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	
	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
3 3 2 5	3 h 7 a
$t/^{\circ}C$ $10^{3}g(1)/dm^{3}a$ $10^{5}mol(1)/dc$	10'x(1)
25 7.92 2.8	5.06
a. Calculated by F. W. Getzen.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi- tated for a period of 4-48 hours (the exact	C ₆ H ₄ BrI: Commercial reagent (Aldrich or Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The	H ₂ 0: Source and purity not specified.
saturated solution was diluted and assayed spectrophotometrically. At least two inde-	2
pendent determinations were carried out.	
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES:

COMP	ONENTS:	EVALUATOR:
(1)	1,2-Dibromobenzene; C ₆ H ₄ Br ₂ ; [583-53-9]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ O; [7732-18-5]	May 1979

The experimental results consist of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison With previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1,2-dibromobenzene in water is tentative:

т/к	10^4 mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
298.15	3.16	7.48	5.71

REFERENCES

1. Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,2-Dibromobenzene; C ₆ H ₄ Br ₂ ; [583-53-9]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
	· · · · · · · · · · · · · · · · · · ·
$t/^{\circ}C$ $10^{2}g(1)/dm^{3}$ a $10^{4}mol(1)/dc$	$m^{3} b 10^{6} x(1)^{a}$
25 7.455 3.16	5.710
a. Calculated by compiler. b. Reported.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,2-dibromobenzene in water was agitated at room temperature for	C ₆ H ₄ Br ₂ : Aldrich commercial grade, used as received.
a period of about 24 hours and then fil- tered. The saturated filtrate solution was	H ₂ 0: Deionized.
diluted and assayed spectrophotometrically. Two independent determinations were carried	
out.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES:

COMPONENTS:		EVALUATOR:	
(1)	1,3-Dibromobenzene; C ₆ H ₄ Br ₂ ; [108-36-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2)	Water, H ₂ 0; [7732-18-5]	May 1979	

The solubility of 1,3-dibromobenzene in water was reported by Hine et al. (1) at 308.15 K and by Yalkowsky et al. (2) at 298.15 K. Considering the small temperature difference of 10 degrees between the two measurements, one can judge that the data show a significant discrepancy.

The reported experimental errors $\pm 5.6\%$ (1) and $\pm 10\%$ (2) are probably greater than the authors anticipated. A creful study of the experimental procedures described in both articles reveals no obvious shortcomings. Despite the long time allowed for equilibrium by Hine et al., their measurements resulted in a lower solubility value than that obtained by Yalkowsky et al. The most likely explanation for the discrepancy could be attributed to the analytical determinations used.

It is well known that the solubility curve for halogenated benzenes in water has a positive slope above room temperature. However, the two measured points reported are against this trend, that is, the values decrease with rising temperature. In order to choose between the two solubility data values, use was made of the theoretical relationship between the molar solubilities and the molar volumes of the solutes at 298.15 K as described in the Introduction. The following solubility value for 1,3-dibromobenzene in water is tentative:

т/к	10^4 mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
298.15	2.70	6.40	4.89

- 1. Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. 1963, 85(10), 1473-6.
- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,3-Dibromobenzene; C ₆ H ₄ Br ₂ ; [108-36-1]	Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. <u>1963</u> , 85(10), 1473-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{4}mol(1)/dm^{3}$	$m^{3 b} 10^{6} x(1)^{c}$
30.0 6.747 2.86	5.184
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of 1,3-dibromobenzene was added to a 4 ml of water in an ampoule which was	C ₆ H ₄ Br ₂ : Eastman Kodak Co.; redistilled before use.
then sealed. The sealed ampoule was ro- tated in a constant temperature water bath for a week to establish saturation. The 1,3-dibromobenzene solubility was deter- mined by ultraviolet spectrophotometric measurements using a Beckman DU spectro- photometer.	H ₂ 0: Not specified.
	ESTIMATED ERROR: Solubility: ±5.6% S.D.
	Temperature: ±0.1 K.
	REFERENCES :

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 1,3-Dibromobenzene; C₆H₄Br₂; [108-36-1]</pre>	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
-	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{2}g(1)/dm^{3}$ a $10^{4}mol(1)/d$	$m^{3 b} 10^{6} x(1)^{a}$
25 9.838 4.17	7.536
a. Calculated by compiler.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,3-dibromobenzene in water was agitated at room temperature for	C ₆ H ₄ Br ₂ : Aldrich commercial grade, used as received.
a period of about 24 hours and then fil-	
tered. The saturated filtrate solution was diluted and assayed spectrophotometrically.	H ₂ 0: Deionized.
Two independent determinations were carried	
out.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES:

0	COMP	ONENTS:	EVALUATOR:
	(1)	1,4-Dibromobenzene; C ₆ H ₄ Br ₂ ; [106-37-6]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
	(2)	Water; H ₂ 0; [7732-18-5]	January 1979

There are only a few reported measurements dealing with the solubility of 1,4dibromobenzene in water. In the published papers on the subject, the determination of solubility is not the primary purpose of the work, a fact that may affect the precision of the reported results. Of the values reported, only that of Andrews and Keefer (1) refers to 298.2 K. The measured data reported by Hine, Haworth, and Ramsay (2) at 308.2 K lend some support to the value assigned at 298.2 K.

The experimental procedure employed by Andrews and Keefer is adequate for the determination in question. However, the time of equilibrium (about 20 hours as given in (3)) seems quite short in the absence of tests for solution equilibrium. It is reasonable to attribute most of the uncertainty in the reported solubility of 8.5 x $10^{-5} \text{ mol}(1)/\text{dm}^3$ (7.0 x 10^{-5} molar at an ionic strength of 1.0 molar, a figure incorrectly cited for solubility in pure water in some papers). Support for the reliability of the value at 298.2 K is provided by the value of 1.12 x $10^{-4} \text{ mol}(1)/\text{kg}(2)$ given by Hine et al. for 308.2 K. This value was obtained by employing a long equilibration (at least a week) and a direct spectrophotometric measurement. In fact, it seems reasonable that the latter measurements should be more reliable than those of Andrews and Keefer although the agreement between the two measurements is quite satisfactory. The increase in the solubility is about 30% for a temperature rise of 10 degrees from 298.2 K. This value as that for 1,4-dichlorobenzene in water in the same range of temperature as measured by Wauchope and Getzen (4).

The following is the tentative solubility of 1,4-dibromobenzene in water at 298.15 K:

т/к	10^{5} mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
298.15	8.5	2.0	1.5

REFERENCES

1. Andrews, L. M.; Keefer, R. M. J. Am. Chem. Soc. 1950, 72(7), 3113-6.

2. Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. 1963, 85(10), 1473-6.

3. Andrews, L. M.; Keefer, R. M. J. Am. Chem. Soc. 1949, 71(11), 3644-7.

4. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data 1972, 17(1), 38-41.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,4-Dibromobenzene; C ₆ H ₄ Br ₂ ; [106-37-6]	Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1950</u> , 72(7), 3113-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3} = 10^{5}mol(1)/dm^{3}$	$dm^{3 b}$ 10 ⁶ x(1) ^b
25.0 2.0 8.48	1.53
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The saturation process was done in sealed Erlenmeyer flasks. The time used for the equilibration was 20 hours. Measured vol-	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ Br ₂ : Commercial product (Eastman Kodak Co.), used as received.
umes of the saturated solutions were then extracted using measured volumes of hexane to remove the aromatic matter from the	H ₂ 0: Source and purity not specified.
aqueous layer. The solute concentration was measured spectrophotometrically using a Beckman DU spectrophotometer (1).	
	ESTIMATED ERROR:
	Solubility: >10% (evaluator). Temperature: ±0.2 K (evaluator).
	REFERENCES :
	 Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1949</u>, 71(11), 3644-7.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,4-Dibromobenzene; C ₆ H ₄ Br ₂ ; [106-37-6]	Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. <u>1963</u> , 85(10), 1473-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES :	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ² g(1)/dm ^{3 a} 10 ⁴ mol(1)/d	$m^{3 b} 10^{6} x(1)^{a}$
35.0 2.642 1.12	2.030
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of the aromatic compound was com- bined with 4.0 ml of water in a 5.0 ml	C ₆ H ₄ Br ₂ : Source not specified; presumably commercial material, recrystal-
ampoule which was then sealed. The sealed	lized to constant melting point
ampoule was rotated in a constant tempera- ture water bath for at least a week to estab-	in agreement with literature values.
lish saturation. The undissolved solid was removed by filtration and the concentration	H ₂ 0: Source and purity not specified.
of the solute in the saturated filtrate	
solution was determined by ultraviolet spectrophotometric measurements using a	
Beckman DU spectrophotometer. Several mea- surements were taken at intervals of at	
least a day in order to be sure that an equilibrium had been established.	ESTIMATED ERROR:
equilibrium had been established.	Solubility: ±7% (standard deviation re-
	ported by authors). Temperature: ±0.1 K (authors).
	REFERENCES:

COMPONENTS:		EVALUATOR:
(1)	2,4-Dibromophenol; C ₆ H ₄ Br ₂ O; [615-58-7]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979

Only a rather old value, that of Werner (1), has been found in the literature for the solubility of 2,4-dibromophenol in water. Considering the incomplete information on the purity of the reagents used and the equilibration method itself coupled with the fact that the instrumentation used was relatively simple, one must regard the reported value as doubtful.

It must be noted that the solubility of this compound, like any protolytic solute, is considerably dependent upon the acidity of the solution. Here it is assumed that the PH value is that for the saturated solution without the addition of other protolytic solutes. Thus, the solubility value reported is for such a saturated solution.

Based upon an assumed density of 1.0 g/cm^3 for the solution (an estimate which has no effect on the accuracy of the reported value), the doubtful solubility of 2,4-dibromophenol in water at 298 K is as follows:

т/к	10^{3} mol(1)/dm ³	g(1)/kg	$10^4 x(1)$
298	8.	2.	1.4

REFERENCES

1. Werner, E. Ann. Chim. Phys. Ser. 6 1884, 571-2.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 2,4-Dibromophenol; C ₆ H ₄ Br ₂ O; [615-58-7]	Werner, E. Ann. Chim. Fhys. Ser. 6 <u>1884</u> , Vol. 3, 571-2.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES :	PREPARED BY:
One temperature	A. Vesala
· • •••	
EXPERIMENTAL VALUES:	
t/°C g(1)/dm ^{3 a} 10 ³ mol(1)/dm ^{3 b}	$10^4 x(1)^{b}$
15 1.94 7.701	1.391
a. Reported. b. Calculated by F. W. Getzen.	
b. Calculated by r. w. Getzen.	
3	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturation procedure was not specified.	$C_{6}H_{4}Br_{2}O$: Synthesized product, melting
The analysis of the saturated solution was done by titration with a standardized bro-	o 4 2 point 40°C, boiling point 154°C at 17 mmHg.
mine solution.	
	H ₂ 0: Source and purity not specified.
}	ESTIMATED ERROR:
}	REFERENCES :
]	

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COMPONENTS:		EVALUATOR:
(1)	1-Chloro-2-iodobenzene; C ₆ H ₄ ClI; [615-41-8]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ O; [7732-18-5]	Мау 1979

The experimental results consist of a single value at 298.15 K published by 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1-chloro-2-iodobenzene in water is tentative:

$T/K = 10^4 mol(1)/dm^3$		10 ² g(1)/kg	$10^{6}x(1)$	
298.15	2.88	6.89	5.21	

REFERENCES

 Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:		
<pre>(1) 1-Chloro-2-iodobenzene; C₆H₄ClI; [615-41-8]</pre>	Yalkowsky, S. H., Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.		
(2) Water; H ₂ 0; [7732-18-5]			
2			
VARIABLES:	PREPARED BY:		
One temperature	A. L. Horvath		
EXPERIMENTAL VALUES:			
$t/^{\circ}C = 10^{2}g(1)/dm^{3} a = 10^{4}mol(1)/dm^{3} b$	$dm^{3 b} 10^{6} x(1)^{a}$		
25 6.868 2.88	5.204		
a. Calculated by compiler.b. Reported.			
AUXILIARY	INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:		
A small excess of 1-chloro-2-iodobenzene	$C_{6}H_{4}$ ClI: Aldrich commerical grade, used		
in water was agitated at room temperature for a period of about 24 hours and then	as received.		
filtered. The saturated filtrate solution was diluted and assayed spectrophotometri-	H ₂ 0: Deionized.		
cally. Two independent determinations were			
carried out.			
	ESTIMATED ERROR:		
	Solubility: ±10%.		
]	Temperature: ±1 K.		
	REFERENCES :		

COMPONENTS:		EVALUATOR:	
(1)	1-Chloro-3-iodobenzene; C ₆ H ₄ ClI; [625-99-0]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2)	Water, H ₂ O; [7732-18-5]	May 1979	

The experimental results consist of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1-chloro-3-iodobenzene in water is tentative:

T/K 10 ⁴ mo1(1)/dm		10 ² g(1)/kg	$10^{6}x(1)$	
298.15	2.82	6.74	5.09	

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS :	ORIGINAL MEASUREMENTS:	
	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.	
<pre>(1) 1-Chloro-3-iodobenzene; C₆H₄ClI; [625-99-0]</pre>	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.	
(2) Water; H ₂ O; [7732-18-5]		
VARIABLES:	PREPARED BY:	
One temperature	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 10 ² g(1)/dm ^{3 a} 10 ⁴ mol(1)/	$dm^{3} b 10^{6} x(1)^{a}$	
25 6.725 2.82	5.096	
a. Calculated by compiler.		
b. Reported.		
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
A small excess of 1-chloro-3-iodobenzene	C ₆ H ₄ ClI: Aldrich commercial grade, used	
in water was agitated at room temperature for a period of about 24 hours and then	6 4 as received.	
filtered. The saturated filtrate solution	H ₂ 0: Deionized.	
was diluted and assayed spectrophotometri- cally. Two independent determinations were		
carried out.		
	ESTIMATED ERROR: Solubility: ±10%.	
	Temperature: ±1 K.	
	REFERENCES:	

COMPONENTS :		EVALUATOR:	
<pre>(1) 1-Chloro-4-iodobenzene; C₆H₄ClI; [637-87-6]</pre>		A. Vesala, Department of Chemistry and Biochemistry, University of Turku.	
(2) Water; H ₂ O; [7732-18-5]		November 1979	

A single solubility measurement of 1-chloro-4-iodobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. All measurements were done at room temperature $(25 \pm 1^{\circ}C)$. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurement is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of ± 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1-chloro-4-iodobenzene in water is reported here as a tentative value:

T/K	$10^4 mol(1)/dm^3$	10 ² g(1)/kg	$10^{6}x(1)$	
298.15	1.3	3.11	2.35	

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

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COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) 1-Chloro-4-iodobenzene; C ₆ H ₄ ClI; [637-87-6]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.	
(2) Water; H ₂ 0; [7732-18-5]		
VARIABLES:		
One temperature	PREPARED BY: A. Vesala	
EXPERIMENTAL VALUES:		
$t/°C = 10^2 g(1)/dm^3 a = 10^4 mol(1)/dm^3 a$	$dm^{3}b$ $10^{6}x(1)^{a}$	
25 3.10 1.3	2.35	
a. Calculated by F. W. Getzen. b. Reported.		
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
	C ₂ H ₂ ClI: Commercial reagent (Eastman or	
tated for a period of 4-48 hours (the exact time for equilibration was not reported in	Aldrich), used as received.	
single cases) and then filtered. The saturated solution was diluted and assayed spectrophotometrically. At least two	H ₂ 0: Source and purity not specified.	
independent determinations were carried out.		
	ECTIVATED EDDOD	
	ESTIMATED ERROR: Solubility: ±10% (authors).	
	Temperature: ±1 K (authors).	
	REFERENCES:	

COMPONENTS:		EVALUATOR:	
(1) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]		A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2) Water; H ₂ 0; [7732-18-5]		January 1983	

Data for the solubility of 1,2-dichlorobenzene in water have been reported in several papers. The approximate solubility at 298.15 K was reported by Booth and Everson (1) while, somewhat earlier, Klemenc and Low (2) produced measurements between 293 and 333 K. More recent data have been published only between 276 and 307 K (8-11).

The approximate value provided by Booth and Everson is not given further consideration here. The remaining available data from (2, 8-11) have been correlated relative to absolute temperature using a normal polynomial equation:

$$S_1(g(1)/kg) = 19.2314 - 1.81140 \times 10^{-1}T$$

+ 5.6509 x $10^{-4}T^2 - 5.77683 \times 10^{-7}T^3$ [1]

The significance of this equation is that the curve representing the solubility data passes through a minimum at 283.7 K. This behavior is consistent with the theory discussed by Gill et al. (12) for the solubility of aromatic compounds in water.

Recommended solubility values between 273 and 333 K have been calculated from equation [1] and presented in Table 1. The solubility behavior described by equation [1] is shown also in Figure 1 which contains the reported experimental values for the solubility of 1,2-dichlorobenzene in water.

The solubility of water in 1,2-dichlorobenzene has also been investigated and reported by a number of workers (3-7) in the 298 to 318 K temperature range. In general, these solubility values show some irregularity. While the agreement between the measurements of Jones and Monk (5) and those of Wing and Johnston (7) is reasonable, the deviation is considerable in relation to the measurements of Goldman (4) and of Kirchnerova and Cave (6) at 298.15 K. Identical measured values at 298.15 K were reported from the same department of McGill University by Goldman and by Kirchnerova and Cave.

From the description of the experimental measurements and the reliability if the results reported, it is not reasonable to exclude any of the data points except that quoted by Dreisbach (3) from further analysis. Therefore, all the water in 1,2-dichlorobenzene solubility data except that from Dreisbach were combined and incorporated in a regression equation. However, because of the long equilibration times allowed in Goldman and in Kirchnerova and Cave, their data points were given a weight twice as large as those values provided in Jones and Monk and in Wing and Johnston. The correlation equation is given by:

$$\log_{10} x(2) = 2.23134 - 1448.67/T$$
 [2]

In this equation, x(2) is the mole fraction solubility of water in the 1,2-dichlorobenzenewater system and T is the Absolute temperature. The calculated solubility values in the 273 and 333 K range are shown in Figure 2 along with the reported values.

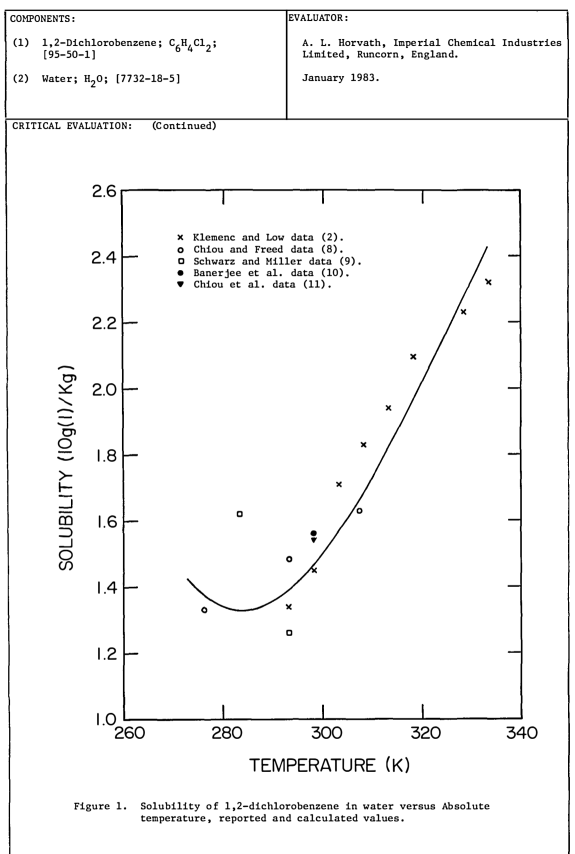
The calculated mole fraction values for the solubility of water in 1,2- dichlorobenzene from equation [2] are given in Table 2 together with the corresponding molarities and g(2)/kg values in the temperature range between 293 and 328 K.

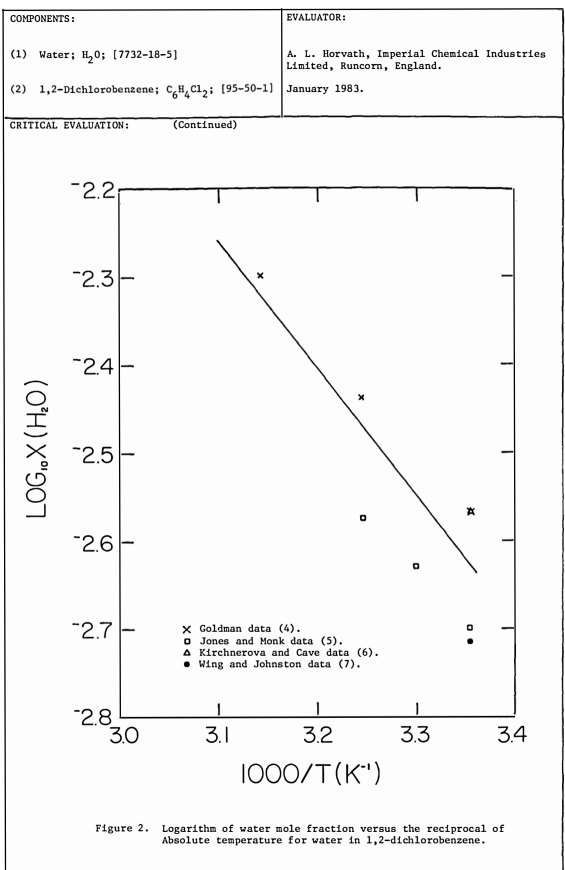
COMPONENTS:			EVALUATOR:	
<pre>(1) 1,2-Dichlorobenzene; C₆H₄Cl₂; [95-50-1]</pre>			A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2) Water; H ₂ 0;	[7732-18-5]	Janua	ry 1983	
CRITICAL EVALUATION	N: (Continued)			
Table 1. Solubili	ty of 1,2-Dichloro	benzene in Wate	r.	
т/к	10 ⁴ mol(1)/dm ³	10g(1)/kg	$10^5 x(1)$	
273.15	9.64	1.42	1.74	
278.15	9.20	1.35	1.66	
283.15	9.04	1.33	1.63	
288.15	9.13	1.34	1.65	
293.15	9.44	1.39	1.70	
298.15	9.94	1.47	1.80	
303.15	10.6	1.57	1.92	
308.15	11.4	1.69	2.07	
313.15	12.3	1.82	2.23	
318.15	13.2	1.97	2.41	
323.15	14.2	2.12	2.60	
328.15	15.3	2.28	2.79	
333.15	16.2	2.43	2.98	
Table 2. Solubili	lty of Water in 1,2	-Dichlorobenzen	1e.	
T/K	$10^2 mol(2)/dm^3$	10g(2)/kg	10x(2)	
293.15	1.73	2.39	1.95	
298.15	2.09	2.90	2.36	
303.15	2.50	3.48	2.84	
308.15	2.98	4.17	3.39	
313.15	3.53	4.96	4.03	
318.15	4.16	5.86	4.76	
323.15	4.87	6.90	5.60	
328.15	5.68	8.08	6.56	

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COMPO	DNENTS:	EVALUATOR:	
(1)	1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2)	Water; H ₂ 0; [7732-18-5]	January 1983	
CRIT	ICAL EVALUATION: (Continued)		
	REFEI	RENCES	
1.	Booth, H. S.; Everson, H. E. Ind. Eng.	Chem. <u>1948</u> , 40(8), 1491-3.	
2.	Klemenc, A.; Löw, M. Rec. Trav. Chim. H	Pays-Bas <u>1930</u> , 49(4), 629-40.	
3.	Dreisbach, R. R. "Physical Properties of Chemical Compounds," Advances in Chemistry Series No. 15; American Chemical Society: Washington, D.C., <u>1955</u> ; p. 135.		
4.	Goldman, S., Ph.D. Dissertation, McGill University, Montreal, 1969, p 84.		
5.	Jones, J. R.; Monk, C. B. J. Chem. Soc. <u>1963</u> , Part III, 2633-5.		
6.	Kirchnerova, J.; Cave, G.C.B. Can. J. C	Chem. <u>1976</u> , 54(24), 3909-16.	
7.	Wing, J.; Johnston, W. H. J. Am. Chem.	Soc. <u>1957</u> , 79(4), 864-5.	
8.	Chiou, C. T.; Freed, V. H. "Chemodynami National Technical Information Service:	lc Studies on Bench Mark Industrial Chemicals"; Springfield, Virginia, <u>1977;</u> PB-274263.	
9.	Schwarz, F. P.; Miller, J. Anal. Chem.	<u>1980</u> , <i>52(13)</i> , 2162-4.	
10.	Banerjee, S.; Yalkowsky, S. H.; Valvani, 1227-9.	, S. C. Environ. Sci. Technol. <u>1980</u> , 14(10),	
11.	Chiou, C. T.; Schmedding, D. W.; Manes,	M. Environ. Sci. Technol. <u>1982</u> , 16(1), 4-10.	
12.	Gill, S. J.; Nichols, N. F.; Wadso, I.	J. Chem. Thermodyn. <u>1976</u> , 8(5), 445-52.	

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COMPONENTS:				ORIGINAL MEASUREMENTS:	
(1)	(1) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]			Klemenc, A.; Löw, M. <i>Rec. Trav. Chim.</i> <i>Pays-Bas.</i> <u>1930</u> , 49(4), 629-40.	
(2) 6	(2) Water; H ₂ 0; [7732-18-5]				
	2				
VARIABI Temper				PREPARED BY: A. L. Horvath	
Temper	ature			A. L. HOIVALII	
EXPERIN	ÆNTAL VALU	JES:			
	t/°C	10g(1)/kg(2) ^a	10 ³ mol(1)	$10^{5} kg^{b} = 10^{5} x(1)^{c}$	
	20	1.34	0.9114	1.642	
	25	1.45	0.9862	1.777	
	30	1.71	1.163	2.096	
	35	1.83	1.245	2.243	
	40	1.94	1.319	2.377	
	45	2.03	1.381	2.488	
	55	2.23	1.517	2.733	
	60	2.32	1.578		
	00		1.570	2.045	
		ced. lated by F. W. Ged lated by compiler.			
			AUXILIARY	INFORMATION	
METHOD	APPARATUS	/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:	
based the me	upon volum easurement	on of the solubil: metric principles of excess solute as described by	applied to in a cali-	C ₆ H ₄ Cl ₂ : Kahlbaum reagent, used as received. H ₂ O: Distilled.	
				ESTIMATED ERROR: Solubility: ±10% (compiler).	
				Temperature: ±1 K (compiler).	
				REFERENCES: 1. Rex, A. Z. Phys. Chem. <u>1906</u> , 55, 355-70.	
				1. Ker, R. <i>D. 116</i> 0. Oren. <u>1700</u> , <i>D.</i> , 10.	

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COMPONENTS:	ORIGINAL MEASUREMENTS:
	Booth, H. S.; Everson, H. E., Ind. Eng.
(1) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	Chem. <u>1948</u> , 40, 1491-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{-1}ml(1)/dm^{3}(2)^{a}$ $10mo$	$1(1)/dm^{3}$ $10^{3}x(1)$ c
25.0 < 2 <	1.7 < 3.2
	*
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
AUXILIARY INFORMATION	
METHOD / AP PARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
The measurements were made with samples con-	
tained in a stoppered Goetz tube placed in a constant temperature water bath. Equili-	used as received.
brium was assured through repeated shaking and centrifuging the mixture in the stop-	H ₂ 0: Distilled.
pered tube while it was temporarily removed	
from the water bath. The amount of solute dissolved was determined as the difference	
between total amount added and amount re- maining in excess upon saturation. The	
determination of the excess amount of solute added has been described by Hanslick (1).	
	ESTIMATED ERROR:
	Solubility: <100%.
	Temperature: ±1 K (compiler).
	REFERENCES:
	 Hanslick, R. S., Dissertation, Columbia University, 1935.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Dreisbach, R. R. "Physical Properties of
-	Chemical Compounds", Advances in Chemistry
(2) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	Series No. 15; American Chemical Society: Washington, D. C., <u>1955</u> ; p 135.
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10 ⁻¹ g(1)/kg(2) ^a mol(1)	$(l_{r_2} \stackrel{b}{=} 10m(1) \stackrel{c}{=}$
25 2.1 1.14	1.46
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS/PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
	ISOUNCE AND FUNITE OF PATENTALS:
EVDERIMENTSI METHODE WORD NOT described	H Or Distilled
Experimental methods were not described.	H ₂ 0: Distilled.
Experimental methods were not described.	H ₂ O: Distilled. C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before use.
Experimental methods were not described.	C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before
Experimental methods were not described.	C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before
Experimental methods were not described.	C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before
Experimental methods were not described.	C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before use.
Experimental methods were not described.	C ₆ H ₄ Cl ₂ : Dow Chemical Co., 99.85% pure, purified by distillation before
Experimental methods were not described.	<pre>Z C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler).</pre>
Experimental methods were not described.	<pre>2 C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler). Temperature: ±1 K (compiler).</pre>
Experimental methods were not described.	<pre>Z C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler).</pre>
Experimental methods were not described.	<pre>2 C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler). Temperature: ±1 K (compiler).</pre>
Experimental methods were not described.	<pre>2 C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler). Temperature: ±1 K (compiler).</pre>
Experimental methods were not described.	<pre>2 C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler). Temperature: ±1 K (compiler).</pre>
Experimental methods were not described.	<pre>2 C₆H₄Cl₂: Dow Chemical Co., 99.85% pure, purified by distillation before use. ESTIMATED ERROR: Solubility: <100% (compiler). Temperature: ±1 K (compiler).</pre>

COMPONENTS :	ORIGINAL MEASUREMENTS:
 Water; H₂0; [7732-18-5] 1,2-Dichlorobenzene; C₆H₄Cl₂; [95-50-1] 	Wing, J.; Johnston, W. H. J. Am. Chem. Soc. 1957, 79(4), 864-5.
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10m1(1)/dm ^{3 a} 10 ² mol(1),	$/dm^{3 b}$ 10 $^{3}x(1)^{c}$
25.0 3.09 1.710	1.930
 a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler. 	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Tritiated water was equilibrated with 20 ml 1,2-dichlorobenzene by stirring in a flask in a constant temperature water bath for	H ₂ 0: Tracerlab Inc., tritiated water, used as received.
two hours. The concentration of the tri- tiated water in the organic phase was de- termined by isotopic dilution. The tritium activities in the tritiated water samples were determined by the acetylene method (1,2). At least four independent experi- ments were done. The article was based upon work reported	C ₆ H ₄ Cl ₂ : Source not specified, chemical grade, redistilled before use.
in a Ph.D. dissertation (2).	ESTIMATED ERROR:
	Solubility: ±1.5%.
	Temperature: ±0.02 K.
	REFERENCES: 1. Wing, J.; Johnston, W. H. Science <u>1955</u> , 121, 674-5.
	 Wing, J., Ph. D. Dissertation, Purdue University, Lafayette, <u>1956</u>.
METHOD/APPARATUS/PROCEDURE: Tritiated water was equilibrated with 20 ml 1,2-dichlorobenzene by stirring in a flask in a constant temperature water bath for two hours. The concentration of the tri- tiated water in the organic phase was de- termined by isotopic dilution. The tritium activities in the tritiated water samples were determined by the acetylene method (1,2). At least four independent experi- ments were done.	 ² used as received. C₆H₄Cl₂: Source not specified, chemical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±1.5%. Temperature: ±0.02 K. REFERENCES: 1. Wing, J.; Johnston, W. H. Science 195 121, 674-5. 2. Wing, J., Ph. D. Dissertation, Purdue

COMPONENTS :		ORIGINAL MEASUREMENTS:
(1) Water;	H ₂ 0; [7732-18-5]	Jones, J. R.; Monk, C. B. J. Chem. Soc.
(2) 1,2-Dic	hlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	<u>1963</u> , Part III, 2633-5.
VARIABLES:		PREPARED BY:
Temperature		A. L. Horvath
EXPERIMENTAL	VALUES:	
t/°C	10m1(1)/dm ³ (2) ^a 10 ² mol(1	$1)/dm^{3} b 10^{3}x(1)^{c}$
25	3.2 1.77	2.00
30	3.75 2.07	2.348
35	4.25 2.34	44 2.667
a. Rep b. Cal	orted. culated by F. W. Getzen.	
	culated by compiler.	
	AUXILIARY	INFORMATION
METHOD/APPAR	ATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
	ater was shaken with 1,2-	H ₂ 0: Tritiated.
4 hours in a	zene in 1:10 volume ratios for a flask in a water thermostat	C ₆ H ₄ Cl ₂ : Source not known, laboratory
	vater content was determined by y. The count rates were deter-	grade, dried over CaCl ₂ and fractionally distilled before
mined using solution tec	a typical liquid scintillator	use.
		ESTIMATED EDDOD.
		ESTIMATED ERROR: Solubility: ±5%.
		Temperature: ±0.5 K (compiler).
		REFERENCES:
		ALE ENERGES .

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Goldman, S., Ph.D. Dissertation, McGill
(2) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	University, Montreal, <u>1969</u> , p. 84.
(2) 1,2-bichiorobenzene; $6^{\circ}4^{\circ}2^{\circ}$, $[55-50^{\circ}1]$	
VARIABLES :	PREPARED BY:
Temperature	A. L. Horvath
Temperature	
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ² mol(1)/dm	$10^{3} b 10^{3} x(1)^{c}$
25.0 4.342 2.41	2.719
35.1 5.783 3.21	3.650
45.06 7.909 4.39	5.028
45.00 7.909 4.55	5.020
a. Calculated by F. W. Getzen.	
b. Reported.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	COURCE AND DUDITY OF MATERIALS.
Equilibrium experiments were carried out in	SOURCE AND PURITY OF MATERIALS: H ₂ O: Distilled.
a constant temperature water bath using a	2
stirrer. An equilibration period of at least 5 days was allowed. The total water	$C_6H_4Cl_2$: Reagent grade, washed with conc. H ₂ SO ₄ and with 1 M NaHCO ₃ and
content in the organic phase was determined	then fractionally distilled
by a Karl Fischer titration. Each reported water solubility was obtained as an average	over silica gel.
of at least two independent determinations.	
	ESTIMATED ERROR:
	Solubility: ±4%.
	Temperature: ±0.1 K.
	REFERENCES:
1	

COMPONENTS :	ORIGINAL MEASUREMENTS:
 Water; H₂0; [7732-18-5] (2) 1,2,-Dichlorobenzene; C₆H₄Cl₂; [95-50-1] 	Kirchnerova, J.; Cave, G.C.B. Can. J. Chem. 1976, 54(24), 3909-16.
VARIABLES: One temperature	PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES:	
/ t/°C 10g(1)/dm ^{3 a} 10 ² mo1(1)/dm	3 b $10^{3}x(1)$ c
25 4.324 2.40	2.707
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: A mixture of 50 ml 1,2-dichlorobenzene and 6 ml water in a bottle was submerged in a water thermostat bath for 2 days. The con- centration of the water in the organic phase was determined by a conventional Karl Fischer dead stop back titration. Determinations were done in triplicate. A detailed description of the complete ex- perimental procedure has been included in a Ph.D. dissertation (1).	SOURCE AND PURITY OF MATERIALS: H ₂ O: Distilled and deionized. C ₆ H ₄ Cl ₂ : Fisher - B255, washed with cc. H ₂ SO ₄ and K ₂ SO ₃ solution and distilled water. Dried over silica gel and fractionally distilled, purity 99.8%.
	ESTIMATED ERROR:
	Solubility: ±1%. Temperature: ±0.1 K.
	REFERENCES: 1. Kirchnerova, J., Ph.D. Dissertation, McGill University, Montreal, Quebec, <u>1974</u> .

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COMPONENT	`S :	ORIGINAL MEASUREMENTS:
[95	2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; 5-50-1]	Chiou, C. T.; Freed, V. H. "Chemodynamic Studies on Bench Mark Industrial Chemicals"; National Technical Information Service:
(2) Wat	ter; H ₂ 0; [7732-18-5]	Springfield, Virginia, <u>1977</u> ; PB-274263.
VARIABLES	5:	PREPARED BY:
Temperat	ture	A. L. Horvath
EXPERIMEN	NTAL VALUES:	I
t/°	°C 10g(1)/dm ³ (2) ^a 10 ³ mol(1)	$dm^{3 b}$ 10 ⁵ x(1) ^c
3	3 1.33 0.904	46 1.630
20	0 1.48 1.00	7 1.817
34	4 1.62 1.102	2 1.997
c.	Calculated by compiler.	
	AUXILIARY	INFORMATION
METHOD/A	PPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
was equi water in temperat was remo by gas c equipped column w Chromoso	as of 1,2-dichlorobenzene, 5-10 g, Llibrated with 100 ml distilled h a bottle for 24 hours at constant cure. The saturated aqueous phase by the container and analyzed chromatography using a chromatograph with a Ni ⁶³ EC detector. The GLC was packed with porous polymer by 101. details on the determinations and	C ₆ H ₄ Cl ₂ : Commercial reagent, used as received. H ₂ O: Distilled.
	ons of experimental procedures in included in (1).	ESTIMATED ERROR:
		Solubility: ±10% (compiler).
		Temperature: ±0.5 K.
		REFERENCES: 1. Chiou, C. T.; Schmedding, D. W. Test. Protoc. Environ. Fate Mov. Toxicants, Proc. Symp., <u>1980</u> (Publ. 1981), pp 28-42.
L		

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) 1,2-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [95-50-1]	Schwarz, F. P.; Miller, J. Anal. Chem. 1980, 52(13), 2162-4.	
(2) Water; ¹¹ 2 ⁰ ; [7732-18-5]		
-		
VARIABLES: Temperature	PREPARED BY: A. L. Horvath	
EXPERIMENTAL VALUES:		
Experimentally determined values:		
10g(1)/kg		
t/°C Elution Chromatography UV	/ Absorption	
10.0 1.69 ± 0.03 1.	.56 ± 0.17	
20.0 1.28 ± 0.06 1.	.24 ± 0.04	
Values derived from average measured solubilities:		
	^b $10^{5}x(1)$ ^c	
10.0 1.62 ± 0.07 1.102 ± 0.048	3 1.986 ± 0.086	
20.0 1.26 ± 0.02 0.8571± 0.013	36 1.544 ± 0.025	
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.		
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Both elution chromatography and UV absorp- tion methods were used to determine the	C ₆ H ₄ Cl ₂ : Commercial, 98 wt. %.	
aqueous solubilities. The agreement was	H ₂ 0: Distilled.	
within an experimental error of 4% between the two methods. The average deviations		
were determined from several measurements made on different samples.		
The analytical procedures for determining organic liquid solubilities in water based on liquid phase elution chromatography has been described in (1).		
	ESTIMATED ERROR: Solubility: ±4%.	
	Temperature: ±0.5 K.	
	REFERENCES :	
	 Schwarz, F. P. Anal. Chem. <u>1980</u>, 52(1), 10-15. 	

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2-dichlorobenzene; C ₆ H ₄ Cl ₂ : [95-50-1]	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environm. Sci. Techn. <u>1980</u> , 14(10), 1227-9.
(2) Water; H ₂ 0; [7732-18-5]	1227-7.
1	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm	$10^{3} b 10^{5} x(1)^{c}$
25 1.5583 1.060	1.9156
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of 1,2-dichlorobenzene was added to water in a stainless steel centrifuge tube which was then sealed. The equilibrium was established by allowing the sample to stand, with intermittent shaking, for a week at constant temperature. The mixture was then centrifuged and aliquots of the solution were removed either by a pipet or syringe for analysis. Liquid scintillation counting with ¹⁴ C-labelled solute was em- ployed in the solubility determinations. The entire procedure was carried out at	C ₆ H ₄ Cl ₂ : New England Nuclear, used as received. H ₂ O: Distilled.
least twice and each analysis was also con- ducted in duplicate.	ESTIMATED ERROR: Solubility: ±0.9% S.D.
	Temperature: ±0.3 K.
	REFERENCES :

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,2-Dichlorobenzene; C₆H₄Cl₂; [95-50-1]</pre>	Chiou, C. T.; Schmedding, D. W.; Manes, M. Environ. Sci. Technol. <u>1982</u> , 16(1), 4-10.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm	$10^{5} x(1)^{c}$
25 1.5392 1.047	1.8921
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: An excess of 1,2-dichlorobenzene was equili- brated with water in screwcapped bottles in a reciprocal shaker for 24 hours. After two days settling, samples were taken from the solution for analysis by gas chromato- graphy. The chromatograph was equipped with a Ni ⁶³ EC detector. Analyses were continued until a constant concentration was observed.	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ Cl ₂ : Not specified. H ₂ O: Distilled. ESTIMATED ERROR: Solubility: ±5% (compiler). Temperature: ±0.5 K.
	REFERENCES :

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COMPO	ONENTS:	EVALUATOR:
(1)	1,3-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [541-73-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ O; [7732-18-5]	January 1983.

Seven experimental determinations have been published on the solubility of 1,3-dichlorobenzene in water in various temperature intervals (1-7), see Figure 1. Some rather serious discrepancies between the various solubility measurements are evident in the figure. The discrepancy between the two determinations (3) and (6) is about 47 percent at ambient temperature. The solubility data of Klemenc and Low (1) cover the temperature range between 293 and 333 K, while the measurements of Vesala (2,3) fall in the temperature range between 283 and 308 K. The remaining measurements reported (4-7) fall in the room temperature range.

The solubility measurements of Vesala (2,3) have been taken as most reliable and, therefore, these values were heavily weighed in the correlation of solubility versus Absolute temperature by means of a normal, three degree polynomial equation:

$$S_1(g(1)/kg) = 27.6827 - 2.61597 \times 10^{-1} T +$$

8.19706 x $10^{-4} T^2 - 8.4698 \times 10^{-7} T^3$ [1]

The significance of this equation representing the solubility data is that the curve passes through a minimum at 298.2 K. This observation is consistent with the theory discussed by Gill et al. (8) for the solubility of aromatic compounds in water.

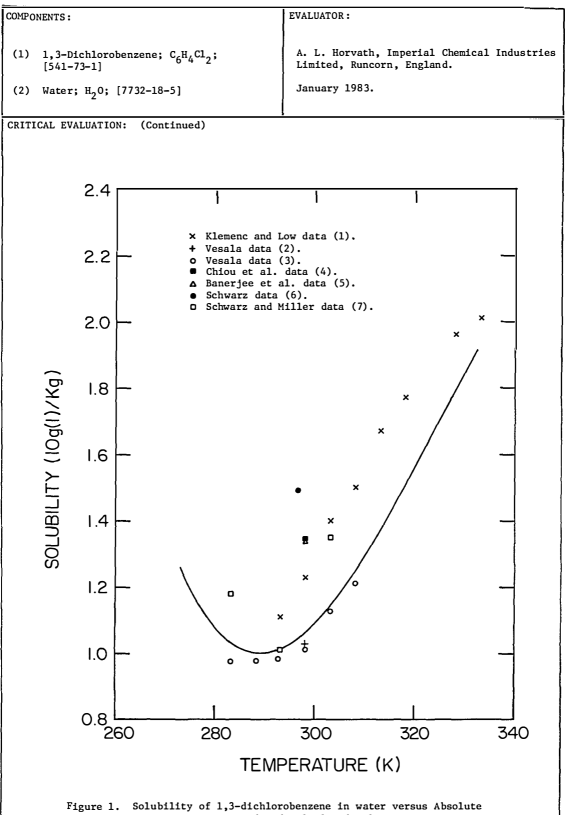
The errors in the recommended solubility values given below could be as large as ± 10 percent, particularly at the higher temperatures. Recommended solubility values calculated from equation [1] together with corresponding molarity and mole fraction values are listed in Table 1. Also, the solubility values calculated from equation [1] are shown in Figure 1 as a solid line along with the measured values.

Table 1. Solubility of 1,3-Dichlorobenzene in Water.

т/к	10^{4} mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
283.15	7.01	1.03	1.26
288.15	6.79	0.999	1.22
293.15	6.86	1.01	1.24
298.15	7.19	1.06	1.30
303.15	7.73	1.14	1.40
308.15	8.43	1.25	1.53
313.15	9.24	1.37	1.68
318.15	10.14	1.51	1.84
323.15	11.07	1.65	2.02
328.15	11.98	1.79	2.19
333.15	12.84	1.92	2.35

The recommended molar concentrations and mole fractions have been calculated from the g(1)/kg and the densities of the two components at the equilibrium temperatures indicated.

-		
COMP	ONENTS:	EVALUATOR:
(1)	1,3-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [541-73-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ 0; [7732-18-5]	January 1983.
CRIT	ICAL EVALUATION: (Continued)	
	REFERENC	
	KEFERENC	,25
1.	l. Klemenc, A.; Löw, M. Rec. Trav. Chim. Pays-Bas <u>1930</u> , 49(4), 629-40.	
2.	2. Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.	
3.	3. Vesala, A., Ph.D. Dissertation, University of Turku, Turku, 1973.	
4.	. Chiou, C. T.; Schmedding, D. W.; Maines, M. Environ. Sci. Technol. <u>1982</u> , 16(1), 4-10	
5.	 Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environ. Sci. Technol. <u>1980</u>, 14(10), 1227-9. 	
6.	. Schwarz, F. P. Anal. Chem. <u>1980</u> , 52(1), 10-15.	
7.	. Schwarz, F. P., Miller, J. Anal. Chem. <u>1980</u> , 52(13), 2162-4.	
8.	. Gill, S. J.; Nichols, N. F.; Wadso, I. J. Chem. Thermodyn. <u>1976</u> , 8(5), 445-52.	



2		1,3-Dichloi	robenzene
COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) 1,3-D [541-	Dichlorobenzene; C ₆ H ₄ Cl 73-1]	2;	Klemenc, A.; Löw, M. <i>Rec. trav. chim.</i> <i>Pays-Bas <u>1930</u>, 49(4), 629-40.</i>
(2) Water	; H ₂ 0; [7732-18-5]		
ARIABLES:			PREPARED BY:
Temperatur	e		A. L. Horvath
XPERIMENTA	AL VALUES:		
t/°C	10g(1)/kg(2) ^a	10 ³ mol(1)/	kg^{b} 10 ⁵ $x(1)^{c}$
20	1.11	0.7550	1.360
25	1.23	0.8366	1.507
30	1.40	0.9522	1.716
35	1.50	1.020	1.838
40	1.67	1.136	2.047
45	1.77	1.204	2.169
55	1.96	1.333	2.402
60	2.01	1.367	2.463
		AUXILIARY	INFORMATION
The determ based upon to the mea	NRATUS/PROCEDURE: ination of the solubil volumetric principles surement of excess sol apparatus as describe	applied ute in a	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ Cl ₂ : Kahlbaum reagent, purified by the method of Friedel and Crafts before use. H ₂ O: Distilled.
			ESTIMATED ERROR:
			Solubility: ±10% (compiler).
			Temperature: ±1 K (compiler).
			REFERENCES :
			 Rex, A. Z. Phys. Chem. <u>1906</u>, 55(A), 355-70.

COMPONENTS :		ORIGINAL MEASUREMENTS:
(1) 1,3-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [541-73-1]		Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u> .
(2) Water; H ₂ O; [7732-18-5]		
VARIABLES:		
Temperature		PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES	:	
t/°C 10g	(1)/kg ^a 10 ⁴ mol(1)/kg((2) ^b $10^5 x(1)^a$
10.0 0.9	7632 6.642 ± 0.088	3 1.1966
15.2 0.9	7764 6.651 ± 0.091	1.1982
19.6 0.9	8323 6.689 ± 0.096	1.2051
25.1 1.0	6.889 ± 0.079	1.2411
30.0 1.1	274 7.670 ± 0.061	1.3818
35.0 1.2	113 8.241 ± 0.159	1.4847
b. Reported.		
	AUXILIARY	INFORMATION
in a sealed flask magnetic stirrer for bath at constant to solution was filter plug, the solute was trimethylpentene. each sample was def metrically (2). Me	OCEDURE: nzene was mixed with water (1) with the aid of a or 48 hours in a water emperature. After the red through a glass-wool as extracted with 2,2,4- The optical density of cermined spectrophoto- can and standard devia- ted from three measure-	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ Cl ₂ : Fluka AG, puriss, >99% GLC, used as received. H ₂ O: Distilled, deionized, and degassed.
		ESTIMATED ERROR: Solubility: ±1.93%. Temperature: ±0.05 K. REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J. Chem. Soc. <u>1963</u> , Part III, 2716-23. 2. Wauchope, R. D.; Getzen, F. U. J. Chem. Eng. Data <u>1972</u> , 17(1), 38-41.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,3-Dichlorobenzene; C₆H₄Cl₂; [541-73-1]</pre>	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ⁴ mol(1)/kg(2	2) b $10^5 x(1)^{a}$
25 1.029 7.00	1.261
a. Calculated by compiler. b. Reported.	
AUXILIARY	INFORMATION
METHOD /APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established between the water and the 1,3-dichlorobenzene in a sealed flask (1) with the aid of a magnetic stirrer during 48 hours under isothermal conditions. After the sample was filtered through a glass-wool plug, the 1,3- dichlorobenzene was extracted with 2,2,4- trimethylpentene. Sample optical den- sities were determined spectrophotometri- cally (2). Five parallel determinations were done.	<pre>C₆H₄Cl₂: Commercial reagent of analytical grade distilled through a column resulting in a more than 99% pure sample. H₂O: Distilled, deionized, and degassed.</pre>
The reported work was based upon a Ph.D. dissertation (3).	ESTIMATED ERROR: Solubility: ±1.0%.
	Temperature: ±0.05 K.
	 REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J. Chem. Soc. <u>1963</u>, Part III, 2716-23. 2. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u>, 17(1), 38-41. 3. Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u>.

COMPONENTS :		ORIGINAL MEASUREMENTS:	
(1) 1,3-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [541-73-1]		Schwarz, F. P. Anal. Chem. <u>1980</u> , 52(1), 10-15.	
(2) Water; H ₂ 0; [7732-18-5]			
_			
VARIABLES:		PREPARED BY:	
One temperature		A. L. Horvath	
EXPERIMENTAL VALUES:			
t/°C 10g(1)/kg ^a 10	³ mol(1)/kg	^b $10^5 x(1)$ ^c	
23.5 1.49 ± 0.11	1.014	1.826	
23.5 1.44 ± 0.03	0.9796	1.765	
a. Reported. b. Calculated by F. W. Getzer c. Calculated by compiler.	n.		
	AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:	
Solubilities were determined by use elution chromatography analytical of The procedure involved the use of a solid packing coated with a known a solute in a transparent tube. Wate forced through the packed tube to a amounts of solute. The solubility calculated from the length of the depleted zone (as observed from the difference between the depleted pac and that in the remaining portion of tube) and the volume of water passe	technique. an inert amount of er was remove was solute e color cking of the ed	C ₆ H ₄ Cl ₂ : Commercial, 98 wt %. H ₂ O: Distilled.	
through the tube. Chemisorb P was the inert solid packing.	used as	ESTIMATED ERROR:	
		Solubility: ±7% S.D. Temperature: ±1.5 K.	
		REFERENCES :	
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COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,3-Dichlorobenzene; C₆H₄Cl₂; [541-73-1]</pre>	Schwarz, F. P.; Miller, J. Anal. Chem. 1980, 52(13), 2162-4.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
Temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
Experimentally determined values:	
10g(1)/kg	
t/°C Elution Chromatography U	V Absorption
10.0 1.16 ± 0.06	1.20 ± 0.06
20.0 0.89 ± 0.04	1.13 ± 0.05
30.0 1.39 ± 0.07	1.32 ± 0.05
Values derived from average measured so	lubilities
$t/^{\circ}C$ 10g(1)/kg ^a 10 ⁴ mol(1)/kg	
	5 1.446 ± 0.025
$20.0 1.01 \pm 0.12 6.870 \pm 0.810$	
30.0 1.35 ± 0.03 9.183 ± 0.20	
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Both elution chromatography and UV absorp-	C ₆ H ₄ Cl ₂ : Commercial, 98 wt. %.
tion methods were used to determine the aqueous solubilities. The agreement was	H ₂ 0: Distilled.
within an experimental error of 4% between the two methods. The average deviations	
were determined from several measurements made on different samples.	
The analytical procedures for determining	
organic liquid solubilities in water based on liquid phase elution chromatography	
has been described in (1).	ESTIMATED ERROR:
	Solubility: ±4%.
	Temperature: ±0.5 K.
	REFERENCES :
	 Schwarz, F. P. Anal. Chem. <u>1980</u>, 52(1), 10-15.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,3-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [541-73-1]	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environm. Sci. Techn. <u>1980</u> , 14(10), 1227-9.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10g(1)/dm ³ a 10 ⁴ mol(1)/dm ³	$10^5 x(1)^a$
25 1.335 9.08	1.641
a. Calculated by compiler.b. Reported.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of 1,3-dichlorobenzene was added to water in a stainless steel centrifuge tube which was then sealed. The equilibrium was	C ₆ H ₄ Cl ₂ : New England Nuclear, used as received.
established by allowing the sample to stand, with intermittent shaking, for a week at	H ₂ 0: Distilled.
constant temperature. The mixture was then centrifuged and aliquots of the solution were removed either by a pipet or syringe	
for analysis. Liquid scintillation counting with ^{14}C -labelled solute was employed in the	
solubility determinations. The entire pro- cedure was carried out at least twice and	
each analysis was also conducted in dupli- cate.	ESTIMATED ERROR:
	Solubility: ±1.1% S.D. Temperature: ±0.3 K.
	REFERENCES:

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COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,3-Dichlorobenzene; C₆H₄Cl₂; [541-73-1]</pre>	Chiou, C. T.; Schmedding, D. W.; Manes, M. Environ. Sci. Technol. <u>1982</u> , 16(1), 4-10.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm	$10^5 x(1)^{c}$
25 1.341 9.12	1.648
a Calculated by F. U. Cataoa	
a. Calculated by F. W. Getzen. b. Reported.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of 1,3-dichlorobenzene was equili-	C ₆ H ₄ Cl ₂ : Not specified.
brated with water in screwcapped bottles in	
a reciprocal shaker for 24 hours. After two days settling, samples were taken from	H ₂ 0: Distilled.
the solution for analysis by gas chromato-	
graphy. The chromatograph was equipped	
with a Ni^{63} EC detector. Analyses were con- tinued until a constant concentration was	
observed.	
	ESTIMATED ERROR:
	Solubility: ±5% (compiler).
	Temperature: ±0.5 K.
	REFERENCES :

COMPONEN	NTS:	EVALUATOR:
(1) 1,4	4-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; 06-46-7]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Wa	ter; H ₂ 0; [7732-18-5]	Мау 1979.

Two sets of experimental data have been reported in the literature for the solubility of liquid 1,4-dichlorobenzene in water (1,2). Klemenc and Low (1) reported measurements in the temperature range between 328 and 333 K while Wauchope and Getzen (2) reported data in the narrow temperature interval between 332 and 346 K, see Figure 1. There is, however, a considerable discrepancy between the two results. For example, at 333 K, the difference is about 25 percent between the reported solubility values.

While the early solubility measurements of Klemenc and Low (1) in 1930 used the volumetric determination of the excess solute, the more recent determination of Wauchope and Getzen (2) in 1972 employed the extraction method followed by spectrophotometric analysis. The reagents used in the latter investigation were of higher purity.

Klemenc and Low (1) did not state the accuracy or the reliability of their method; neither duplicate nor triplicate samples were taken. Their temperature control during the equilibration periods of their measurements was about ± 1 K. Therefore, a conservative estimate of the possible errors in their solubility determinations is about ± 10 percent. The investigation of Wauchope and Getzen (2) was done under much more controlled conditions. The time required for completion of the equilibrium was assured in the experiment and replicate samples were always withdrawn and analyzed. The shorter equilibration times in the measurements of Klemenc and Low (1) may well account for the low solubility values at 328 and 333 K.

The solubility data of Wauchope and Getzen (2) have been assigned a higher weight in the establishment of recommended solubility values. The data have been correlated against Absolute temperature using the following second degree polynomial equation:

$$S_1(g(1)/kg) = 13.974 - 8.5829 \times 10^{-2} T + 1.3365 \times 10^{-4} T^2$$
 [1]

Recommended solubility values calculated from equation [1] together with corresponding molarity and mole fraction values are listed in Table 1. Also, the solubility values calculated from equation [1] are shown in Figure 1 as a solid line along with the measured values.

Table 1. Solubility of 1,4-Dichlorobenzene in Water.

т/к	10^3 mol(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
328.15	1.35	2.01	2.46
333.15	1.43	2.14	2.62
338.15	1.56	2.33	2.86
343.15	1.72	2.59	3.18
348.15	1.94	2.92	3.58

The recommended molar concentrations and mole fractions have been calculated from the g(1)/kg and the densities of the two components at the equilibrium temperatures indicated.

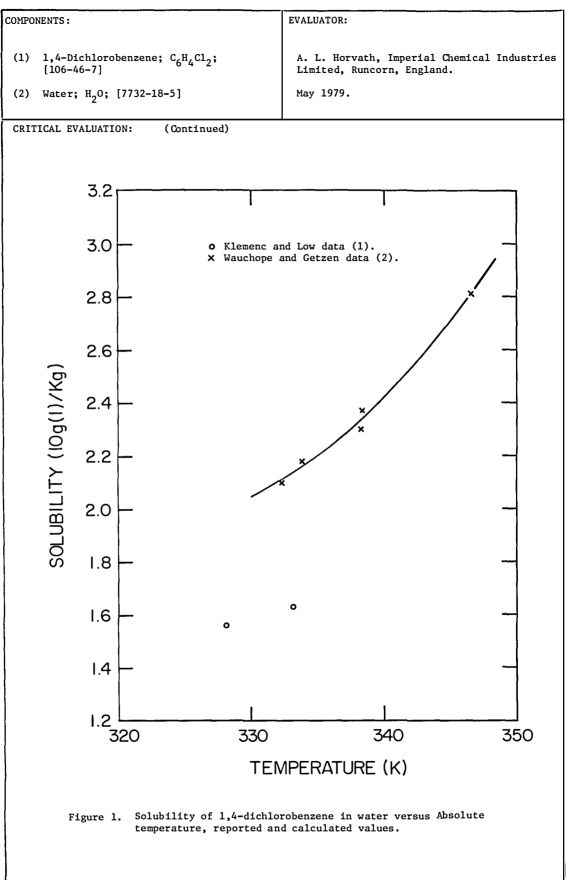
REFERENCES

1. Klemenc, A.; Löw, M. Rec. Trav. Chim. Pays-Bas <u>1930</u>, 49(4), 629-40.

2. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data 1972, 17(1), 38-41.

NOTE:

The critical evaluation for solid 1,4-dichlorobenzene with water appears on page 103.



COMPONENTS :				
		·····	ORIGINAL MEASUREMENTS:	
(1) 1,4-Dichlord [106-46-7]	obenzene; C ₆ H ₄ Cl ₂ ;	:	Klemenc, A.; Löw, M. <i>R</i> Pays-Bas <u>1930</u> , 49(4), 6	
(2) Water; H ₂ O;	[7732-18-5]			
				l
VARIABLES:		<u>2277</u>	PREPARED BY:	
Temperature			A. L. Horvath	
EXPERIMENTAL VALU	IES:			
t/°C 10)g(1)/kg(2) ^a	10 ³ mo1(1)/1	$10^{5}x(1)^{c}$	
55	1.56	1.061	1.912	
60	1.63	1.109	1.998	
	ed by compiler.			
		AUXILIARY	INFORMATION	
based upon volume the measurement o	PROCEDURE: of the solubilit stric principles a f excess solute i as described by R	y was pplied to n a cali-	SOURCE AND PURITY OF MAT C ₆ H ₄ Cl ₂ : Prepared by chl zene using iodi The preparate of	lorination of ben- lne as catalyst. was recrystallized In ethanol before

COMPONENTS :				ORIGINAL MEASUREMENTS:
(1) 1,4-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [106-46-7]			Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u> , 17(1), 38-41.	
(2) Water; H ₂ 0; [7732-18-5]				
VARIA	BLES:		<u></u>	PREPARED BY:
	rature			A. L. Horvath
}				
EXPER	IMENTAL VAL	JES:		
	t/°C 1	0g(1)/kg ^a 10	³ mol(1)/kg ¹	$10^5 x(1)^{c}$
	59.2	2.10	1.429	2.574
	60.7	2.18	1.483	2.672
1	65.1	2.30	1.565	2.819
	65.2	2.37	1.612	2.905
	73.4	2.81	1.911	3.445
	c. Calcula	ted by compiler.		
			AUXILIARY	INFORMATION
METHO	DD/APPARATUS	/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:
An excess of 1,4-dichlorobenzene, 20 g, in water was shaken gently for one week in a water bath at constant temperature. Repli- cate samples were filtered through a glass- wool plug. Then, the aqueous solution was extracted with cyclohexane which was then analyzed spectrophotometrically for organic solute using a Cary 14 spectrometer. The article was based upon a Ph.D. disser- tation (1).		ek in a . Repli- a glass- tion was as then r organic r.	<pre>C₆H₄Cl₂: Recrystallized three times and vacuum-sublimed twice of Matheson, Coleman, and Bell reagent. H₂O: Distilled and deionized.</pre>	
1				ESTIMATED ERROR:
				Solubility: ±3%.
				Temperature: ±0.5 K.
[REFERENCES:
				 Wauchope, R. D., Ph.D. Dissertation, North Carolina State Univ., Raleigh, <u>1970</u>.

COMPONENTS :	EVALUATOR:
<pre>(1) 1,4-Dichlorobenzene; C₆H₄Cl₂; [106-46-7]</pre>	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2) Water; H ₂ 0; [7732-18-5]	September 1982.

There have been eight published sets of data on the solubility of 1,4-dichlorobenzene in water at ordinary temperatures. Accordingly, the evaluation of the solubility data for this system lies on a firmer base than in the cases of the other solid halogenated benzenes in water.

The oldest data involving this system, determined by Klemenc and Low (1) in 1930, appear to be low and are imprecise because of the method used for saturation of water. There is, however, some evidence that the method allows the separation of the solubilities of the two crystallographic forms of 1,4-dichlorobenzene although the experimental values give no distinct temperature for the conversion from the α -form to the β -form. The solubility measured by Gross and Saylor (2) at 308.2 K is probably too low because of the short time periods involved in the equilibrations. For the same reason, the value determined by Andrews and Keefer (3) must be rejected as doubtful. Booth and Everson (4) have applied a residue-volume method of Vaughn and Nutting (5) and found the solubility of 1,4dichlorobenzene in water to be less than 0.5 g(1)/dm³(2). Their method is rapid but insensitive giving only rough values for substances having low solubilities.

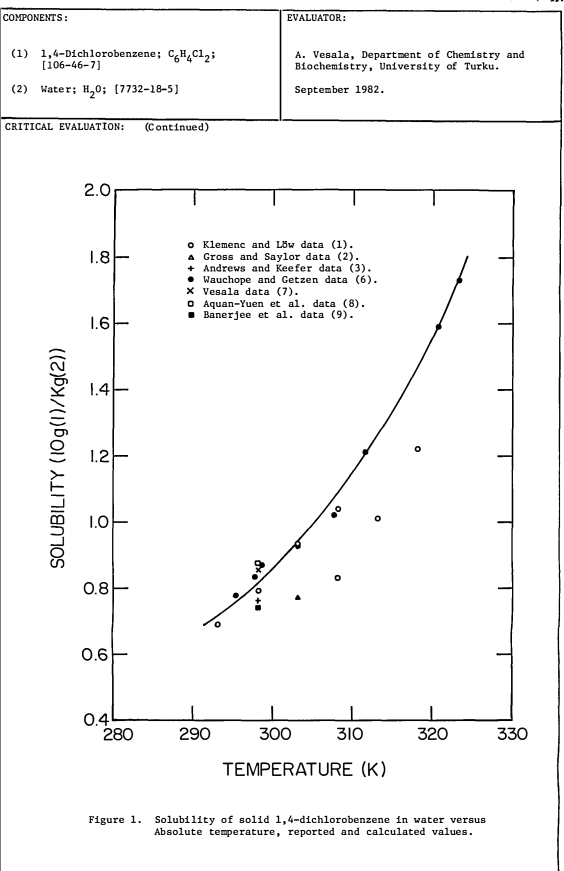
More reliable data have been produced by Wauchope and Getzen (6) who reported errors to be in the range of 2 percent. These data are supported by the solubility values obtained by Vesala (7) as part of a study of the transfer free energies of certain nonelectrolytes from H₂O to D₂O. The recent data of Aquan-Yuen et al. (8) of the solubilities of 1,4dichlorobenzene in aqueous electrolyte solutions produce a value for the solubility in water that agrees well with the above values. Banerjee et al. (9) report a solubility that is somewhat different from the former ones. The reason for the differences may be attributed to the radiochemical method of analysis used. For instance, the radiochemical purity of the substrate and the quenching of the samples in the scintillation analysis remain open to question. Their value can thus be regarded only as a slight support to the other data - even in cases of a full agreement.

The recommended solubility values are calculated on the basis of the data given by Wauchope and Getzen (6). However, instead of their smoothed equation, the calculations are done using an equation of simpler form. This, in turn, suggests that the magnitude of the reported errors is too optimistic. (In the work of Wauchope and Getzen, the meaning of the term "av.% dev.obsd. smoothed solubility" is somewhat unclear.) The values calculated from the simpler equation and the experimental data allow a fairly good estimate of precision. A standard deviation for a single value established in this fashion is of the order of 4.0 mg(1)/kg(2) or 0.027 mmol(1)/kg(2). The existence of the two crystallographic forms of the solute in the range of temperatures where the solubilities were measured is probably one reason for the deviations. However, this brings about no greater effect in the lower temperature range which suggests that the value for 298.2 K, for instance, is quite reliable. The simplified equation for concentration, g(1)/kg(2), in terms of Absolute temperature, T, is as follows:

$$\log_{10}(S_1(g(1))/kg(2)) = 2.86294 - 1176/T$$
 [1]

The observed values from the seven relevant data are shown in Figure 1 together with the calculated behavior (shown as a solid line) from the simpler equation discussed above. The densities of pure water and the saturated solutions were assumed to be equal for the determination of the reported molarity values from the calculated g(1)/kg(2) values.

The recommended g(1)/kg solubility values for solid 1,4-dichlorobenzene in water calculated from values obtained from equation [1] together with corresponding molarity and mole fraction values are listed in Table 1.



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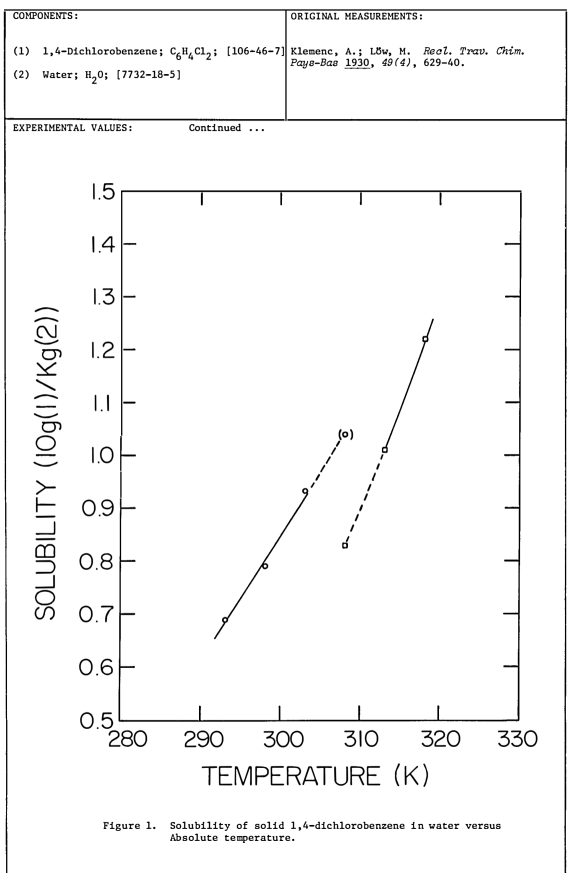
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	1,4-0101		
COMPONENTS:		EVALUATOR:	
(1) 1,4-Di [106-4	chlorobenzene; C ₆ H ₄ Cl ₂ ; 6-7]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.	
(2) Water;	H ₂ 0; [7732-18-5]	September 1982.	
CRITICAL EV	ALUATION: (Continued)		
Table 1. S	olubility of 1,4-Dichlorobenzene	in Water.	
Т	/K 10 ⁴ mo1(1)/dm ³ 10 ²	$g(1)/kg$ $10^5 x(1)$	
283	.15 3.48	5.12 0.628	
288	.15 4.11	6.05 0.741	
293	.15 4.82	7.10 0.870	
298	.15 5.62	8.29 1.016	
303	.15 6.52	9.63 1.18	
308	.15 7.53	11.13 1.36	
313	.15 8.65	12.8 1.57	
318	.15 9.88	14.7 1.80	
323	.15 11.25	16.7 2.05	
328	.15 12.4	19.0 2.33	
	librium temperatures indicated.	values and the densities of the two components	
	REFEREN	ICES	
1. Klemen	c, A.; Löw, M. <i>Rec. Trav. Chim</i> .	Pays-Bas 1930, 49(4), 629-40.	
2. Gross,			
3. Andrew	3. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1950</u> , 72(7), 3113-6.		
4. Booth,	Booth, H. S.; Everson, H. E. Ind. Eng. Chem. <u>1948</u> , 40(8), 1491-3.		
. Vaughn, T. H.; Nutting, E. G. Ind. Eng. Chem. Anal. Ed. <u>1942</u> , 14(6), 454-6.			
6. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u> , 17(1), 38-41.			
7. Vesala	V. Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.		
8. Aquan-	Aquan-Yuen, M.; Mackay, D.; Shiu, W. Y. <i>J. Chem. Eng. Data</i> <u>1979</u> , 24(1), 30-4.		
	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environ. Sci. Technol. <u>1980</u> , 14(10), 1227-9.		

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,4-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [106-46-7]	Andrews, L. J.; Keefer, R. M. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1950</u> , <i>72(7)</i> , 3113-6.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm ³	$^{3 b} 10^{6} x(1)^{b}$
25.0 7.6 5.17	9.34
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Water was saturated with 1,4-dichlorobenzene in a sealed Erlenmeyer flask by rotating the	C ₆ ^H ₄ Cl ₂ : Commercial product (Eastman Kodak Co.), used as received.
flask in a constant temperature bath for 20 hours. Measured volumes of the saturated	H ₂ 0: Distilled.
solution were extracted with measured volumes of n-hexane for analysis. The optical den- sity of the extract was measured against a	
n-hexane standard at 234 nm using a Beckman spectrophotometer (1).	
	ESTIMATED ERROR:
	Solubility: >10% (compiler).
	Temperature: ±0.2 K (compiler).
	REFERENCES: 1. Andrews, L. J.; Keefer, R. M. J. Am. <i>Chem. Soc.</i> <u>1949</u> , 71(11), 3644-7.

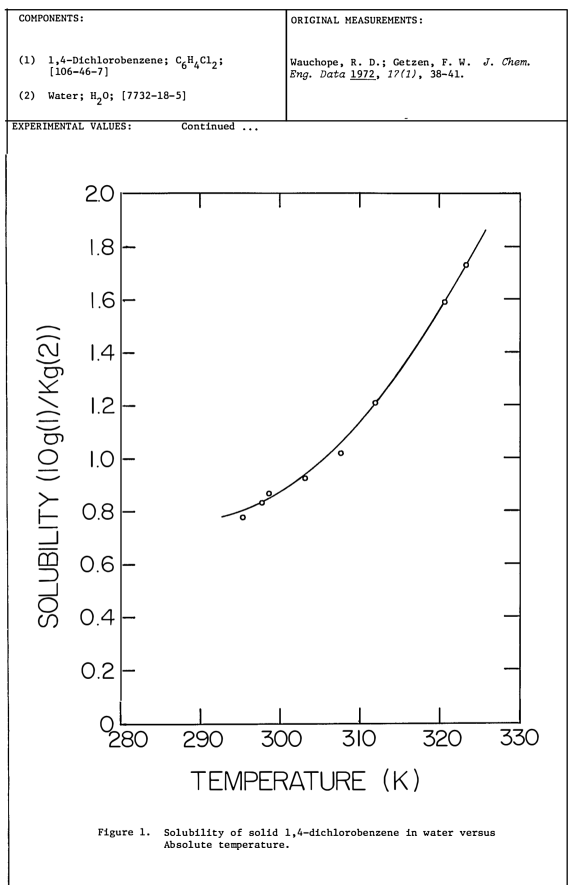
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I,4-DICNIO	robenzene 107
COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,4-Dichlorobenzene; C₆^{II}₄Cl₂; [106-46-7]</pre>	Klemenc, A.; Low, M. <i>Recl. Trav. Chim.</i> <i>Pays-Bas <u>1930</u>, 49(4), 629-40.</i>
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
Temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C $10^{2}g(1)/kg(2)^{a}$ $10^{4}mol(1)$	$/kg^{b}$ 10 ⁶ x(1) ^b
20 6.89 4.687	8.444
25 7.91 5.380	9.694
30 9.33 6.346	11.04
35 ^c 10.4 8.30 7.074 5	.646 12.75 10.17
40 10.1 6	.870 12.38
45 12.2 8	.298 14.95
	Continued
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
The equilibrations were done over periods from 2 to 17 days (average 7-10 days). Gra- vimetric analysis was the basis for the con- centration determinations. The solute was placed in a spiral tube and weighed. Then, a known amount of water was passed through the tube and it was re-weighed. The weight loss and the mass of the water passed through the tube gave the solubility value directly.	<pre>C₆H₄Cl₂: Synthesized from benzene and chlorine, recrystallized from abs. ethanol, m.p. 53°C. The purity of the product was checked by elemental analysis.</pre> H ₂ O: Source and purity not specified.
	ESTIMATED ERROR: Solubility: >10% (evaluated on the basis of deviations from the averages). Temperature: ±0.5 K. REFERENCES:



COMPONENTS :		ORIGINAL MEASUREMENTS:
<pre>(1) 1,4-Dichlorobenzene; ([106-46-7]</pre>	6 ^H 4 ^{C1} 2;	Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u> , 17(1), 38-41.
(2) Water; H ₂ 0; [7732-18-5	5]	
2		
VARIABLES: Temperature		PREPARED BY: A. Vesala
Temperature		A. Veodia
EXPERIMENTAL VALUES:		
t/°C 10g(1)/kg(2)	a 10 ⁴ mol(1)/	$kg^{b} 10^{5}x(1)^{b}$
22.2 0.778	5.292	0.9535
24.6 0.834	5.673	1.022
25.5 0.869	5.911	1.065
30.0 0.926	6.298	1.135
34.5 1.02	6.938	1.250
38.4 1.21	8.230	1.483
47.5 1.59	10.81	1.949
50.1 1.73	11.77	2.120
a. Reported. b. Calculated by F. W Measurements are shown		gure 1. Continued
	AUXILIARY	INFORMATION
		SOURCE AND PURITY OF MATERIALS:
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by allowing the mixtures to stand in glass stoppered flasks, with occasional shaking, for 1-3 week periods at constant temperature. Samples of the saturated aqueous solutions were withdrawn with pipettes through glass-wool plugs and transferred to separatory funnels for weighing. An extraction with cyclohexane was then performed after which the extracts were analyzed spectrophotometrically at 273 nm. The concentrations were established from previously obtained absorbance values of standard reference solutions (which were observed to obey Beer's law).		<pre>Source And Former of Markans. C₆H₄Cl₂: Commercial product (Matheson, Coleman and Bell), recrystallized three times from abs. ethanol, vacuum sublimed twice, UV-spectrum and melting point were determined and were identical to published data. H₂O: Distilled and deionized water.</pre>

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,4-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [106-46-7]	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.
(2) Water; H ₂ 0; [7732-18-5]	
£	
VARIABLES: One temperature	PREPARED BY: H. Lonnberg
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{2}g(1)/kg(2)^{a}$ $10^{4}mol(1)$	$/kg^{b} = 10^{5} x(1)^{b}$
25.0 8.55 5.816	1.048
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Saturation of the 1,4-dichlorobenzene in water was established by shaking the mixture	C ₆ H ₄ Cl ₂ : Commercial product (E. Merck AG), recrystallized twice from abs.
in a sealed tube suspended in a thermostat water bath for 1-2 weeks. The sealed tube	ethanol.
was allowed to stand for two to three days without shaking before analysis. Samples	H ₂ 0: Distilled, deionized and degassed.
from the saturated solution were extracted with 2,2,4-trimethylpentene. Then, the	
optical densities of the liquid extracts were determined spectrophotometrically. The	
concentration was established from optical	
density using a previously prepared standard calibration curve.	ESTIMATED ERROR:
	Solubility: ±1.5% (author).
	Temperature: ±0.05 K (author).
	REFERENCES :

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COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,4-Dichlorobenzene; C₆H₄Cl₂; [106-46-7]</pre>	Gross, P. M.; Saylor, J. H. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1931</u> , <i>53(5)</i> , 1744-51.
(2) Water; H ₂ 0; [7732-18-5]	
_	
VARIABLES:	PREPARED BY: A. Vesala
One temperature	A. VESALA
EXPERIMENTAL VALUES:	
LA DRIIDRIAL VALUED.	
t/°C 10 ² g(1)/kg(2) ^a 10 ⁴ mol(1),	$u_{\rm kg}^{\rm b}$ 10 ⁶ $x(1)^{\rm b}$
30 7.7 5.24	9.44
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturated solution was prepared by shaking the substances in a thermostat bath for at least 12 hours. An interferometric method was used for the analysis of the	C ₆ ^{II} ₄ Cl ₂ : Commercial product (Eastman Kodak Co.), recrystallized twice from ethanol, m.p. 52.84°C.
saturated solution (1). Samples were with- drawn from several saturation flasks and read against a pure water sample in an interferometer. The concentration was estab- lished from a calibration obtained from pre-	H ₂ O: Distilled water "of good quality".
vious measurements made with a reference solution.	
	ESTIMATED ERROR:
	Solubility: ±5% (authors).
	Temperature: ±0.01 K (authors).
	REFERENCES:
	1. Gross, P. M. J. Am. Chem. Soc. <u>1929</u> , 51(8), 2362-6.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 1,4-Dichlorobenzene; C₆II₄Cl₂; [106-46-7]</pre>	Aquan-Yuen, M.; Mackay, D.; Shiu, W. Y. J. Chem. Eng. Data <u>1979</u> , 24(1), 30-4.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ² g(1)/dm ^{3 a} 10 ⁴ mol(1)/d	$m^{3 b} 10^{5} x(1)^{b}$
25 8.715 5.9283	1.0713
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Saturated solutions were prepared by adding an excess of solute to water in a closed flask with standing for 24 hours. Then, the sample was allowed to settle at 25°C in a thermostat bath for at least 48 hours before analysis. The solubility was determined by solvent extraction followed by gas chromato- graphic analysis using a Hewlett Packard Model 5750 apparatus (equipped with a flame ionization detector and a 10 ft stainless steel column packed with 10% SE 30 ultra- phase on high performance Chromosorb P, 60/80 mesh).	C ₆ H ₄ Cl ₂ : Commercial product (BDH), evi- dently used as received. H ₂ O: Source and purity not specified. ESTIMATED ERROR: Solubility: ±3% (authors, the value is based
	on the std. error of the least squares fit on the solubility of 1,4-dichlorobenzene in water and aqueous electrolyte solutions. Temperature: ±0.2 K (compiler). REFERENCES:

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 1,4-Dichlorobenzene; C₆^{II}₄Cl₂; [106-46-7]</pre>	Booth, H. S.; Everson, H. E. <i>Ind. Eng.</i> <i>Chem.</i> <u>1948</u> , 40(8), 1491-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	• • • • • • • • • • • • • • • • • • •
$t/^{\circ}C$ 10g(1)/kg(2) ^a 10 ³ mol(1)/kg(2)	$(g^{b} 10^{5}x(1)^{b})$
25.4 <5 <3.4	<6.1
a. Reported. b. Calculated by F. W. Getzen.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The composition analysis was based upon a volumetric method described in (1). In this	C ₆ H ₄ Cl ₂ : Commercial reagents of highest grade, probably used as received.
method, further addition of solute to a vol- ume of solution yielded, upon saturation,	H_20 : Source and purity not specified.
proportional amounts of undissolved solute	2
residue which could be measured volumetri- cally. A plot of added solute mass versus	
volume of residue gave a straight line the intercept of which gave the solubility.	
Stoppered tubes with capillaries graduated in	
steps of 0.05 ml were used for the measure- ments. A known volume of solvent (50 ml) was	ESTIMATED ERROR:
added to the tube in a constant temperature bath and weighted quantities of the solid	Solubility: Within 0.1 g as reported in
solute were added to the liquid. The mixture	
was then shaken for 5 minutes, returned to the bath for a minimum of 10 minutes, and	Temperature: ±0.2 K (compiler).
centrifuged for 5 minutes. Then, the volume of residue was read. The procedures were repeated to ensure that the equilibrium had	REFERENCES: 1. Vaughn, T.H.; Nutting, E. G. Ind. Eng. Chem., Anal. Ed. 1942, 14(6), 454-6.
been reached.	5,500,5 1,140, Due <u>1772</u> , 13107, 434-0.

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	1,4-Dichlo	robenzene 115
COMPO	DNENTS:	ORIGINAL MEASUREMENTS:
	1,4-Dichlorobenzene; C ₆ H ₄ Cl ₂ ; [106-46-7] Water; H ₂ O; [7732-18-5]	Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. <i>Environm. Sci. Techn.</i> <u>1980</u> , 14(10), 1227-9.
VARIA	ABLES:	PREPARED BY:
	One temperature	A. Vesala
EXPER	RIMENTAL VALUES:	
	$t/^{\circ}C = 10^{2}g(1)/dm^{3}a = 10^{4}mol(1)/d$	$^{3 b}$ 106 6 $^{(1)}$ a
	25 7.380 5.02	9.071
	a. Calculated by F. W. Getzen. b. Reported.	
5a		
	AUXILIARY	INFORMATION
METH	OD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
stai: stan	equilibrium was performed in sealed nless steel centrifugre tubes with con- t or intermittent shaking. The equili- m was generally complete within 1 week.	C ₆ H ₄ Cl ₂ : Commercial reagent. the ¹⁴ C- labeled compound was purchaed by NEN, the nonlabeled one by Aldrich.
The r	mixture was then centrifugred for 60 tes after which aliquots of the solution	H20: Distilled.
were	removed either by pipet or syringe for ysis. Liquid scintillation counting with	
¹⁴ C-	labelled solute was employed in the solu-	
was o	ty determinations. The entire procedure carried out at least twice and each anal-	
ysis	was also conducted in duplicate.	ESTIMATED ERROR:
		Solubility: ±6.0% (std. deviation estimated by authors).
		Temperature: ±0.2 K (equilibration), ±0.3 K (centrifugation).
		REFERENCES :

COMPONENTS:		EVALUATOR:
(1)	2,4-Dichlorophenol; C ₆ H ₄ Cl ₂ O; [120-83-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

The solubility of 2,4-dichlorophenol in water was determined many decades ago by Mosso (1). While the measurements may be subject to question because of their age and obscurity, the reported melting point strongly indicates that the "m-dichlorophenol" does refer to 1,2-dichlorophenol. Undoubtedly, the experimental procedure was based upon a classical gravimetric analysis. Because of the lack of other data, the value recommended here must be classified as doubtful.

The following solubility value for 2,4-dichlorophenol in water is based solely upon the work reported by Mosso.

T/K	10^2 mol(1)/dm ³	g(1)/kg	$10^4 x(1)$
293.15	2.7	4.5	4.9

REFERENCES

1. Mosso, U. Jahresber. Fortschr. Chem. 1887, 1800.

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 2,4-Dichlorophenol; C ₆ ^H ₄ Cl ₂ O; [120-83-2]	Mosso, U. Jahresber. Fortschr. Chem. <u>1887</u> , 1800.
(2) Water; H ₂ 0; [7732-18-5]	
_	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C g(1)/kg(2) ^a 10 ² mol(1)/kg	^b $10^4 x(1)^{b}$
20 4.47-4.66 2.730-2.845	4.938-5.148
a. Reported. b. Calculated by F. W. Getzen.	
- AUXILI ARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS;
Experimental methods were not described.	C ₆ ^H ₄ Cl ₂ O: Synthesized from dry chlorine and phenol, distilled and re- crystallized from benzene, re- ported melting point 43 - 44°C.
	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR:
	REFERENCES :

COMPONENTS :	EVALUATOR:
<pre>(1) 1,2-Difluorobenzene; C₆H₄F₂; [367-11-3]</pre>	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ 0; [7732-18-5]	May 1979.

The experimental results consist of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonble observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1,2-difluorobenzene in water is tentative:

T/K $10^2 \text{mol}(1)/\text{dm}^3$ g(1)/kg $10^4 x(1)$ 298.15 1.0 1.14 1.81

REFERENCES

 Yalkowsky, S.H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

2. Yalkowsky, S.H., Personal Communication, 1979.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,2-Difluorobenzene; C ₆ H ₄ F ₂ ; [367-11-3]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.
	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/dm ³ a 10 ² mol(1)/dm ³	^b $10^4 x(1)$ ^c
25 1.141 1.00	1.809
a. Calculated by F. W. Getzen. b. Reported.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
_	· · · · · · · · · · · · · · · · · · ·
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,2-difluorobenzene in water was agitated at room temperature for	C ₆ H ₄ F ₂ : Aldrich commercial grade, used as received.
a period of about 24 hours and then fil-	as received.
tered. The filtrate was diluted and	H ₂ O: Deionized.
assayed spectrophotometrically. The deter- mination was done in duplicate.	
•	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES:

COMPONENTS :		EVALUATOR:
(1) 1,3-D: [372-]	ifluorobenzene; C ₆ H ₄ F ₂ ; 18-9]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water	; H ₂ 0; [7732-18-5]	May 1979.

The experimental results consist of a single value at 298.15 K published in 1979 by Yalkowsky et al. (1) of the Upjohn Company, Kalamazoo, Michigan. According to the authors (2), the accuracy of the experimental determinations was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure.

The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1,3-difluorobenzene in water is tentative:

т/к	10^2 mol(1)/dm ³	g(1)/kg	$10^4 x(1)$
298.15	1.0	1.14	1.81

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,3-Difluorobenzene; C ₆ H ₄ F ₂ ; [372-18-9]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.
3 · -	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
	. ,
$t/^{\circ}C$ g(1)/dm ³ a 10 ² mol(1)/dm ³	$10^4 x(1)^{c}$
25 1.141 1.00	1.809
a. Calculated by F. W. Getzen.	
b. Reported.c. Calculated by compiler.	
c. calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,3-difluorobenzene in	C ₆ H ₄ F ₂ : Aldrich commercial grade, used
water was agitated at room temperature for	as received.
a period of about 24 hours and then fil- tered. The filtrate was diluted and	H ₂ 0: Deionized.
assayed spectrophotometrically. The deter- mination was done in duplicate.	2
mination was done in dupitcate.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES :
	ELENCED,

1,4-Difluorobenzene

COMP	ONENTS:	EVALUATOR:
(1)	1,4-Difluorobenzene; C ₆ H ₄ F ₂ ; [540-36-3]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ O; [7732-18-5]	May 1979.

CRITICAL EVALUATION:

Two experimental measurements have been reported on the solubility of 1,4-difluorobenzene in water (1,2). Jockers (1) has determined the solubility at high temperatures between 533 K and 553 K and high pressures only in connection with studies of the parameters that influence phase separation. The single solubility measurement reported by Yalkowsky et al. (2) cannot be easily compared with the results of Jockers. Both results are included in the table below.

According to the authors (3), the accuracy of the experimental determinations reported by Yalkowsky et al. was \pm 10 percent. This is a reasonable observation considering the details available on the experimental procedure. The manufacturer's reagent (Aldrich and Eastman) was neither further purified nor degassed before use. The time required for equilibration was indicated as between 4 and 48 hours. The saturated solutions were assayed spectrophotometrically.

The solubility value was the average of at least two independent determinations according to the investigators. The experimental result was expressed in Briggsian logarithms only with three significant figures.

The objective of the solubility measurements was to extend the correlation technique of aqueous solubilities to a broader group of planar nonelectrolytes by involving the melting points and total molecular surface areas as dependent variables. Using such methods, the authors also showed that branched and cyclic compounds have greater solubilities in water than corresponding linear compounds.

To compare, relate, and correlate the solubility data for liquid halogenated benzenes in water, use was made of the theoretical relationship between the molar solubility and solute molar volumes at 298.15 K as discussed in the Introduction. A data comparison with previously selected solubilities shows that new measurements are urgently required in order to resolve the anomalous trend in the correlated experimental data reported by Yalkowsky et al.

The uncertainty in the single reported solubility value may be as large as 10 percent or even larger. The following solubility value for 1,4-difluorobenzene in water is tenta-tive:

$P/P_{\theta} = 1.0$				
т/к	10^2 mol(1)/dm ³	g(1)/kg	$10^4 x(1)$	
298.15	1.07	1.22	1.94	

	$P/P_{0} = 80 - 100 \times 10^{5}$		
T/K	$mol(1)/dm^3$	10 ⁻² g(1)/kg	$10^{2}x(1)$
533.15-553.15	3.47	3.50	7.84

REFERENCES

- 1. Jockers, R., Ph.D. Dissertation, University of Bochum, Bochum, 1976.
- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.

3. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:		12
	ORIGINAL ME	ASUREMENTS :
 (1) 1,4-Difluorobenzene; C₆H₄F₂; [540-36-3] (2) Water; H₂O; [7732-18-5] 		R., Ph.D. Dissertation, University , Bochum, <u>1976</u> , pp 94-5.
VARIABLES:	PREPARED BY	:
Temperature and pressure	A. L. Hory	vath
-		
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ P/bar $10^{-2}g(1)/kg^{a}$	mol(1)/kg ^b	$10^2 x(1)^{c}$
260.0 80 3.501	3.069	7.84
270.0 92 3.501	3.069	7.84
280.0 100 3.501	3.069	7.84
b. Calculated by F. W. Getzen. c. Reported.		
AUXILIARY		
	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND	PURITY OF MATERIALS:
		PURITY OF MATERIALS: Fluka AG., Buchs, Schweiz, 99.5% pure, redistilled before use. Boiled with KMnO ₄ and redis- tilled before use.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 1,4-Difluorobenzene; C ₆ H ₄ F ₂ ; [540-36-3]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.
• • • •	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/dm ^{3 a} 10 ² mol(1)/dm ³	^b $10^4 x(1)$ ^c
25 1.221 1.07	1.935
a. Calculated by F. W. Getzen. b. Reported.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A small excess of 1,4-difluorobenzene in	$C_6H_4F_2$: Aldrich commercial grade, used
water was agitated at room temperature for a period of about 24 hours and then fil-	as received.
tered. The filtrate was diluted and assayed spectrophotometrically. The deter-	H ₂ 0: Deionized.
mination was done in duplicate.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	REFERENCES :
1	

COMPONENTS:	EVALUATOR:
(1) 1,2-Diiodobenzene; C ₆ H ₄ I ₂ ; [615-42-9]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2) Water; H ₂ 0; [7732-18-5]	November 1979.

A single solubility measurement of 1,2-diiodobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,2-diiodobenzene in water is reported here as a tentative value:

т/к	10^{5} mol(1)/dm ³	10 ² g(1)/kg	$10^{6}x(1)$
298.15	5.8	1.92	1.05

REFERENCES

- 1. Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS:	ORIGINAL MEASUREMENTS:
 1,2-Diiodobenzene; C₆H₄I₂; [615-42-9] Water; H₂0; [7732-18-5] 	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
VARIABLES: One temperature	PREPARED BY: A. Vesala
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{2}g(1)/dm^{3}a = 10^{5}mol(1)/dm^{3}$	$10^{3} b$ $10^{6} x(1)^{a}$
25 1.91 5.8	1.05
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: A small excess of solute in water was agi- tated for a period of 4-48 hours (the exact time for equilibration was not reported in single cases) and then filtered. The sat- urated solution was extracted, diluted with solvent, and assayed spectrophotometrically. At least two independent determinations were carried out.	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ I ₂ : Commercial reagent (Aldrich or Eastman), used as received. H ₂ O: Source and purity not specified.
	ESTIMATED ERROR: Solubility: ±10% (authors). Temperature: ±1 K (authors). REFERENCES:

COMPO	ONENTS:	EVALUATOR:
(1)	1,3-Diiodobenzene; C ₆ H ₄ I ₂ ; [626-00-6]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ O; [7732-18-5]	November 1979.

A single solubility measurement of 1,3-diiodobenzene in water has been reported by Yalkowsky, Orr, and Valvani (1). The principal motivation for their study was to obtain certain correlations for solubilities of halogenated benzenes in water.

In these measurements, a conventional experimental procedure was employed. Commercial reagents were used without further purification. All measurements were done at room temperature (25 \pm 1°C). The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. The saturated sample analyses were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (2), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The Possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The solubility of 1,3-diiodobenzene in water is reported here as a tentative value:

т/к	10^5 mol(1)/dm ³	10 ³ g(1)/kg	$10^{7}x(1)$
298.15	2.7	8.93	4.87

REFERENCES

- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 2. Yalkowsky, S. H., Personal Communication, 1979.

COMPONENTS: ORIGINAL MEASUREMENTS: (1) 1,3-Diiodobenzene; C₆H₄I₂; [626-00-6] Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. 1979, 18(4), 351-3. (2) Water; H₂O; [7732-18-5] VARIABLES: PREPARED BY: One temperature A. Vesala EXPERIMENTAL VALUES: $t/^{\circ}C$ 10³g(1)/dm³ a 10⁵mol(1)/dm³ b 10⁷x(1) a 2.7 4.88 25 8.91 a. Calculated by F. W. Getzen. b. Reported. AUXILIARY INFORMATION METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS: Commercial reagent (Aldrich or A small excess of solute in water was agi-C6H412: Eastman), used as received. tated for a period of 4-48 hours (the exact time for equilibration was not reported in H₂0: single cases) and then filtered. The saturated solution was extracted, diluted Source and purity not specified. with solvent, and assayed spectrophotometrically. At least two independent determinations were carried out. ESTIMATED ERROR: Solubility: ±10% (authors). Temperature: ±1 K (authors). **REFERENCES:**

COMPONENTS:	EVALUATOR:
(1) 1,4-Diiodobenzene; C ₆ H ₄ I ₂ ; [624-38-4]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2) Water; H ₂ 0; [7732-18-5]	September 1979.

Two experimental values have been reported for the solubility of 1,4-diiodobenzene in water, that determined by Andrews and Keefer (1) and a recent one measured by Yalkowsky, Orr, and Valvani (2). The former study deals with the argentation of substituted benzenes and the latter work presents some useful correlations for the solubilities of halogenated benzenes in water.

The analytical procedure used by Andrews and Keefer is, in principle, quite adequate for the purpose; however, the time for saturation equilibration seems to have been too short and no tests for reaching saturation equilibria were presented (3). Furthermore, the melting point of 360-361 K reported for 1,4-diiodobenzene deviates considerably from reported literature values of 402-403 K (4a,4b). Whether this difference resulted from impurities or from some other cause is difficult to establish.

The saturation equilibrium periods for the measurements of Yalkowsky et al. also seem rather short. The time for sample equilibration varied from 4 to 48 hours. The exact time was not reported in single cases. A conventional experimental procedure was employed in their measurements. Commercial reagents were used without further purification. All measurements were done at room temperature ($25 \pm 1^{\circ}$ C). The saturated sample analyses Were carried out spectrophotometrically either after dilution or after concentration by extraction with methylene chloride. At least two independent determinations were made. However, the separate measurements were not reported and the precision of the measurements is difficult to estimate. According to the authors in a private communication (5), the error in their solubility measurement may be as great as 10 percent. The estimate seems reasonable for the following reasons. First, the variation of temperature was considerable. Second, the sample equilibration time was sufficiently long to assure an accuracy of \pm 5 percent but hardly better. The filtering and extraction procedures may well have produced further errors, so the total 10 percent error seems reasonable. The Possible systematic errors should have no effect on the correlations obtained. Therefore, solubilities of comparable accuracy for many purposes may be calculated from the reported solubility correlations.

The value reported by Yalkowsky et al. appears to be more reliable than that of Andrews and Keefer. Therefore, the value of Yalkowsky et al. is reported here as a tentative solubility of 1,4-diiodobenzene in water:

T/K 10 ⁶ mo1(1	10 ⁶ mo1(1)/dm ³	1)/dm ³ 10 ³ g(1)/kg	
298.15	5.6	1.85	1.01

REFERENCES

- 1. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. 1950, 72(7), 3113-6.
- Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u>, 18(4), 351-3.
- 3. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. 1949, 71(11), 3644-7.
- 4. a. Paulsen, O. Monatsh. Chem. 1939, 72, 254.
 - b. "Handbook of Tables for Organic Compound Identification", 3rd ed.; Rappoport, Z., Compiler; CRC Press: Cleveland, Ohio, <u>1967</u>; p. 76.
- 5. Yalkowsky, S. H., Personal Communication, 1979.

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,4-Diiodobenzene; C ₆ H ₄ I ₂ ; [624-38-4]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ O; [7732-18-5]	
-	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ³ a 10 ⁶ mol(1)/	
25 1.85 5.6	1.01
a. Calculated by F. W. Getzen. b. Reported.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
A small excess of solute in water was agi- tated for a period of 4 - 48 hours (the exact	C ₆ H ₄ I ₂ : Commerical reagent (Aldrich or Eastman), used as received.
time for equilibration was not reported in single cases) and then filtered. The satu- rated solution was diluted and assayed spec- trophotometrically. At least two independent determinations were carried out.	H ₂ O: Source and purity not reported.
	ESTIMATED ERROR:
	Solubility: ±10% (authors).
	Temperature: ±1 K (authors).
	REFERENCES:

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 1,4-Diiodobenzene; C ₆ H ₄ I ₂ ; [624-38-4]	Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1950</u> , 72(7), 3113-6.
(2) Water; H ₂ O; [7732-18-5]	
-	
VARIABLES: One temperature	PREPARED BY: A. Vesala
	A. VEBALA
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁶ mol(1)/d	$m^{3 b} 10^{8} x(1)^{b}$
25.0 1.4 4.24	7.67
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The saturation process was done in sealed	C ₆ H ₄ I ₂ : Commercial product (Eastman Kodak
Erlenmeyer flasks. The time used for the equilibration was 20 hours. Measured vol-	^{b 4 2} Co.), recrystallized twice from ethanol, reporting melting point
umes of the saturated solution were then	360-361 K.
extracted using measured volumes of hexane to remove the aromatic compound from the	H ₂ O: Source and purity not specified.
aqueous layer. The solute concentration was measured spectrophotometrically (1).	2
····	
	ESTIMATED ERROR:
	Solubility: >10% (compiler).
	Temperature: ±0.2 K.
	REFERENCES:
	 Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1949</u>, 71(11), 3644-7.

COMPONENTS:	EVALUATOR:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ O; [7732-18-5]	January 1983.

The solubility of bromobenzene in water has been reported in eight published sets of data (1-6,14,15). The solubility behavior as a function of temperature is illustrated in Figure 1. Some rather serious discrepancies between the various solubility measurements are evident in the figure.

The solubility of water in bromobenzene has been measured by several investigators between 1932 and 1963 and reported in five published sets of data (7-10,14). As shown in Figure 2, which gives the solubility behavior relative to Absolute temperature, the agreement among experimental values is reasonable.

Of the more recent works, that of Nelson and Smit (4) reported the solubility data for temperatures in the range between 278 and 318 K. However, despite the equilibration period of 24 hours (which might not have been long enough), the solubility values obtained are substantially lower than those found by the earlier investigators. It is not possible to establish any shortcomings of their experimental procedure from the very brief description. No information was provided on the source and purity of materials used. Also, it was not indicated whether or not a water stripper had been employed for the analysis of the very dilute aqueous solutions by gas chromatography, or whether or not an internal standard had been used for the calibration of the gas chromatograph which employed a flame ionization detector. However, the authors agreed to re-examine their raw data in order to verify the reported values (13). Consequently, for the present evaluation, their results have not been considered for inclusion in the selected solubility values.

The reported solubility value of Yalkowsky et al. is too low. This could be due to an insufficient time allowed for saturation equilibrium. According to the authors, the accuracy of the experimental determination was \pm 10 percent (11). The remaining reported data, that of Andrews and Keefer (1), Gross and Saylor (2), Hine et al. (3), and Vesala (5,15), have been correlated against Absolute temperature using a normal polynomial equation of second degree. In the regression, twice as much weight was assigned to the data of Vesala as was assigned to the remaining data. The equation below represents the solubility of bromobenzene in water between 283 and 308 K:

$$S_1(g(1)/kg) = 1.8293 - 1.35675 \times 10^{-2} T$$

+ 2.99322 x $10^{-5} T^2$ [1]

The values calculated from equation [1] for the saturation of bromobenzene in water in the range of temperatures between 283 and 313 K together with corresponding molarities and mole fractions are given in Table 1. Also, the solubility values calculated from equation [1] are shown in Figure 1 as a solid line along with the measured values.

The solubility of water in bromobenzene has been reported in the temperature range between 288 and 308 K in five investigations. The original report from Columbia University by Hutchinson and Lyon (8) has been lost, and all details reported here on that work have been taken from a secondary source (12). The single value from Hutchinson and Lyon is too high in relation to other reported determinations at the same temperature so it was not included in the correlation of the data against temperature. On the other hand, the solubility reported by Donahue and Bartell (14) appears too low and this value also was not used in the correlation. The reported values of Bell (7), Jones and Monk (9), and Wing and Johnston (10) are represented by the equation:

$$\log_{10} x(2) = 2.43149 - 1514.54/T$$
[2]

In this equation, x(2) is the mole fraction solubility of water in the water-bromobenzene system and T is the Absolute temperature. The calculated solubility values in the 283 to 308 K range are shown in Figure 2 as a solid line along with the reported values.

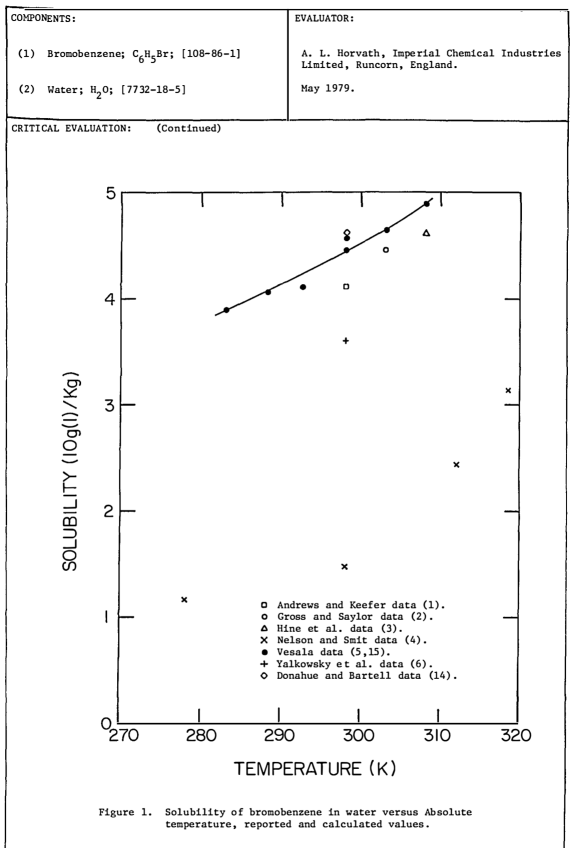
The calculated mole fraction values for the solubility of water in bromobenzene from equation [2] are included in Table 2 together with the corresponding molarities and g(2)/kg values in the temperature range between 283 and 313 K.

[
COMPONENTS:			EVALUATOR:	
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]				A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ O; [7732-18-5]				January 1983.
	-			
CRIT	ICAL EVALUATION	: (Continued)		
Table 1. Solubility of Bromobenzene in Water.				
	т/к	10 ³ mol(1)/dm ³	10g(1)/kg $10^5 x(1)$
	283.15	2.47	3.8	7 4.45
	288.15	2.58	4.0	5 4.65
	293.15	2.70	4.2	4 4.87
	298.15	2.83	4.4	5 5.11
	303.15	2.96	4.6	7 5.36
ł	308.15	3.11	4.9	1 5.63
	313.15	3.26	5.1	6 5.92
Tabl	e 2. Solubili	ty of Water in Brom	ohenzene	
140.		-		
	т/к	10 ² mo1(2)/dm ³	10g(2)/kg $10^3 x(2)$
	283.15	1.16	1.3	9 1.21
	288.15	1.43	1.7	2 1.50
	293.15	1.76	2.1	2 1.84
	298.15	2.13	2.5	8 2.25
	303.15	2.58	3.14	4 2.73
	308.15	3.09	3.78	8 3.29
	313.15	3.69	4.5	3 3.94
		ום	EFERENCE	s
1.	1. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1950</u> , 72(7), 3113-6.			
2.	Gross, P. M.; Saylor, J. H. J. Am. Chem. Soc. <u>1931</u> , 53(5), 1744-51.			
3.	Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. <u>1963</u> ,85(10), 1473-6.			
4.	Nelson, H. D.; Smit, J. H. SAfr. Tydskr. Chem. <u>1978</u> , 31(2), 76.			
5.	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.			8), 839-45.
6.	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C. Ind. Eng. Chem. Fundam. 1979, 18(4), 351-3.			
7.	Bell, R. P. J. Chem. Soc. <u>1932</u> , Part II, 2905-11.			
8.	Hutchinson, C.	. A.; Lyon, A. M. (Columbia	University Report A-745, July 1, <u>1943</u> .
9.	Jones, J. R.;	Monk, C. B. J. Che	em. Soc.	<u>1963</u> , Part III, 2633-5.
10.	Wing, J.; Johr	nston, W. H. J. Am.	. Chem. S	Soc. <u>1957</u> , 79(4), 864-5.
11.	Yalkowsky, S.	H., Personal Commun	nication,	, <u>1979</u> .
12.	Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M.; Urey, H. C.; Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II, Chapter 7, pp 129-44.			

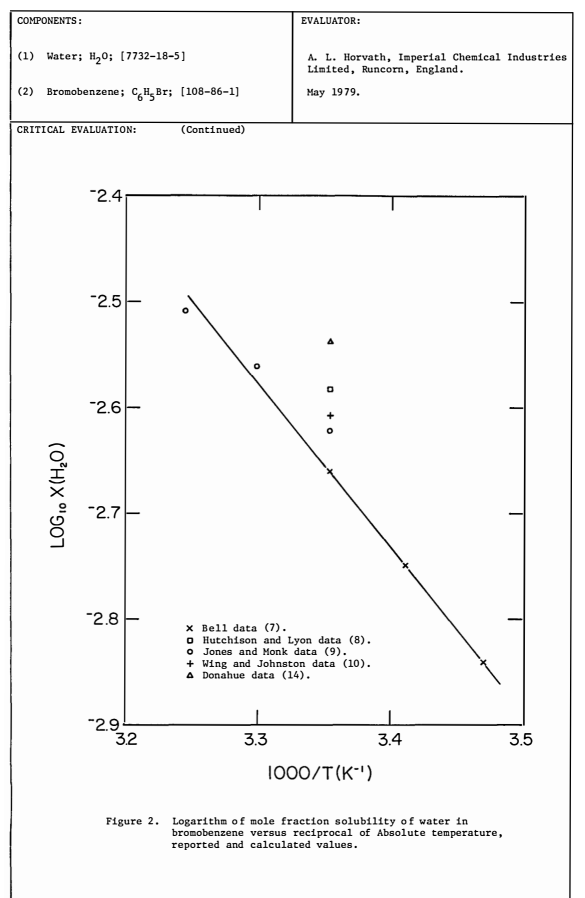
COMPONENTS:		EVALUATOR:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]		A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ 0; [7732-18-5]		January 1983.
-		,
CRITICAL EVALUATION: (Continued)		
13.	Nelson, H. D., Personal Communication, J	1979.

14. Donahue, D. J.; Bartell, F. E. J. Phys. Chem. 1952, 56(4), 480-4.

15. Vesala, A., Ph.D. Dissertation, University of Turku, Turku, 1973.



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COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Gross, P. M.; Saylor, J. H. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1931</u> , <i>53(5)</i> , 1744-51.
(2) Water; H ₂ O; [7732-18-5]	
VAR ANY DO	
VARI ABLES :	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10g(1)/kg(2)^{a} = 10^{3}mol(1)/kg(2)^{a}$	b 10 ⁵ $x(1)$ ^c
30 4.46 2.839	5.117
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of bromobenzene in 500 g water was shaken for 12 hrs. in a thermostat bath.	C ₆ H ₅ Br: Eastman Kodak Co., was purified by fractional distillation before
Samples were then withdrawn and read against water in an interferometer made by Zeiss (1).	use.
A detailed description of the complete pro-	H ₂ 0: Distilled.
cedure is given in a Ph.D. dissertation (2).	
	ESTIMATED ERROR:
	Solubility: ± 2%.
	Temperature: ± 0.02 K.
	REFERENCES:
	 Gross, P. M. J. Am. Chem. Soc. <u>1929</u>, 51(8), 2362-6.
	 Saylor, J. H., Ph.D. Dissertation, Duke University, Durham, <u>1930</u>.

OMPONENTS				
COMPONENTS :		1 contract of the second se	ORIGINAL MEASUREMENTS:	
(1) Water; H ₂ O; [7732-18-5]			Bell, R. P. J. Chem. Soc. <u>1932</u> , Part II, 2905-11.	
(2) Bromob	enzene; C ₆ H ₅ Br; [108-	-86-1]		
VARIABLES :		<u> </u>	PREPARED BY:	
Temperature			A. L. Horvath	
EXPERIMENTAL	VALUES:			
	10g(1)/dm ³ (2) ^a	2	3 b 10 ³ (1) C	
t°C				
15	2.49	1.382	1.443	
20	3.06	1.698	1.781	
25	3.74	2.075	2.186	
			ΙΝΈΩΡΜΑΤΙΩΝ	
			INFORMATION	
	ATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS;	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abse NaOH solution	ATUS/PROCEDURE: 5 ratio solute/solved in a thermostat bad equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. To orbed in water and to on (1). Successive ere carried out with	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS; H ₂ O: Distilled (compiler).	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abse NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS; H ₂ O: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade, redistilled before use.	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxyy HC1 was abse NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS; H ₂ O: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade,	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abse NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS: H ₂ O: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR:	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abse NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS: H ₂ 0: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±2.6%. Temperature: ±0.02 K.	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abse NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS; H ₂ O: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±2.6%. Temperature: ±0.02 K. REFERENCES:	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abso NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	SOURCE AND PURITY OF MATERIALS: H ₂ 0: Distilled (compiler). C ₆ H ₅ Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±2.6%. Temperature: ±0.02 K.	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abso NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	<pre>SOURCE AND PURITY OF MATERIALS: H₂0: Distilled (compiler). C₆H₅Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±2.6%. Temperature: ±0.02 K. REFERENCES: 1. Bell, R. P. J. Chem. Soc. <u>1932</u>, Part</pre>	
About 1 to were rotated hrs. After samples were cottonwool. content was α -naphthoxy, HCl was abso NaOH solution	5 ratio solute/solver d in a thermostat bar equilibrium was esta e taken and filtered The determination of based on the reaction dichlorophosphine. orbed in water and to on (1). Successive	nt mixtures th for 12 ablished, through of the water on with The evolved itrated with (2-4) deter-	<pre>SOURCE AND PURITY OF MATERIALS; H₂0: Distilled (compiler). C₆H₅Br: Merck reagent, analytical grade, redistilled before use. ESTIMATED ERROR: Solubility: ±2.6%. Temperature: ±0.02 K. REFERENCES: 1. Bell, R. P. J. Chem. Soc. <u>1932</u>, Part</pre>	

COMPONENTS :	ORIGINAL MEASUREMENTS:
, 2, -	Hutchison, C. A.; Lyon, A. M. Columbia University Report A-745, July 1, 1943.
(2) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	
VARIABLES:	
One temperature	PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ² mol(1)/kg ^b	$10^{3}x(1)^{c}$
25 3.009 1.67	2.616
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
ME THOD /APPARATUS/PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
A mixture of 1 to 15 volume ratio of solute	H ₂ 0: Distilled.
to solvent was placed in an equilibration flask and then lowered into a thermostat water bath. The assembly was shaken mechani- cally for about 90 min. at constant tempera- ture. The amount of water in the organic phase was determined by a modified Karl Fisher titration. The determination was done in triplicate. The description of the procedure was taken from a secondary source (1). The original report is no longer avail- able.	C ₆ H ₅ Br: Source is not known, purified and dried before use.
	ESTIMATED ERROR:
	Solubility: ± 1%.
	Temperature: ± 0.05 K.
	REFERENCES: Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M.; Urey, H. C.; Kirshenbaum, I., Eds.; McGraw-Hill: New York, 1955; Part II, Chapter 7, pp 129-44.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Andrews, L. J.; Keefer, R. M. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1950</u> , 72(7), 3113-6.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm ³	^b $10^{5}x(1)^{c}$
25.0 4.1 2.61	4.72
December 1	
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
AUXILI ARY	INFORMATION
METHOD /APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
In a constant temperature bath, water was	C ₆ H ₅ Br: Eastman Kodak Co., commercial reagent, b.p. 156.3°C, fraction-
mixed with bromobenzene through rotation in a glass-stoppered Erlenmeyer flask for 30	ated before use.
hrs. The saturated solution was extracted with n-hexane. The optical density of the	H ₂ 0: Not specified.
extract was measured against a n-hexane blank using a Beckman spectrophotometer (1).	2
blank using a beckman spectrophotometer (1).	
	ESTIMATED ERROR:
	Solubility: ± 10% (compiler).
	Temperature: ± 0.1 K (compiler).
	REFERENCES:
	 Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1949</u>, 71, 3644-7.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Donahue, D. J.; Bartell, F. E. <i>J. Phys.</i> <i>Chem.</i> <u>1952</u> , 56(4), 480-4.
(2) Water; H ₂ O; [7732-18-5]	
-	
VARIABLES:	PREPARED BY:
	A. L. Horvath
One temperature	
EXPERIMENTAL VALUES:	
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10g(1)/kg ^a 10 ³ mol(1)/kg ¹	$10^{5}x(1)^{c}$
25 4.62 2.94	5.3
25 4.02 2.54	
t/°C 10g(2)/kg ^a 10 ² mol(2)/kg ^b	$10^3 x(2)$ c
25 3.34 1.85	2.9
a. Calculated by compiler.	
b. Calculated by F. W. Getzen.c. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A mixture of water and bromobenzene was	C ₆ H ₅ Br: Reagent grade, purified by
placed in a glass stoppered flask and shaken intermittently for at least three days in a	fractional distillation.
constant temperature water bath. The water	H ₂ O: Purified by distillation.
content of the organic phase was established by the Karl Fisher method and the organic	
content of the aqueous phase was established	
interferometrically.	
	ESTIMATED ERROR:
	Solubility: ± 5% (compiler).
	Temperature: ± 0.1 K.
	REFERENCES :

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Wing, J.; Johnston, W. H. J. Am. Chem.
1 ²	Soc. 1957, 79(4), 864-5.
(2) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10m1(1)/dm^{3} a = 10^{2}mo1(1)/dr$	$3 b 10^{3} (1) c$
$t/^{\circ}C = 10m1(1)/dm^{3/a} = 10^{2}mo1(1)/dr$	10 x(1)
25.0 4.24 2.347	2.470
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
1	
1	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Tritiated water was equilibrated with 20 ml	H ₂ 0: Tracerlab, Inc., tritiated water,
bromobenzene by stirring in a flask in a	used as received.
constant temperature water bath for two hrs. The concentration of the tritiated water in	C.H.Br: Source not specified, chemical
the organic phase was determined by isoto-	C ₆ H ₅ Br: Source not specified, chemical grade, redistilled before use.
pic dilution. The tritium activities for	
the tritiated water samples were determined by the acetylene method (1,2). At least	
four independent experiments were done.	
The article describes work reported in a	
Ph.D. dissertation (2).	
	ESTIMATED ERROR:
	Solubility: ± 2.3%.
	Temperature: ± 0.02 K.
	DEFEDENCES -
	REFERENCES:
	 Wing, J.; Johnston, W. H. Science <u>1955</u>, 121, 674-5.
	2. Wing, J., Ph.D. Dissertation, Purdue
	University, Lafayette, IN, <u>1956</u> .

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Hine, J.; Haworth, H. W.; Ramsay, O. B. J. Am. Chem. Soc. <u>1963</u> , 85(10), 1473-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm	3 b 10 ⁵ (1) C
35.0 4.585 2.92	5.294
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of bromobenzene was combined with 4 ml of water in an ampoule and sealed. The	C ₆ H ₅ Br: Eastman Kodak Co., redistilled before use.
sealed ampoule was then rotated in a water bath for a week. The bromobenzene solubi- lity was determined by ultraviolet spectro-	H ₂ O: Not specified.
photometric measurements using a Beckman DU spectrophotometer.	
	ESTIMATED ERROR:
	Solubility: ±2% S.D. Temperature: ±0.1 K.
	REFERENCES:
L	

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Jones, J. R.; Monk, C. B. <i>J. Chem. Soc.</i> 1963, Part III, 2633-5.
(2) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	<u>1905</u> , Fart 111, 2055-5.
VARIABLES:	PREPARED BY:
Temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{4} dm^{3}(1)/dm^{3}(2)$ ^a $10^{2} mol($	
25 4.1 2	.27 2.39
30 4.7 2	.60 2.75
35 5.3 2	.92 3.10
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Tritiated water was shaken with bromobenzene	H ₂ 0: Tritiated.
in 1:10 volume ratios for 4 hours in a flask in a water thermostat bath. The water con- tent was determined by tritium assay. The count rates were determined using a typical liquid scintillator solution technique.	C ₆ H ₅ Br: Source not known, laboratory grade, dried over CaCl ₂ and fractionally distilled before use.
	ESTIMATED ERROR:
	Solubility: ±5%.
	Temperature: ±0.5 K (compiler).
	REFERENCES :

COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]			Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u> .
(2) Water	; H ₂ 0; [7732-18-5]		
VARIABLES:			PREPARED BY:
Temperatur	e		A. L. Horvath
EXPERIMENTA	AL VALUES:		
t/°C	10g(1)/kg ^a	10 ³ mol(1)/kg(2)	^b $10^{5}x(1)^{a}$
10.0	3.8957	2.482 ± 0.008	4.4714
15.2	4.0605	2.587 ± 0.020	4.6605
19.6	4.1107	2.619 ± 0.020	4.7182
25.0	4.5719	2.913 ± 0.033	5.2478
30.0	4.6456	2.960 ± 0.040	5.3325
35.0	4.8919	3.117 ± 0.040	5.6153
<u> </u>		AUXILIARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :			SOURCE AND PURITY OF MATERIALS:
The bromobenzene was mixed with water in a sealed flask (1) with the aid of a magnetic stirrer for 48 hours in a water bath at constant temperature. After the solution was filtered through a glass-wool plug, the solute was extracted with 2,2,4-trimethyl-pentene. The optical density of each sample was determined spectrophotometrically (2). Mean and standard deviations were calculated from three measurements.		d of a magnetic ter bath at the solution wool plug, the 2,4-trimethyl- of each sam- ptometrically (2).	C ₆ H ₅ Br: Merck AG., >99% GLC, used as received. H ₂ O: Distilled, deionized, and degassed.
			ESTIMATED ERROR: Solubility: ±1.35%.
			Temperature: ±0.05 K.
			 REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J. Chem. Soc. <u>1963</u>, Part III, 2716-23. 2. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u>, 17(1), 38-41.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10 ³ mol(1)/kg(2) ^a 10g(1)/kg	$b 10^5 x(1)^{c}$
25 2.84 4.457	5.116
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established between the water and the bromobenzene in a sealed flask (1) with the aid of a magnetic stir- rer during 48 hours under isothermal con- ditions. After the sample was filtered through a glass-wool plug, the bromobenzene was extracted with 2,2,4-trimethylpentene.	<pre>C₆H₅Br: Commercial reagent of analytical grade distilled through a column resulting in a more than 99% pure sample.</pre> H ₂ 0: Distilled, deionized, and degassed.
Sample optical densities were determined spectrophotometrically (2). Five parallel determinations were done.	
The reported work was based upon a Ph.D. dissertation (3).	ESTIMATED ERROR:
	Solubility: ±1.1%. Temperature: ±0.05 K.
	REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J. <i>Chem. Soc.</i> <u>1963</u> , Part III, 2716-23.
	 Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u>, 17(1), 38-41.
	 Vesala, A., Ph.D. Dissertation, Univer- sity of Turku, Turku, <u>1973</u>.

ORIGINAL MEASUREMENTS:
Nelson, H. D.; Smit, J. H. SAfr. Tydskr.
Chem. 1978 , $31(2)$, 76.
PREPARED BY: A. L. Horvath
A. L. HOIVILI
_
$b^{b} 10^{5}x(1)^{c}$
1. 34
1.7
2.8
3.6
-
ARY INFORMATION
SOURCE AND PURITY OF MATERIALS:
ase C ₆ H ₅ Br: Not specified.
H ₂ 0: Not specified.
th gth
,
ESTIMATED ERROR:
Solubility: ±14.9% (compiler).
Temperature: ±0.1 K (compiler).
REFERENCES :
1. Nelson, H. D.; de Ligny, C. L. <i>Rec.</i> <i>Trav. Chim.</i> 1968, <i>87</i> , 528-44.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	Yalkowsky, S. H.; Orr, R. J.; Valvani, S. C.
	Ind. Eng. Chem. Fundam. <u>1979</u> , 18(4), 351-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10g(1)/dm^{3} = 10^{3}mo1(1)/dm^{3} b$	$10^5 x(1)^{a}$
t/°C 10g(1)/dm ⁻ 10 ⁻ mol(1)/dm ⁻	$10^{-}x(1)$
25 3.596 2.29	4.139
a. Calculated by compiler. b. Reported.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Samples of excess bromobenzene in water	C _c H ₅ Br: Aldrich commercial grade, used as
were agitated at room temperature for about	C ₆ H ₅ Br: Aldrich commercial grade, used as received.
24 hours and then filtered. The filtrate	
was diluted and assayed spectrophotometri-	H ₂ O: Deionized.
cally. Determinations were carried out in duplicate.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±1 K.
	•
	REFERENCES:
4	
1	

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water-d ₂ ; D ₂ 0; [7789-20-0]	Hutchison, C. A.; Lyon, A. M. Columbia
(2) Bromobenzene; C ₆ H ₅ Br; [108-86-1]	University Report A-745, July 1, <u>1943</u> .
B	
VARIABLES:	PREPARED BY:
One temperature	G. Jancso
EXPERIMENTAL VALUES:	
	2
$t/^{\circ}C = 10g(1)/kg^{a} = 10^{2}mol(1)/kg^{b}$	$10^{3}x(1)^{a}$
25.0 2.8261 1.411	2.2113
a. Calculated by F. W. Getzen.	
b. Reported (average of two experimental	l values).
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Between 25 and 100 ml bromobenzene and 1 to	C ₆ H ₅ Br: Carefully purified and dried before
2 ml D ₂ O were placed in a flask and shaken for about 90 min. The thermostat water	use. Source and method not given.
bath temperature was maintained within	100% D ₂ 0: Source not specified.
$\pm 0.05^{\circ}$ C. Then, a sample was removed and the amount of D ₀ O dissolved was determined	2
by a modified Karl Fischer titration (1).	
The original report was unavailable; how- ever, the method and the results were de-	
scribed in sufficient detail in (1). The	
solubility of H ₂ O in bromobenzene was also	
determined and found to be 0.0167 mol(1)/kg. The average deviation for two experiments	ESTIMATED ERROR:
was ±0.0001 mol(1)/kg.	Solubility: ±1 x 10 ⁻⁶ mol (std. dev.)
	$D_20/100$ g solution.
	Temperature: ±0.05 K.
	REFERENCES :
	 Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series
	Division III-Vol. 4F, Murphy, G. M.;
	Urey, H. C.; Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II,
	Chapter 7, pp 129-44.

COMPONENTS :		EVALUATOR:
(1)	4-Bromophenol; C ₆ H ₅ Br0; [106-41-2]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

The only two available reported measurements of the solubility of 4-bromophenol in water were published many years apart and the information concerning the methods of analysis provided in the earlier of these works is incomplete. The older one by Werner (1) appeared in 1884 and the other more recent work was published in 1971. Therefore, certain difficulties have been encountered in the evaluation of these combined data. The solubility measurements reported by Parsons, Rochester, and Wood (2) for 4-bromophenol in water appear quite reliable when the saturation equilibrium, the method of analysis, and the reported standard deviation have been considered. On the basis of these criteria, the value reported by Werner should be rejected. However, if one can accept a satisfactory state of purity for the synthesized material, the value of Werner, which refers to a temperature of 288 K, can be regarded as fairly good. Considering the difference in the temperatures of the reported solubility values, the agreement between the two values is quite satisfactory in spite of the long time interval between their determinations. However, the solubility value reported by Parsons et al. must be considered as only a tentative value.

It should be noted that the solubility of 4-bromophenol in water is considerably dependent upon the acidity of the solution. Here, it is assumed that the solubility established refers to that of a solution having a pH value prevailing for saturated 4bromophenol in water. An entire series of measurements of solubilities at various pH values is required to complete the picture of the solubility behavior for protolyzing compounds such as 4-bromophenol in water.

The solubility of 4-bromophenol in water is reported here as a tentative value:

т/к	10mo1(1)/dm ³	10 ⁻¹ g(1)/kg	$10^{3}x(1)$
298.15	1.07	1.86	1.97

REFERENCES

1. Werner, E. Ann. Chim. Phys. Series 6(III) 1884, 567-74.

2. Parsons, G. H.; Rochester, C. H.; Wood, C.E.C. J. Chem. Soc. B 1971, 533-6.

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 4-Bromopheno1; C₆H₅Br0; [106-41-2] (2) Water; H₂0; [7732-18-5]</pre>	Parsons, G. H.; Rochester, C. H.; Wood, C. E. C. <i>J. Chem. Soc. B</i> <u>1971</u> , 533-6.
VARIABLES: One temperature	PREPARED BY: A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁻¹ g(1)/kg ^a 10mo1(1)/kg(2) ^b $10^{3}x(1)^{a}$
25 1.8261 1.075	1.9330
a. Calculated by F. W. Getzen. b. Reported.	
AUXI LIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS:	
The experimental procedure was probably similar to that reported in (1). Excess solid was shaken with water at $25 \pm 0.1^{\circ}$ C for 24 hours. The saturated solution was then filtered at thermostat temperature and, with appropriate dilution, analyzed spectrophotometrically. Linear calibration graphs of absorbance at a wave length of maximum absorbance of the neutral phenols against concentration confirmed the appli- cability of the Beer-Lambert law for the analyses.	C ₆ H ₅ Br0: Commercial product (manufacturer not reported), recrystallized to a constant melting point. H ₂ 0: Source and purity not specified. ESTIMATED ERROR: Solubility: ±2% (standard deviation of the
	average value from five deter- minations as reported by the authors). REFERENCES:
	 Parsons, G. H.; Rochester, C. H.; Rostron, A.; Sykes, P. C. J. Chem. Soc. Perkin Trans. II(1) <u>1972</u>, 136-8.

COMPONENTS:	EVALUATOR:
(1) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ 0; [7732-18-5]	January 1983.
CRITICAL EVALUATION:	

Thirteen experimental determinations of the solubility of chlorobenzene in water between 278 and 363 K have been reported in the literature (1-10,23-25), see Figure 1. The solubility at higher temperatures, between 513 and 533 K, have been reported also by Vorozhtsov and Kobelev (11) in graphical form only without further details concerning the source of the original measurements.

The solubility of water in chlorobenzene has been reported in eleven published works (5,9,12-20) in the temperature range between 288 and 322 K, see Figure 2.

Some of the more recent data for the solubility of chlorobenzene in water is that of Nelson and Smit (8) in the temperature range between 278 and 318 K. However, despite equilibrium periods of 24 hours (which might not have been long enough), the measured solubilities are substantially lower than those found by earlier investigators. It is not possible to establish any shortcomings of the experimental procedures from the very brief description. No information was provided on the source and purity of materials used. Also, it was not indicated whether or not a water stripper had been employed for the analysis of the very dilute aqueous solutions by gas chromatography, or whether or not an internal standard had been used for the calibration of the gas chromatograph which employed a flame ionization detector. However, the authors agreed to re-examine their raw data in order to verify the reported values (21). Consequently, for the present evaluation, their results have not been considered for inclusion in the selected solubility values.

The reported solubilities of Othmer et al. (9) and those of Newman et al (23) are too high in relation to the other reported solubility data. Consequently, these values have not been included in the correlation procedure. The approximate value given by Booth and Everson (3) has been excluded also from further consideration. The solubility data at high temperatures by Vorozhtsov and Kobelev (11) could not be correlated with the other data given. These remaining data given in (1,2,4-7,10,24,25) were correlated with Absolute temperature using polynomial equations of various degrees. The equation given below represents the solubility of chlorobenzene in water between 283 and 363 K within an estimated 10 percent. Equal weight was given to each of the data points in the regression procedure for the following equation:

$$S_1(g(1)/kg) = 11.3351 - 3.0290 \times 10^{-2} T$$

- 1.8716 x $10^{-4} T^2 + 0.559466 \times 10^{-6} T^3$ [1]

The significance of this equation is that the curve representing the solubility data passes through a minimum at 286.1 K. This behavior is consistent with the theory discussed by Gill et al. (27) for the solubility of aromatic compounds in water.

Recommended solubility values between 283 and 363 K have been calculated from equation [1] and presented in Table 1. The solubility behavior described by equation [1] is shown also in Figure 1 which contains the reported experimental values for the solubility of chlorobenzene in water.

The solubility of water in chlorobenzene has also been investigated and reported by a number of workers (9,13,17,18,19,20,26) in the 298 to 318 K temperature range. In general, these solubility values show some irregularity, but those showing reasonable agreement can be correlated with respect to Absolute temperature. The reported solubilities of Zielinski (20) and of Othmer et al. (9) are too high while those of Jones and Monk (17) are too low. Consequently, these data are not considered further. Also, the solubility expressed as a distribution coefficient by an equation only (26) cannot be included in the correlation. The good agreement of the values at 298.15 K reported in (13,18,19) are heavily weighed in the correlation. The remaining data were all assigned equal weight to produce the following equation:

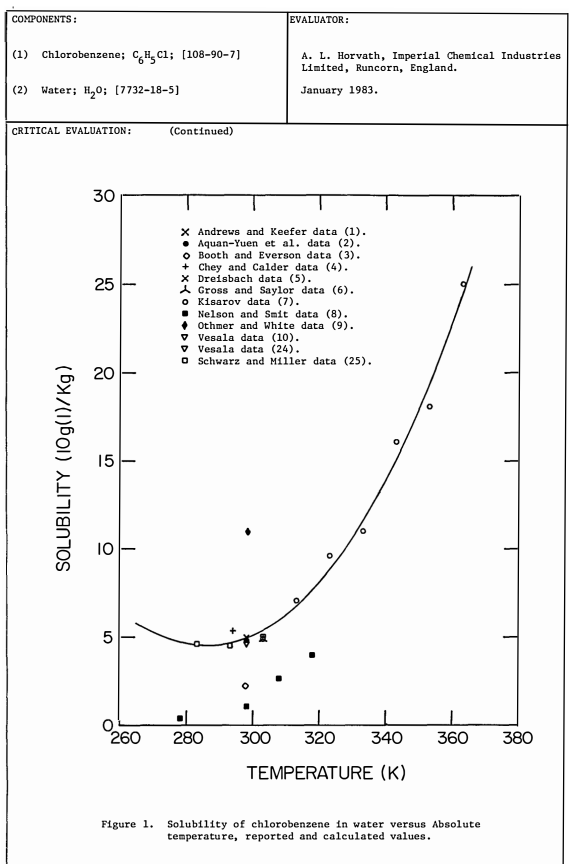
$$\log_{10} x(2) = 2.99105 - 1668.56/T$$

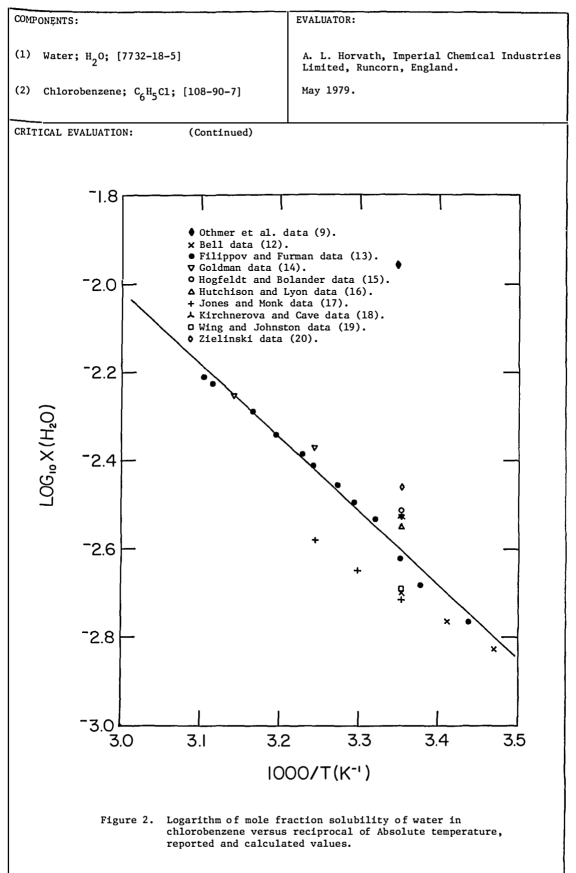
[2]

In this equation, x(2) is the mole fraction solubility of water in the chlorobenzenewater system and T is the Absolute temperature. The calculated solubility values in the 273 to 333 K range are shown in Figure 2 along with the reported values.

· · ·			
COMPONENTS :		EVALUAT	FOR:
(1) Chlorobenzene;	с ₆ н ₅ с1; [108-90-7] A. L. Limite	Horvath, Imperial Chemical Industries d, Runcorn, England.
(2) Water; H ₂ 0; [7	732-18-5]	Januar	y 1983.
	-		-
CRITICAL EVALUATION:	(Continued)		
The calculated mole fraction values for the solubility of water in chlorobenzene from equation [2] are given in Table 2 together with the corresponding molarities and g(2)/kg values in the temperature range between 283 and 333 K.			
The azeotrope temperature (363.35 K) and composition (284 g(2)/kg) at 1.0133 x 10^5 Pa pressure were also reported (22).			
Table 1. Solubilit	y of Chlorobenzene	in Water.	
T/K	10 ³ mo1(1)/dm ³	10g(1)/kg	$10^{5}x(1)$
283.15	4.03	4.54	7.27
288.15	4.02	4.52	7.24
293.15	4.13	4.66	7.46
298.15	4.38	4.95	7.92
303.15	4.77	5.39	8.63
308.15	5.30	6.00	9.60
313.15	5.97	6.77	10.84
318.15	6.78	7.71	12.34
323.15	7.74	8.82	14.13
328.15	8.85	10.11	16.20
333.15	10.12	11.58	18.56
338.15	11.53	13.24	21.22
343.15	13.11	15.09	24.18
348.15	14.84	17.13	27.46
353.15	16.73	19.37	31.06
363.15	20.98	24.47	39.24
	y of Water in Chlo		3
Т/К	$10^2 mol(2) / dm^3$	10g(2)/kg	$10^{3}x(2)$
283.15	1.24	2.01	1.25
288.15	1.57	2.54	1.59
293.15	1.96	3.19	1.99
298.15	2.43	3.98	2.48
303.15	2.99	4.92	3.07
308.15	3.66	6.05	3.77
313.15	4.44	7.39	4.60
318.15	5.36	8.97	5.58
323.15	6.43	10.82	6.72
328.15	7.67	12.99	8.06
333.15	9.10	15.50	9.61

Chlorobenzene 15			
COMP	ONENTS:	EVALUATOR:	
(1)	Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2)	Water; H ₂ O; [7732-18-5]	January 1983.	
CRIT	ICAL EVALUATION: (Continued)		
	REFERENC	ES	
1.	Andrews, L. J.; Keefer, R. M. J. Am.	Chem. Soc. <u>1950</u> , 72(7), 3113-6.	
2.	• Aquan-Yuen, M.; Mackay, D.; Shiu, W. Y. J. Chem. Eng. Data <u>1979</u> , 24(1), 30-4.		
3.	Booth, H. S.; Everson, H. E. Ind. Eng.	Chem. <u>1948</u> , 40(8), 1491-3.	
4.	Chey, W.; Calder, G. V. J. Chem. Eng.	Data <u>1972</u> , 17(2), 199-200.	
5.	Dreisbach, R. R. "Physical Properties o Series No. 15; American Chemical Society	f Chemical Compounds", Advances in Chemistry y: Washington, D. C., <u>1955;</u> p 134.	
6.	Gross, P. M.; Saylor, J. H. J. Am. Cher	n. Soc. <u>1931</u> , 53(5), 1744-51.	
7.	Kisarov, V. M. Zh. Prikl. Khim. 1962, 3	35(10), 2347-9.	
8.	Nelson, H. D.; Smith, J. H. SAfr. Typ	dskr. Chem. <u>1978</u> , 31(2), 76.	
9.	Othmer, D. F.; White, R. C.; Truegar, E. Ind. Eng. Chem. 1941, 33(12), 1513.		
10.	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A	(8), 839-45.	
11.	Vorozhtsov, N. N.; Kobelev, V. A. Zh. (Dbs. Khim. <u>1938</u> , 8(12), 1106–19.	
12.	2. Bell, R. P. J. Chem. Soc. <u>1932</u> , Part II, 2905-11.		
13.	3. Filippov, T. S.; Furman, A. A. Zh. Prikl. Khim. <u>1952</u> , 25, 895-7.		
14.	Goldman, S., Ph.D. Dissertation, McGill University, Montreal, 1969, 84.		
15.	5. Högfeldt, E.; Bolander, B. Ark. Kemi <u>1963</u> , 21(16), 161-86.		
16.	6. Hutchinson, C. A.; Lyon, A. M. Columbia University Report A-745, July 1, <u>1943</u> .		
17.	7. Jones, J. R.; Monk, C. B. J. Chem. Soc. <u>1963</u> , Part III, 2633-5.		
18.	3. Kirchnerova, J.; Cave, G.C.B. Can. J. Chem. <u>1976</u> , 54(24), 3909-16.		
19.	• Wing, J.; Johnston, W. H. J. Am. Chem. Soc. <u>1957</u> , 79(4), 864-5.		
20.	- Zielinski, A. Z. Chem. Stosowana <u>1959</u> , 3, 377-84.		
21.	Nelson, H. D., Personal Communication, 1	<u>979</u> .	
22.	Prahl, W.; Hathes, W. Angew. Chem. 1934	, 47, 11-13.	
23.	Newman, M.; Hayworth, C. B.; Treybal, R.	E. Ind. Eng. Chem. <u>1949</u> , 41(9), 2039-43.	
24.	Vesala, A., Ph.D. Dissertation, Universi	ty of Turku, Turku, <u>1973</u> .	
25.	Schwarz, F. P.; Miller, J. Anal. Chem.	<u>1980</u> , <i>52(13)</i> , 2162-4.	
26.	 Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. Khim. Tekhnol. Inst. 1974, 81, 55-6. 		
27.	Gill, S. J.; Nichols, N. F.; Wadso, I.	J. Chem. Thermodyn. <u>1976</u> , 8(5), 445-52.	





COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Gross, P. M.; Saylor, J. H. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1931</u> , <i>53(10)</i> , 1744-51.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg(2) ^a 10 ³ mol(1)/kg	^b $10^5 x(1)$ ^c
30 4.88 4.333	7.810
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of chlorobenzene in 500 g water was shaken for 12 hours in a thermostat bath. Samples were then withdrawn and read against water as a reference using	C ₆ H ₅ Cl: Eastman Kodak Co., was purified by fractional distillation before use.
an interferometer made by Zeiss (1). A detailed description of the complete	H ₂ O: Distilled.
procedure has been described in a Ph.D. dissertation (2).	
	ESTIMATED ERROR: Solubility: ±1.0%.
	Temperature: ±0.02 K.
	REFERENCES: 1. Gross, P. M. J. Am. Chem. Soc. <u>1929</u> , 51(8). 2362-6.
	 Saylor, J. H., Ph.D. Thesis, Duke University, Durham, 1930.

COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) Water;	н ₂ 0; [7732-18-5]		Bell, R. P. J. Chem. Soc. <u>1932</u> , Part II, 2905-11.
(2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]			2905-11.
VARIABLES:			PREPARED BY:
Temperature			A. L. Horvath
EXPERIMENTAL	VALUES:	·····	
t/°C	10g(1)/dm ³ (2) ^a	10 ² mol(1)/d	$\lim_{m}^{3} b 10^{3} x(1)^{c}$
15	2.64	1.471	1.488
20	3.05	1.691	1.719
25	3.54	1.953	1.995
	culated by F. W. Get culated by compiler.		
	<u></u>		
METHOD /ADDADAD	TUS/PROCEDURE:	AUXILIARI	INFORMATION
	solute and solvent	in about l	SOURCE AND PURITY OF MATERIALS: H ₂ 0: Distilled (compiler).
bath for 12 attained, sa tered throug tion of the the reaction	were rotated in a t hours. After equil amples were withdraw gh cotton-wool. The water content was b h of water with α -na sphine. The evolved	ibrium was n and fil- determina- ased upon phthoxy-	C ₆ H ₅ Cl: Merck reagent, analytical grade, redistilled before use.
absorbed in dard NaOH so	water and titrated volution (1). From 2 Inations were made w	with stan- -4 succes-	
			ESTIMATED ERROR:
			Solubility: ±1.6%.
			Temperature: ±0.02 K.
			REFERENCES: 1. Bell, R. P. J. Chem. Soc. <u>1932</u> , Part II, 2903-5.

COMPONENTS :		ORIGINAL MEASUREMENTS:
 Chlorobenzene; C₆H₅C1; [108-90-7] Water; H₂O; [7732-18-5] 		Vorozhtsov, N. N.; Kobelev, V. A. <i>Zh. Obs.</i> <i>Khim.</i> <u>1938</u> , <i>8(12)</i> , 1106-19.
VARIABLES:		PREPARED BY:
Temperature		A. L. Horvath
EXPERIMENTAL VALUES:		
	1)/kg ^a 10mol(1)/kg ^b	
240 5.		
250 6.	2 5.51	10.5
260 6.	8 6.04	11.5
a. Values obta b. Calculated c. Calculated	ined from graph plot (Fig by F. W. Getzen. by compiler.	. 4).
<u> </u>		
		INFORMATION
METHOD/APPARATUS/PRO	CEDURE:	SOURCE AND PURITY OF MATERIALS:
Not specified.		Not specified.
		ESTIMATED ERROR: Solubility: ±10% (compiler).
		Temperature: ±1 K (compiler).
		REFERENCES :

			001	
Сомро	NENTS:			ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]			4 ₅ C1; [108-90-7]	Othmer, D. F.; White, R. E.; Trueger, E.
(2) Water; H ₂ O; [7732-18-5]			-18-5]	Ind. Eng. Chem. <u>1941</u> , 33(12), 1513.
		2		
VARIA	BLES:			PREPARED BY:
One	tempera	ture		A. L. Horvath
			· · · · · · · · · · · · · · · · · · ·	
EXPER	RIMENTAL	VALUES:		
	t/°C	g(1)/kg	10 ³ mol(1)/kg	$10^4 x(1)$
	25.5	1.1	9.77	1.76
	t/°C	a(2)/ka	10 ² mol(2)/kg	$10^2 x(2)$
	25.5	1.8	9.99	1.11
	a. Rep	orted.		
	b. Cal	culated by 1	. W. Getzen.	
	c. Cal	culated by o	compiler.	
			·····	
			AUXILIA	RY INFORMATION
		ATUS / PROCEDU		SOURCE AND PURITY OF MATERIALS:
			using about 15 ml meyer flask (1).	C ₆ H ₅ Cl: Source not specified. Purified until it distilled within 1 or 2°C
The	solute	was added to	the water from a	range.
			agitated until bid. The amounts	H ₂ 0: Distilled.
of	reagent	added were o	converted from	
		ass using the components.	e known densities	
	ene pure	componenter		
				ESTIMATED ERROR: Solubility: ±10% (compiler).
				Temperature: ±0.5 K.
				REFERENCES: 1. Othmer. D. F.: White. R. E.: Trueger. E.
				 Othmer, D. F.; White, R. E.; Trueger, E. Ind. Eng. Chem. <u>1941</u>, 33(10), 1240-8.

COMPONENTS :	ORIGINAL MEASUREMENTS:
 Water; H₂0; [7732-18-5] Chlorobenzene: C₆H₅Cl; [108-90-7] 	Hutchison, C. A.; Lyon, A. M. Columbia University Report A-745, July 1, <u>1943</u> .
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10g(1)/kg^{a} = 10^{2}mol(1)/kg^{b}$	$10^3 x(1)$ c
25 4.504 2.50	2.807
b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Mixtures of 1 to 15 volume ratios of solute to solvent were introduced into an equili- bration flask and placed in a water ther- mostat bath. The assembly was shaken mechanically for about 90 minutes at con- stant temperature. The amount of water in the organic phase was determined by a modi- fied Karl Fischer titration. The determi- nations were done in triplicate. The ex- perimental procedure was taken from a secondary source (1). The original re- port is no longer available.	H ₂ O: Distilled. C ₆ H ₅ Cl: Source is not known, purified and dried before use. ESTIMATED ERROR: Solubility: ±1%.
	Temperature: ±0.05 K.
	REFERENCES: 1. Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series Division III-Vol. 4F, Murphy, G. M., Urey, H. C., Kirshenbaum, I., Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II, Chapter 7, pp 129-44.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Booth, H. S.; Everson, H. E. Ind. Eng.
	Chem. <u>1948</u> , 40(8), 1491-3.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10m1(1)/dm ³ (2) ^a 10 ³ mo1(1)	$/dm^{3 b} 10^{5} x(1)^{c}$
25.0 < 2 < 2.	0 < 3.6
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by r. w. Getzen.	
AUXILIARY	INFORMATION
METHOD /APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The measurements were made with samples	C ₆ H ₅ Cl: Commercial reagent, C.P. grade,
contained in a stoppered Goetz tube placed in a constant temperature water bath. E-	used as received.
quilibrium was assured through repeated	H ₂ 0: Distilled.
shaking and centrifuging the mixture in the stoppered tube while it was temporarily	
removed from the water bath. The amount of solute dissolved was determined as the	
difference between total amount added and	
amount remaining in excess upon saturation. The determination of the excess amount of	
solute added has been described by Hanslick (1).	POTIMATED PROD-
	ESTIMATED ERROR: Solubility: <100%.
	Temperature: ±1 K (compiler).
	REFERENCES: 1. Hanslick, R. S., Ph.D. Dissertation.
	 Hanslick, R. S., Ph.D. Dissertation, Columbia University, New York, <u>1935</u>.

164	Chlorobenzene			
COMPONENTS:			ORIGINAL MEASUREMENTS:	
	obenzene; C ₆ H ₅ Cl; [10 ; H ₂ O; [7732-18-5]	8-90-7]	Newman, M.; Hayworth, C. B.; Treybal, R. E. Ind. Eng. Chem. <u>1949</u> , 41(9), 2039-43.	
VARIABLES:				
One tempera	ature		PREPARED BY: A. L. Horvath	
EXPERIMENTAL	VALUES:			
t/°C	g(1)/kg ^a 1	0 ² mol(1)/kg ^b	$10^4 x(1)$ c	
25	4.0	3.55	6.42	
t/°C	10g(1)/kg ^d 1	0 ³ mol(1)/kg ^d	$10^5 x(1)^{d}$	
25	4.0	3.55	6.40	
<u> </u>		AUXILIARY	INFORMATION	
Equilibrium was establi bath with s position an	ATUS/PROCEDURE: a between the solute shed in a constant t sufficient agitation. alysis was by standa with specific gravity malysis.	emperature The com - rd proce-	SOURCE AND PURITY OF MATERIALS: C ₆ H ₅ Cl: Not specified. H ₂ O: Not specified. ESTIMATED ERROR:	
			Solubility: Too large by a factor of 10 (compiler). Temperature: ±0.3 K (compiler). REFERENCES: 1. Smith, J. C. Ind. Eng. Chem. <u>1942</u> , 34(2), 234-7.	
			53(2), 234-1.	

COMPONENTS:	ORIGINAL MEASUREMENTS:
 (1) Chlorobenzene; C₆H₅Cl; [108-90-7] (2) Water; H₂O; [7732-18-5] 	Andrews, L. J.; Keefer, R. M. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1950</u> , 72(9), 3113-6.
VAPTARY DC.	
VARIABLES: One temperature	PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm ³	^b $10^5 x(1)$ ^c
25.0 5.0 4.44	8.03
b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Water was saturated with chlorobenzene in a glass stoppered Erlenmeyer flask by ro- tating the flask in a constant temperature bath for 20 hours. The saturated solution was extracted with n-hexane for analysis. The optical density of the extract was measured against a n-hexane standard using a Beckman spectrophotometer (1).	C ₆ H ₅ Cl: Eastman Kodak Co., commercial reagent, b.p. 131.9-132°C, fractionated before use. H ₂ O: Not specified.
	ESTIMATED ERROR:
	Solubility: ±10% (compiler).
	Temperature: ±0.1 K (compiler).
	REFERENCES: 1. Andrews, L. J.; Keefer, R. M. J. Am. <i>Chem. Soc.</i> <u>1949</u> , 71(11), 3644-7.

Chlorobenzene

166	Chlorob	penzene		
COMPONENTS:		ORIGINAL MEASUREMENTS:		
(1) Water; H ₂ 0; [7732-18	-5]	Filippov, T. S.; Furman, A. A. <i>2h. Prikl.</i> <i>Khim</i> . 1952, <i>25</i> , 895-7.		
(2) Chlorobenzene; C ₆ H ₅ C	1; [108-90-7]			
VARIABLES:		PREPARED BY:		
Temperature		A. L. Horvath		
EXPERIMENTAL VALUES:				
t/°C 10g(1)/kg ^a	10^2 mol(1)/kg ^b	$10^{3}x(1)$ c		
17.7 2.75	1.526	1. 716		
22.9 3.33	1.848	2.077		
25.2 3.82	2.120	2.382		
28.0 4.70	2.609	2.929		
30.4 5.12	2.842	3.190		
32.4 5.60	3.108	3.489		
35.3 6.21	3.447	3.867		
36.6 6.60	3.663	4.109		
39.8 7.31	4.058	4.550		
42.7 8.25	4.579	5.132		
47.9 9.55	5.301	5.937		
49.0 9.90	5.495	6.153		
b. Calculated by F. c. Calculated by com				
	AUXILIARY	INFORMATION		
METHOD/APPARATUS/PROCEDURE	:	SOURCE AND PURITY OF MATERIALS:		
Measured amounts of water		H ₂ 0: Double distilled.		
were introduced into a ta was then sealed. The am shaken in a thermostat at ulated temperatures until cloudy. Variations in te clouding and clearing cor more than 0.2-0.3°C. Thi solubility measurements h to as the cloud point met	ooule was constantly a series of reg- it appeared emperature for aditions were not s procedure for as been referred	times over freshly calcined cal- cium oxide.		
		ESTIMATED ERROR:		
		Solubility: ±5% (compiler).		
		Temperature: ±0.1 K (compiler).		
		REFERENCES :		
		1. Alexejew, Wladimir Ann. d. Phys. u. Chem. <u>1886</u> , 28, 305-38.		

		Chiorod	benzene		16
COMPONENTS:			ORIGINAL	MEASUREMENTS:	
 Chlorobenezene; C₆H₅C1; [108-90-7] Water; H₂O; [7732-18-5] 			Chemical Series N	ch, R. R. "Physical Properties of Compounds", Advances in Chemist No. 15; American Chemical Society ton, D. C., <u>1955</u> ; p 134.	ry
VARIABLES :			PREPARED	RY •	
One temperatur	e		A. L. Ho		
EXPERIMENTAL VAL	LUES:	·····			
t/°C 1	.0g(1)/kg(2) ^a	10 ³ mol(1)/k	g z	$10^{5}x(1)^{c}$	
25	5.0	4.44		8.00	
t/°C 10) ⁻¹ g(2)/kg(1) ^a	mo1(2)/k	.g D	10x(2) ^c	
25	4.4	2.34		2.16	
		AUXILIARY	1		
METHOD/APPARATU				ND PURITY OF MATERIALS;	
No details are	e available.		с ₆ н ₅ с1:	Dow Chemical Co., 99.98% pure, purified by distillation before use.	
			^н 2 ^{0:}	Distilled.	
			ESTIMATEI Solubili	Lty: (1) in (2) ±10% (compiler). (2) in (1) <100% (compiler)	
			Temperat REFERENCI	ture: ±1 K (compiler).	

COMPONENTS:	ORIGINAL MEASUREMENTS:
 Water; H₂0; [7732-18-5] Chlorobenzene: C₆H₅C1; [108-90-7] 	Wing, J.; Johnston, W. H. <i>J. Am. Chem. Soc.</i> <u>1957</u> , 79(4), 864-5.
VARIABLES: One temperature	PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10m1(1)/dm ^{3 a} 10 ² mo1(1)/dm	$^{3 b} 10^{3} x(1)^{c}$
25.0 3.60 1.992	2.035
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Tritiated water was equilibrated with 20 ml chlorobenzene through stirring in a flask in a constant temperature water bath for two hours. The concentration of the tritiated water in the organic phase was determined by isotopic dilution. For the tritiated water samples, the tritium acti- vities were determined by the acetylene method (1,2). At least four independent determinations were made.	H ₂ 0: Tracerlab Inc., tritiated water, used as received. C ₆ H ₅ Cl: Source not specified, chemical grade, redistilled before use.
The article was based upon work reported in a Ph.D. dissertation (2).	ESTIMATED ERROR:
	Solubility: ±2.8%.
	Temperature: ±0.02 K.
	 REFERENCES: 1. Wing, J.; Johnston, W. H. Science <u>1955</u>, 121, 674-6. 2. Wing, J., Ph.D. Dissertation, Purdue University, Lafayette, <u>1956</u>.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ O; [7732-18-5]	Zieleinski, A. Z. Chem. Stosowana <u>1959</u> ,
(2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	3, 377-84.
6576	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ² mo1(1)/dm ³	^b $10^{3}x(1)$ ^b
25 6.1 3.39	3.45
a. Reported. b. Calculated by compiler.	
,,,	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Not specified.	Not specified.
	ESTIMATED ERROR: Solubility: ±10% (compiler).
	Temperature: ±1 K (compiler).
	REFERENCES:

	COMPONENTS :	ORIGINAL MEASUREMENTS:
Temperature A. L. Horvath EXPERIMENTAL VALUES: $t/^{c}$ log(1)/kg ^a 10 ³ mol(1)/kg ^b 10 ⁴ x(1) ^c 30 4.90 4.353 40 7.05 6.263 40 7.05 6.263 50 9.60 8.529 50 9.60 8.529 50 9.60 8.529 70 16.05 14.259 80 18.05 16.036 90 25.00 22.210 4.0098 a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler. METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS: Calculated by compiler. SOURCE AND PURITY OF MATERIALS: Calculated by compiler. Ch5C1: Source not specified, $n_D^{20} = 1.5248$ bcvr2 1 liter aqueous solutions in a flask. The equilibrium chlorobenzene vapor twas obtained from the gata law. The amount of water vapor was obtained from the gata law. The exhipt increase of the absorbel lass the weight of the chlorobenzene absorbed. The chlorobenzene vapor pressure was directly proportional to its content in water. Dupit retailed before use. ESTIMATED ERROR: Solubility: ±43. Temperature: ±0.1 K.	(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Kisarov, V. M. Zh. Prikl. Khim. <u>1962</u> , 35(10)
$\frac{t/^{\circ}C}{30} = \frac{10^{3}mol(1)/kg}{30} = $		
$\frac{30}{4.90} + \frac{4.353}{6.263} + \frac{1.129}{1.538}$ $\frac{40}{50} + \frac{9.60}{9.60} + \frac{8.529}{8.529} + \frac{1.538}{1.7623}$ $\frac{50}{70} + \frac{16.05}{14.259} + \frac{1.7623}{2.5724} + \frac{1.60}{2.8934}$ $\frac{90}{90} + \frac{25.00}{22.210} + \frac{2.934}{4.0098}$ $\frac{1.8}{1.805} + \frac{16.036}{1.6036} + \frac{2.8934}{2.8934}$ $\frac{90}{90} + \frac{25.00}{22.210} + \frac{2.098}{4.0098}$ $\frac{1.8}{1.805} + \frac{1.60}{1.6036} + \frac{1.80}{1.808} + 1.$	EXPERIMENTAL VALUES:	
$\frac{40}{100} 7.05 \qquad 6.263 \qquad 1.129$ $\frac{50}{50} 9.60 \qquad 8.529 \qquad 1.538$ $\frac{60}{11.00} \qquad 9.7726 \qquad 1.7623$ $\frac{70}{16.05} \qquad 14.259 \qquad 2.5724$ $\frac{80}{90} \qquad 18.05 \qquad 16.036 \qquad 2.8934$ $\frac{90}{25.00} \qquad 22.210 \qquad 4.0098$ a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler. $\frac{400}{100} \frac{100}{100} \frac$	t/°C 10g(1)/kg ^a 10 ³ mol(1)/kg ^b	$10^4 x(1)^{c}$
509.608.5291.5386011.009.77261.76237016.0514.2592.57248018.0516.0362.89349025.0022.2104.0098a. Reported.b. Calculated by F. W. Getzen.c. Calculated by compiler.LINER MATIONMUXILIARY INFORMATIONMUXILIARY INFORMATION <td>30 4.90 4.353</td> <td>0.7846</td>	30 4.90 4.353	0.7846
	40 7.05 6.263	1.129
7016.0514.2592.57248018.0516.0362.89349025.0022.2104.0098a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler	50 9.60 8.529	1.538
80 18.05 16.036 2.8934 90 25.00 22.210 4.0098 a. Reported. b. Calculated by F. W. Getzen. c. c. Calculated by compiler. Value of the compiler. SURCE AND PURITY OF MATERIALS: METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS: Temperatures were maintained using a thermostat bath with continuous stirring. Partial pressures of chlorobenzene vere determined over 2 liter aqueous solutions in a flask. SOURCE AND PURITY OF MATERIALS: The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor was obtained from the total weight increase of the absorber less the weight of the chlorobenzene absorbed. H20: Redistilled before use. H20: Redistilled before use. ESTIMATED ERROR: Solubility: ±4%. Temperature: ±0.1 K. Temperature: ±0.1 K.	60 11.00 9.7726	1.7623
 90 25.00 22.210 4.0098 a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler. AUXILIARY INFORMATION METROD/APPARATUS/PROCEDURE: Temperatures were maintained using a thermostat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The total weight of the chlorobenzene absorbed. The chlorobenzene apor pressure was directly proportional to its content in water. Duplicate measurements were made at each temperature: Source and the total of the vapor mass directly proportional to its content in water. Duplicate measurements were made at each temperature: ESTIMATED ERROR: Solubility: ±4%. Temperature: ±0.1 K.	70 16.05 14.259	2.5724
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler. AUXILIARY INFORMATION METHOD/APPARATUS/PROCEDURE: Temperatures were maintained using a thermo- stat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor ware obtained from the total weight of the chlorobenzene absorbed. The chlorobenzene vapor pressure was directly proportional to its content in water. Dupli- cate measurements were made at each tempera- ture. Solubility: $\pm 4\%$. Temperature: $\pm 0.1 K$.	80 18.05 16.036	2.8934
b. Calculated by F. W. Getzen. c. Calculated by compiler. AUXILIARY INFORMATION METHOD/APPARATUS/PROCEDURE: Temperatures were maintained using a thermo- stat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor was obtained from the total weight increase of the absorber less the weight of the chlorobenzene was directly proportional to its content in water. Dupli- cate measurements were made at each tempera- ture. ESTIMATED ERROR: Solubility: $\pm 4\%$. Temperature: $\pm 0.1 K$.	90 25.00 22.210	4.0098
METHOD/APPARATUS/PROCEDURE:SOURCE AND PURITY OF MATERIALS:Temperatures were maintained using a thermo- stat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor was obtained from the total weight increase of the absorber less the weight of the chlorobenzene absorbed. The chlorobenzene vapor pressure was directly proportional to its content in water. Dupli- cate measurements were made at each tempera- ture.SOURCE AND PURITY OF MATERIALS: Cate Source not specified, $n_D^{20} = 1.5248$ $b.p. = 132°C$ H20: Redistilled before use.ESTIMATED ERROR: Solubility: $\pm 4\%$. Temperature: ± 0.1 K.		
Temperatures were maintained using a thermo- stat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor was obtained from the total weight increase of the absorber less the weight of the chlorobenzene absorbed. The chlorobenzene vapor pressure was directly proportional to its content in water. Dupli- cate measurements were made at each tempera- ture. ESTIMATED ERROR:Solubility: ±4%. Temperature: ±0.1 K.	AUXILIARY	INFORMATION
	Temperatures were maintained using a thermo- stat bath with continuous stirring. Partial pressures of chlorobenzene were determined over 2 liter aqueous solutions in a flask. The equilibrium chlorobenzene content of the vapor was calculated from the gas laws. The amount of water vapor was obtained from the total weight increase of the absorber less the weight of the chlorobenzene absorbed. The chlorobenzene vapor pressure was directly proportional to its content in water. Dupli- cate measurements were made at each tempera-	C ₆ H ₅ Cl: Source not specified, n _D ²⁰ = 1.5248 b.p. = 132°C H ₂ O: Redistilled before use. ESTIMATED ERROR: Solubility: ±4%. Temperature: ±0.1 K.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Högfeldt, E.; Bolander, B. Ark. Kemi 1963,
(2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	21(16), 161-86.
65	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10g(1)/dm^{3} = 10^{2}mol(1)/dc$	$\lim^{3} b$ 10 ³ x(1) c
25 5.40 3.0	3.06
a. Calculated by F. W. Getzen. b. Reported.	
c. Calculated by compiler.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE: A mixture of 15 ml chlorobenzene with 10 ml	SOURCE AND PURITY OF MATERIALS: H _n O: Distilled.
distilled water was shaken in a glass-	2
stoppered bottle overnight at room tempera- ture. After centrifugation, duplicate sam-	b J purity: 99.8%,
ples were taken for the determination of the water content in the organic phase. Titra-	impurity: 0.11% benzene.
tion of the samples was carried out by a modified Karl Fischer method (1). A small	
correction was made for water in the vapor	
phase and adsorbed on the pipet.	
	ESTIMATED ERROR:
	Solubility: ±10%.
	Temperature: ±0.3 K.
	REFERENCES:
	1. Johansson, A. Sven. Papperstidn. <u>1947</u> ,
	<i>50(11B)</i> , 124–33.
	1

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ O; [7732-18-5] (2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	Jones, J. R.; Monk, C. B. <i>J. Chem. Soc.</i> <u>1963</u> , Part III, 2633-5.
VARIABLES:	PREPARED BY:
Temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10m1(1)/dm ³ (2) ^a 10 ² mol($1)/dm^{3 b}$ $10^{3}x(1)^{c}$
25 3.4 1.8	38 1.92
30 3.95 2.1	82 2.239
35 4.6 2.5	2.62
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Tritiated water was shaken with chlorobenzene in 1:10 volume ratios for 4 hours in flasks placed in a water thermostat bath. The water content was determined by tritium assay of samples taken from the flasks. The count rates were determined using a typical liquid scintillator solution technique.	<pre>H₂0: Tritiated. C₆H₅Cl: Source not known, laboratory grade, dried over CaCl₂ and fractionally distilled before use. ESTIMATED ERROR: Solubility: ±5%. Temperature: ±0.5 K (compiler). REFERENCES:</pre>

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ 0; [7732-18-5]	Goldman, S., Ph.D. Dissertation, McGill
(2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	University, Montreal, <u>1969</u> , p. 84.
VARIABLES:	PREPARED BY:
Temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ² mol(1)/d	$dm^{3 b}$ 10 ³ x(1) ^c
25.0 5.225 2.90	2.959
35.1 7.405 4.11	4.234
45.06 9.639 5.35	5.567
a. Calculated by F. W. Getzen. b. Reported.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibration experiments were carried out	H ₂ 0: Distilled.
in a constant temperature water bath using a stirrer. An equilibration period of at	C ₆ H ₅ Cl: Reagent grade, washed with
least 5 days was allowed. The total water content in the organic phase was determined	conc. H ₂ SO ₄ and with 1 M NaHCO ₃ and then fractionally distilled
by a Karl Fischer titration. Each reported water solubility was obtained as an average	over silica gel.
of at least two independent determinations.	
	ESTIMATED ERROR:
	Solubility: ±2%.
	Temperature: ±0.1 K.
	REFERENCES :

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7] (2) Water; H ₂ O; [7732-18-5]	Chey, W.; Calder, G. V. J. Chem. Eng. Data 1972, 17(2), 199-200.
VARIABLES: One temperature	PREPARED BY: A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg(2) ^a 10 ³ mo1(1)	$1/kg^{b}$ $10^{5}x(1)^{c}$
21 5.34 4.742	8.546
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
A 1 to 50 volume ratio solute/solvent mix- ture was stirred in a separatory funnel for 20 hr in a thermostat bath. Then, the solu- tion was passed through a separation column of nonpolar resin. The solute was eluted with isopropyl ether. Traces of water were removed using molecular sieve pellets, and the sample was injected into a gas chroma- tograph fitted with a stainless steel column packed with Chromosorb having a liquid phase of Carbowax for analysis. The determination was done in duplicate.	C ₆ H ₅ Cl: Source not specified, Analytical Reagent grade. H ₂ O: Distilled.
was uone in dupitate.	ESTIMATED ERROR: Solubility: ±2%. Temperature: ±1 K.
	REFERENCES :

COMPONENTS:	ORIGINAL MEASUREMENTS:
 (1) Chlorobenzene; C₆H₅C1; [108-90-7] (2) Water; H₂O; [7732-18-5] 	Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u> .
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ³ mol(1)/k	$g(2)^{b}$ $10^{5}x(1)^{a}$
25.0 4.622 4.108 ± 0.	062 7.4004
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The solute was equilibrated with water by stirring in a sealed flask (1) with a magne- tic stirrer for 48 hours at constant tem- perature. A sample was withdrawn and fil- tered through a glass-wool plug and the solute was extracted with 2,2,4-trimethyl- pentene. The optical densities were deter- mined spectrophotometrically (2). The mean and standard deviation were obtained from eight independent measurements.	<pre>SOURCE AND PURITY OF MATERIALS: C₆H₅Cl: Merck AG., >99% GLC, used as received. H₂O: Distilled, deionized, and degassed.</pre>
	ESTIMATED ERROR:
	Solubility: ±1.51%.
	Temperature: ±0.05 K.
	 REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J. Chem. Soc. <u>1963</u>, Part III, 2716-23. 2. Wauchope, R. D.; Getzen, F. W. J. Chem. Eng. Data <u>1972</u>, 17(1), 38-41.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8), 839-45.
(2) Water; H ₂ ⁰ ; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ³ mol(1)/k	$g(2)^{b}$ 10 ⁵ $x(1)^{a}$
25.0 4.624 4.11	7.404
a. Calculated by compiler.	
b. Reported.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium between the water and chloro- benzene was established by stirring the sam-	C ₆ H ₅ Cl: Commercial reagent of analytical grade distilled through a column
ple in a sealed flask (1) with a magnetic stirrer for 48 hours. The sample was fil-	resulting in a more than 99% pure sample.
tered through a glass-wool plug and the	-
chlorobenzene was extracted with 2,2,4- trimethylpentene. The optical density was	H ₂ O: Distilled, deionized, and degassed.
determined spectrophotometrically (2). Five parallel determinations were made.	
The reported work was based upon a Ph.D.	
dissertation (3).	ESTIMATED ERROR:
	Solubility: ±1.7%.
	Temperature: ±0.05 K.
	REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. J.
	Chem. Soc. <u>1963</u> , Part III, 2716-23. 2. Wauchope, R. D.; Getzen, F. W. J. Chem.
	Eng. Data <u>1972</u> , 17(1), 38-41.
	 Vesala, A., Ph.D. Dissertation, Univer- sity of Turku, Turku, <u>1973</u>.

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COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H ₂ 0; [7732-18-5]	Prosyanov, N. N.; Shalygin, V. A;	
(2) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Zel'venskii, Ya. D. Tr. Mosk. Khim. Tekhnol. Inst. <u>1974</u> , 81, 55-6.	
VARIABLES:	PREPARED BY:	
Temperature: 298-361 K	A. L. Horvath	
EXPERIMENTAL VALUES:		
1222.37		
$\log_{10} \alpha = \frac{1222.37}{T(K)} - 1.3225$		
where α = distribution coefficient.		
At the normal boiling point of $C_{\beta}H_{5}C1$: $\alpha = 49.0$.		
0.	6 5	
AUXILIARY INFORMATION		
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The water concentration in the saturated solutions was determined radiometrically	H ₂ 0: Not specified.	
using tritium labelled water.	C ₆ H ₅ Cl: Not specified.	
The method of investigation has been de- scribed in greater detail in (1).		
	ESTIMATED ERROR: Solubility: ±10% (compiler).	
	Temperature: ±0.5 K (compiler).	
	REFERENCES:	
	1 Programow N N · Shalvein V A ·	

 Prosyanov, N. N.; Shalygin, V. A.; Zel'venskii, Ya. D. Tr. Mosk. Khim. Tekhnol. Inst. <u>1973</u>, 75, 100-2.

HBTPW-G*

COMPONENTS :	ORIGINAL MEASUREMENTS:
 Water; H₂0; [7732-18-5] Chlorobenzene; C₆H₅C1; [108-90-7] 	Kirchnerova, J.; Cave, G.C.B. Can. J. Chem. 1976, 54(24), 3909-16.
(-, -, -, -, -, -, -, -, -, -, -, -, -, -	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)dm ^{3 a} 10 ² mol(1)/d	$m^{3 b} 10^{3} x(1)^{c}$
25 5.225 2.90	2.959
a. Calculated by F. W. Getzen. b. Reported. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: A mixture of 50 ml chlorobenzene and 6 ml water in a bottle was submerged in a water thermostat bath for 2 days. The concentra- tion of the water in the organic phase was determined by a conventional Karl Fischer dead stop back titration. Determinations were done in triplicate. A detailed description of the complete ex- perimental procedure has been included in a Ph.D. dissertation (1).	<pre>SOURCE AND PURITY OF MATERIALS: H₂O: Distilled and deionized. C₆H₅Cl: Fischer - B255, washed with cc. H₂SO₄ and K₂CO₃ solutions and distilled water. Dried over silica gel and fractionally dis- tilled, purity: 99.8%.</pre>
	ESTIMATED ERROR:
	Solubility: ±1%. Temperature: ±0.1 K.
	REFERENCES:
	 Kirchnerova, J., Ph.D. Dissertation, McGill University, Montreal, Quebec, <u>1974</u>.

COMPONENTS:		ORIGINAL MEASUREMENTS:
 Chlorobenzene; Water; H₂0; [7 	C ₆ H ₅ Cl; [108-90-7] 732-18-5]	Nelson, H. D.; Smit, J. H. SAfr. Tydskr. Chem. <u>1978</u> , 31(2), 76.
VARIABLES:		PREPARED BY:
Temperature		A. L. Horvath
EXPERIMENTAL VALUES	:	
t/°C 10	lg(1)/kg ^a 10 ³ mol(1)/k	$x_{g}^{b} 10^{5}x(1)^{c}$
5	0.400 0.355	0.64
25	1.068 0.9492	1.71
35	2.674 2.376	4.28
45	4.005 3.558	6.41
	ed by compiler. ed by F. W. Getzen.	
AUXILIARY INFORMATION		INFORMATION
METHOD/APPARATUS/PR	OCEDURE:	SOURCE AND PURITY OF MATERIALS:
	with chlorobenzene from a special flask (1) placed	C ₆ H ₅ Cl: Not specified.
in a thermostat bat	h using a shaker for a	H ₂ 0: Not specified.
tographic analysis tion into a 1.5 m s having a 5% Apiezon operated at 120°C. fitted with a flame	on period. A gas chroma- was done by sample injec- tainless steel column M coating on Celite The chromatograph was ionization detector. each flask were analyzed.	
		ESTIMATED ERROR:
		Solubility: ±2.9% (compiler).
		Temperature: ±0.1 K (compiler).
		REFERENCES :
		 Nelson, H. D.; de Ligny, C. L. Rec. Trav. Chim. <u>1968</u>, 87, 528-44.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	Aquan-Yuen, M.; Mackay, D.; Shiu, W. Y.
(2) Water; H ₂ O; [7732-18-5]	J. Chem. Eng. Data 1979, 24(1), 30-4.
(-,, -2, ,	
VARIABLES:	PREPARED BY: A. L. Horvath
One temperature	A. L. HOFVALII
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10g(1)/dm ³ a 10 ³ mol(1)/	$\dim^{3} b$ $10^{5}x(1)$ c
25 4.717 4.1907	7.5753
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
An excess of solute was added to pure water in a flask which was then placed in a con-	C ₆ H ₅ Cl: Fisher Scientific Inc., certified grade, used as received.
stant temperature bath for equilibration	
for at least 48 hours before analysis. The concentration of the chlorobenzene was de-	H ₂ 0: Specified as "pure".
termined by solvent extraction, followed by	
gas chromatographic analysis. The chromato- graph was equipped with a flame ionization	
detector as described in (1).	
	ESTIMATED ERROR:
	Solubility: ±3.8%.
	Temperature: ±0.5 K (evaluator).
	REFERENCES:
	1. Mackay, D.; Shiu, W. Y. Bull. Environ. Contam. Toxicol. <u>1976</u> , 15, 101-12.

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Chlorobenzene; C ₆ H ₅ C1; [108-90-7]	Schwarz, F. P.; Miller, J. Anal. Chem. 1980, 52(13), 2162-4.
(2) Water; H ₂ O; [7732-18-5]	<u>1960</u> , <i>52(13)</i> , 2102-4.
_	
VARIABLES :	PREPARED BY:
Temperature	A. L. Horvath
-	
EXPERIMENTAL VALUES:	
Experimentally determined values:	
10g(1)/kg	
t/°C Elution Chromatography	UV Absorption
10.0 4.4 ± 0.2	4.9 ± 0.3
20.0 4.2 ± 0.2	4.8 ± 0.4
30.0 4.9 ± 0.1	5.0 ± 0.2
Values derived from average measured s	olubilities:
t/°C 10g(1)/kg ^a 10 ³ mol(1)/k	b 10 ⁵ x(1) ^c
10.0 4.6 ± 0.2 4.09 ± 0.18	
20.0 4.5 \pm 0.4 4.00 \pm 0.36	
$30.0 5.0 \pm 0.1 4.44 \pm 0.09$	
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by r. w. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Both elution chromatography and UV absorption	C ₆ H ₅ Cl: Commercial, 98 wt %.
methods were used to determine the aqueous solubilities. The agreement was within an	H ₂ 0: Distilled.
experimental error of 4% between the two methods. The average deviations were deter-	-
mined from several measurements made on dif- ferent samples.	
The analytical procedures for determining	
organic liquid solubilities in water based upon liquid phase elution chromatography has	
been described in (1).	ESTIMATED ERROR:
	Solubility: ±4%.
	Temperature: ±0.5 K.
	REFERENCES:
	 Schwarz, F. P. Anal. Chem. <u>1980</u>, 52(1), 10-15.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water-d ₂ ; D ₂ 0; [7729-20-0]	Hutchison, C. A.; Lyon, A. M. Columbia
	University Report A-745, July 1, <u>1943</u> .
(2) Chlorobenzene; C ₆ H ₅ Cl; [108-90-7]	
VARIABLES:	PREPARED BY:
One temperature	G. Jancso
EXPERIMENTAL VALUES:	
	b 3 a
t/°C 10g(1)/kg ^a 10 ² mol(1)/k	$g^{b} 10^{3}x(1)^{a}$
25.0 4.386 2.19	2.460
a. Calculated by F. W. Getzen.	
b. Reported (average of two experimen	tal measurements).
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	
Between 25 and 100 ml of chlorobenzene and	SOURCE AND PURITY OF MATERIALS:
1 to 2 ml of D_2O were introduced into a flask	
and shaken for about 90 min. The water bath temperature was maintained within ±0.05°C.	dried before use. Source and method not given.
Then, a sample was taken and the dissolved	
D_2O content was determined by a modified Karl Fischer titration as described in (1).	100% D ₂ 0: Source not specified.
The original report was unavailable, but the	
method and results have been described in	
sufficient detail in (1). The solubility of H ₂ O in chlorobenzene was also determined	
and found to be $0.00250 \pm 0.00003 \text{ mol H}_20/$	ESTIMATED ERROR:
100 g solution from three separate experi- ments.	Solubility: av. dev. $\pm 1 \times 10^{-5}$ mol D ₂ 0/100 g
	solution.
	Temperature: ±0.05 K.
	REFERENCES:
	 Eidinoff, M. L. In "Production of Heavy Water", National Nuclear Energy Series
	Division III-Vol. 4F, Murphy, G. M.;
	Urey, H. C.; Kirshenbaum, I.; Eds.; McGraw-Hill: New York, <u>1955</u> ; Part II,
	Chapter 7, pp 129-44.

COMPONENTS:	EVALUATOR:
(1) 2-Chlorophenol; C ₆ H ₅ C10; [95-57-8]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2) Water; H ₂ O; [7732-18-5]	September 1982.

CRITICAL EVALUATION:

The mutual solubility of 2-chlorophenol and water was investigated by Sidgwick and Turner (1) in 1922. While the measurements seem to be performed with care and the method, which involved classical synthetic procedures based upon gravimetry, gives relatively accurate results, there is some difficulty in resolving a solubility value at 298 K from the data. This circumstance arises because of the range of temperatures over which the measurements were made which makes the interpolation of the data inaccurate. Also, some of the data values scatter considerably in the low temperature range.

Kuroda (2) found the solubility of 2-chlorophenol in water to be in the order of 2 percent at room temperature (291 K). Better agreement is obtained when the data of Mulley and Metcalf (3) are compared with the values of Sidgwick and Turner. The 24.65 g(1)/dm³ value of Mulley and Metcalf corresponds to a value of 25 g(1)/kg (assuming a density of 1.0 g/cm³ for the solution).

The more recent work of Banerjee et al. (4) gives a value of $88.3 \text{ mmol}(1)/\text{dm}^3$ corresponding to approximately 11 g(1)/kg, a value that does not agree well with the earlier values. One possible reason for this lack of agreement may be associated with the use of a radiochemical method of analysis by Banerjee et al. For instance, a systematic error could have resulted from radiochemical impurities of the substrate. In any event, the lack of relevant information makes an evaluation here too speculative. The available experimental data are shown in Figure 1.

It should be mentioned that the pH of the solutions exerts a considerable influence on the solubility values of those solutes such as 2-chlorophenol capable of protolytic action. Here it is assumed that the reported solubilities refer to the pH values which prevail in the saturated solutions of 2-chlorophenol in water.

No more than a tentative value can be assigned to the solubility of 2-chlorophenol in water. The following value has been obtained from the data shown in Figure 1 by a graphical interpolation:

T/K	10mo1(1)/dm ³	$10^{-1}g(1)/kg$	$10^{3}x(1)$
298.15	1.6	2.	2.9

REFERENCES

- 1. Sidgwick, N. V.; Turner, S. L. J. Chem. Soc. 1922, 121, Part II, 2256-63.
- 2. Kuroda, T. cf. Chem. Zentralbl. 1926, I, 3610.
- 3. Mulley, R. A.; Metcalf, A. D. Sci. Pharm. 1966, 2, 481-8.
- 4. Banerjee, S.; Yalkowsky, S. H.; Valvani, S. C. Environ. Sci. Technol. <u>1980</u>, 14(10), 1227-9.

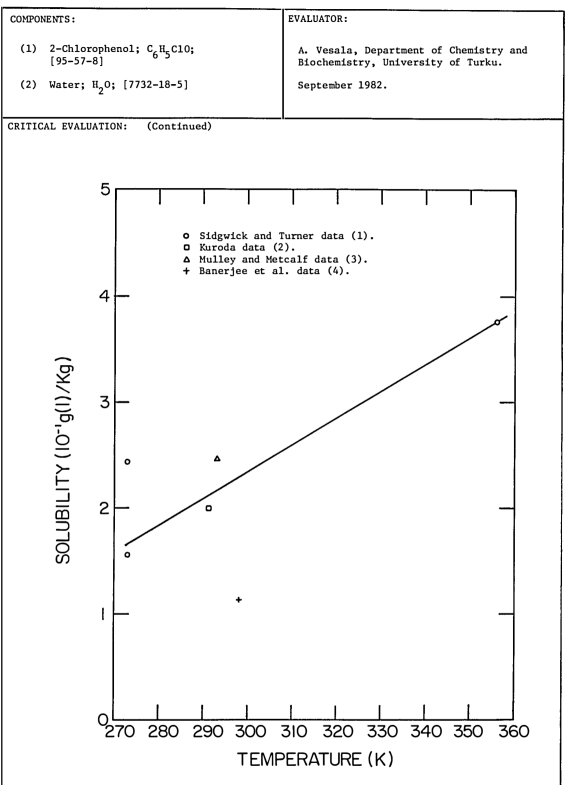
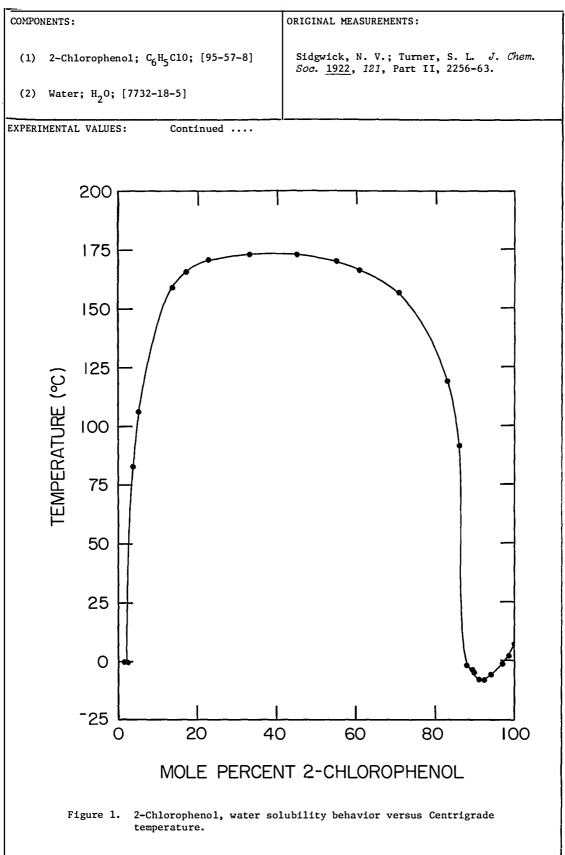


Figure 1. Solubility of 2-chlorophenol in water versus Absolute temperature.

COMPONENTS :			ORIGINAL	MEASUREMENTS:
(1) 2-Chlorophenol; C ₆ H ₅ Cl0; [95-57-8]			, N. V.; Turner, S. L. J. Chem.	
Û Û			2, 121, Part II, 2256-63.	
(2) Water; H ₂ 0; [7732-18-5]				
VARIABLES:			PREPARED	RY •
Temperature: -8.	.2 - 173.0°C		A. Vesala	
EXPERIMENTAL VAL				
t/°C	$10^{-2}g(1)/kg^{a}$	mol(1)/	'kg ^D	10x(1)
-8.2	9.220	7.1	.719	6.2357
-8.0	9.087	7.0	685	5.8243
-6.0	9.393	7.3	065	6.8440
-5.0	8.962	6.9	712	5.4750
-4.0	8.925	6.9	424	5.3778
-2.0	8.773	6.8	242	5.0050
-1.5	9.676	7.5	266	8.0714
-0.30	0.244	0.1898	0.	.03493
-0.20	0.156	0.1213	0.	.02216
2.0	9.839	7.6	534	8.9544
7.0	10.00	7.7	787	10.0
82.9	0.376	0.2925	0.	.05445
91.5	8.590	6.6	819	4.6056
1				Continued
AUXILIARY INFORMATION METHOD/APPARATUS/PROCEDURE: SOURCE AND PURITY OF MATERIALS:				
]	versus temperature	measure-	C ₆ H ₅ C10:	Synthesized by the chlorination
	by the so called volved the mixing		0 3	of phenol in carbon tetrachloride, purified as described in Ref. 1,
masses of solut	e and solvent and	the testing		no criterions of the purity given.
ing procedure w	with temperature. as applied. In th	is method,	н, о:	Source and purity not specified.
	thermostats, one a the solubility ter			
were used. The	contents of sample	e tubes in	}	
maintain one ho	the two thermostat baths were observed to maintain one homogeneous and the other			
heterogeneous as the temperature interval between the two thermostats were reduced by successive steps.				
		ESTIMATE) ERROR:	
			REFERENCE	7 C •
				eben, W. J. Ber. Dtsch. Chem. Ges.
		<u>1909</u> ,	42, 4370.	
]		
]	
L				

COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) 2-Chlorophenol; C ₆ H ₅ Cl0; [95-57-8] (2) Water; H ₂ 0; [7732-18-5]			Sidgwick, N. V.; Turner, S. L. <i>J. Chem.</i> <i>Soc.</i> <u>1922</u> , <i>121</i> , Part II, 2256-63.
EXPERIMENTA	L VALUES: conti	nued	
t/°C	10 ⁻² g(1)/kg ^a	mol(1)/kg	$b^{b} = 10x(1)^{b}$
106.3	0.512	0.3983	0.07506
118.9	8.282	6.4423	4.0319
156.6	7.062	5.4933	2.5197
159.1	1.358	1.0563	0.21547
165.8	1.695	1.3185	0.27806
166.2	6.072	4.7232	1.7806
170.1	5.495	4.2744	1.4598
170.7	2.259	1.7572	0.39289
172.9	4.504	3.5035	1.0301
173.0	3.300	2.5670	0.64568
	rted. ulated by F. W. Ge ments are shown gra		ure l.



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COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 2-Chlorophenol; C ₆ H ₅ C10; [95-57-8]	Mulley, B. A.; Metcalf, A. D. Sci. Pharm.
	<u>1966</u> , 2, 481-8.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
LA LATENTAL VALUES.	
$t/^{\circ}C$ $10^{-1}g(1)/dm^{3}a$ $10mol(1)$	$/dm^{3 b} 10^{3}x(1)^{b}$
20 2.465 1.917	4 3.5357
a. Reported.	
b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The sample preparation and equilibration	C ₆ H ₅ ClO: Commercial reagent, manufacturer not specified, suitably purified
procedures were not specified. Saturated solutions were prepared and analyzed spec-	until physical constants corres-
trophotometrically. Also, duplicate deter-	sponded with literature values.
minations were probably made by a synthetic	The reagent was dried in a desic-
method described in (1).	cator for two days before use.
	H ₂ 0: Source and purity not specified.
	2
	ESTIMATED ERROR:
	Solubility: <4% (estimated here on the
	basis of the deviations in the reported values).
	REFERENCES :
	1. Mulley, B. A.; Metcalf, A. D. J. Pharm.
	Pharmacol. <u>1956</u> , 8, 774.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 2-Chlorophenol; C ₆ H ₅ C10; [95-57-8]	Banerjee, S.; Yalkowsky, S. H.; Valvani,
(2) Water; H ₂ 0; [7732-18-5]	S. C. Environm. Sci. Techn. <u>1980</u> , 14(10), 1227-9.
(1) "acci, "20, [//32 10 5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{-1}g(1)/dm^{3}a$ $10^{2}mol(1)$	$)/dm^{3 b}$ $10^{3}x(1)^{a}$
25 1.135 8.83	1.611
a. Calculated by F. W. Getzen.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The equilibrations were performed in sealed	C ₆ H ₅ C10: Commercial reagent, the ¹⁴ C-
stainless steel centrifuge tubes with con- stant or intermittent shaking. The equili-	labeled compound was purchased by NEN, the nonlabeled one by
bration was generally complete within 1 week.	Aldrich.
The mixture was then centrifuged for 60 minutes after which aliquots of the solution	H ₂ 0: Distilled.
were removed for analysis either by a pipet	
or a syringe. Liquid scintillation counting with $^{14}\mathrm{C-labeled}$ solute was employed in	
solubility determinations. The entire pro-	
cedure was carried out at least twice and	
each analysis was also conducted in dupli- cate.	ESTIMATED ERROR:
	Solubility: ±0.5% (std. deviation estimated by authors).
	Temperature: ±0.2 K (equilibration) ±0.3 K (centrifugation):
	REFERENCES:

COMPONENTS:	EVALUATOR:	
(1) 3-Chlorophenol; C ₆ H ₅ ClO; [108-43-0]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.	
(2) Water; H ₂ O; [7732-18-5]	November 1979.	

CRITICAL EVALUATION:

The earliest and most thorough investigation of the mutual solubilities of 3chlorophenol and water was done in 1922 by Sidgwick and Turner (1). Their measurements of solubilities were accomplished using the so called synthetic method, i.e., the determination of solubilities from weighed masses of 3-chlorophenol and water at the appropriate temperatures. They reported no directly measured values for the solubilities at 298 K, the nearest measured values were obtained at temperatures relatively far from room temperature (276 K and 359 K), which makes an accurate interpolation difficult.

The solubilities of three isomers of monochlorophenols in water have been determined by Kuroda (2) who reported the magnitude of this solubility to be 2 percent. More support for the work of Sidgwick and Turner is provided by the value measured by Mulley and Metcalf (3) at 293.15 K. Their result of 22.4 g(1)/dm³ (which corresponds to 22.4 g(1)/kg assuming a density of 1.0 for the solution) agrees quite satisfactorily with the values of Sidgwick and Turner as shown in Figure 1. There are not, however, enough data to allow a more rigorous evaluation to be made; therefore, the solubility value reported here must be tentative.

It should be mentioned that the pH of the solutions exerts a considerable influence on the solubility values of those solutes such as 3-chlorophenol capable of protolytic action. Here it is assumed that the reported solubilities refer to the pH values which prevail in the saturated solutions of 3-chlorophenol in water.

No more than a tentative value can be assigned to the solubility of 3-chlorophenol in water 298.15 K as follows:

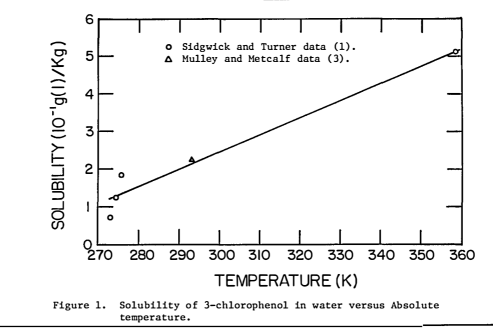
т/к	10mo1(1)/dm ³	10 ⁻¹ g(1)/kg	$10^{3}x(1)$
298.15	1.7	2.2	3.2

REFERENCES

1. Sidgwick, N. V.; Turner, S. L. J. Chem. Soc. <u>1922</u>, 121, Part II, 2256-63.

2. Kuroda, T. cf. Chem. Zentralbl. 1926, I, 3610.

3. Mulley; B. A.; Metcalf, A. D. Sci. Pharm. 1966, 2, 481-8.

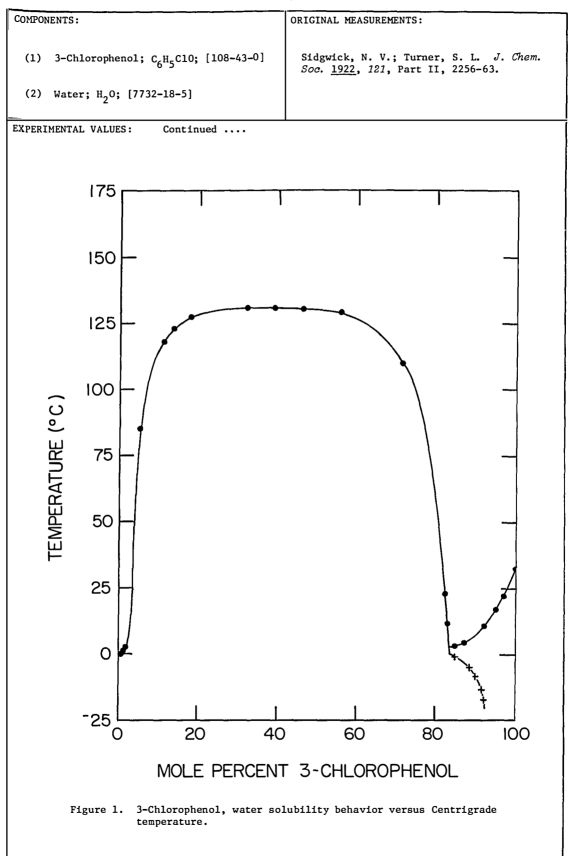


COMPONENTS:		<u></u>	ORIGINAL	MEASUREMENTS:
(1) 3-Chlorophenol; C ₆ H ₅ Cl0; [108-43-0]		Sidgwic	k, N. V.; Turner, S. L. J. Chem.	
(2) Water; H ₂ 0; [7732-18-5]		Soc. <u>19</u>	<u>22</u> , <i>121</i> , Part II, 2256-63.	
(
VARIABLES:			PREPARED	BY:
Temperature: -17	7.0 - 130.8°C		A. Vesa	la
EXPERIMENTAL VALU		- (-)	., b	10x(1) b
t/°C	$10^{-2}g(1)/kg^{a}$			
-17.0	9.223		. 1743	6.2455
-13.2	9.173		.1354	6.0852
-8.2	9.011		.0093	5.6080
-4.8	8.866	6.	. 8966	5.2282
-0.9	8.487	6.	.6017	4.4012
-0.18	0.073	0.0568		0.0103
1,2	0.125	0.09723		0.01771
2.5	0.185	0.1439		0.02635
3.2	8.487	6	.6017	4.4012
4.5	8.719	6.	. 7822	4.8819
10.8	9.223	7.	1742	6.2455
11.8	8,290	6.	. 4485	4.0455
17.0	9.510	7.	. 3975	7.3117
				Continued
	AUXILIARY			ION
METHOD/APPARATUS/PROCEDURE: The solubility versus temperature measure- ments were made by the so called synthetic method which involved the mixing of known		SOURCE A C6H5C10	ND PURITY OF MATERIALS: : Synthesized from 1-chloro-3- nitrobenzene by reduction and diazotisation. Melting point	
	and solvent and vith temperature.			32.5°C.
ing procedure wa	as applied. In th hermostats, one a	is method,	н ₂ 0:	No specifications given.
	the solubility ter contents of sample			
	at baths were obs			
maintain one homogeneous and the other heterogeneous as the temperature interval between the two thermostats was reduced by successive steps.				
		ESTIMATE	ED ERROR:	
		DEFEDENC	νες.	
			REFERENC	
I			I	

5-chlorophenol			
COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) 3-Chlorophenol; C ₆ H ₅ Cl0; [108-43-0] (2) Water; H ₂ 0; [7732-18-5]			Sidgwick, N. V.; Turner, S. L. <i>J. Chem.</i> <i>Soc.</i> <u>1922</u> , <i>121</i> , Part II, 2256-63.
EXPERIMENTA	L VALUES: Cont	inued	
t/°C	$10^{-2}g(1)/kg^{a}$	mol(1)/kg	b 10x(1) b
22.2	9.711	7.553	8 8.2484
23.1	8.230	6.401	8 3.9453
32.5	10.00	7.778	7 10.0
85.25	0.512	0.3983	0.07506
109.8	7.123	5.540	7 2.5759
118.0	1.113	0.86576	0.17248
123.0	1.356	1.0548	0.21511
127.5	1.784	1.3877	0.29531
129.1	5.565	4.328	8 1.4955
130.5	4.612	3.587	5 1.0711
130.7	3.889	3.025	1 0.81882
130.8	3.202	2.4907	0.61922

a. Reported.b. Calculated by F. W. Getzen.

Measurements are shown graphically in Figure 1.



3-Chlorophenol

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 3-Chlorophenol; C ₆ H ₅ C10; [108-43-01]	Mulley, B. A.; Metcalf, A. D. Sci. Pharm.
(2) Water; H ₂ 0; [7732-18-5]	1966, 2, 481-8.
(2) water, n_2^{0} ; $(7752-10-5)$	
VARIABLES:	PREPARED BY:
	A. Vesala
One temperature	A. VESALA
EXPERIMENTAL VALUES:	
-1 3 2	3 h 3 h
t/°C 10 ⁻¹ g(1)/dm ³ a 10mol(1),	$/dm^{3}$ 10 ³ x(1)
20 2.242 1.7439	3.2095
a. Reported.	
b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The sample preparation and equilibration procedures were not specified. Saturated	C ₆ H ₅ ClO: Commercial reagent, suitably purified, until physical constants
solutions were prepared and analyzed spectro-	corresponded with literature val-
photometrically. Also, duplicate determina-	ues. The reagent was dried in a
tions were probably made by a synthetic method described in (1).	desiccator for two days before use.
method described in (i).	
	H ₂ 0: Source and purity not specified.
	ESTIMATED ERROR: Solubility: <4% (estimated here on the
	basis of the deviations in the
	values reported for the solubi-
	lity of this phenol in cetoma- grol solns).
	REFERENCES: 1. Mulley, B. A.; Metcalf, A. D. J. Pharm.
	Pharmacol. 1956, 8, 774.
1	

COMPONENTS :		EVALUATOR:
(1)	4-Chlorophenol; C ₆ H ₅ ClO; [106-48-9]	A. Vesala, Department of Chemistry and Biochemistry, University of Turku.
(2)	Water; H ₂ 0; [7732-18-5]	November 1979.

The earliest of the published works on the solubility of 4-chlorophenol in water is that of Sidgwick and Turner (1). Even though this work has been available for more than 50 years, it still serves as a useful reference for comparison of later results. No solubility value at 298.15 K has been reported as a measured value; the nearest measurements reported have been relatively far from room temperature (274 K and 339 K by Sidgwick and Turner) making an interpolation difficult.

Kuroda (2) reported an approximate solubility of the three isomeric monochlorophenols in water in the range of 2 percent. More support for the work of Sidgwick and Turner is given by the work of Blackman et al. (3) and that of Mulley and Metcalf (4). The data reported in these two works are in good agreement. For an assumed solution density of 1.0 g/cm^3 , a value of 27 g(1)/kg results from the work of Blackman et al. and a value of 26 g(1)/kg results from the work of Mulley and Metcalf.

In reviewing the work of Blackman et al. it should be stressed that the solubility was determined at a pH of 5.1, a condition being controlled by the addition of a phosphate buffer (a KH_2PO_4 solution). However, the effect of the buffer solution can be minimized since the pH of a saturated 4-chlorophenol solution can be assumed to be of the same magnitude as that of the buffer and the ionic effects can be assumed to be quite low (as reported by Robinson and Stokes (5), the pK for 4-chlorophenol in water is 9.4). Even so, one should remember that the pH does exert some influence on the solubility of protolytic solutes such as 4-chlorophenol and attention should be paid to agents in the solution which change its acidity. The data are shown in Figure 1.

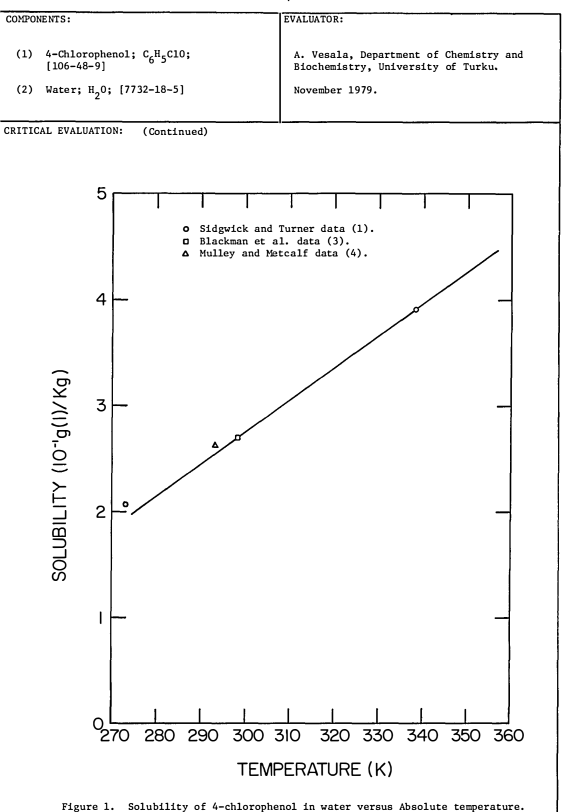
The solubility of 4-chlorophenol in water at normal temperatures has a reliable value as deduced from the three references. However, the mutual solubility of 4-chlorophenol and water as reported by Sidgwick and Turner over its entire temperature range must be considered tentative because of the lack of other supporting data.

The following solubility of 4-chlorophenol in water is tentative and based primarily on the reported value of Blackman et al.:

T/K	10mo1(1)/dm	10 ⁻¹ g(1)/kg	$10^{3}x(1)$
298.15	2.1	2.7	3.9

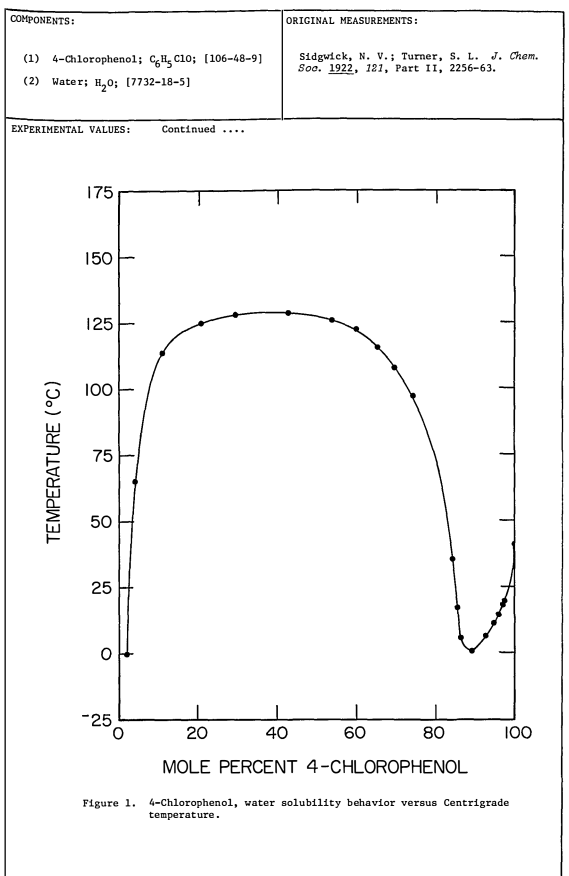
REFERENCES

- 1. Sidgwick, N. V.; Turner, S. L. J. Chem. Soc. 1922, 121, Part II, 2256-63.
- 2. Kuroda, T. cf. Chem. Zentralbl. 1926, I, 3610.
- 3. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.
- 4. Mulley, B. A.; Metcalf, A. D. Sci. Pharm. 1966, 2, 481-8.
- Robinson, R. A.; Stokes, R. H. "Electrolyte Solutions"; Butterworths: London, <u>1970</u>; p 533.



COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) 4-Chlorop	henol; C ₆ H ₅ ClO; [1 0; [7732–18–5]	06-48-9]	Sidgwick, N. V.; Turner, S. L. <i>J. Chem.</i> <i>Soc.</i> <u>1922</u> , <i>121</i> , Part II, 2256-63.
VARIABLES: Temperature: -(0.2 - 128.7°C		PREPARED BY: A. Vesala
EXPERIMENTAL VAL	-	- (-> (-	
t/°C		mol(1)/k	
-0.2			0.02953
0.5	8.892		68 5.2934
5.5	8.619	6.704	44 4.6656
6.2	9.248	7.193	37 6.3282
11.0	9.448	7.349	93 7.0576
14.2	9.570	7.444	42 7.5722
17.0	8.542	6.644	45 4.5086
18.0	9.682	7.531	13 8.1013
19.5	9.729	7.567	78 8.3419
35.5	8.402	6.535	56 4.2424
41.0	10.00	7.778	87 10.0
65.0	0.391	0.3041	0.05670
97.0	7.403	5.758	35 2.8545
			Continued
AUXILIARY		AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The solubility versus temperature measure- ments were made by the so called synthetic method which involved the mixing of known masses of solute and solvent and the testing of miscibility with temperature. A bracket- ing procedure was applied. In this method, two adjustable thermostats, one above and the other below the solubility temperature, were used. The contents of sample tubes in the two thermostat baths were observed to maintain one homogeneous and the other heterogeneous as the temperature interval between the two thermostats was reduced by successive steps.		·······	SOURCE AND PURITY OF MATERIALS:
		synthetic of known	C ₆ H ₅ ClO: Synthesized from 4-chloroaniline, melting point of the product 40.9°C.
		A bracket- his method, above and emperature, le tubes in served to other	H ₂ 0: Source and purity not specified.
			ESTIMATED ERROR:
			REFERENCES:

COMPONENTS: ORIGINAL MEASUREMENTS: (1) 4-Chlorophenol; C ₆ H ₅ Cl0; [106-48-9] Sidgwick, N. V.; Turner, S. L. J. Chem. (2) Water; H ₂ 0; [7732-18-5] Sidgwick, N. V.; Turner, S. L. J. Chem. t/°C 10 ⁻² g(1)/kg mol(1)/kg 10x(1) 107.7 6.936 5.3953 2.4083 113.8 1.066 0.82920 0.16446 115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. Measurements are shown graphically in Figure 1.
Soc. 1922 , 121 , Part II, $2256-63$.EXPERIMENTAL VALUES:Continued $t/^{\circ}C$ $10^{-2}g(1)/kg$ mol(1)/kg107.7 6.936 5.3953 113.8 1.066 0.82920 115.5 6.505 5.0600 122.4 5.962 4.6376 125.0 2.050 1.5946 125.8 5.349 4.1608 128.2 2.916 2.2683 128.7 4.257 3.3114 0.94104 a. Reported.b. Calculated by F. W. Getzen.
(2) Water; H_2O ; [7732-18-5] Soc. <u>1922</u> , 221, Part II, 2256-63. EXPERIMENTAL VALUES: Continued $t/^{\circ}C$ $10^{-2}g(1)/kg$ mol(1)/kg $10x(1)$ 107.7 6.936 5.3953 2.4083 113.8 1.066 0.82920 0.16446 115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Cetzen. 5.0000 5.0000
EXPERIMENTAL VALUES: Continued $t/^{\circ}C$ $10^{-2}g(1)/kg$ mol(1)/kg $10x(1)$ 107.7 6.936 5.3953 2.4083 113.8 1.066 0.82920 0.16446 115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. External contraction of the second contraction of the
t/°C $10^{-2}g(1)/kg$ mol(1)/kg $10x(1)$ 107.7 6.936 5.3953 2.4083 113.8 1.066 0.82920 0.16446 115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. $V. Getzen.$
t/°C $10^{-2}g(1)/kg$ mol(1)/kg $10x(1)$ 107.7 6.936 5.3953 2.4083 113.8 1.066 0.82920 0.16446 115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. $V. Getzen.$
113.81.0660.829200.16446115.56.5055.06002.0687122.45.9624.63761.7144125.02.0501.59460.34876125.85.3494.16081.3880128.22.9162.26830.54540128.74.2573.31140.94104a. Reported.b. Calculated by F. W. Getzen.
115.5 6.505 5.0600 2.0687 122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. F. W. Getzen. Ketzen.
122.4 5.962 4.6376 1.7144 125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. Example 1.200
125.0 2.050 1.5946 0.34876 125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. Image: Calculated by F. W. Getzen.
125.8 5.349 4.1608 1.3880 128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen. Example 100 (2000)
128.2 2.916 2.2683 0.54540 128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen.
128.7 4.257 3.3114 0.94104 a. Reported. b. Calculated by F. W. Getzen.
a. Reported. b. Calculated by F. W. Getzen.
b. Calculated by F. W. Getzen.
b. Calculated by F. W. Getzen.
Measurements are shown graphically in Figure 1.
Continued



COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 4-Chlorophenol; C ₆ H ₅ ClO; [106-48-9]	Mulley, B. A.; Metcalf, A. D. <i>Sci. Pharm.</i> <u>1966</u> , 2, 481-8.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁻¹ g(1)/dm ³ a 10mol(1)	$/dm^{3}b$ $10^{3}x(1)^{b}$
20 2.625 2.041	8 3.7705
- December 1	
a. Reported. b. Calculated by F. W. Getzen.	
AUXILIARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
The sample preparation and equilibration	C ₆ H ₅ C10: Commercial reagent, manufacturer
procedures were not specified. Saturated solutions were prepared and analyzed spec-	not specified, suitably purified until physical constants corre-
trophotometrically. Also, duplicate deter- minations were probably made by a synthetic	ponded with literature values. The reagent was dried in a desic-
method described in (1).	cator for two days before use.
	H ₂ 0: Source and purity not specified.
	_
	ESTIMATED ERROR:
	Solubility: <4% (estimated here on the basis of the deviations in
	the reported values).
	REFERENCES:
	1. Mulley, B. A.; Metcalf, A. D. J. Pharm.
	Pharmacol. <u>1956</u> , 8, 774.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 4-Chlorophenol; C ₆ H ₅ Cl0; [106-48-9]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	Aren. Blochem. Blophys. <u>1755</u> , 04(17, 55-71.
2	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	A. Vesala
EXPERIMENTAL VALUES:	
t/°C 10 ⁻¹ g(1)/dm ^{3 a} 10mol(1)	$/dm^{3 b}$ 10 ³ x(1) ^a
25 2.70 2.1	3.89
a. Calculated by F. W. Getzen.	
b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The samples were equilibrated in a thermostat	
bath with temporary shaking over periods of $3-4$ weeks. During the equilibration time,	source and purity not specified.
the pH values of the solutions were con-	H ₂ 0: Distilled.
trolled by dropwise addition of phosphate buffer solution. The analysis of the solute	_
concentration in the saturated samples was	
done spectrophotometrically either directly or by using proper colorizing agents.	
or by using proper cororizing agence.	
	ESTIMATED ERROR:
	Solubility: <5% (evaluated on the basis of
	the reported results of the two techniques of analysis).
	REFERENCES:

COMPONENTS:	EVALUATOR:	
(1) Fluorobenzene; C ₆ H ₅ F; [462-06-6]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.	
(2) Water; H ₂ O; [7732-18-5]	May 1979	

Four experimental measurements of the solubility of fluorobenzene in water have been reported in the literature (1-4), see Figure 1. However, only one single measurement of the solubility of water in fluorobenzene has been reported (5).

Some of the more recent data for the solubility of fluorobenzene in water is that of Nelson and Smit (4) in the temperature range between 278 and 318 K. However, despite equilibrium periods of 24 hours (which might not have been long enough), the measured solubilities are substantially lower than those found by earlier investigators. It is not possible to establish any shortcomings of the experimental procedures from the very brief description. No information was provided on the source and purity of materials used. Also, it was not indicated whether or not a water stripper had been employed for the analysis of the very dilute aqueous solutions by gas chromatography, or whether or not an internal standard had been used for the calibration of the gas chromatograph which employed a flame ionization detector. However, the authors agreed to re-examine their raw data in order to verify the reported values (6). Consequently, for the present evaluation, their results have not been considered for inclusion in the selected solubility values.

Jockers (3) reported the solubility of fluorobenzene in water at high temperatures and high pressures only. The objective of these measurements was to establish the phase equilibria and thermodynamics for fluid mixtures in which fluorobenzene and water are miscible in all proportions. The critical curve was also determined (7,8). There are no other solubility measurements in the temperature range between 546 and 562 K and in the pressure range between 96 x 10^5 and 114×10^5 Pa for comparison.

The two experimental determinations by Andrews and Keefer (1) at 298.15 K and Gross et al. (2) at 303.15 K are in good agreement with respect to the errors estimated by the authors. The recommended solubility has been established as the arithmetic mean of the two solubilities and temperatures.

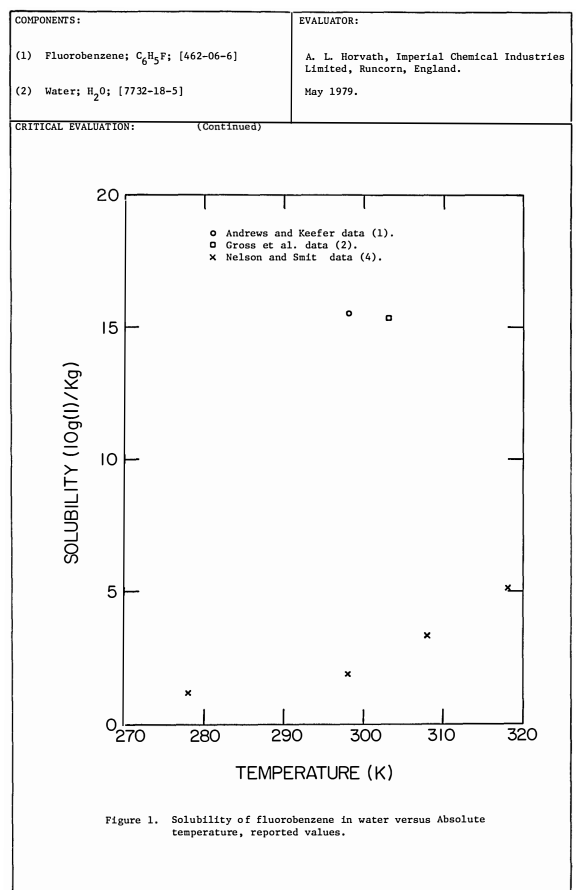
Only a single solubility measurement for the solubility of water in fluorobenzene has been reported in the literature (5). This experimental determination was done by Wing in 1956 (9) at Purdue University in partial fulfillment of the requirements for the Doctor of Philosophy degree. This measurement was made by an isotopic dilution technique using tritiated water as the tracer. There are no other reported values for comparison and selection.

Recommended solubility values are as follows (that for water in fluorobenzene is tentative):

т/к	$10^2 mol(1)/dm^3$	g(1)/kg	$10^4 x(1)$
300.65	1.59	1.54	2.89
T/K	10^2 mol(2)/dm ³	10g(2)/kg	$10^{3}x(2)$
298.15	1.75	3.08	1.65

COMP	COMPONENTS: EVALUATOR:	
(1)	Fluorobenzene; C ₆ H ₅ F; [462-06-6]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2)	Water; H ₂ 0; [7732-18-5]	May 1979
CRIT	ICAL EVALUATION: (Continued)	
REFERENCES		
1.	Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1950</u> , 72(7), 3113-6.	
2.	Gross, P. M.; Saylor, J. H.; Gorman, M. A. J. Am. Chem. Soc. <u>1933</u> , 55(2), 650-2.	
3.	Jockers, R., Ph.D. Dissertation, University of Bochum, Bochum, 1976.	
4.	Nelson, H. D.; Smit, J. H. SAfr. Tydskr. Chem. <u>1978</u> , 31(2), 76.	
5.	Wing, J.; Johnston, W.H. J. Am. Chem. Soc. <u>1957</u> , 79(4), 864-5.	
6.	Nelson, H. D.; Smit, J. H., Personal Communication, <u>1979</u> .	
7.	Götze, G.; Jockers, R.; Schneider, G. M., 4th Internat. Conf. Chem. Thermodyn. IUPAC, Montpellier, Aug. 26-30, <u>1975</u> , Part IV/9, 57-64.	
8.	Jockers, R.; Schneider, G. M. Ber. Bunsenges, Phys. Chem. <u>1978</u> , 83, 576-82.	
9.	Wing, J., Ph.D. Dissertation, Purdue University, Lafayette, 1956.	

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COMP ONENTS:	ORIGINAL MEASUREMENTS:
 (1) Fluorobenzene; C₆H₅F; [462-06-6] (2) Water; H₂O; [7732-18-5] 	Gross, P. M.; Saylor, J. H.; Gorman, M. A. J. Am. Chem. Soc. <u>1933</u> , 55(1), 650-2.
-	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/kg(2) ^a 10 ² mol(1)/	kg^{b} 10 ⁴ x(1) ^c
30 1.54 1.600	2.886
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: The saturated solution was prepared in a flask placed in a water thermostat bath. The samples were analyzed by means of a Zeiss combination liquid and gas interferometer as described in (1). A detailed description of the complete procedure has been included in a M.A. dis- sertation (2).	range 84.55° - 84.63°C. H ₂ 0: Distilled.
	<pre>ESTIMATED ERROR: Solubility: ±1.5%. Temperature: ±0.02 K. REFERENCES: 1. Gross, P. M.; Saylor, J. H. J. Am. Chem. Soc. <u>1931</u>, 53(5), 1744-51. 2. Gorman, M. A., M.A. Thesis, Duke Uni- versity, Durham, <u>1932</u>.</pre>

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Fluorobenzene; C ₆ H ₅ F; [462-06-6]	Andrews, L. J.; Keefer, R. M. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1950</u> , 72(7), 3113-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C g(1)/dm ^{3 a} 10 ² mol(1)/dm	$10^4 x(1)^{c}$
25.0 1.55 1.613	2.918
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
The water was saturated with fluorobenzene in a glass-stoppered Erlenmeyer flask ro- tated in a constant temperature bath for 20 hours. The saturated solution was extracted with n-hexane. The optical density of the extract was measured against a standard n- hexane blank with a Beckman spectrophotome- ter (1).	C ₆ H ₅ F: Eastman Kodak Co., commercial reagent, b.p. 84.4°C, fractionated before use. H ₂ O: Not specified.
	ESTIMATED ERROR:
	Solubility: ±10% (compiler). Temperature: ±0.1 K (compiler).
	<pre>REFERENCES: 1. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. 1949, 71(11), 3644-7.</pre>

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) Water; H ₂ O; [7732-18-5]	Wing, J.; Johnston, W. H. J. Am. Chem. Soc.
(2) Fluorobenzene; C ₆ H ₅ F; [462-06-6]	<u>1957</u> , <i>79(4)</i> , 864–5.
(27) Filorobenzene, 665F, [402-00-0]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10ml(1)/dm ^{3 a} 10 ² mol(1)	$/dm^{3 b} 10^{3}x(1)^{c}$
25.0 3.16 1.749	1.651
a. Reported. b. Calculated by F. W. Getzen.	
c. Calculated by compiler.	
AUXI LI ARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Tritiated water was equilibrated with 20 ml	H ₂ 0: Tracerlab Inc., tritiated water, used as received.
fluorobenzene for two hours in a flask placed in a constant temperature bath. The sample	Í
was stirred throughout the equilibration period. The concentration of the tritiated	C ₆ H ₅ F: Source not specified, chemical grade, redistilled before use.
water in the organic phase was determined	
by isotopic dilution. The tritium activities in the tritiated water were determined by	
the acetylene method (1,2). At least four	
independent experiments were done.	
The article was based upon work reported in a Ph.D. dissertation (2).	
	ESTIMATED ERROR: Solubility: ±2.8%.
	Temperature: ±0.02 K.
	REFERENCES :
	 Wing, J.; Johnston, W. H. Science 1955, 121, 674-6.
	2. Wing, J., Ph.D. Dissertation, Purdue
	University, Lafayette, <u>1956</u> .

Fluorobenzene

Fluoron	Jenzene	
COMPONENTS:	ORIGINAL MEASUREMENTS:	
 (1) Fluorobenzene; C₆H₅F; [462-06-6] (2) Water; H₂O; [7732-18-5] 	Jockers, R., Ph.D. Dissertation, University of Bochum, Bochum, <u>1976</u> , pp. 91-2.	
VARIABLES:	PREPARED BY:	
Temperature and pressure	A. L. Horvath	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ P/bar $10^{-2}g(1)/kg^{a}$	$mol(1)/kg^{b}$ $10x(1)^{c}$	
284.0 100 4.9130	5.1123 1.533	
289.5 114 5.6684	8.8985 1.970	
279.0 103 5.6684	8.8985 1.970	
273.5 96 5.6684	8.8985 1.970	
a. Calculated by compiler. b. Calculated by F. W. Getzen. c. Reported.		
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
The measurements were made using a high temperature, high pressure optical cell made from stainless steel with an internal stirrer. Predetermined mixtures of fluoro- benzene in water were introduced into the cell and homogenized at various tempera- tures and pressures. The phase transitions were observed through a sapphire window at the appropriate temperatures and pressures. Further details have been included in (1,2).	C ₆ H ₅ F: Fluka AG., Buchs, Schweiz, 99.5% pure, redistilled before use. H ₂ O: Boiled with KMnO ₄ and redis- tilled before use.	
	ESTIMATED ERROR: Solubility: ±1%. Temperature: ±0.5 K. Pressure: ±2 bar. REFERENCES: 1. Götze, G.; Jockers, R.; Schneider, G. M. 4th Inter. Conf. Chem. Thermodyn. IUPAC,	

2	0	9

FILUIU	Senzene 20
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Fluorobenzene; C ₆ H ₅ F; [462-06-6]	Nelson, H. D.; Smit, J. H. SAfr. Tydskr.
(2) Water: H ₂ 0; [7732-18-5]	Chem. <u>1978</u> , 31(2), 76.
2	
VARIABLES:	DDEDADED DV.
Temperature	PREPARED BY: A. L. Horvath
-	
 EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ³ mol(1)/kg	^b $10^{5}x(1)$ ^c
5 1.184 1.232	2.22
25 1.893 1.970	3.55
35 3.36 3.50	6.3
	9.6
45 5.12 5.33	9.0
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Water was saturated through the vapor phase	C ₆ H ₅ F: Not specified.
with fluorobenzene in a special flask (1) using a shaker in a thermostat bath for 24	H ₂ 0: Not specified.
hours. A gas chromatographic analysis of the samples was done by injection into a 5% Apiezon M stainless steel column with Celite as the supporter. The column length was 1.5 m and the temp. was 120°C. The chromatograph was equipped with a flame ionization detector. Three samples were analyzed from each flask.	2
	ESTIMATED ERROR:
	Solubility: ±1.9% (compiler).
	Temperature: ±0.1 K (compiler).
	REFERENCES :
	REFERENCES: 1. Nelson, H. D.; de Ligny, C. L. <i>Rec.</i> <i>Trav. Chim.</i> <u>1968</u> , <i>87</i> , 528-44.
	1. Nelson, H. D.; de Ligny, C. L. Rec.

COMPONENTS :	EVALUATOR:
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]	A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ 0; [7732-18-5]	January 1983

Five experimental determinations have been published on the solubility of iodobenzene in water in the temperature interval between 278 and 318 K (1-4,8), see Figure 1. The serious discrepancies between the various solubility measurements for iodobenzene in water are evident in Figure 1. The solubility of water in iodobenzene between 298 and 308 K has been reported in two publications (5,6), see Figure 2.

Some of the more recent data for the solubility of iodobenzene in water is that of Nelson and Smit (3) in the temperature range between 278 and 318 K. However, despite equilibrium periods of 24 hours (which might not have been long enough), the measured solubilities are substantially lower than those found by earlier investigators. It is not possible to establish any shortcomings of the experimental procedures from the very brief description. No information was provided on the source and purity of materials used. Also, it was not indicated whether or not a water stripper had been employed for the analysis of the very dilute aqueous solutions by gas chromatography, or whether or not an internal standard had been used for the calibration of the gas chromatograph which employed a flame ionization detector. However, the authors agreed to re-examine their raw data in order to verify the reported values (7). Consequently, for the present evaluation, their results have not been considered for inclusion in the selected solubility values.

The solubility value for iodobenzene in water reported by Gross et al. (2) is too high in relation to other reported values (1,4). While the equilibration time was 20 hours for measurements made by Andrews and Keefer (1), the time was 48 hours for those of Vesala (4). This equilibration time difference may have affected the reported results. For this reason, the higher solubility value given by Vesala is recommended, see Table 1. The remaining available data have been correlated relative to Absolute temperature using a normal polynomial equation:

$$S_1(g)1)/kg$$
 = 0.59293 - 4.8616 x $10^{-3}T$
+ 1.21754 x $10^{-5}T^2$ [1]

The values calculated from equation [1] for the saturation of iodobenzene in water in the range of temperatures between 283 and 318 K together with corresponding molarities and mole fractions are given in Table 1. Also, the solubility values calculated from equation [1] are shown in Figure 1 as a solid line along with the measured values.

For the solubility of water in iodobenzene, the agreement between the two solubilities reported by Jones and Monk (5) and by Wing and Johnston (6) at 298.15 K is quite good. At higher temperatures, measurements were made only by Jones and Monk (5); therefore, there are no other data for comparison. The data from both investigations (5,6) were correlated using the following equation:

$$\log_{10} x(2) = 0.792324 - 985.707/T$$
[2]

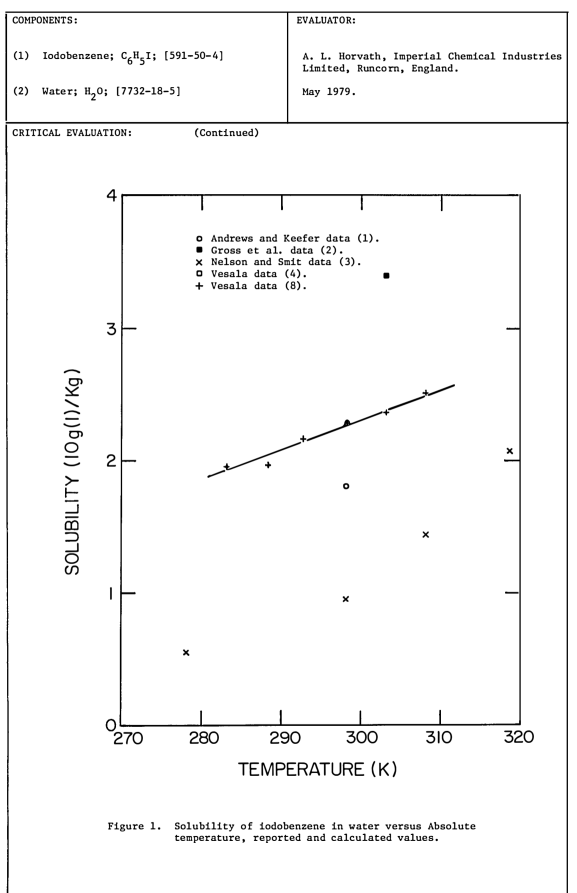
In this equation, x(2) is the mole fraction solubility of water in the water-iodobenzene system and T is the Absolute temperature. The calculated solubility values in the 283 to 308 K range are shown in Figure 2 along with the reported values.

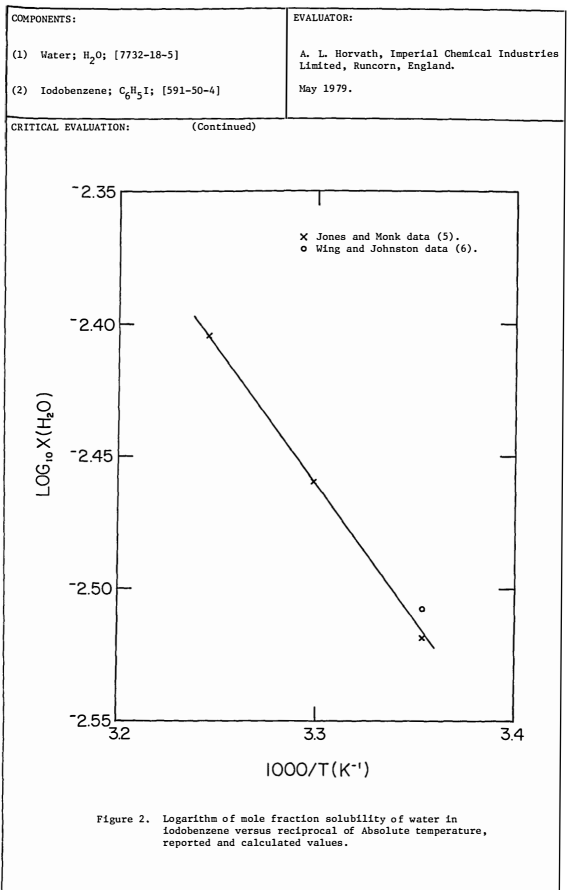
The calculated mole fraction values for the solubility of water in iodobenzene from equation [2] are included in Table 2 together with the corresponding molarities and g(2)/kg values in the temperature range between 293 and 313 K.

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COMPONENTS :		E	VALUATOR:
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]			A. L. Horvath, Imperial Chemical Industries Limited, Runcorn, England.
(2) Water; H ₂ O;	[7732-18-5]	j	January 1983
CRITICAL EVALUATIO	N: (Continued)		
Table 1. Solubil:	lty of Iodobenzene i	n Water.	
T/K	10 ⁴ mol(1)/dm ³		$x_{g} = 10^{5} x(1)$
283.15	9.43	1.93	1.70
288.15	9.94	2.03	1.79
293.15	10.5	2.14	1.89
298.15	11.0	2.26	1.99
303.15	11.6	2.38	2.10
308.15	12.2	2.51	2.22
313.15	12.9	2.64	2.34
318.15	13.5	2.79	2.46
Table 2. Solubility of Water in Iodobenzene. T/K $10^2 \text{mol}(2)/\text{dm}^3$ $10g(2)/\text{kg}$ $10^3 x(2)$			$k_{g} = 10^{3} x(2)$
293.15	2.42	2.38	2.69
298.15	2.74	2.71	3.06
303.15	3.10	3.08	3.47
308.15	3.49	3.48	3.92
313.15	3.90	3.91	4.41
		REFERI	ENCES
1. Andrews, L	J.; Keefer, R. J. J	. Am. Cher	m. Soc. <u>1950</u> , 72(7), 3113-6.
2. Gross, P. M.	Savlor, J. H.; Gor	man. M. A.	. J. Am. Chem. Soc. 1933, 55(2), 650-2.
 Gross, P. M.; Saylor, J. H.; Gorman, M. A. J. Am. Chem. Soc. <u>1933</u>, 55(2), 650-2. Nelson, H. D.; Smit, J. H. SAfr. Tydskr. Chem. <u>1978</u>, 31(2), 76. 			
,			
			y of Turku, Turku, <u>1973</u> .
	· · · · · · · · · · · · · · · · · · ·	-	
1			

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lodobenzene





COMPONENTS:	ORIGINAL MEASUREMENTS:		
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]	Gross, P. M.; Saylor, J. H.; Gorman, M. J.		
(2) Water; H ₂ O; [7732-18-5]	J. Am. Che	em. Soc. <u>1933</u> , 55(2),	650-2.
(2) water, "20, [//32-10-5]			
VARIABLES:	PREPARED B		
One temperature	A. L. Hory		
EXPERIMENTAL VALUES:			
2 3	Ъ	5 0	1
t/°C 10g(1)/kg(2) ^a 10 ³ mo1(1)/	kg 1	$10^{5}x(1)^{-1}$	
30 3.4 1.67		3.00	
a. Reported.			
b. Calculated by F. W. Getzen.c. Calculated by compiler.			
c. calculated by complicit.			
AUXILIARY	INFORMATIO	N	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND	D PURITY OF MATERIALS	:
The saturated solution was prepared in a	С6Н21:	Eastman Kodak Co.,	
thermostat water bath. The samples were analyzed by means of a Zeiss combination		fractionally befor range 84.55° - 84.	e use, b.p.
liquid and gas interferometer described in		Tange 04.55 - 04.	05 0.
(1). A detailed description of the complete	^H 2 ⁰ :	Distilled.	
procedure has been included in a M.A. thesis (2).			
	ESTIMATED	ERROR:	
	Solubility		
	-		
	Temperatur	re: ±0.02 K.	
	REFERENCES		
		, P. M.; Saylor, J. H 1931, 53(5), 1744-51,	
	Soc. <u>1931</u> , 53(5), 1744-51. 2. Gorman, M. A., M. A. thesis, Duke		
		rsity, Durham, <u>1932</u> .	_,

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]	Andrews, L. J.; Keefer, R. M. <i>J. Am. Chem.</i> <i>Soc.</i> <u>1950</u> , 72(7), 3113-6.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10g(1)/dm^{3} = 10^{4}mol(1)/dm^{3}$	10^{3} b $10^{5}x(1)$ c
25.0 1.8 8.82	1.59
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Water was saturated with iodobenzene in a glass-stoppered Erlenmeyer flask by ro- tating the flask in a constant temperature bath for 20 hours. The saturated solution was extracted with n-hexane for analysis. The optical density of the extract was measured against a n-hexane standard using a Beckman spectrophotometer (1).	SOURCE AND PURITY OF MATERIALS: C ₆ H ₅ I: Eastman Kodak Co., commercial reagent, b.p. 188.3°C, frac- tionated before use. H ₂ O: Not specified.
	ESTIMATED ERROR: Solubility: ±10% (compiler). Temperature: ±0.1 K (compiler). REFERENCES: 1. Andrews, L. J.; Keefer, R. M. J. Am. Chem. Soc. <u>1949</u> , 71(11), 3644-7.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H ₂ O; [7732-18-5]	Wing, J.; Johnston, W. H. J. Am. Chem.	
(2) Iodobenzene; C ₆ H ₅ I; [591-50-4]	Soc. <u>1957</u> , 79(4), 864-5.	
657		
VARIABLES:	PREPARED BY:	
One temperature	A. L. Horvath	
EXPERIMENTAL VALUES:		
t/°C 10m1(1)/dm ^{3 a} 10 ² mol(1)/d	$1m^{3 b} 10^{3}x(1)^{c}$	
25.0 5.03 2.784	3.107	
a. Reported.		
b. Calculated by F. W. Getzen.c. Calculated by compiler.		
AUXILIARY	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
Tritiated water was equilibrated with 20 ml	H ₂ 0: Tracerlab Inc., tritiated water,	
iodobenzene through stirring in a flask in a constant temperature water bath for	² used as received.	
two hours. The concentration of the tri-	C ₆ H ₅ I: Source not specified, chemical	
tiated water in the organic phase was de- termined by isotopic dilution. For the	grade, redistilled before use.	
tritiated water samples, the tritium acti-		
vities were determined by the acetylene method (1,2). At least four independent		
determinations were made. The article was		
based upon work reported in a Ph.D. dis- sertation (2).		
	ESTIMATED ERROR:	
	Solubility: ±1.6%.	
	Temperature: ±0.02 K.	
	REFERENCES :	
	1. Wing, J.; Johnston, W. H. Science	
1	<u>1955</u> , <i>121</i> , 674–5.	
	 Wing, J., Ph.D. Dissertation, Purdue University, Lafayette, <u>1956</u>. 	
1		

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Water; H_20 ; [7732-18-5] (2) Jodobonzeno: C.H.J: [591-50-6]	Jones, J. R.; Monk, G. B. J. Chem. Soc. <u>1963</u> , Part III, 2633-5.
(2) Iodobenzene; C ₆ H ₅ I; [591-50-4]	
VARIABLES:	PREPARED BY:
Temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10m1(1)/dm ³ (2) ^a 10 ² mol($1)/dm^{3}b$ $10^{3}x(1)^{c}$
25 4.9 2.7	1 3.03
30 5.6 3.0	9 3.47
35 6.35 3.5	3.940
a. Reported. b. Calculated by F. W. Getzen. c. Calculated by compiler.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE: Tritiated water was shaken with iodobenzene in 1:10 volume ratios for 4 hours in flasks placed in a water thermostat bath. The water content was determined by tritium assay of samples taken from the flasks. The count rates were determined using a typical liquid scintillatior solution technique.	SOURCE AND PURITY OF MATERIALS: H ₂ O: Tritiated. C ₆ H ₅ I: Source not known, laboratory grade, dried over CaCl ₂ and fractionally distilled before use.
	ESTIMATED ERROR: Solubility: ±5%. Temperature: ±0.5 K (compiler). REFERENCES:

lodobenzene

COMPONENTS :		ORIGINAL MEASUREMENTS:				
 Iodobenzene; C₆H₅I; [591-50-4] Water; H₂0; [7732-18-5] 		Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u> ,				
		-				
VARIA	BLES:			PREPARED BY	Y:	
	Temperat	ure		A. L. Horv	vath	
EXPER	IMENTAL V	ALUES:				
	t/°C	10g(1)/kg ^a	10 ³ mol(1)/kg	(2) ^b	$10^{5}x(1)^{a}$	
	10.0	1.9529	0.9574 ± 0.0	093 1	1.7248	
	15.2	1.9656	0.9636 ± 0.0	084 1	1.7360	
	19.6	2.1642	1.061 ± 0.0	33 1	1.9115	
	25.1	2.2845	1.120 ± 0.0	32 2	2.0177	
	30.0	2.3620	1.158 ± 0.0	26 2	2.0862	
	35.0	2.5109	1.231 ± 0.0	07 2	2.2177	
			AUXILIARY	INFORMATION	I	
METHOD/APPARATUS/PROCEDURE: The iodobenzene was mixed with water in a sealed flask (1) with the aid of a magnetic stirrer for 48 hours in a water bath at constant temperature. After the solution was filtered through a glass-wool plug, the solute was extracted with 2,2,4- trimethylpentene. The optical density of each sample was determined spectrophoto- metrically (2). Mean and standard devia- tions were calculated from three measure- ments.		SOURCE AND C ₆ H ₅ I: H ₂ O:	PURITY OF MAT Flika AG, pur used as recei Distilled, de degassed.	riss, >99% GLC, .ved.		
				ESTIMATED	ERROR:	
				Solubility	≠3.11%.	
				Temperatur	re: ±0.05 K.	
		J. Che 2. Waucho	s, F.; Gent, M 2m. Soc. <u>1963</u> ,	1.; Johnson, H. H. Part III, 2716-23. Stzen, F. W. J. Chem. 1), 38-41.		

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]	Vesala, A. Acta Chem. Scand. <u>1974</u> , 28A(8),
0.5	839-45.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	A. L. Horvath
EXPERIMENTAL VALUES:	
t/°C 10g(1)/kg ^a 10 ³ mol(1)/kg	(2) ^b $10^{5}x(1)^{a}$
25.0 2.2845 1.120	2.0177
a. Calculated by compiler.	
b. Reported.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
The iodobenzene was mixed with water in a	C ₆ H ₅ I: Commercial reagent of analytical grade distilled through a column
sealed flask (1) with the aid of a magnetic stirrer for 48 hours in a water bath at	resulting in a more than 99% pure
constant temperature. After the solution	sample.
was filtered through a glass-wool plug, the solute was extracted with 2,2,4-trimethyl-	H ₂ 0: Distilled, deionized, and
pentene. The optical density of each sam-	2 degassed.
ple was determined spectrophotometrically (2). Mean and standard deviations were	
established from five parallel determina-	
tions.	
The reported work was based upon a Ph.D.	ESTIMATED ERROR:
dissertation (3).	Solubility: ±2.7%.
	Temperature: ±0.05 K.
	REFERENCES: 1. Franks, F.; Gent, M.; Johnson, H. H. <i>J. Chem. Soc.</i> <u>1963</u> , Part III, 2716-23.
	2. Wauchope, R. D.; Getzen, F. W. J. Chem.
	Eng. Data <u>1972</u> , $17(1)$, 38-41.
	 Vesala, A., Ph.D. Dissertation, University of Turku, Turku, <u>1973</u>.

lodobenzene

COMPONENTS:			ORIGINAL MEASUREMENTS:
(1) Iodobenzene; C ₆ H ₅ I; [591-50-4]		-50-4]	Nelson, H. D.; Smit, J. H. SAfr. Tydskr.
(2) Water	; H ₂ 0; [7732-18-5]		Chem. <u>1978</u> , 31(2), 76.
	2		
VARIABLES:			PREPARED BY:
Temperatur	e		A. L. Horvath
EXPERIMENTAL	L VALUES:		
t/°C	10g(1)/kg ^a	10 ⁴ mol(1)/kg	^b $10^6 x(1)$ ^c
5	0.5492	2.692	4.85
25	0.9500	4.657	8.39
35	1.438	7.048	12.7
45.5	2.072	10.16	18.3
ъ. С	alculated by compile alculated by F. W. G		
c. R	eported.		
		AUXILIARY	INFORMATION
METHOD/APPA	RATUS/PROCEDURE:		SOURCE AND PURITY OF MATERIALS:
Water was saturated through the vapor phase with iodobenzene in a special flask (1)			C ₆ H ₅ I: Not specified.
using a sha	aker in a thermostat	bath for 24	H ₂ 0: Not specified.
the samples	as chromatographic was done by inject	ion into a 5%	
	tainless steel colu the supporter. The		
was 1.5 m a	and the temp. was 12 uph was equipped wit	0°C. The	
ionization	detector. Three sa		
anaryzeu II	com each flask.		
			ESTIMATED ERROR:
			Solubility: ±2% (compiler).
			Temperature: ±0.1 K (compiler).
			REFERENCES:
			 Nelson, H. D.; de Ligny, C. L. Rec. Trav. Chim. <u>1968</u>, 87, 528-44.

COMPONENTS:		EVALUATOR:	
(1) (2)	2,4,5,6-Tetrachloro-3-methylphenol; C ₇ H ₄ Cl ₄ O; [10460-33-0] Water; H ₂ O; [7732-18-5]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA July 1983.	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4,5,6-tetrachloro-3-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of KH_2PO_4 buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentration. A standardized curve obtained from measurements using known phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4,5,6-tetrachloro-3-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4,5,6-tetrachloro-3-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4,5,6-tetrachloro-3-methylphenol in water is tentative:

t/°C	$10^{3}g(1)/dm^{3}$	$10^5 mol(1)/dm^3$	$10^{7}x(1)$
25	6.15	2.5	4.52

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 2,4,5,6-Tetrachloro-3-methylphenol; C ₇ H ₄ Cl ₄ O; [10460-33-0]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature	
One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)/dm	$10^{3 b} 10^{7} x(1)^{a}$
25 6.15 2.5	4.52
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with	C ₇ H ₄ Cl ₄ O: Probably a commercial reagent.
KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated	H ₂ 0: Distilled water.
for an additional 2 weeks to assure satura- tion.	
Concentration was determined either directly by spectrophotometric techniques or by a	ESTIMATED ERROR:
colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to	Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques).
yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem.
	<u>1912,</u> 12, 239. 3. Folin, 0.; Ciocalteu, V. J. Biol. Chem. <u>1927</u> , 73, 627.

COMPONENTS:		EVALUATOR:	
(1)	2,4,6-Trichloro-3-methylphenol; C ₇ H ₅ Cl ₃ O; [551-76-8]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA	
(2)	Water; H ₂ 0; [7732-18-5]	July 1983.	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4,6-trichloro-3-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of KH_2PO_4 buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4,6-trichloro-3-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4,6-trichloro-3-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4,6-trichloro-3-methylphenol in water is tentative:

t/°C	10g(1)/dm ³	$10^4 mol(1)/dm^3$	$10^{6}x(1)$
25	1.12	5.3	9.58

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4,6-Trichloro-3-methylphenol; C₇H₅Cl₃0; [551-76-8]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm ³	$10^{6}x(1)^{a}$
25 1.12 5.3	9.58
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	C7 ^H 5 ^{Cl} 3 ⁰ : Probably a commerical reagent. H ₂ 0: Distilled water.
Concentration was determined either directly by spectrophotometric techniques or by a	ESTIMATED ERROR:
colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution to to an appropriately diluted solution to wield an appropriately diluted solution to	Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques).
yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, 0.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 3. Folin, 0.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.

COMPONENTS:	ORIGINAL MEASUREMENTS:	
(1) Water; H ₂ O; [7732-18-5]	Dreisbach, R. R. "Physical Properties of	
-	Chemical Compounds", Advances in Chemistry	
(2) 3,4-Dichlorotoluene; C ₇ H ₆ Cl ₂ ; [95-75-0]	Series No. 15; American Chemical Society: Washington, D. C., <u>1955</u> ; p. 140.	
	······································	
VARIABLES:	PREPARED BY:	
One temperature	F. W. Getzen	
EXPERIMENTAL VALUES:		
t/°C 10 ² g(1)/kg(2) ^a 10 ³ mol(1)	$/kg^{b}$ 10 ⁴ x(1) ^b	
30 2.6 1.44	2.32	
a. Reported.		
b. Calculated by compiler.		
	INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:	
No experimental details were available.	C7H6C12: Dow Chemical Co., purified by distillation before use.	
	H ₂ O: Distilled.	
·		
	ESTIMATED ERROR: Solubility: <100% (compiler).	
1		
	Temperature: ±1 K (compiler).	
	REFERENCES :	

COMPONENTS :	EVALUATOR:
 (1) 2,6-Dichloro-4-methylphenol; C₇H₆Cl₂0; [2432-12-4] (2) Water; H₂0; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA July 1983.

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,6-dichloro-4-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2PO_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,6-dichloro-4-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,6-dichloro-4-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,6-dichloro-4-methylphenol in water is tentative:

t/°C	10g(1)/dm ³	10^{3} mol(1)/dm ³	$10^{5}x(1)$
25	6.73	3.8	6.87

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

2,0-Dichioro-4	
COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,6-Dichloro-4-methylphenol; C₇H₆Cl₂0; [2432-12-4]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mo1(1)/dm ³	$10^5 x(1)^a$
25 6.73 3.8	6.87
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
	INFORMATION SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with KH_2PO_4 at a pH of 5.1 in the glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura-	C ₇ H ₆ Cl ₂ O: Probably a commercial reagent. H ₂ O: Distilled water.
tion.	

COMPONENTS:	EVALUATOR:
 (1) 2,4-Dichloro-6-methylphenol; C₇H₆Cl₂0; [1570-65-6] (2) Water; H₂0; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4-dichloro-6-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\mathrm{KH}_2\mathrm{PO}_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4-dichloro-6-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4-dichloro-6-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4-dichloro-6-methylphenol in water is tentative:

t/°C	$10g(1)/dm^{3}$	10^{3} mol(1)/dm ³	$10^{5}x(1)$
25	2.83	1.6	2.89

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch, Biochem. Biophys. 1955, 54(1), 55-71.

2,4-Dichioro-0	
COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 2,4-Dichloro-6-methylphenol; C ₇ H ₆ Cl ₂ O: [1570-65-6]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ³ mol(1)/dm	$10^{5}x(1)^{a}$
25 2.83 1.6	2.89
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with KH_2PO_4 at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure saturation.	
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	<pre>ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem. 1912, 12, 239.</pre>

COMPONENTS:	ORIGINAL MEASUREMENTS:
 (1) 4-Bromotoluene; C₇H₇Br; [106-38-7] (2) Water; H₂0; [7732-18-5] 	Dreisbach, R. R. "Physical Properties of Chemical Compounds", Advances in Chemistry Series No. 15; American Chemical Society: Washington, D. C., <u>1955</u> ; p. 153.
VARIABLES:	PREPARED BY:
One temperature	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C $10g(1)/kg(2)^{a}$ $10^{4}mol(1)/kg^{b}$ $10^{5}x(1)^{b}$	
25 1.1 6.43	1.16
a. Reported. b. Calculated by compiler.	
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
No experimental details were available.	C ₇ H ₇ Br: Dow Chemical Co., purified by distillation before use.
	H ₂ O: Distilled.
	2
	ESTIMATED ERROR:
	Solubility: <100% (compiler).
	Temperature: ±1 K (compiler).
	REFERENCES:

COMPONENTS:	EVALUATOR:
<pre>(1) 4-Chloro-2-methylphenol; C7H7Cl0; [1570-64-5]</pre>	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA
(2) Water; H ₂ 0; [7732-18-5]	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-2-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of KH_2PO_4 buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-2-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-2-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-2-methylphenol in water is tentative:

t/°C	g(1)/dm ³	10^2 mol(1)/dm ³	$10^4 x(1)$
25	6.84	4.8	8.73

REFERENCES

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 4-Chloro-2-methylphenol; C ₇ H ₇ C10; [1570-64-5]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C g(1)/dm ^{3 a} 10 ² mol(1)/dm ³	b $10^4 x(1)^{a}$
25 6.84 4.8	8.73
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by mixing the solute with distilled water buffered with KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion. Concentration was determined either directly by spectrophotometric techniques or by a	
colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	<pre>ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorime- tric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, 0.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 3. Folin, 0.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.</pre>

COMPONENTS :	EVALUATOR:
 4-Chloro-3-methylphenol; C₇H₇Cl0; [59-50-7] Water; H₂O; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-3-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2PO_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-3-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-3-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-3-methylphenol in water is tentative:

t/°C	g(1)/dm ³	$10^2 mol(1) / dm^3$	$10^{4}x(1)$
25	3.99	2.8	5.08

REFERENCES

1. Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. 1955, 54(1), 55-71.

HBTPW-I

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COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 4-Chloro-3-methylphenol; C7H7Cl0; [59-50-7]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	
One temperature	PREPARED BY:
One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C g(1)/dm ^{3 a} 10 ² mol(1)/dm ^{3 1}	$10^4 x(1)^{a}$
25 3.99 2.8	5.08
a. Calculated by compiler.	
b. Reported value measured at pH 5.1.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the	C7H7C10: Probably a commercial reagent.
solute with distilled water buffered with KNI_PO, at a pH of 5.1 in a glass-stoppered	H ₂ O: Distilled water.
bottle suspended in a thermostat bath at	
25°C. The sample was shaken periodically for	
2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually	
about 1 week), the sample was equilibrated	
for an additional 2 weeks to assure satura-	
Concentration was determined either directly	
by spectrophotometric techniques or by a colorimetric method based upon that given by	ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of
(1) in which suitable colorizing agents (2,3)	the reported results of the two
were added either to the saturated solution or to an appropriately diluted solution to	analytical techniques).
yield an optical density proportional to the	
solute concentration. Concentration was ob-	REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric
tained as the average of three replicate measurements.	Methods of Analysis", Vol. II; Chapman
	and Hall: London, England, <u>1936</u> .
	2. Folin, O.; Denis, W. J. Biol. Chem. 1912, 12, 239.
	3. Folin, 0.; Ciocalteu, V. J. Biol Chem.
	<u>1927</u> , 73, 627.

COMPONENTS:	EVALUATOR:
(1) 2-Chloro-6-methylphenol; C ₇ H ₇ ClO; [87-64-9]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA
(2) Water; H ₂ O; [7732-18-5]	July 1983.

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2-chloro-6-methylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2-chloro-6-methylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2-chloro-6-methylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2-chloro-6-methylphenol in water is tentative:

t/°C	g(1)/dm ³	10 ² mol(1)/dm ³	$10^4 x(1)$
25 3.56	2.5	4.53	
		PEFFRENCES	

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 2-Chloro-6-methylphenol; C ₇ H ₇ C10: [87-64-9]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/dm ³ a 10 ² mol(1)/dm ³	$10^4 x(1)^{a}$
25 3.56 2.5	4.53
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXTLTARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by mixing the solute with distilled water buffered with KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	SOURCE AND PURITY OF MATERIALS: C7 ^H 7 ^{ClO:} Probably a commercial reagent. H ₂ O: Distilled water.
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 3. Folin, O.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.

COMPONENTS :	ORIGINAL MEASUREMENTS:	
(1) 1,3-bis(trifluoromethyl)benzene; C ₆ H ₄ (CF ₃) ₂ ; [402-31-3]	Strachan, A. N.; Field, J. P.; Fleming, K. A. Int. Solvent Extr. Conf., (Proc.), Vol. 2, <u>1980</u> , page paper 80-2, 6 pp.	
(2) Water; H ₂ 0; [7732-18-5]	, <u></u> , page paper of 1 , o pp.	
VARIABLES:	PREPARED BY:	
One temperature	F. V. Getzen	
EXPERIMENTAL VALUES:		
$t/^{\circ}C$ $10^{2}g(1)/dm^{3}a$ $10^{4}mol(1)/dm^{3}a$	$dm^{3 b}$ 10 ⁶ x(1) ^a	
25 4.090 1.91	3.451	
a. Calculated by compiler. b. Reported.		
	INFORMATION	
METHOD/APPARATUS/PROCEDURE: An excess of 1,3-bis(trifluoromethyl)ben- zene was stirred with 100 ml water in a closed flask in a thermally regulated bath (±0.1°C) for 30 min. After phase separa- tion and standing for several hours, two samples of known amounts were removed from the aqueous phase and extracted with hexane. The absorbances of the extracted samples were measured (265 nm) with a Pye Unicam SP 30 spectrophotometer. Hexane solutions of the solute in known concentrations were used to establish the extinction coeffi- edeet (266 the 3-biller 2). The extender	SOURCE AND PURITY OF MATERIALS: C ₆ H ₄ (CF ₃) ₂ : Fluorochem Limited 97-100%; redistilled and middle cut used. H ₂ O: Distilled.	
cient (246 dm ³ mol ⁻¹ cm ⁻¹). The procedure was repeated to give a total of four con-	ESTIMATED ERROR:	
centration measurements.	Solubility: ±3% (authors).	
	Temperature: ±0.1 K (authors).	
	REFERENCES :	

COMPONENTS:	EVALUATOR:
<pre>(1) 2,4,6-Trichloro-3,5-dimethylphenol; C₈H₇Cl₃0; [6972-47-0]</pre>	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA
(2) Water; H ₂ O; [7732-18-5]	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4,6-trichloro-3,5-dimethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2PO_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4,6-trichloro-3,5-dimethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4,6-trichloro-3,5-dimethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4,6-trichloro-3,5-dimethylphenol in water is tentative:

t/°C	$10^{3}g(1)/dm^{3}$	10^5 mol(1)/dm ³	$10^{7}x(1)$
25	4.96	2.2	3.98

REFERENCES

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4,6-Trichloro-3,5-dimethylphenol; C₈H₇Cl₃0; [6972-47-0]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C = 10^{3}g(1)/dm^{3}a = 10^{5}mol(1)/dm^{3}a$	$m^{3 b} 10^{7} x(1)^{a}$
25 4.96 2.2	3.98
a Calculated by compiler	
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD /APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with	C ₈ H ₇ Cl ₃ 0: Probably a commercial reagent.
KII ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at	H ₂ 0: Distilled water.
25°C. The sample was shaken periodically for	
2 days with appropriate adjustment of pll to 5.1. After the pH became stabilized (usually	
about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura-	
tion.	
Concentration was determined either directly	
by spectrophotometric techniques or by a colorimetric method based upon that given by	ESTIMATED ERROR:
(1) in which suitable colorizing agents (2,3) were added either to the saturated solution	Solubility: <5% (evaluated on the basis of the reported results of the
or to an appropriately diluted solution to	two analytical techniques).
yield an optical density proportional to the solute concentration. Concentration was ob-	REFERENCES:
tained as the average of three replicate measurements.	 Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman
	and Hall: London, England, 1936. 2. Folin, O.; Denis, W. J. Biol. Chem.
	<u>1912, 12, 239.</u>
	 Folin, 0.; Ciocalteu, V. J. Biol. Chem. 1927, 73, 627.

COMPONENTS :	EVALUATOR:
 (1) 2,4-Dichloro-6-ethylphenol; C₈H₈Cl₂O; [24539-94-4] (2) Water; H₂O; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4-dichloro-6-ethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4-dichloro-6-ethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4-dichloro-6-ethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4-dichloro-6-ethylphenol in water is tentative:

t/°C	10g(1)/dm ³	10^{3} mol(1)/dm ³	$10^{5}x(1)$
25	2.48	1.3	2.35

REFERENCES

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4-Dichloro-6-ethylphenol; C₈H₈Cl₂O; [24539-94-4]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
 VARIABLES:	PREPARED BY:
One temperature	R. W. Ostan
One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ 10g(1)/dm ³ a 10 ³ mol(1)/dm	$3 b 10^5 x(1)^a$
25 2.48 1.3	2.35
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the	C ₈ H ₈ Cl ₂ O: Probably a commerical reagent.
solute with distilled water buffered with KH_2PO_4 at a pH of 5.1 in a glass-stoppered	H ₂ 0: Distilled water.
bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	2 2
Concentration was determined either directly	
by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to	ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques).
yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem. 1912, 12, 239.
	 Folin, 0.; Ciocalteu, V. J. Biol. Chem. 1927, 73, 627.

COMPONENTS:	EVALUATOR:
 (1) 4-Chloro-2,5-dimethylphenol; C₈H₉Cl0; [1124-06-7] (2) Water; H₂0; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-2,5-dimethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-2,5-dimethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-2,5-dimethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-2,5-dimethylphenol in water is tentative:

t/°C	g(1)/dm ³	10^{2} mol(1)/dm ³	$10^{3}x(1)$
25	8.93	5.7	1.04

REFERENCES

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COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 4-Chloro-2,5-dimethylphenol; C ₈ H ₉ ClO; [1124-06-7]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/dm ^{3 a} 10 ² mol(1)/dm ^{3 b}	$10^{3}x(1)^{a}$
25 8.93 5.7	1.04
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD / APPARATUS / PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with KH2P04 at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	C ₈ H ₉ ClO: Probably a commercial reagent. H ₂ O: Distilled water.
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	<pre>ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, 1936. 2. Folin, O.; Denis, W. J. Biol. Chem. 1912, 12, 239. 3. Folin, O.; Ciocalteu, V. J. Biol. Chem. 1927, 73, 627.</pre>

COMP	ONENTS:	EVALUATOR:
(1)	4-Chloro-2,6-dimethylphenol; C ₈ H ₉ ClO; [1123-63-3]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North
(2)	Water; H ₂ 0; [7732-18-5]	Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-2,6-dimethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentration. A standardized curve obtained from measurements using known phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-2,6-dimethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-2,6-dimethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-2,6-dimethylphenol in water is tentative:

t/°C	10g(1)/dm ³	10^{3} mo1(1)/dm ³	$10^{5}x(1)$
25	5.17	3.3	5.97

REFERENCES

COMPONENTS :	ORIGINAL MEASUREMENTS:
(1) 4-Chloro-2,6-dimethylphenol; C ₈ H ₉ ClO; [1123-63-3]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10g(1)/dm^{3}$ a $10^{3}mol(1)/dm$	1^{3} b $10^{5}x(1)$ a
25 5.17 3.3	5.97
a. Calculated by compiler. b. Reported value measured by pH 5.1.	
AU XILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE :	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the	C ₈ H ₉ Cl0: Probably a commercial reagent.
solute with distilled water buffered with KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure saturation.	H ₂ 0: Distilled water.
Concentration was determined either directly	
by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the	ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques).
solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 3. Folin, O.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.

COMPONENTS :	EVALUATOR:
 (1) 4-Chloro-3,5-dimethylphenol; C₈H₉Cl0; [88-04-0] (2) Water; H₂0; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-3,5-dimethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentration. A standardized curve obtained from measurements using known phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-3,5-dimethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-3,5-dimethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-3,5-dimethylphenol in water is tentative:

t/°C	g(1)/ém ³	10^2 mol(1)/dm ³	$10^4 x(1)$
25	3.45	2.2	3.99

REFERENCES

COMPONENTS:	ORIGINAL MEASUREMENTS:
(1) 4-Chloro-3,5-dimethylphenol; C ₈ H ₉ ClO; [88-04-0]	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ g(1)/dm ³ a 10 ² mol(1)/dm ³	^b $10^4 x(1)^{a}$
25 3.45 2.2	3.99
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXI LI ARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by mixing the solute with distilled water buffered with Kll_2PO_4 at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	<pre>ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, 0.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 3. Folin, 0.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.</pre>

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СОМ	PONENTS:	EVALUATOR:
(1) (2)	2,4-Dichloro-6-propylphenol; C ₉ H ₁₀ Cl ₂ 0; [91399-12-1] Water; H ₂ 0; [7732-18-5]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4-dichloro-6-propylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of KH_2PO_4 buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4-dichloro-6-propylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4-dichloro-6-propylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4-dichloro-6-propylphenol in water is tentative:

t/°C	10g(1)/dm ³	$10^4 mol(1)/dm^3$	$10^{6}x(1)$
25	1.00	4.9	8.85

REFERENCES

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4-Dichloro-6-propylphenol; C9^H10^{C1}2⁰; [91399-12-1]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10g(1)/dm ^{3 a} 10 ⁴ mol(1)/dm	1^{3} b $10^{6}x(1)$ a
25 1.00 4.9	8.85
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by mixing the solute with distilled water buffered with KH ₂ PO ₄ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	 ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u>. 2. Folin, O.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. 2. Folin, O.; Ciocalteu, V. J. Biol. Chem. <u>1927</u>, 73, 627.

COMPO	DNENTS:	EVALUATOR:
(1)	4-Chloro-3-methyl-5-ethylphenol; C ₉ H ₁₁ ClO; [1125-66-2] Water; H ₂ O; [7732-18-5]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-3-methyl-5-ethylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2P0_4$ buffer. Concentration determinations were accomplished both by direct spectrophotometric techniques and, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentration. A standardized curve obtained from measurements using known phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. The agreement between the measurements of concentration using the two methods of analysis was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-3-methyl-5-ethylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-3-methyl-5-ethylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-3-methyl-5-ethylphenol in water is tentative:

t/°C	10g(1)/dm ³	10^3 mol(1)/dm ³	$10^{5}x(1)$
25	3.58	2.1	3.80

REFERENCES

COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 4-Chloro-3-methyl-5-ethylphenol; C9^H11^{Cl0}; [1125-66-2]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES: 10 ³ mol(1)/dm ³ a	
t/°C Spectrophotometry Colorim	etry
25 2.0 2.2	
Values derived from average measured solu	ubilities: ^b
$t/^{\circ}C$ $10g(1)/dm^{3}$ $10^{3}mol(1)/dm^{3}$	$10^5 x(1)$
25 3.58 2.1	3.80
a. Reported value measured at pH 5.1 b. Calculated by compiler.	
	INFORMATION
METHOD/APPARATUS/PROCEDURE: Equilibrium was established by mixing the solute with distilled water buffered with KH ₂ PO ₄ at a pII of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pII to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion. Concentration was determined both by direct	SOURCE AND PURITY OF MATERIALS: C ₉ H ₁₁ ClO: Probably a commercial reagent. H ₂ O: Distilled water.
Spectrophotometric techniques and by a colori- metric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was obtained as the average of three replicate measurements. Agreement between the two methods was within an experimental error of 5%.	Solubility: <5% (evaluated on the basis of

COMPONENTS:	EVALUATOR:
 2,4-Dichloro-6-butylphenol; C₁₀H₁₂Cl₂O; [91399-13-2] Water; H₂O; [7732-18-5] 	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4-dichloro-6-butylphenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\mathrm{KH}_2\mathrm{P0}_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentration. A standardized curve obtained from measurements using known phenol concentrations was required for the determinations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4-dichloro-6-butylphenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4-dichloro-6-butylphenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4-dichloro-6-butylphenol in water is tentative:

t/°C	$10^2 g(1) / dm^3$	10^4 mol(1)/dm ³	$10^{6}x(1)$
25	5.26	2.4	4.34

REFERENCES

COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4-Dichloro-6-butylphenol; C₁₀^{II}₁₂Cl₂O; [91399-13-2]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pII: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10 ² g(1)/dm ^{3 a} 10 ⁴ mol(1)/d	$m^{3 b}$ 10 ⁶ x(1) ^a
25 5.26 2.4	4.34
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with KH_2PO_4 at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for 2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure saturation.	C ₁₀ H ₁₂ Cl ₂ O: Probably a commercial reagent. H ₂ O: Distilled water.
Concentration was determined either directly by spectrophotometric techniques or by a colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) Were added either to the saturated solution or to an appropriately diluted solution to yield an optical density proportional to the solute concentration. Concentration was ob- tained as the average of three replicate measurements.	<pre>ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques). REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Hall: London, England, 1936. 2. Folin, 0.; Denis, W. J. Biol. Chem. 1912, 12, 239. 3. Folin, 0.; Ciocalteu, V. J. Biol. Chem. 1927, 73, 627.</pre>

COMPO	DNENTS:	EVALUATOR:
(1)	2,4-Dichloro-6-(phenylmethyl)-phenol; C ₁₃ H ₁₀ Cl ₂ O; [19578-81-5]	F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA
(2)	Water; H ₂ 0; [7732-18-5]	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 2,4-dichloro-6-(phenylmethyl)-phenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2PO_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 2,4-dichloro-6-(phenylmethyl)-phenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 2,4-dichloro-6-(phenylmethyl)-phenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 2,4-dichloro-6-(phenylmethyl)-phenol in water is tentative:

t/°C	$10^{3}g(1)/dm^{3}$	10^{5} mol(1)/dm ³	$10^{7}x(1)$
25	5.82	2.3	4.16

REFERENCES

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COMPONENTS:	ORIGINAL MEASUREMENTS:
<pre>(1) 2,4-Dichloro-6-(phenylmethyl)-phenol; C₁₃H₁₀Cl₂O; [19578-81-5]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ 0; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
t/°C 10 ³ g(1)/dm ^{3 a} 10 ⁵ mol(1)/d	$m^{3 b} 10^{7} x(1)^{a}$
25 5.82 2.3	4.16
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY INFORMATION	
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the	
solute with distilled water buffered with KH ₂ PO ₄ at a pH of 51. in a glass-stoppered	C ₁₃ H ₁₀ Cl ₂ O: Probably a commercial reagent.
bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for	H ₂ 0: Distilled water.
2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually	
about 1 week), the sample was equilibrated	
for an additional 2 weeks to assure satura- tion.	
Concentration was determined either directly	
by spectrophotometric techniques or by a colorimetric method based upon that given by	ESTIMATED ERROR: Solubility: <5% (evaluated on the basis of
(1) in which suitable colorizing agents (2,3) were added either to the saturated solution	the reported results of the two analytical techniques).
or to an appropriately diluted solution to yield an optical density proportional to the	
solute concentration. Concentration was ob- tained as the average of three replicate	REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric
measurements.	Methods of Analysis", Vol. II; Chapman and Hall: London, England, <u>1936</u> .
	 Folin, 0.; Denis, W. J. Biol. Chem. 1912, 12, 239.
	 Folin, 0.; Ciocalteu, V. J. Biol. Chem. 1927, 73, 627.

COMPONENTS:		EVALUATOR:
(1) 4-Chloro-3,5-dimethyl- phenol; C ₁₃ H ₁₅ Cl0; [18		F. W. Getzen, Chemistry Department, North Carolina State University, Raleigh, North Carolina, USA
(2) Water; H ₂ O; [7732-18-5	1	

Only one measurement, that of Blackman et al. (1), was found in the literature for the solubility of 4-chloro-3,5-dimethyl-2-(phenylmethyl)-phenol in water. The determination was part of a study to provide some insight into the mode of biological action of substituted phenols as related to differences in physical properties.

Saturation was established at 298.15 K over a 3-4 week period, a reasonably sufficient time interval for saturation equilibrium. The system pH was adjusted to 5.1 by addition of small amounts of $\rm KH_2PO_4$ buffer. Concentration determinations were accomplished either by direct spectrophotometric techniques or, indirectly, by colorimetry using added colorizing reagents. The colorimetry method was based upon a standard procedure in which suitable colorizing agents were added either to the saturated phenol solution or to appropriately diluted solutions to yield an optical density proportional to the phenol concentrations.

The concentration was obtained from three replicate measurements as an average value. Where both analytical procedures were used for the same solute, the agreement between the measurements of concentration was within 5 percent.

No mention was made of further purification of the solute reagent (probably a commercial product). The water was probably distilled. The ionic effects of the buffer upon the saturation value can be taken as minimal. However, it must be noted that pH does affect the solubility of protolytic solutes such as 4-chloro-3,5-dimethyl-2-(phenylmethyl)-phenol. In this work, it can be assumed that the reported solubility refers, within experimental error, to the saturation value at the pH which exists for the saturated solution of 4-chloro-3,5-dimethyl-2-(phenylmethyl)-phenol alone, in water.

The uncertainty in the single reported value can be assumed to be as large as \pm 5 percent. The following solubility value for 4-chloro-3,5-dimethyl-2-(phenylmethyl)-phenol in water is tentative:

t/°C	$10^2 g(1) / dm^3$	10^{5} mol(1)/dm ³	$10^{7}x(1)$
25	1.11	5.0	9.03

REFERENCES

4-Chloro-3,5-dimethyl-	2-(phenylmethyl)phenol 257
COMPONENTS :	ORIGINAL MEASUREMENTS:
<pre>(1) 4-Chloro-3,5~dimethyl~2-(phenylmethyl)- phenol; C₁₃H₁₅Cl0; [1867-85-2]</pre>	Blackman, G. E.; Parke, M. H.; Garton, G. Arch. Biochem. Biophys. <u>1955</u> , 54(1), 55-71.
(2) Water; H ₂ O; [7732-18-5]	
VARIABLES:	PREPARED BY:
One temperature One pH: 5.1	F. W. Getzen
EXPERIMENTAL VALUES:	
$t/^{\circ}C$ $10^{2}g(1)/dm^{3}$ a $10^{5}mol(1)/dm^{3}$	$m^{3 b}$ 10 ⁷ x(1) ^a
25 1.11 5.0	9.03
a. Calculated by compiler. b. Reported value measured at pH 5.1.	
AUXILIARY	INFORMATION
METHOD/APPARATUS/PROCEDURE:	SOURCE AND PURITY OF MATERIALS:
Equilibrium was established by mixing the solute with distilled water buffered with	C ₁₃ H ₁₅ ClO: Probably a commercial reagent.
$\rm KH_2PO_4$ at a pH of 5.1 in a glass-stoppered bottle suspended in a thermostat bath at 25°C. The sample was shaken periodically for	H ₂ 0: Distilled water.
2 days with appropriate adjustment of pH to 5.1. After the pH became stabilized (usually about 1 week), the sample was equilibrated for an additional 2 weeks to assure satura- tion.	
Concentration was determined either directly by spectrophotometric techniques or by a	ESTIMATED ERROR:
colorimetric method based upon that given by (1) in which suitable colorizing agents (2,3) were added either to the saturated solution or to an appropriately diluted solu- tion to yield an optical density proportional	Solubility: <5% (evaluated on the basis of the reported results of the two analytical techniques).
to the solute concentration. Concentration was obtained as the average of three repli- cate measurements.	REFERENCES: 1. Snell, F. D.; Snell, C. T. "Colorimetric Methods of Analysis", Vol. II; Chapman and Wall: London England 1936
	 and Hall: London, England, <u>1936</u>, Folin, O.; Denis, W. J. Biol. Chem. <u>1912</u>, 12, 239. Folin, O.; Clocalteu, V. J. Biol. Chem.
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