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DATA ACQUISITION FOR ENVIRONMENTAL TRANSPORT AND FATE SCREENING FOR COMPOUNDS OF INTEREST TO THE OFFICE OF EMERGENCY AND REMEDIAL RESPONSE

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Office of Health and Environmental Assessment Washington DC 20460 DATA ACQUISITION FOR ENVIRONMENTAL TRANSPORT AND FATE SCREENING FOR COMPOUNDS OF INTEREST TO THE OFFICE OF EMERGENCY AND REMEDIAL RESPONSE

Ву

H. M. Jaber, W. R. Mabey, A. T. Liu,T. W. Chou, H. L. Johnson, T. Mill,R. T. Podoll, and J. S. Winterle

SRI International 333 Ravenswood Avenue Menlo Park, CA 94025

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Project Officer
Lee A. Mulkey
Environmental Research Laboratory
U.S. Environmental Protection Agency
Athens, GA 30613

Technical Project Monitor
Gregory Kew
Exposure Assessment Group
Office of Health and Environmental Assessment
Office of Research and Development
U.S. Environmental Protection Agency
Washington, DC 20460

OFFICE OF HEALTH AND ENVIRONMENTAL ASSESSMENT
OFFICE OF RESEARCH AND DEVELOPMENT
U.S. ENVIRONMENTAL PROTECTION AGENCY
WASHINGTON, DC 20460

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FOREWORD

The Exposure Assessment Group of EPA's Office of Research and Development has three main functions: 1) to conduct exposure assessments; 2) to review assessments and related documents; and 3) to develop guidelines for Agency exposure assessments. The activities under each of these functions are supported by and respond to the needs of the various EPA program offices. This project was undertaken to gather data for use in assessing environmental transport and fate of chemicals for the Office of Emergency and Remedial Response (OERR) and therefore falls under the first function.

The Office of Emergency and Remedial Response is required to determine "reportable quantities" (RQs) for various chemicals under Sections 101(14) and 102 of the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA). After promulgation of regulations listing RQs, releases to the environment of amounts of these chemicals in excess of the corresponding RQ must be reported to a designated Federal organization. The toxicity of each chemical is one of several factors considered in determining an appropriate RQ. Since the knowledge of the transport and fate of these chemicals in the environment is important in determining their potential toxicity it is imperative to gather basic data to evaluate transport and fate.

James W. Falco, Director Exposure Assessment Group

ABSTRACT

Physical properties, equilibrium, and kinetic constants for evaluating the transformation and transport in aquatic systems for organic chemicals of interest to the Environmental Protection Agency have been obtained from the literature and calculated from theoretical or empirical relations. Values for selected physical properties such as melting point, boiling point, vapor pressure, water solubility, and octanol/water partitioning, and for rate constants such as hydrolysis, microbial degradation, photolysis, and oxidation are listed for each chemical along with the source of the data. Values are reported in units suitable for use in a current aquatic fate model. A discussion of the empirical relationships between water solubility, octanol/water partition coefficients, and partition coefficients for sediment and biota is presented.

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SECTION 1

INTRODUCTION

PURPOSE

Decisions on possible regulatory action or cost-effective remedial measures for toxic chemicals require an understanding of environmental and human risks associated with the manufacture, use, and disposal of chemicals. Part of the risk assessment requires the best scientific information about what the concentration of a chemical is in the environment. In the absence of reliable and extensive monitoring data, the concentration of a chemical can be estimated using one of several fate models and data for the individual processes that may be dominant for that chemical. These data may be measured in the laboratory, obtained from literature sources, or estimated using appropriate structure-activity relationships (SARs) or correlation methods. These data used with environmental parameters in a mathematical model constitute the process modeling approach (Baughman and Burns, 1980; Baughman and Lassiter, 1978; Smith et al., 1977; Mill, 1978).

The Environmental Protection Agency's Office of Emergency and Remedial Repsonse (OERR) is required to determine "reportable quantities" (RQs) for various chemicals under Sections 101(14) and 102 of the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA). After promulgation, releases to the environment of amounts of these chemicals in excess of the corresponding RQ must be reported to a designated Federal organization (usually the U.S. Coast Guard). The toxicity of each chemical is one of several factors considered in determining an appropriate RQ. Earlier, scientists in the Office of Health and Environmental Assessment (OHEA) summarized the toxicological literature on these compounds for OERR. Since the

transport and fate of these chemicals and their persistence in the environment are also important factors in determining their potential toxicity, Work Assignment 14 provides for a literature search for 10 fundamental physical and chemical parameters for 82 compounds not previously investigated in this fashion by the Agency. This information will be analyzed to predict potential environmental transport and fate.

These data are to be used in a screening assessment to decide what chemicals clearly would not persist in aquatic environments because of exceptional reactivity and what environments may be of particular concern because of dominant volatilization or sorption processes. This information will also be used to decide what data gaps exist and what particular process data need to be obtained for subsequent and more detailed assessments.

The data are made available in this report with the expectation that they may be of interest in other assessment efforts. Use of the data in the context of other assessments requires that each user understand the sources and limitations of the data. Each user must decide what additional data are required for the particular assessment. Any user of these data must particularly recognize that some values were estimated by SRI staff with expertise in the process of interest, and that considerable subjective judgment was applied for some of the estimates. Such judgments based on even crude analogies are indeed valuable and acceptable in screening level evaluations. In cases where even expert judgment cannot be used to provide a value, no value was entered. Users of these data are encouraged to conduct more intensive literature searches or to consult other knowledgeable scientists to augment or supplant data in this report.

In this report, "process data" are defined as data relating to rate constants, equilibrium constants, or physical properties that describe the intrinsic processes the chemicals may undergo independent of environmental influences. "Environmental parameter" in this report refers to properties or data that describe (or are a function of) the environment.

BACKGROUND

The processes that can be important for transforming or transporting a chemical in an aquatic environment are shown in Figure 1.1. The following discussion summarizes the mathematical basis for the process modeling approach applied to aquatic systems: (1) the evaluation of rates of loss of chemical due to transformations and volatilization processes, (2) the influence of sorption processes on the rates of loss of chemical, and (3) the prediction of concentration and half-life of chemical in the aquatic environment, including terms for input of chemical, dilution, and flow out of the environment. This discussion assumes that sorption to particulates in the environment is not kinetically controlled (i.e., sorption equilibrium is attained instantaneously).

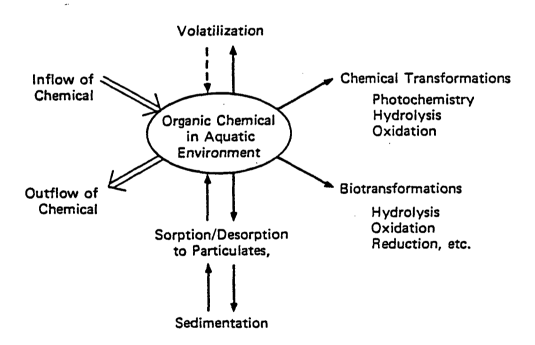


FIGURE 1.1 TRANSPORT AND TRANSFORMATION PROCESSES IN AQUATIC ENVIRONMENTS

EVALUATION OF CHEMICAL LOSS RATES

The rate of loss of a chemical due to the above transformation processes and volatilization, $R_{\rm T}$, is given by the sum of the rates of the individual processes, $R_{\rm i}$, according to the equation

$$R_{T} = \Sigma R_{i} = \Sigma k_{i}'[E_{i}][C]$$
 (1.1)

where k_i ' is the rate constant for the i-th process, $[E_i]$ is an environmental parameter that is kinetically important for the i-th process, and [C] is the concentration of the chemical. The calculations of R_i for individual processes from environmental parameter and process data are discussed in subsequent sections. The important environmental parameters for each process have been reviewed, and the use of the parameters in the calculations of environmental transformation rates has been discussed in detail by Baughman and Burns (1980), Mill (1978), and Smith et al. (1977).

The above expression for R_T assumes that the loss of chemical is first order in the chemical concentration, as certainly must be the case at the highly dilute concentrations expected in the environment. Equation (1.1) also requires that the rate of loss of chemical due to any one process R_1 is first order in the environmental parameter term E_1 ; R_1 is then considered as following overall second-order kinetic behavior. If it is assumed that the low concentration of chemical in the environment has no significant effect on the environment (for example, does not change pH, biomass, dissolved oxygen, etc.) and that the environmental parameter, E_1 , is constant over a specific region and time period, the term k_1 ' $[E_1]$ can be expressed as a simple pseudofirst-order rate constant, k_1 , and then

$$R_{T} = [\Sigma k_{i}][C] = k_{T}[C]$$
 (1.2)

or

$$k_{T} = \Sigma k_{i} \tag{1.3}$$

where \mathbf{k}_{T} is the overall pseudo-first-order rate constant for loss of chemical due to transformation and volatilization. The half-life for

loss of chemical due to these processes is then given by

$$t_{1/2} = \ln 2/k_T$$
 (1.4)

INFLUENCE OF SORPTION

In addition to losses of chemicals due to these transformation and volatilization processes, sorption to particulates can also reduce the concentrations of chemicals in aquatic systems. These particulates may be either suspended sediments or biotic in origin and the particulates may eventually be deposited into benthic sediments. The suspended or benthic sediment may later serve as a source of chemical from sorptiondesorption equilibrium as the chemical in solution volatilizes or undergoes transformation in the water column. If biotransformation does not occur in biota (such as bacteria, algae, and fish), the chemical may be released back into solution when the organism dies and decomposes. The understanding of chemical transformation when the chemical is sorbed onto particulates is inadequate to predict or measure the rates of such reactions for use in modeling. Therefore, the following discussion assumes that no transformations occur on particulates and that sorption is completely reversible and rapid in comparison with transformations that occur in solution.

The partitioning of a chemical between particulates (sediment or biota) and water at the low concentrations of chemicals usually found in the environment can be expressed as a partition coefficient $K_{\rm B}$:

$$k_p = C_s/C_w \tag{1.5}$$

where C_s and C_w are the equilibrium concentrations of chemical on sediment and in water, respectively (Baughman and Lassiter, 1978; Smith and Bomberger, 1982).

By convention, K_p is unitless when C_s is in units that are equivalent to C_w (i.e., C_s is in g chemical/g particulate and C_w is in g chemical/g water). In this discussion, $[C_w]$ will be defined in these weight units and [C] will be defined in molecular units (moles L^{-1}); because 1 g water is approximately 1 mL, it follows that $[C] = 10^3 [C_w][MW]^{-1}$

where MW is the molecular weight of chemical. Note that [C] and [C_W] can be used interchangeably in expressions such as equation (1.2) because first-order rate constants are concentration independent, but the rate of loss term, R, is of course defined in units corresponding to $[C_W]$ or [C].

For a chemical in aqueous solution containing particulates, the chemical is equilibrated between the water and particulate P according to the relation

$$C + P \longrightarrow C-P$$
 (1.6)

and the partition coefficient can be rewritten as

$$K_{p} = \frac{[C-P]}{[C_{w}][P]}$$
 (1.7)

where [C-P] is the mass of sorbed chemical per unit solution volume and [P] is the mass of sorbing particulate per unit solution volume. The mass balance of chemical in the solution-sediment system is given by

$$[C_T] = [C-P] + [C_w]$$
(1.8)

where $[C_T]$ is the total mass of chemical in a unit solution volume of water containing [P] grams of particulate. Combining equations (1.7) and (1.8) then gives the fraction of the total chemical dissolved in solution:

$$\frac{[C_w]}{[C_T]} = \frac{1}{K_p[P] + 1} \tag{1.9}$$

Baughman and Lassiter (1978) have pointed out that, given the relationship shown in equation (1.9), the fraction of chemical in solution may be quite high in spite of a large K_p value because the sediment or biota loading, [P], is often low in aquatic systems (i.e., $K_p[P] < 1$).

The concentration of chemical in solution $[C_w]$ in the presence of a particulate-water system is then given by

$$\begin{bmatrix} C_{\mathbf{w}} \end{bmatrix} = \frac{\begin{bmatrix} C_{\mathbf{T}} \end{bmatrix}}{\begin{bmatrix} P \end{bmatrix} K_{\mathbf{p}} + 1} \tag{1.10}$$

Substituting equation (1.10) into equation (1.2) for the rate of loss of chemical then gives

$$R_{T} = \frac{k_{T}C_{T}}{\lceil P \rceil K_{p} + 1} \tag{1.11}$$

This relationship shows that unless transformation on particulate is as fast as or faster than that in solution, the net effect of sorption will be to reduce the overall rate of loss of chemical from the aquatic system. From equation (1.11), it also follows that the half-life of the chemical is given by

$$t_{1/2} = \frac{(\lceil P \rceil K_p + 1) \ln 2}{k_T}$$
 (1.12)

The process modeling approach is then a valuable tool in risk assessments. Although values of $t_{1/2}$ or $C_{\rm w}$ can be manually calculated, the calculations are more easily done using computer programs. One such computer model is EXAMS, which allows the user to choose environmental parameters and is able to accommodate chemicals when several processes compete to be the important fate pathway. Computer models also allow for sophisticated and environmentally realistic dynamic models to be used rather than assuming steady-state conditions.

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SECTION 2

DEFINITIONS OF PROCESSES AND SOURCES OF DATA

BASIS FOR DERIVATION OF DATA

The data on chemicals given in this report were obtained from the literature and from calculations based on theory, SARs, or empirical calculations. In general, the physical properties of a chemical are functions of the molecular structure as an entity; that is, the elemental composition, spatial relationships and size, molecular weight, and functional groups of the molecule all may contribute to the property of the chemical. In contrast, the chemical or biological reactivity of a molecule is usually caused by selected functional groups in the molecular structure, and the functional group may undergo transformation with sometimes only minor changes in the total structure of the molecule.

The individual processes that chemicals may undergo can then be classified and evaluated according to specific physical properties or the reactive functional groups that these chemicals may have in common. The basis for the empirical correlations between $K_{\rm ow}$ and $S_{\rm w}$ is discussed in Section 4. These constants describe equilibrium processes for the chemical between water and a second (organic) phase. Similarly, the volatilization of a chemical can be evaluated in terms of Henry's constants, which are functions of vapor pressure and water solubility.

The reactivity of a chemical can be classified according to select functional groups in the molecular structure. For evaluations of hydrolysis reactions, chemicals are classified as carboxylic acid esters $(-CO_2R)$, carboxylic acid amides $(-CONH_2)$, alkyl halides (R-X), and phosphoric acid esters $((RO)_3PO)$, to name only a few. Data for hydrolysis of a chemical can often be estimated by analogy to another chemical with a similar functional group or calculated by more formal

procedures using linear free-energy relationships such as the Taft equation, the Hammett equation, or other such correlations (Mill, 1979; Wolfe et al., 1978 and 1980).

Chemical oxidation rate constants can be calculated by evaluating the reaction of an oxidant at a particular type of carbon-hydrogen bond (i.e., hydrogen abstraction process) or at an olefinic bond. No SAR or correlation method is available for predicting a direct photolysis rate constant except by analogy to other chemicals, which is often unreliable because of the complex chemistry of photoexcited states.

For this report, data obtained from calculations involving theory, SARs, or empirical correlations have been clearly identified so that the user can recognize the source of such data and can recalculate data using current or improved procedures.

The following briefly describes the environmental processes and the process data important in aquatic fate assessments. The process data are discussed in the order that they appear on the data sheets. The sources of the process data are also discussed.

CHEMICAL NAME, CHEMICAL ABSTRACTS SERVICE REGISTRY NUMBER, AND MOLECULAR WEIGHT

The names of the chemicals used on the data sheets are those as given on the original EPA list. The Chemical Abstracts Service (CAS) number has been obtained mainly from the original EPA list. Handbooks, catalogs, or the CAS were used when necessary. The molecular weight (MW) has been calculated from the molecular formula. Although the MW is not used for environmental assessments, it is required for conversion of units from ppm to molar units (M). The MW has also been used to calculate the molecular weight/oxygen ratio.

WATER SOLUBILITY

Water solubility (S_w) data are required for calculating Henry's constant and for calculating other partition coefficients using the correlation equations discussed in Section 4. Values of S_w (ppm or mg L^{-1}) were calculated from K_{OW} using a correlation equation developed by Yalkowsky and Valvani (1980).

For organic pollutants that are liquid in their pure state at 25°C

$$\log S_w = -1.08 \log K_{ow} + 3.70 + \log MW$$
 (2.1)

where MW is the molecular weight of the pollutant (g mole⁻¹). For organic pollutants that are solid in their pure state at 25°C

$$\log S_{w} = -1.08 \log K_{ow} + 3.70 + \log MW - (\frac{\Delta S_{p}}{1360}) (mp - 25)$$
 (2.2)

where mp is the melting point of the pollutant (°C) and ΔS_F is the entropy of fusion of the pollutant (cal mol⁻¹deg⁻¹). If ΔS_F is not known, it may be approximated by

$$\Delta S_{R} \sim 13.6 + 2.5 (n - 5)$$
 (2.3)

where n is the number of flexible atoms (i.e., atoms not involved in double bonds, triple bonds, or part of a ring structure) in the pollutant molecule, other than hydrogen. If n is less than 5, n-5 is set equal to zero.

For solids that had no literature melting points available, $S_{\rm w}$ was calculated using the equation for liquids. This results in a maximum $S_{\rm w}$ and should be used only for a screening risk assessment.

MELTING AND BOILING POINT

These data are not used directly in aquatic fate assessments, but they show in which phase (gas, liquid, solid) the pure chemical is found under environmental conditions. Boiling point data are cited for 760 torr (1 atmosphere) unless otherwise noted. The melting point should be used in the calculation of water solubility from octanol/water partition coefficient (K_{OW}) data for compounds that are solids above 25°C.

VAPOR PRESSURE

The vapor pressure P_{v} (torr) of an organic chemical is, in itself, a qualitative or relative measure of the volatility of the chemical in

its pure state and can be used to calculate the Henry's constant used in volatilization rate constant calculations. Unless otherwise specified, the P_{ν} values listed are at 25°C.

Vapor pressure data not found in the literature were calculated using procedures described by Grain (1982). The method uses a modification of the Watson correlation to express the temperature dependence of ΔH_{ν} such that

$$\Delta H_{v} \approx \Delta H_{vb} \left[3 - 2(T/T_{B})\right]^{m}$$
 (2.4)

where ΔH_{v} is the heat of vaporization at temperature T, ΔH_{vb} is the heat of vaporization at the normal boiling point, and m is a constant that depends upon the physical state. Substitution in the Clausius-Clapeyron equation and integration results in an expression with adjustable parameters that depend on the molecular structure, functional groups, and physical state at the temperature of interest. With further modification, the method can also be used to extrapolate vapor pressures from one temperature to another.

This procedure has an estimated maximum error of 7.1% over the pressure range 10-760 torr, 50% over 10^{-3} -10 torr, and 200% below 10^{-3} torr. The average error is <50%, which is often less than the range of $P_{\rm w}$ s found in the literature.

MOLECULAR WEIGHT TO OXYGEN RATIO

The molecular weight to oxygen ratio is used to estimate the volatilization rate constant for a chemical (Mabey et al. 1983). The ratio was calculated from the molecular weights of the chemical and molecular oxygen (the latter is 32 g/mole).

OCTANOL/WATER PARTITION COEFFICIENT

The octanol/water partition coefficient $K_{\rm ow}$ has been used in medical and environmental science as a measure of the hydrophobicity/hydrophilicity of chemicals (Hansch and Leo, 1979; Kenaga and Goring, 1978). The $K_{\rm ow}$ values in this report were used to calculate $S_{\rm w}$; $K_{\rm ow}$ values also are useful for estimating sediment and biota partitioning

coefficients (see Section 4). The calculation of K_{OW} from structural features of the molecule is also discussed in Section 4.

HYDROLYSIS RATE CONSTANTS

Hydrolysis refers to reaction of a chemical with water, usually resulting in the introduction of a hydroxyl function into a molecule and loss of a leaving group -X:

$$R-X + H_2O \longrightarrow ROH + HX$$
 (2.5)

The hydrolyses of some classes of compounds are catalyzed by acid or base, and therefore the hydrolysis rates of these chemicals in the environment can be pH dependent. The subject of hydrolysis in aquatic systems has been reviewed in detail by Mill et al. (1982), and an extensive compilation of hydrolysis data was published in a review by Mabey and Mill (1978).

The rate of hydrolysis of a compound at a specific pH value is given by the equation

$$R_{H} = k_{h}[C] = (k_{A}[H^{+}] + k_{N} + k_{B}[OH^{-}])[C]$$
 (2.6)

where $k_{\rm h}$ is the first-order rate constant for hydrolysis at the pH, $k_{\rm A}$ and $k_{\rm B}$ are second-order rate constants for acid- and base-promoted hydrolyses respectively, and $k_{\rm N}$ is the first-order rate constant for the pH-independent, neutral hydrolysis process. Using the autoprotolysis equilibrium expression

$$[H^+][OH^-] = K_w$$
 (2.7)

equation (2.6) can be rewritten as

$$k_h = k_A[H^+] + k_N + \frac{k_B^K w}{[H^+]}$$
 (2.8)

Equation (2.8) shows that k_h will depend on the pH of the aquatic system and on the relative values of k_A , k_B , and k_N . At present, no reliable information shows that hydrolysis rates in aquatic environments will be catalyzed by species other than $[H^+]$ or $[OH^-]$.

The hydrolysis rate constants k_A , k_B , and k_N used to calculate k_h as a function of pH are described below along with the source codes for calculating or estimating the values of the rate constants.

ACID-PROMOTED HYDROLYSIS RATE CONSTANT

The acid-promoted hydrolysis rate constant k_A (M⁻¹h⁻¹) is for the acid-promoted hydrolysis of a chemical. In regions where only k_A contributes to hydrolysis (i.e., $k_A[H^+] >> k_N = k_B[OH^-]$), k_h will decrease by a factor of 10 for each 1-unit increase in pH.

BASE-PROMOTED HYDROLYSIS RATE CONSTANT.

The base-promoted hydrolysis rate constant k_B (M^{-l}h^{-l}) is for the base-promoted (OH⁻) hydrolysis of a chemical. In regions where only k_B contributes to hydrolysis, k_h will increase by a factor of 10 for each l-unit increase in pH.

NEUTRAL-HYDROLYSIS RATE CONSTANT.

The neutral-hydrolysis rate constant k_N (h^{-1}) is for the pH-independent hydrolysis of a chemical. Data or sources pertaining to the hydrolysis of the organic chemicals have been entered in the data sheets in several ways. When a chemical structure had no hydrolyzable functional groups, NHFG was entered. When chemical hydrolysis occurs only at extreme pH values or temperatures or with catalysts not available in aquatic environments, HNES was entered (hydrolysis not environmentally significant). Other data for hydrolysis are referenced or are based on analogy to similar chemicals.

MICROBIAL DEGRADATION RATE CONSTANT

Biotransformations are undoubtedly important processes for degradation of chemicals in aquatic environments, resulting in hydrolysis, oxidation, and reduction of the chemical structure to ultimately produce carbon dioxide and water. The complex factors influencing the biotransformation of a chemical include pH, temperature, dissolved oxygen, available nutrients, other organic chemicals (synthetic or naturally occurring) that may serve as cometabolites or

alternative energy sources, and the populations and types of organisms capable of transforming the chemical. For most assessments, the initial biotransformation step is of prime importance (i.e., removal of the specific chemical from the environment). However, the biotransformation process is still too complex to be used to reliably predict a biotransformation rate constant using theoretical approaches such as those available for chemical and physical processes.

Maki et al. (1980) recently reviewed some of the aspects of the measurement of biotransformation rates and the use of such data. The rates of biotransformation are complex functions of chemical concentration and microbial biomass. However, at the typical concentrations of a chemical in the environment (< 1 ppm), the rates may be expected to follow second-order kinetics because they are first order in chemical kinetics and first order in biomass kinetics. Furthermore, the microorganism growth due to consumption of the chemical may not be significant; therefore, the rates of biotransformation are pseudo-first-order as a function of the chemical concentration.

The biotransformation data given in this report were estimated for the approach described by Baughman et al. (1980), in which the rate of biotransformation of a chemical, $R_{\rm B}$, is given by the expression

$$R_B = -k_b[B][C]$$
 (2.9)

where k_b is a second-order rate constant for biotransformation of a chemical by bacteria of population [B] in the solution phase of the water column. When k_b is given in mL cell⁻¹ h^{-1} , the units of [B] are in cell mL⁻¹. Because data for k_b were not available for most chemicals covered by this report, the rate constants were estimated solely for use in aquatic fate modeling by EPA. These data were estimated using on relative rates of biodegradation of the chemicals as reported in literature, structural analogies, and judgment of SRI staff with expertise in biotransformation studies. These data have been estimated and appropriate caution should be exercised in the use of the data.

PHOTOLYSIS RATE CONSTANT

The direct photolysis rate constants k_p (h^{-1}) for most of the chemicals cannot be estimated because of insufficient spectral and quantum yield data. For chemicals where no light absorption occurs above the solar cutoff (300 nm), the rate constant can be considered as zero, and therefore photolysis is not environmentally relevant (NER).

In cases where the chemical is expected to photolyze in the environment but no data are available, no value is entered. Similarly, no value is entered if nothing is known about a chemical's photolytic reactivity. No data for indirect photolysis of chemicals is provided in this report except that which results from oxidation processes (see next section).

OXIDATION RATE CONSTANTS

Chemical oxidation of organic chemicals in aquatic environments may be caused by several different oxidants, among which are singlet oxygen $(^{1}O_{2})$, alkyl peroxyl radical $(RO_{2} \cdot)$, alkoxy radical $(RO \cdot)$, or hydroxyl radical $(\cdot OH)$. The source of these oxidants is primarily photochemical, but because the oxidants react with chemicals in their ground state, and oxidation therefore does not involve the photochemistry of the chemical itself, oxidations are reasonably considered as discrete processes apart from photochemistry. Each oxidant has a unique reactivity toward organic moieties, and the relative and absolute concentrations of these oxidants will vary with environmental parameters, such as concentration and origin of humic-fulvic materials and sunlight intensity.

Literature information classifies reported data on oxidation of organic chemicals by oxy radicals such as RO_2 • and 1O_2 . The laboratory study conducted by Mill et al. (1982) using natural waters indicates that RO_2 • radical concentrations of $\sim 1 \times 10^{-9}$ M may be present in the surface waters of sunlit water bodies. Oxidation reactions initiated by RO_2 • include the following:

$$RO_2 \cdot + -C-H \longrightarrow RO_2H + -C \cdot$$
 (2.10)

$$RO_2 \cdot + C = C \longrightarrow RO_2 - C - C \cdot$$
 (2.11)

$$RO_2$$
 + ArOH RO_2 H + ArO • (2.12)

$$RO_2 \cdot + ArNH_2 \longrightarrow RO_2H + ArNH$$
 (2.13)

Of these reactions, the last two are quite rapid in aquatic environments $(t_{1/2} < \text{several days})$, whereas the others are slower and usually will not be important for most chemicals.

Zepp et al. (1978) have shown that $^{1}O_{2}$ can be formed at $\approx 1 \times 10^{-12}$ M concentrations in sunlit natural waters. The most important reactions for $^{1}O_{2}$ with organic chemicals are those involving reaction with olefinic moieties (Ranby and Rabek, 1978).

$$^{1}O_{2} + C = C \xrightarrow{CH_{2}^{-}} - \overset{!}{C^{-}C^{-}C^{-}C^{+}}$$

$$^{-}C^{-}C = CH$$

$$^{-}CH_{2}^{-} - CH_{2}^{-} - CH_{2}^{$$

Some rate constants for ${}^{1}\mathrm{O}_{2}$ and $\mathrm{RO}_{2}^{}\cdot$ are listed in a review by Mill (1980).

The rate of loss of organic chemicals \mathbf{R}_{OX} by oxidation is

$$R_{OX} = k_{RO_2} \cdot [RO_2 \cdot] [C] + k_1 \begin{bmatrix} 1 & 0 \\ 0 & 2 \end{bmatrix} [C] + k_{OX} [OX] [C] \qquad (2.15)$$
 where k_{OX} and $[OX]$ are the rate constants and concentration values for other unspecified oxidants. Only data for the second-order rate constants $k_{RO_2} \cdot \text{ and } k_1$ have been estimated for this report. When two rate constants are given on the data sheets, the second-order rate

constants should be multiplied by their respective oxidant concentrations to determine which of the first-order rate constant values is larger, and that rate constant should be used for an assessment.

Apart from a direct measurement of a rate constant at a specific temperature (which is rare), most rate constants in this report were obtained either from extrapolation of a rate constant for the organic chemical measured at another temperature or from a correlation of structure with reactivity as discussed below.

RATE CONSTANT FOR OXIDATION BY ALKYL PEROXYL RADICAL

Because many chemicals on the list of chemicals of concern have several kinds of reactive centers for oxidation by RO2., the overall rate constant k_{RO_2} $(M^{-1}h^{-1})$ was obtained by first calculating the individual rate constants for each reactive site and then summing these rate constants. For example, acrolein has two reactive sites: (1) addition to the double bond and (2) H-atom transfer from the carbonyl

$$RO_{2} \cdot + CH_{2} = CHCHO \xrightarrow{k_{1}} RO_{2}C - \dot{C} - CHO$$

$$RO_{2} \cdot + CH_{2} = CHCHO \xrightarrow{k_{2}} CH_{2} = CH\dot{C}O + RO_{2}H$$

$$(2.16)$$

$$RO_2$$
 + CH_2 = $CHCHO$ CH_2 = CH_2 + RO_2 H (2.17)

$$k_{RO_2} = k_1 + k_2$$
 (2.18)

When one oxidation process was found to be fast, the important oxidant was listed and the other reactions were ignored. When there were more than one -CH bond of a given kind, the rate constant was multiplied by the number of similar -CH bonds to give the correct total rate constant for oxidation of that CH-bond.

Two procedures were used to calculate individual $k_{\mbox{\scriptsize OX}}$ values for $\mathrm{RO}_2^{}$ reactions. In the first, when a structure was analogous to another chemical structure with a measured rate constant at a similar temperature, the measured rate constant was used directly (Hendry et al., 1974). (The -CHO bond in acrolein is an example.) The second procedure, used most often, is based on SARS established by Howard and coworkers for H-atom transfer (Korcek et al., 1972) and addition to double bonds (Howard, 1972), as shown below.

For the H-atom transfer reaction

$$\log k_{RO_2} = 18.96 - 0.2[D(R-H)]$$
 (2.19)

where D(R-H) is the bond dissociation energy of the CH-bond.

For the RO2 addition to double bonds

$$\log k_{RO_2} = [16.54 - 0.2D(XCR_2-H)]/0.75$$
 (2.20)

where $D(XCR_2-H)$ is the bond dissociation energy of a species that gives the radical formed by RO_2 addition and where RO_2 is assumed to have the same effect as methyl (Me) on D(C-H). Thus for oxidation of vinyl chloride

$$RO_2 \cdot + C = C = C1$$
 $RO_2 \cdot -C \cdot C1$
 $RO_2 \cdot -C \cdot C1$
(2.21)

the closest analog would be $MeCH_2$ CHCl, and the value of $D(MeCH_2CHCl-H)$ would be used in equation (2.20). Bond dissociation energies were taken from Furuyama et al. (1969).

RATE CONSTANT FOR OXIDATION BY SINGLET OXYGEN

Only a few of the chemicals tabulated in this report are reactive toward $^1\mathrm{O}_2$; these include some polycyclic aromatic and a few olefinic double bond or diene systems. All reactive chemicals were assigned rate constants by analogy with similar structures that have shown rate constants for reaction with singlet oxygen. For cyclic olefins, the values of Matsuura et al. (1973) were used. For alicyclic olefins and other structures, the rate data summarized by Gollnick (1978) were used.

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SECTION 3

DATA SHEETS FOR CHEMICALS OF INTEREST

This section contains a list of data sheets showing the Chemical Abstract Services registry number and compound name, a list of source codes for the data sheets, and the data sheets and references for this work assignment.

LIST OF DATA SHEETS

	CAS	
Number	Registry Number	Compound Name
1	62-44-2	Acetamide, N-(ethoxyphenyl)-
2	53-96-3	Acetamide, N-9H-fluoren-2-y1-
_		(2-Acetamidofluorene)
3	1402-68-2	Aflatoxin
4	148-82-3	Alanine, [p-bis(2-chloroethyl)amino]phenyl-, L-
5	92-67-1	4-Aminobiphenyl
6	61-82-5	Amitrole (3-Amino-1,2,4-triazole)
7	7778-39-4	Arsenic acid
8	492-80-8	Auramine
9	115-02-6	Azaserine
10	151-56-4	Aziridine
		(Ethyleneimine)
11	50-07-7	Azirino(2',3':3,4)pyrrolo(1,2-a)indole-4,7-dionen,6-amino-8-[((aminocarbonyl)oxy)methyl]-1,1a,2,8,8a,8b-hexahydro-8a-methoxy-5-methyl-(Porifomycin)
12	56-49-5	Benz[j]aceanthrylene, 1,2-dihydro-3-methyl-
13	225-51-4	Benz[c]acridine
14	57-97-6	1,2-Benzanthracene, 7,12-dimethyl-
15	3165-93-3	Benzenamine, 4-chloro-2-methyl-, hydrochloride
16	60-11-7	Benzenamine, N,N-dimethyl-4-phenylano-
17	101-14-4	Benzenamine, 4,4'-methylenebis(2-chloro-)-
18	636-21-5	Benzenamine, 2-methyl-, hydrochloride
19	99-55-8	Benzenamine, 2-methyl-5-nitro-
		(2-Amino-4-nitrotoluene)
20	510-15-6	Benzeneacetic acid, 4-chloro-alpha-
		<pre>(4-chlorophenyl)-alpha-hydroxy-, ethyl ester (Chlorobenzilate)</pre>
21	94-59-7	Benzene, 1,2-methylenedioxy-4-allyl-(Safrole)
22	120-58-1	Benzene, 1,2-methylenedioxy-4-propenyl- (Isosafrole)
23	94-58-6	Benzene, 1,2-methylenedioxy-4-propyl- (Dihydrosafrole)
24	82-68-8	Benzene, pentachloronitro-
25	13597-99-4	Beryllium nitrate

LIST OF DATA SHEETS (continued)

	CAS	
Number	Registry Number	Compound Name
26	1464-53-5	2,2'-Bioxirane
27	119-93-7	(1,1'-Bipheny1)-4,4'-diamine, 3,3'-dimethoxy-
28	119-90-7	(1,1'-Biphenyl)-4,4'-diamine, 3,3'-dimethyl- (o-Tolidine)
29	305-03-3	Butanoic acid, 4-[bis(2-chloroethyl)amino] benzene-
30	8001-35-2	Camphene, octachloro-
31	51-79-6	Carbamic acid, ethyl ester (Ethyl carbamate urethane)
32	615-53-2	Carbamic acid, methylnitroso-, ethyl ester
33	759-73-9	Carbamide, N-ethyl-N-nitroso-
34	684-93-5	Carbamide, N-methyl-N-mitroso-
35	62-56-6	Carbamide, thio- (Thiourea)
36	79-44-7	Carbamoyl chloride, dimethyl-
37	494-03-1	Chloronaphazine
38	106-89-8	1-Chloro-2,3-epoxypropane (Epichlorohydrin)
39	50-18-0	Cyclophosphamide
40	20830-81-3	Daunomycin
41	2303-16-4	Diallate
42	302-01-2	Diamine
		(Hydrazine)
43	95-80-7	2,4-Diaminotoluene
44	189-55-9	1,2:7,8-Dibenzopyrene
45	96-12-8	1,2-Dibromo-3-chloropropane
46	692-42-2	Diethylarsine
47	123-91-1	1,4-Diethylene dioxide (p-Dioxane)
48	57-14-7	1,1-Dimethylhydrazine
49	1615-80-1	N, N'-Diethylhydrazine
50	56-53-1	Diethylstilbestrol
51	77-78-1	Dimethyl sulfate
52	25321-14-6	Dinitrotoluenes
53	602-01-7	2,3-Dinitrotoluene
54	619-15-8	2,5-Dinitrotoluene
55	610-39-9	3,4-Dinitrotoluene

LIST OF DATA SHEETS (continued)

Number	CAS Registry Number	Compound Name
	100 20 /	
56 57	122-39 - 4	N, N-Diphenylamine
	55-18-5	Ethanamine, N-ethyl-N-nitroso-
58 59	62-55-5	Ethanethioamide
60	1116-54-7	Ethanol, 2,2'-(nitrosoimino)bis-
90	4549-40-0	Ethenamine, N-methyl-N-nitroso-
61	106-93-4	Ethylene dibromide
62	75-21-8	Ethylene oxide
63	62-50-0	Ethyl methanesulfonate
64	9004-66-4	Ferric dextran
65	18883-66-4	D-Glucopyranose, 2-deoxy-2-(3-methy1-3-
		nitrosoureido)-
66	765-34-4	Glycidylaldehyde
67	70-25-7	Guanidine, N-nitroso-N-methyl-N'-nitro-
68	143-50-0	Kepone
69	303-34-4	Lasiocarpine
70	75-55 - 8	2-Methylaziridine
		(Propyleneimine)
71	505-60-2	Mustard Gas
72	72-57-1	2,7-Napththalenedisulfonic acid, 3,3'-[(3,3'-dimethyl-(1,1'-biphenyl)-4,4'-diyl)-bis(azo)]
		bis(5-amino-4-hydroxy)-tetrasodium salt
73	134-32-7	l-Naphthylamine
74	91-59-8	2-Naphthylamine
75	13463-39-3	Nickel carbonyl
76	100-75-4	N-Nitrosopiperidine
77	930-55-2	N-Nitrosopyrrolidine
78	1120-71-4	1,2-Oxathiolane, 2,2-dioxide
79	50-06-6	Phenobarbital
80	126-72-7	1-Propanol, 2,3-dibromo-, phosphate (3:1)
		<pre>[tris(2,3-Dibromopropyl)phosphate]</pre>
81	75-86-5	Propanenitrile, 2-hydroxy-2-methyl- (Acetone cyanohydrin)
82	72-54-8	TDE
92	66 _7 E_1	(1,1-Dichloro-2,2-bis(p-chlorophenyl)ethane)
83	66-75-1	<pre>Uracil, 5-[bis(2-chloroethy1)amino]- (Uracil mustard)</pre>

LIST OF SOURCE CODES

Calc	Molecular weight/oxygen ratio was calculated directly.
CC-Kow	Value of the octanol/water partition coefficient (K_{ow}) was obtained by computer calculation using FRAGMENT calculation procedure (see Section 4.4).
C-Sw f Kow	The water solubility (S_w) was calculated from the octanol/water partition coefficient (K_{OW}) using the equation of Yalkowsky and Valvani (1980); the calculation of S_w values is discussed in Section 2.
C-vp f bp	Vapor pressure (vp) was calculated from the boiling point (bp) using the method discussed by Grain (1982); the method is discussed in Section 2.
E-A-Aziridine	Estimate by analogy to aziridine; hydrolysis data for aziridine from Mabey and Mill (1978) or from Earley et al. (1958).
E-A-Dibromopropane	Estimate by analogy to dibromopropane; hydrolysis data for dibromopropane from Vogel (1983).
E-A-Glycirdol	Estimate by analogy to glycirdol; hydrolysis data for glycirdol from Mabey and Mill (1978).
E-A-Naled	Estimate by analogy to Naled; hydrolysis data for Naled from Jentzsch and Fischer (1978).
E-A-NM	Estimated by analogy to nitrogen mustards; hydrolysis data for nitrogen mustards from Ross (1949) indicates a maximum half-life of 8 hours in aquatic environments.
E-KB	Estimate of biotransformation rate constant (k_b) is based on relative rates of transformation reported in literature or on structure-reactivity analogies.
HF-NBD	Hydrolysis is too fast for biotransformation studies to be conducted. Therefore, no biotransformation data are available.
HNES	Hydrolysis is not environmentally significant. Chemical hydrolysis occurs only at extreme pHs or temperatures or with catalysts not available in aquatic environments.

INERT Oxidation reactions at ambient oxygen levels have half-life

greater than 2 years and are therefore considered

unimportant fate processes.

M-OX Oxidation rate constants were modelled using functional

group reactivity toward alkyl peroxyl radical (RO2) and

singlet oxygen (SO).

NBD No biotransformation data are available.

NHFG No hydrolyzable functional groups in molecule.

OX Oxidation rate constants are experimental values.

partial Partial notation indicates that the computer calculated

octanol/water partition coefficient has not accounted for each functional moiety of the molecule. This occurs when a substituent fragment is not represented in the data base or

when there are possible hydrogen bonding interactions.

PNER Photolysis is not environmentally relevant.

RO2 Alkyl peroxyl radical, RO2.

Singlet oxygen, ¹0₂

VF-NBD Volatilization is too fast for biotransformation studies to

be conducted. No biotransformation data are therefore

available.

EPA CONTRACT 68-03-2981 WORK ASSIGNMENT NO. 14

Compound Name: Acetamide, N-(ethoxyphenyl)-			
CAS Registry Number: 62-44-2	Molecular Weight	(g): <u>179.22</u>	
Parameters:		Reference	
Water Solubility (ppm)	578	C-Sw f Kow	
Melting Point (°C)	134-135	Merck (1976)	
Boiling Point (°C)			
Vapor Pressure (torr)	6.9×10^{-7}	Wiedemann (1972)	
Molecular Weight/Oxygen	5.60	Calc	
Log (Octanol/Water Partition Coefficient)	1.76	CC-Kow	
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES		
Acid Hydrolysis Rate Constant (M hr-1)	HNES		
Neutral Hydrolysis Rate Constant (hr 1)			
Microbial Degradation Rate Constant (ml cell-lhr-1)	3 × 10 ⁻⁹	E-KB	
Photolysis Rate Constant (hr ⁻¹)			
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO	

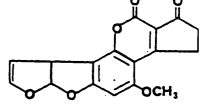
These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

EPA CONTRACT 68-03-2981 WORK ASSIGNMENT NO. 14

Compound Name: Acetamide, N-9H-flu	oren-2-yl- (2-Ace	etamidofluorene)
CAS Registry Number: 53-96-3	Molecular Weight	223.28 <u>223.28</u>
Parameters:		Reference
Water Solubility (ppm)	6.5	C-Sw f Kow
Melting Point (°C)	192-196	Aldrich (1982)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	6.98	Calc
Log (Octanol/Water Partition Coefficient)	3.28	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M hr 1)	HNES	
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell -1hr -1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

EPA CONTRACT 68-03-2981 WORK ASSIGNMENT NO. 14



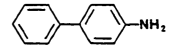
Compound Name: Aflatoxin		
CAS Registry Number: 1402-68-2	Molecular Weight	(g): 312
·		
Parameters:		Reference
Water Solubility (ppm)	993	C-Sw f Kow
Melting Point (°C)	<u> 268-269</u>	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	9.75	Calc
Log (Octanol/Water Partition Coefficient)	0.71 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr 1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M-1hr-1)	2 x 10 ¹⁰	M-OX SO

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

Compound Name: Alanine, [p-bis(2	-chloroethyl)amino]	phenyl-, L-
CAS Registry Number: 148-82-3	Molecular Weight	(g): <u>305.20</u>
Parameters:		Reference
Water Solubility (ppm)	22	C-Sw f Kow
Melting Point (°C)	180-181	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	9.54	Calc
Log (Octanol/Water Partition Coefficient)	1.20	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		EA-NM
Acid Hydrolysis Rate Constant (M hr-1)		EA-NM
Neutral Hydrolysis Rate Constant (hr ⁻¹)		EA-NM
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr^{-1})		-
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	unknown	

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Aryldialkyl amines react with RO2 by electron-, not H-transfer.



Compound Name: 4-Aminobiphenyl		
CAS Registry Number: 92-67-1	— Molecular Weight	(g): 169.23
Parameters:		Reference
Water Solubility (ppm)	842	C-Sw f Kow
Melting Point (°C)	53	Verschueren (1977)
Boiling Point (°C)	302	Weast (1981)
Vapor Pressure (torr)	6.0×10^{-5}	C-VP f bp
Molecular Weight/Oxygen	5.29	Calc
Log (Octanol/Water Partition Coefficient)	2.78	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr 1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻⁹	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁶	M-OX RO2



Compound Name: _Amitrole (3-Amino-1,2,4-triazole)		
CAS Registry Number: 61-82-5	Molecular Weight	(g): 84.08
Parameters:		Reference
Water Solubility (ppm)	$2.8 \times 10^{5}/25^{\circ}C$	Spencer (1973)
Melting Point (°C)	159	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	2.66	Calc
Log (Octanol/Water Partition Coefficient)	-2.08	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr 1)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁶	M-OX RO2

H3 AS O4 . 1/2 H2 O

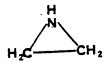
Compound Name: Arsenic acid		
CAS Registry Number: 7778-39-4	Molecular Weight((g): 150.9
Parameters:		Reference
Water Solubility (ppm)	1	
Melting Point (°C)	35.5	Hawley (1977)
Boiling Point (°C)	loses H ₂ O at 160°	Hawley (1977)
Vapor Pressure (torr)	non-volatile	Spencer (1973)
Molecular Weight/Oxygen	4.72	Calc
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Neutral Hydrolysis Rate Constant (hr)	hydrated in solution	
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)		NBD
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	

Compound Name: <u>Auramine</u>		
CAS Registry Number: 492-80-8	Molecular Weight(g): 267.38
	,	
Parameters:		Reference
Water Solubility (ppm)	2.1	C-Sw f Kow
Melting Point (°C)	136	Weast (1973)
Boiling Point (°C)		
Vapor Pressure (torr)		-
Molecular Weight/Oxygen	8.34	Calc
Log (Octanol/Water Partition Coefficient)	4.16 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		· · · · · · · · · · · · · · · · · · ·
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	unknown	

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Aryldialkyl amines react with RO2 by electron-, not H-atom transfer.

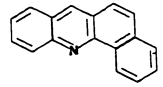
Compound Name: Azaserine		
CAS Registry Number: 115-02-6	Molecular Weight	(g):
Parameters:		Reference
Water Solubility (ppm)	1.36×10^5	C-Sw f Kow
Melting Point (°C)	146-162 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.41	Calc
Log (Octanol/Water Partition Coefficient)	-1.08 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell-lhr-l)	3 x 10 ⁻⁹	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0



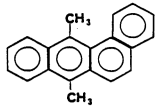
Compound Name: Aziridine (Ethyleneimine)		
CAS Registry Number: 151-56-4	Molecular Weight	(g): 43.07
Parameters:		Reference
Water Solubility (ppm)	2.66×10^6	C-Sw f Kow
Boiling Point (°C)	56-57	Merck (1976)
Vapor Pressure (torr)	255	Osborn and Scott (1980)
Molecular Weight/Oxygen	1.35	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	<u>HNES</u>	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	1.87×10^{-3}	Mabey & Mill (1978)
Neutral Hydrolysis Rate Constant (hr)		Earlevet_al. (1958)
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻⁷	E-KB
Photolysis Rate Constant (hr^{-1})	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	м-Ox RO2, SO

Compound Name: [((aminocarbonyl)o	(xy)methy1]-1,1a,2,8	,8a,8b-hexahydro-8a-methoxy
5-methyl-(Porifomyo CAS Registry Number: 50-07-7		(g): 348.35
	_	
Parameters:		Reference
Water Solubility (ppm)	1.6 x 10 ⁶	C-Sw f Kow
Melting Point (°C)	201-201.5 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	10.89	Calc
Log (Octanol/Water Partition Coefficient)	-2.19	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M hr-1)	2.2×10^{-3}	E-A-Aziridine as cited in Mabey and Mill (1978)
Neutral Hydrolysis Rate Constant (hr)	2.5 x 10 ⁻³	E-A-Aziridine as cited in Mabey and Mill (1978)
Microbial Degradation Rate Constant (ml cell-hr-1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: Benz[j]aceanthryle	ne, 1,2-dihydro-3-m	methyl-
CAS Registry Number: 56-49-5	Molecular Weight	(g): <u>268.34</u>
Parameters:	2	Reference
Water Solubility (ppm)	1.15×10^{-3}	C-Sw f Kow
Melting Point (°C)	179-180	Merck (1976)
Boiling Point (°C)	280/80 torr	Merck (1976)
Vapor Pressure (torr)	3.8×10^{-6}	C-VP f bp
Molecular Weight/Oxygen	8.39	Calc
Log (Octanol/Water Partition Coefficient)	6.97	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁸	M-OX SO



Compound Name: Benz[c]acridine		
CAS Registry Number: 225-51-4	Molecular Weight	(g):
Parameters:		Reference
Water Solubility (ppm)		C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	7.16	Calc
Log (Octanol/Water Partition Coefficient)	4.56	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X RO2, SO



Compound Name: 1,2-Benzanthracene	, 7,12-dimethyl-	
CAS Registry Number: 57-97-6	Molecular Weight	(g): 256.33
Parameters:		Reference
Water Solubility (ppm)	4.40×10^{-3}	C-Sw f Kow
Melting Point (°C)	121-123	Alrich (1982)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	8.01	Calc
Log (Octanol/Water Partition Coefficient)	6.94	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁸	_M-OX SO

Compound Name: Benzenamine, 4-chloro-2-methyl-, hydrochloride		
CAS Registry Number: 3165-93-3	Molecular Weight	(g): <u>176</u>
Parameters:		Reference
Water Solubility (ppm)	1400	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5 50	Calc
Log (Octanol/Water Partition Coefficient)	2.58 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	•
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: Benzenamine, N,N-d	imethyl-4-phenylano	_
CAS Registry Number: 60-11-7	Molecular Weight	(g): 225.28
		•
Parameters:		Reference
Water Solubility (ppm)	13.6	C-Sw f Kow
Melting Point (°C)	114-117	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)	3.3×10^{-7}	Green and Jones (1967)
Molecular Weight/Oxygen	7.04	Calc
Log (Octanol/Water Partition Coefficient)	3.72 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	-3
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M-1hr-1)	unknown	

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Aryldialkyl amines react with RO2 by electron, not H-atom transfer.

EPA CONTRACT 68-03-2981 H₂N — CI NH₂

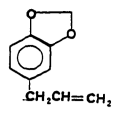
Compound Name: Benzenamine, 4,4'-	methylenebis(2-chlo	ro-)-
CAS Registry Number: 101-14-4	Molecular Weight(g): 232	
Parameters:		Reference
Water Solubility (ppm)	1700	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	7.25	Calc
Log (Octanol/Water Partition Coefficient)	2.62 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	Λ.
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	4 x 10 ⁶	M-OX RO2



Compound Name: Benzenamine, 2-met	hyl-, hydrochloride	
CAS Registry Number: 636-21-5	Molecular Weight(g): 142
Parameters:		Reference
Water Solubility (ppm)		
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	4.44	Calc
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell hr 1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0

Compound Name: Benzenamine, 2-met	hyl-5-nitro- (2-Am:	ino-4-nitrotoluene)
CAS Registry Number: 99-55-8	Molecular Weight	(g): <u>152</u>
Parameters:		Reference
Water Solubility (ppm)	1.3×10^5	C-Sw f Kow
Melting Point (°C)	107-108	Weast (1973)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	4.75	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁶	M-OX RO2

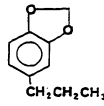
Benzeneacetic acid, Compound Name: hydroxy-, ethyl est		-chlorophenyl)-alpha-
CAS Registry Number: 510-15-6	Molecular Weight((g): 325.20
Parameters:		Reference
Water Solubility (ppm)	21.9	C-Sw f Kow
Melting Point	35-37	Spencer (1973)
Boiling Point (°C)	146-148/0.04 torr	Merck (1976)
	$1.2 \times 10^{-6}/25^{\circ}$ C	C-VP f bp
Vapor Pressure (torr)	$2.2 \times 10^{-6}/20^{\circ}$ C	Merck (1976)
Molecular Weight/Oxygen	10.2	Calc
Log (Octanol/Water Partition Coefficient)	4.51 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell -1hr -1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X RO2, SO



Compound Name: Benzene. 1.2-methy	lenedioxy-4-allyl-	(Safrole)
CAS Registry Number: 94-59-7		
Parameters:	insoluble 1500	Reference Hawley (1977) C-Sw f Kow
Water Solubility (ppm)	11.2	Weast (1973)
Melting Point (°C) Boiling Point (°C)	234.5	Weast (1973)
Vapor Pressure (torr)	9.1 x 10 ⁻²	C-vp f bp
Molecular Weight/Oxygen	5.07	Calc
Log (Octanol/Water Partition Coefficient)	2.53	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr 1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr-1)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ¹⁰	M-OX SO



CH=CHCH3 Compound Name: Benzene, 1,2-methylenedioxy-4-propenyl-(Isosafrole) CAS Registry Number: 120-58-1 162.18 -Molecular Weight(g): _ Reference Parameters: 1.09×10^3 C-Sw f Kow Water Solubility (ppm) 253 Merck (1976) Boiling Point (°C) $1.6 \times 10^{-8} \text{(extrap)}$ Merck (1976) Vapor Pressure (torr) 5.07 Calc Molecular Weight/Oxygen Log (Octanol/Water Partition 2.66 CC-Kow Coefficient) Alkaline Hydrolysis Rate Constant $(M^{-1}hr^{-1})$ NHFG Acid Hydrolysis Rate Constant (M⁻¹hr⁻¹) NHFG Neutral Hydrolysis Rate Constant (hr 1) NHFG Microbial Degradation Rate Constant (ml cell⁻¹hr⁻¹) 1×10^{-10} E-KB Photolysis Rate Constant (hr^{-1}) Oxidation Rate Constant $(M^{-1}hr^{-1})$ 2×10^{10} M-OX SO



Compound Name: Benzene, 1,2-methy	lenedioxy-4-propyl-	(Dihydrosafrole)
CAS Registry Number: 94-58-6	Molecular Weight	(g): <u>164.2</u>
Parameters:		Reference
Water Solubility (ppm)	1500	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.13	Calc
Log (Octanol/Water Partition Coefficient)	2.54	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell hr 1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ¹⁰	M-OX SO

Compound Name: Benzene, pentachlo	ronitro-	
CAS Registry Number: 82-68-8	Molecular Weight	(g): 295.36
Parameters:		Reference
Water Solubility (ppm)	7.11×10^{-2}	C-Sw f Kow
Melting Point (°C)	146	Verschueren (1977)
Boiling Point (°C)	328	Merck (1976)
Vapor Pressure (torr)	1.8×10^{-6} 1.13×10^{-4}	C-Vp f bp Spencer (1973)
Molecular Weight/Oxygen	9.23	Calc
Log (Octanol/Water Partition Coefficient)	5.45 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	<u>M-0X RO2, SO</u>

 $\mathrm{Be}\big(\mathrm{NO_3}\big)_2 \,\cdot\, 3\,\mathrm{H}_2\mathrm{O}$

Compound Name: Beryllium nitrate		
CAS Registry Number: 13597-99-4	Molecular Weight	(g): 187.1
Parameters:		Reference
Water Solubility (ppm)		
Melting Point (°C)	60	Merck (1976)
Boiling Point (°C)	100-200 (dec)	Hawley (1977)
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.85	Calc
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		•
Acid Hydrolysis Rate Constant (M hr-1)	<u>.</u>	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	dissociates in solution	
Microbial Degradation Rate Constant (ml cell hr -1)		NBD
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO



Compound Name: 2,2'-Bioxirane		
CAS Registry Number: 1464-53-5	Molecular Weight	(g): 86.09
Parameters:		Reference
Water Solubility (ppm)	1.07×10^7	C-Sw f Kow
Boiling Point (°C)	138	Merck (1976)
Vapor Pressure (torr)	6.9	C-VP f bp
Molecular Weight/Oxygen	2.69	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M hr 1)	8.9	E-A-Glycirdol
Neutral Hydrolysis Rate Constant (hr)	1.02 x 10 ⁻³	E-A-Glycirdol
Microbial Degradation Rate Constant (ml cell-1hr-1)		NBD
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2

Compound Name: (1,1'-Biphenyl)-4,	4'-diamine, 3,3'-di	imethoxy-
CAS Registry Number: 119-93-7	Molecular Weight	(g): 244.28
Parameters:		Reference
Water Solubility (ppm)	430	C-Sw f Kow
Melting Point (°C)	137-138	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	7.63	Calc
Log (Octanol/Water Partition Coefficient)	1.78	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	.*
Acid Hydrolysis Rate Constant (M hr -1)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell-lhr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	4 x 10 ⁶	M-OX RO2

Compound Name: (1,1'-Bipheny1)-4,	4'-diamine, 3,3'-d	imethyl- (o-Tolidine)
CAS Registry Number: 119-90-7	Molecular Weigh	t(g): 212.28
Parameters:		Reference
Water Solubility (ppm)	73.5	C-Sw f Kow
Melting Point (°C)	129-131	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	6.63	Calc
Log (Octanol/Water Partition Coefficient)	2.88	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	<u> </u>
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-hr-1)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	4 x 10 ⁶	_M=0X_R02

Compound Name: Butanoic acid, 4-[1	ois(2-chloroethyl)am	ino]benzene-
CAS Registry Number: 305-03-3	Molecular Weight((g): 304.23
Parameters:		Reference
Water Solubility (ppm)	1.67×10^3	C-Sw f Kow
Melting Point (°C)	64-66	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	9.51	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		EA-NM
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		EA-NM
Neutral Hydrolysis Rate Constant (hr)		EA-NM
Microbial Degradation Rate Constant (ml cell hr -1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	unknown a	

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Aryldialkyl amines react with RO2 by electron, not H-atom transfer.



Compound Name: Camphene, octachlo	ro-	
CAS Registry Number: 8001-35-2	Molecular Weight(g):
Parameters:	ı	Reference
Water Solubility (ppm)		
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	12.94	Calc
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	(a)	
Acid Hydrolysis Rate Constant (M hr-1)	<u>(a)</u>	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	(a)_	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Some isomers may hydrolyze, but rates will vary with structure.

H₂ N-C-OC₂H₅ || O

Compound Name: Carbamic acid, ethy	yl ester (Ethyl car	bamate urethane)
CAS Registry Number: 51-79-6	Molecular Weight	(g): 89.09
Parameters:		Reference
Water Solubility (ppm)	6.48×10^5	C-Sw f Kow
Melting Point (°C)	48-50	Merck (1976)
Boiling Point (°C)	182-184	Merck (1976)
_	0.44	C-vp f bp
Vapor Pressure (torr)	0.36 (extrap.)	Weast (1973)
Molecular Weight/Oxygen	2.78	Calc
Log (Octanol/Water Partition Coefficient)	-0.15	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr 1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)	1 x 10 ⁻⁷	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: Carbamic acid, met	hylnitroso-, ethyl	ester
CAS Registry Number: 615-53-2	Molecular Weight	(g): <u>132.1</u>
Parameters:	,	Reference
Water Solubility (ppm)	4.43×10^7	C-Sw f Kow
Boiling Point (°C)	62-64/12 torr	Klein (1982)
Vapor Pressure (torr)	1.1	Klein (1982)
Molecular Weight/Oxygen	4.13	Calc
Log (Octanol/Water Partition Coefficient)	-1.69	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		•
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Neutral Hydrolysis Rate Constant (hr. 1)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Ovidation Rate Constant (M-1hr-1)	INERT	M-OX RO2, SO

Compound Name: Carbamide, N-ethyl	-N-nitroso-	
CAS Registry Number: 759-73-9	Molecular Weight	(g): <u>117.1</u>
<pre>Parameters: Water Solubility (ppm)</pre>	3.31 x 10 ⁸	Reference C-Sw f Kow
Melting Point (°C)	103-104	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	3.66	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-lhr-l)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		2
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: <u>Carbamide</u> , N-methy	1-N-nitroso-	
CAS Registry Number: 684-93-5	Molecular Weight	(g): 103.1
Parameters:		Reference
Water Solubility (ppm)	6.89×10^8	C-Sw f Kow
Melting Point (°C)	124	Merck (1976)
Boiling Point (°C)		· · · · · · · · · · · · · · · · · · ·
Vapor Pressure (torr)		
Molecular Weight/Oxygen	3.22	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Neutral Hydrolysis Rate Constant (hr 1)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0

H₂N---C---NH₂

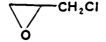
Compound Name: Carbamide, thio-	(Thiourea)	
CAS Registry Number: 62-56-6	Molecular Weight	(g):76.12
Parameters:		Reference
Water Solubility (ppm)	1.72×10^6	C-Sw f Kow
Melting Point (°C)	180-182	Hawley (1977)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	2.38	Calc
Log (Octanol/Water Partition Coefficient)	-2.05 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell hr -1)	1 × 10 ⁻⁷	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	4 × 10 ¹⁰	M-OX SO

npound Name: <u>Carbamovl chloride</u>	, dimethyl-	
CAS Registry Number: 79-44-7	Molecular Weight	(g): <u>107.54</u>
•		
Parameters:		Reference
Water Solubility (ppm)	1.44×10^7	C-Sw f Kow
Boiling Point (°C)	55-57/11 torr	Fluka (1982)
Vapor Pressure (torr)	1.95	Jaber and Gunderson (1983)
Molecular Weight/Oxygen	3.34	Calc
Log (Octanol/Water Partition Coefficient)	-1.32 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr 1)		
Neutral Hydrolysis Rate Constant (hr 1)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant $(M^{-1}hr^{-1})$	INERT	M-0X R02, S0

Compound Name: <u>Chloronaphazine</u>		
CAS Registry Number: 494-03-1	Molecular Weight	(g): 268,20
Parameters:		Reference
Water Solubility (ppm)	165	C-Sw f Kow
Boiling Point (°C)	210	Merck (1976)
Vapor Pressure (torr)	0.30	C-Vp f bp
Molecular Weight/Oxygen	8.38	Calc
Log (Octanol/Water Partition Coefficient)	3.62 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		EA-NM
Acid Hydrolysis Rate Constant (M hr-1)		EA-NM
Neutral Hydrolysis Rate Constant (hr)		EA-NM
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	unknown ^a	

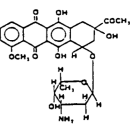
These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) Aryldialkyl amines react with RO2 by electron, not H-atom transfer.



Compound Name: 1-Chloro-2,3-epoxypropane (Epichlorohydrin)		
CAS Registry Number: 106-89-8 Molecular Weight(g): 92.53		
Parameters:		Reference
Water Solubility (ppm)	3.19×10^5	C-Sw f Kow
Boiling Point (°C)	117.9	Merck (1976)
Vapor Pressure (torr)	15.7	Weast (1973)
Molecular Weight/Oxygen	2.89	Calc
Log (Octanol/Water Partition Coefficient)	0.15 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ¹ hr ¹)	HNES	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	2.88.	Mabey & Mill (1978)
Neutral Hydrolysis Rate Constant (hr 1)	3.51×10^{-3}	Mabey & Mill (1978)
Microbial Degradation Rate Constant (ml cell hr -1)		HF-NBD
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: Cyclophosphamide		
CAS Registry Number: 50-18-0	Molecular Weight((g): 261.10
Parameters:		Reference
Water Solubility (ppm) Melting Point (°C) Boiling Point (°C)	1.31 x 10 ⁹ 41-45	C-Sw f Kow Merck (1976)
Vapor Pressure (torr)		
Molecular Weight/Oxygen	8.16	Calc
Log (Octanol/Water Partition Coefficient)	3.22 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX, RO2, SO



Compound Name: Daunomycin	·	
CAS Registry Number: 20830-81-3	Molecular Weight	(g): 527.5
Parameters:		Reference
Water Solubility (ppm)	2.7×10^5	C-Sw f Kow
Melting Point (°C)	188-190 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		·
Molecular Weight/Oxygen	16.48	Calc
Log (Octanol/Water Partition Coefficient)	-1.99	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	*****	•
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
·	8 x 10 ⁹	M-OX SO
Oxidation Rate Constant $(M^{-1}hr^{-1})$	4×10^{7}	M-OX RO2

Compound Name: Diallate		
CAS Registry Number: 2303-16-4	Molecular Weight	(g): 273.5
Parameters:		Reference
Water Solubility (ppm)	14	Spencer (1973)
Boiling Point (°C)	150/9 torr	Merck (1976)
Vapor Pressure (torr)	6.4×10^{-3}	C-Vp f bp
Molecular Weight/Oxygen	8.55	Calc
Log (Octanol/Water Partition Coefficient)	0.73 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	c
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	_M-OX RO2, SO

H₂N ---NH₂

Compound Name: <u>Diamine</u> (Hydrazine))	
CAS Registry Number: 302-01-2	Molecular Weight((g): 32.05
Parameters:	0 .	Reference
Water Solubility (ppm)	3.41×10^8	C-Sw f Kow
Boiling Point (°C)	113.5	Aldrich (1982)
Vapor Pressure (torr)	14	Yaws et al. (1974)
Molecular Weight/Oxygen	1.00	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻⁷	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	labile ^a	

⁽a) Reacts with oxygen directly. Half-life estimated to be less than 10 days.

Compound Name: 2,4-Diaminotoluene	:	
CAS Registry Number: 95-80-7	Molecular Weight	(g): 122.17
Parameters:		Reference
Water Solubility (ppm)	4.77×10^4	C-Sw f Kow
Melting Point (°C)	9.7-98	Aldrich (1982)
Boiling Point (°C)	292	Weast (1981)
Vapor Pressure (torr)	3.8×10^{-5}	C-Vp f bp
Molecular Weight/Oxygen	3.82	Calc
Log (Octanol/Water Partition Coefficient)	0.35	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	-
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	**************************************
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁶	M-OX RO2



Compound Name: 1,2:7,8-Dibenzopyrene		
CAS Registry Number: 189-55-9	Molecular Weight	(g): 305
		•
Parameters:		Reference
Water Solubility (ppm)	0.11	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	9.53	Calc
Log (Octanol/Water Partition Coefficient)	6.62	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ⁸	M-OX SO

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Compound Name: 1,2-Dibromo-3-chloropropane CAS Registry Number: 96-12-8 236.3 -Molecular Weight(g): Reference Parameters: 1000 Spencer (1973) Water Solubility (ppm) 196 Merck (1976) Boiling Point (°C) C-Vp f bp 1.0 0.8/21°C Merck (1976) Vapor Pressure (torr) Calc 7.38 Molecular Weight/Oxygen Log (Octanol/Water Partition CC-Kow 2.29 Coefficient) Alkaline Hydrolysis Rate Constant (M-1hr-1) 21 Burlinson et al. (1982) Acid Hydrolysis Rate Constant (M⁻¹hr⁻¹) **HNES** Neutral Hydrolysis Rate Constant (hr 1) Microbial Degradation Rate Constant (ml cell hr -1) 3×10^{-12} E-KB Photolysis Rate Constant (hr⁻¹) Oxidation Rate Constant $(M^{-1}hr^{-1})$ INERT M-OX RO2, SO

H H₅C₂—As—C₂H₅

Compound Name: Diethylarsine		
CAS Registry Number: 692-42-2	Molecular Weight(g):134.05
Parameters:		Reference
Water Solubility (ppm)	417	C-Sw f Kow
Boiling Point (°C)	105	Weast (1981)
Vapor Pressure (torr)	35	C-VP f bp
Molecular Weight/Oxygen	4.19	Calc
Log (Octanol/Water Partition Coefficient)	2.97 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr 1)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	Е-КВ
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	unknown	



Compound Name: 1,4-Diethylene dioxide (p-Dioxane)		
CAS Registry Number: 123-91-1	Molecular Weight	(g): <u>88.10</u>
Parameters:		Reference
Water Solubility (ppm)	4.31 x 10 ⁵	C-Sw f Kow
Boiling Point (°C)	101.1	Merck (1976)
Vapor Pressure (torr)	39.9	Weast (1973)
Molecular Weight/Oxygen	2.75	Calc
Log (Octanol/Water Partition Coefficient)	0.01	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

H₂N—N—CH₃ CH₃

Compound Name: 1,1-Dimethylhydrazine		
CAS Registry Number: 57-14-7	Molecular Weight	(g): 60.10
Parameters: Water Solubility (ppm)	1.24 x 10 ⁸	C-Sw f Kow
Boiling Point (°C)	63.9	Merck (1976)
Vapor Pressure (torr)	157	Verschueren (1977)
Molecular Weight/Oxygen	1.88	Calc
Log (Octanol/Water Partition Coefficient)	-2.42	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 × 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)		

 H_5C_2 — $\frac{H}{N}$ — $\frac{H}{N}$ — C_2H_5

Compound Name: N,N'-Diethylhydrazine		
CAS Registry Number: 1615-80-1	Molecular Weight	(g): <u>88</u>
Parameters:		Reference
Water Solubility (ppm)	$\frac{2.88 \times 10^7}{}$	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	2.75	Calc
Log (Octanol/Water Partition Coefficient)	-1.68	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	<u>NHFG</u>	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell-hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)		

Compound Name:Diethylstilbestrol		
CAS Registry Number: 56-53-1	Molecular Weight(g	268.34
Parameters:		Reference
Water Solubility (ppm)	9.60×10^{-3}	C-Sw f Kow
Melting Point (°C)	169-172	Merck (1976)
Boiling Point (°C)		-
Vapor Pressure (torr)		
Molecular Weight/Oxygen	8.39	Calc
Log (Octanol/Water Partition Coefficient)	5.46	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10.7	M-OX RO2

(CH₃)₂SO₄

Compound Name: Dimethyl sulfate		
CAS Registry Number: 77-78-1	Molecular Weight	(g): <u>126.13</u>
Parameters:		Reference
Water Solubility (ppm)	3.24×10^5	C-Sw f Kow
	188 (dec)	Verschueren (1977)
Boiling Point (°C)	76/15 torr	Merck (1976)
Vapor Pressure (torr)	0.68/20°C	Weber et al. (1981)
Molecular Weight/Oxygen	3.94	Calc
Log (Octanol/Water Partition Coefficient)	-1.24 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	53	Mabey and Mill (1978)
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Neutral Hydrolysis Rate Constant (hr)	0.6	Mabey and Mill (1978)
Microbial Degradation Rate Constant (ml cell-lhr-l)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)		·
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

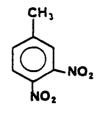
Compound Name: <u>Dinitrotoluenes</u>		
CAS Registry Number: 25321-14-6	Molecular Weight(g):182.14
Parameters:		Reference
Water Solubility (ppm)	3100	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.69	Calc
Log (Octanol/Water Partition Coefficient)	2.29	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	·
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	$(0.2 - 7) \times 10^{-10}$	Spanggord et al. (1981)
Photolysis Rate Constant (hr ⁻¹)	(a)	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

⁽a)Nitrotoluenes with o-nitrotoluene structure have photolysis rate constant $> 0.01~hr^{-1}$ (Tse et al., 1983)

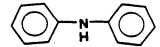
Compound Name: 2,3-Dinitrotoluene		
CAS Registry Number: 602-01-7	Molecular Weight((g): 182.14
Parameters:		Reference
Water Solubility (ppm)	3100	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.69	Calc
Log (Octanol/Water Partition Coefficient)	2.29	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	·
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	5 x 10 ⁻¹⁰	Spanggord et al. (1981)
Photolysis Rate Constant (hr ⁻¹)	0.07	Tse et al. (1983)
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0

Compound Name: 2,5-Dinitrotoluene		
CAS Registry Number: 619-15-8	Molecular Weight	(g): 182.14
David		Reference
Parameters:		
Water Solubility (ppm)	$\frac{1.32 \times 10^3}{}$	C-Sw f Kow
Melting Point (°C)	52.5	Weast (1981)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.69	Calc
Log (Octanol/Water Partition Coefficient)	2.28	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	<u>-</u>
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	<u>.</u>
Microbial Degradation Rate Constant (ml cell hr -1)	7 x 10 ⁻¹⁰	Spanggord et al. (1981)
Photolysis Rate Constant (hr ⁻¹)	0.3	Tse et al. (1983)
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

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Compound Name: 3,4-Dinitrotoluene		
CAS Registry Number: 610-39-9	Molecular Weight	(g): <u>182.14</u>
Parameters:		Reference .
Water Solubility (ppm)	1.08×10^3	C-Sw f Kow
Melting Point	58.3	Weast (1981)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	5.69	
Log (Octanol/Water Partition Coefficient)	2.29	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	4 x 10 ⁻¹⁰	Spanggord et al. (1981)
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO



Compound Name: N,N-Diphenylamine		
CAS Registry Number: 122-39-4	Molecular Weight(g): 169.22
Parameters:		Reference
Water Solubility (ppm)	57.6	C-Sw f Kow
Melting Point (°C)	53-54	Merck (1976)
Boiling Point (°C)	302	Merck (1976)
Vapor Pressure (torr)	3.8 × 10 ⁻⁵	C-vp f bp
Molecular Weight/Oxygen	5.29	Calc
Log (Octanol/Water Partition Coefficient)	3.60	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻⁹	Е-КВ
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	3 x 10 ⁷	M-OX RO2

$$0=N-N C_2H_5$$

Compound Name: Ethanamine, N-ethy	l-N-nitroso-	
CAS Registry Number: 55-18-5	Molecular Weight	(g): 102.14
Parameters:	5	Reference
Water Solubility (ppm)	$\frac{2.2 \times 10^5}{}$	C-Sw f Kow
Boiling Point (°C)	175-177	Merck (1976)
Vapor Pressure (torr)	1.10	Klein (1982)
Molecular Weight/Oxygen	3.19	Calc
Log (Octanol/Water Partition Coefficient)	0.34	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr-1)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	TNERT	M-04 R02 S0

S	
H ₃ CC	NH

Compound Name: Ethanethioamide		
CAS Registry Number: 62-55-5	Molecular Weight	(g):75.14
Parameters:	,	Reference
Water Solubility (ppm)	1.63×10^5	Merck (1976)
Melting Point (°C)	108.5	Verschueren (1977)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	2.35	Calc
Log (Octanol/Water Partition Coefficient)	0.91 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name:Ethanol, 2,2'-(nitrosoimino)bis-			
CAS Registry Number: 1116-54-7	Molecular Weight	(g): <u>134.1</u>	
Parameters:	7 1	Reference	
Water Solubility (ppm)	3.1 x 10 ⁷	C-Sw f Kow	
Boiling Point (°C)	114/1.5 torr	Klein (1982)	
Vapor Pressure (torr)	$\frac{8.3 \times 10^{-4}}{}$	Klein (1982)	
Molecular Weight/Oxygen	4.19	Calc	
Log (Octanol/Water Partition Coefficient)		CC-Kow	
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		•	
Acid Hydrolysis Rate Constant (M hr-1)			
Neutral Hydrolysis Rate Constant (hr 1)			
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	EKB	
Photolysis Rate Constant (hr ⁻¹)			
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO	

N=0 | CH₂= CH-N-CH₃

Compound Name: <u>Ethenamine, N-meth</u>	yl-N-nitroso-	
CAS Registry Number: 4549-40-0	Molecular Weight	(g): 86.1
Parameters:		Reference
Water Solubility (ppm)	$\frac{7.6 \times 10^5}{}$	C-Sw f Kow
Boiling Point (°C)	47-48/30 torr	Klein (1982)
Vapor Pressure (torr)	12.3	Klein (1982)
Molecular Weight/Oxygen	2.69	Calc
Log (Octanol/Water Partition Coefficient)	-0.23	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X RO2, SO

Br-CH2 - CH2- Br

Compound Name: Ethylene dibromide		
CAS Registry Number: 106-93-4	Molecular Weight	(g):
Parameters:		Reference
Water Solubility (ppm)	1.18 × 10 ⁴	C-Sw f Kow
Boiling Point (°C)	131.4	Dreisbach (1959)
Vapor Pressure (torr)	11.7	Dreisbach (1959)
Molecular Weight/Oxygen	5.87	Calc
Log (Octanol/Water Partition Coefficient)	1.76	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	3.7×10^{-5}	E-A-Dibromopropane
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0



Compound Name: Ethylene oxide		
CAS Registry Number: 75-21-8	Molecular Weight	(g): 44.05
Parameters:		Reference
Water Solubility (ppm)	complete	Conway et al. (1983)
Boiling Point (°C)	10.7	Merck (1976)
Vapor Pressure (torr)	1305	Conway et al. (1983)
Molecular Weight/Oxygen	1.38	Calc
Log (Octanol/Water Partition Coefficient)	-0.22	CC-Kow
. Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M hr 1)	33.5	Mabey & Mill (1978)
Neutral Hydrolysis Rate Constant (hr)	2.43 x 10 ⁻³	Mabey & Mill (1978)
Microbial Degradation Rate Constant (ml cell-1hr-1)		VF-NBD
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	OXRO2SO

Compound Name: Ethyl methanesulfor	nate —————————	
CAS Registry Number: 62-50-0	Molecular Weight	(g): 124.16
Parameters:	- 1	Reference
Water Solubility (ppm)	3.69×10^5	C-Sw f Kow
Boiling Point (°C)	85-86/10 torr	Aldrich (1982)
Vapor Pressure (torr)	0.206	Jaber and Gunderson (1983)
Molecular Weight/Oxygen	3.88	Calc
Log (Octanol/Water Partition Coefficient)	0.21	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell hr 1)		HF-NBD
Photolysis Rate Constant (hr-1)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02, S0

Compound Name: Ferric dextran		
CAS Registry Number: 9004-66-4	Molecular Weight(g): 5000-7500
Parameters:	1	Reference
Water Solubility (ppm)		
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen		
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr ⁻¹)		
Microbial Degradation Rate Constant (ml cell -1hr -1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name: D-Glucopyranose, 2-deoxy-2-(3-methyl-3-nitrosoureido)-			
CAS Registry Number: 18883-66-4	Molecular Weight	(g): 265.22	
Parameters:		Reference	
Water Solubility (ppm)	2.7×10^{11}	C-Sw f Kow	
Melting Point (°C)	115 (dec)	Merck (1976)	
Boiling Point (°C)			
Vapor Pressure (torr)			
Molecular Weight/Oxygen	8.29	Calc	
Log (Octanol/Water Partition Coefficient)	-6.20	CC-Kow	
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)			
Acid Hydrolysis Rate Constant (M hr-1)			
Neutral Hydrolysis Rate Constant (hr ⁻¹)			
Microbial Degradation Rate Constant (ml cell hr 1)	1 x 10 ⁻¹⁰	Е-КВ	
Photolysis Rate Constant (hr ⁻¹)			
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO	

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Compound Name: Glycidylaldehyde		
CAS Registry Number: 765-34-4	Molecular Weight	(g): 72.1
Parameters:		Reference
Water Solubility (ppm)	1.70×10^8	C-Sw f Kow
Boiling Point (°C)	112-113	SRI Chemical Handbook
Vapor Pressure (torr)	19.7	C-VP f bp
Molecular Weight/Oxygen	2.25	Calc
Log (Octanol/Water Partition Coefficient)	-1.55	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	8.86	E-A-Glycirdol
Neutral Hydrolysis Rate Constant (hr)	1.02 x 10 ⁻³	E-A-Glycirdol
Microbial Degradation Rate Constant (ml cell ⁻¹ hr ⁻¹)		HF-NBD
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

NO NHH | || | H₃C-N-C-N-NO:

Compound Name:Guanidine, N-nitroso-N-methyl-N'-nitro-		
CAS Registry Number: 70-25-7	Molecular Weight	(g): 147.09
Parameters:		Reference
Water Solubility (ppm)	2.35×10^8	C-Sw f Kow
Melting Point (°C)	118 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	4.60	Calc
Log (Octanol/Water Partition Coefficient)	-3.18 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-lhr-l)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO



Compound Name: <u>Kepone</u>		
CAS Registry Number: 143-50-0	Molecular Weight	(g): 490.6
Parameters:		Reference
Water Solubility (ppm)	9.9×10^{-3}	C-Sw f Kow
Melting Point (°C)	350 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	15.3	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	· · · · · · · · · · · · · · · · · · ·
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

These data were estimated for use in a preliminary assessment to be conducted by the EPA, and should not be used in more detailed assessments.

(a) This compound exists as the gem-dial in aqueous solution.

Compound Name: <u>Lasiocarpine</u>		
CAS Registry Number: 303-34-4	Molecular Weight((g): 411.5
Parameters:		Reference
Water Solubility (ppm)	1600	C-Sw f Kow
Melting Point (°C)	96.4-97	SRI Chemical Handbook
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	13.8	Calc
Log (Octanol/Water Partition Coefficient)	0.99	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		
Acid Hydrolysis Rate Constant (M hr-1)		
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-1hr-1)		NBD
Photolysis Rate Constant: (hr-1)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ¹¹	M-OX SO



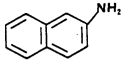
Compound Name: 2-Methylaziridine	(Propyleneimine)	
CAS Registry Number: 75-55-8	Molecular Weight	(g): 57.10
Parameters:		Reference
Water Solubility (ppm)	9.44×10^5	C-Sw f Kow
Boiling Point (°C)	66-67	
Vapor Pressure (torr)	141	C-vp f bp
Molecular Weight/Oxygen	1.78	Calc
Log (Octanol/Water Partition Coefficient)	-0.48	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M hr-1)	1.87×10^{-7}	E-A-Aziridine as cited by Mabey and Mill (1978)
Neutral Hydrolysis Rate Constant (hr 1)	2.5×10^{-3}	E-A-Aziridine as cited by Earley et al. (1958)
Microbial Degradation Rate Constant (ml cell-1hr-1)		HF-NBD
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT.	M-OX RO2, SO

Compound Name: Mustard Gas			
CAS Registry Number: 505-60-2	Molecular Weight	(g): 159.08	
Parameters:		Reference	
Water Solubility (ppm)	800	Franke (1967)	
Melting Point (°C)	13-14	Merck (1976)	
Boiling Point (°C)	215-217	Merck (1976)	
Vapor Pressure (torr)	0.17	Franke (1967)	
Molecular Weight/Oxygen	4.94	Calc	
Log (Octanol/Water Partition Coefficient)	1.37 (partial)	CC-Kow	
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)			
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)			
Neutral Hydrolysis Rate Constant (hr ⁻¹)	·		
Microbial Degradation Rate Constant (ml cell-hr-1)		HF-NBD	
Photolysis Rate Constant (hr-1)	PNER		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	2 x 10 ¹⁰	_M-OX SO	

2,7-Naphthalenedis Compound Name: 4,4'-div1)-bis(azo	ulfonic acid, 3,3'-) bis(5-amino-4-hyd	[(3,3'-dimethyl-(1,1'-bipheroxy)-tetrasodium salt
CAS Registry Number: 72-57-1	Molecular Weight	(g): 960.83
Parameters:		Reference
Water Solubility (ppm)	3.8 x 10 ⁸	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	30.0	Calc
Log (Octanol/Water Partition Coefficient)	1.76 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr 1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	1 x 10 ⁷	M-OX RO2



Compound Name:l-Naphthylamine		
CAS Registry Number: 134-32-7	Molecular Weight	(g): 143.18
Parameters:		Reference
Water Solubility (ppm)	2.35×10^3	C-Sw f Kow
Melting Point (°C)	50.	Verschueren (1977)
Boiling Point (°C)	301	Merck (1976)
Vapor Pressure (torr)	6.5 x 10 ⁻⁵	C-VP f bp
Molecular Weight/Oxygen	4.47	Calc
Log (Octanol/Water Partition Coefficient)	2.07	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	6 x 10 ⁶	OX RO2



Compound Name: 2-Naphthylamine		
CAS Registry Number: 91-59-8	Molecular Weight	(g): 143.18
Parameters:		Reference
Water Solubility (ppm)	586	C-Sw f Kow
Melting Point (°C)	110.2	Verschueren (1977)
Boiling Point (°C)	306	Merck (1976)
Vapor Pressure (torr)	2.56×10^{-4}	Karyakin et al. (1968)
Molecular Weight/Oxygen	4.47	Calc
Log (Octanol/Water Partition Coefficient)	2.07	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M hr-1)	NHFG	
Neutral Hydrolysis Rate Constant (hr 1)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	4 x 10 ⁶	OX RO2

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Compound Name: <u>Nickel carbonyl</u>	· · · · · · · · · · · · · · · · · · ·	
CAS Registry Number: 13463-39-3	Molecular Weight	(g): <u>170.73</u>
Parameters:		Reference
Water Solubility (ppm)		
Boiling Point (°C)	43	Merck (1976)
Vapor Pressure (torr)	~400	Baev and Fedulova (1973)
Molecular Weight/Oxygen	5.34	Calc
Log (Octanol/Water Partition Coefficient)		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	NHFG	
Microbial Degradation Rate Constant (ml cell-1hr-1)		NBD
Photolysis Rate Constant (hr ⁻¹)		r
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

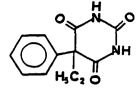


Compound Name: N-Nitrosopiperidine			
CAS Registry Number: 100-75-4	Molecular Weight	(g):114	
		•	
Parameters:		Reference	
Water Solubility (ppm)	1.9 x 10 ⁶	C-Sw f Kow	
Boiling Point (°C)	215/721_torr	Klein (1982)	
Vapor Pressure (torr)	1.4×10^{-1}	Klein (1982)	
Molecular Weight/Oxygen	3.56	Calc	
Log (Octanol/Water Partition Coefficient)	-0.49	CC-Kow	
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG		
Acid Hydrolysis Rate Constant (M hr-1)	NHFG		
Neutral Hydrolysis Rate Constant (hr)	NHFG	1-1	
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	E-KB	
Photolysis Rate Constant (hr ⁻¹)			
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO	



Compound Name: N-Nitrosopyrrolidine					
CAS Registry Number: 930-55-2	Molecular Weight	(g): 100.12			
		•			
Parameters:		Reference			
Water Solubility (ppm)	7.0×10^6	C-Sw f Kow			
Boiling Point (°C)	214	Aldrich (1982)			
Vapor Pressure (torr)	1.1 × 10 ⁻¹	Klein (1982)			
Molecular Weight/Oxygen	3.13	Calc			
Log (Octanol/Water Partition Coefficient)		CC-Kow			
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG				
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG				
Neutral Hydrolysis Rate Constant (hr)	NHFG				
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	E-KB			
Photolysis Rate Constant (hr ⁻¹)					
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2			

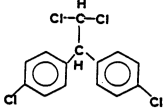
Compound Name: 1,2-0xathiolane, 2	,2-dioxide	-		
CAS Registry Number: 1120-71-4	Molecular Weight(g):122.1			
Parameters:	6 1	Reference		
Water Solubility (ppm)	$\frac{1.7 \times 10^6}{}$	C-Sw f Kow		
Boiling Point (°C)				
Vapor Pressure (torr)				
Molecular Weight/Oxygen	3.82	Calc		
Log (Octanol/Water Partition Coefficient)	-0.40	CC-Kow		
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)				
Acid Hydrolysis Rate Constant (M hr-1)				
Neutral Hydrolysis Rate Constant (hr ⁻¹)				
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	E-KB		
Photolysis Rate Constant (hr^{-1})	PNER			
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-0X R02 S0		



Compound Name: Phenobarbital		
CAS Registry Number: 50-06-6	Molecular Weight	(g): <u>232.23</u>
Parameters:		Reference
Water Solubility (ppm) Melting Point (°C)	1000 174-178	Merck (1976) Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	7.26	Calc
Log (Octanol/Water Partition Coefficient)		CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		,
Acid Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	· · · · · · · · · · · · · · · · · · ·	
Neutral Hydrolysis Rate Constant (hr)		
Microbial Degradation Rate Constant (ml cell-lhr-l)	1 x 10 ⁻¹⁰	E-KB
Photolysis Rate Constant (hr^{-1})		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2. SO

Compound Name: 1-Propanol, 2,3-di tris(2,3-dibromop)		3:1)
CAS Registry Number: 126-72-7		(g):698
		•
Parameters:		Reference
Water Solubility (ppm)	120	C-Sw f Kow
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	21.81	Calc
Log (Octanol/Water Partition Coefficient)	4.12 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	HNES	
Acid Hydrolysis Rate Constant (M ¹ hr ⁻¹)	HNES	
Neutral Hydrolysis Rate Constant (hr ⁻¹)	1.15 x 10 ⁻³	E-A-Naled
Microbial Degradation Rate Constant (ml cell hr -1)	3 x 10 ⁻¹²	Е-КВ
Photolysis Rate Constant (hr ⁻¹)	PNER	
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO

Compound Name:	ropanenitrile,	2-hydroxy-2-methyl-	(Acetone cyanohydrin)
CAS Registry Numbe	r: <u>75-86-5</u>	Molecular Weight((g): 85.11
Parameters:		c 1	Reference
Water Solubility (ppm)	1.52×10^6	C-Sw f Kow
Boiling Point (°C)		95	Weber, et al. (1981)
Vapor Pressure (to	rr)	44.9	C-VP f bp
Molecular Weight/0	xygen	2.66	Calc
Log (Octanol/Water Coefficient)	Partition	-0.51	CC-Kow
Alkaline Hydrolysi Constant (M ⁻¹ hr ⁻¹)	s Rate	HNES	
Acid Hydrolysis Ra Constant (M hr -1)	te	HNES	
Neutral Hydrolysis Constant (hr 1)	Rate	HNES	
Microbial Degradat Rate Constant (ml	cell ⁻¹ hr ⁻¹)	3 x 10 ⁻⁹	E-KB
Photolysis Rate Co	onstant (hr ⁻¹)	PNER	
Oxidation Rate Cor	stant $(M^{-1}hr^{-1})$	INERT	M-OX RO2, SO



Compound Name: TDE (1,1-Dichloro-2,2-bis (4-chlorophenyl)ethane)					
CAS Registry Number: 72-54-8	Molecular Weight(g): 320.1				
Parameters:		Reference			
Water Solubility (ppm)	2.2×10^{-2}	C-Sw f Kow			
Melting Point	109-110	Merck (1976)			
Boiling Point (°C)					
Vapor Pressure (torr)					
Molecular Weight/Oxygen	10.0	Calc			
Log (Octanol/Water Partition Coefficient)	6.21	CC-Kow			
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)	NHFG	· · · · · · · · · · · · · · · · · · ·			
Acid Hydrolysis Rate Constant (M hr-1)	NHFG				
Neutral Hydrolysis Rate Constant (hr 1)	NHFG				
Microbial Degradation Rate Constant (ml cell-1hr-1)	1 x 10 ⁻¹⁰	Е-КВ			
Photolysis Rate Constant (hr^{-1})	PNER				
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2, SO			

Compound Name: <u>Uracil</u> , 5-[bis(2-c	hloroethyl)amino]-(Uracil mustard)
CAS Registry Number: 66-75-1	Molecular Weight	(g): 252.1
Parameters:		Reference
Water Solubility (ppm)	641	C-Sw f Kow
Melting Point (°C)	206 (dec)	Merck (1976)
Boiling Point (°C)		
Vapor Pressure (torr)		
Molecular Weight/Oxygen	7.88	Calc
Log (Octanol/Water Partition Coefficient)	-1.09 (partial)	CC-Kow
Alkaline Hydrolysis Rate Constant (M ⁻¹ hr ⁻¹)		E-A-NM
Acid Hydrolysis Rate Constant (M hr-1)		E-A-NM
Neutral Hydrolysis Rate Constant (hr)		E-A-NM
Microbial Degradation Rate Constant (ml cell-1hr-1)	3 x 10 ⁻¹²	E-KB
Photolysis Rate Constant (hr ⁻¹)		
Oxidation Rate Constant (M ⁻¹ hr ⁻¹)	INERT	M-OX RO2. SO

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SECTION 4

CALCULATION OF PARTITION COEFFICIENTS OF ORGANIC CHEMICALS IN AQUATIC ENVIRONMENTS

This section was taken in whole from W. R. Mabey, J. H. Smith, R. T. Podoll, et al., "Aquatic Fate Process Data for Organic Priority Pollutants," EPA Report No. 440/4-81-014, December 1982.

Section 4

CALCULATION OF PARTITION COEFFICIENT OF ORGANIC CHEMICALS IN AQUATIC ENVIRONMENTS

4.1 BACKGROUND

The partitioning of a chemical between water and sediment and between water and biota will affect the concentration of the chemical in water and the rate of loss of the chemical from aquatic systems. Solubility data, on the other hand, are required for calculation of Henry's constants, which are needed to calculate volatilization rates of chemicals in aquatic systems.

This section discusses the relationships between water solubility, the partition coefficients for a chemical between sediment and biota, and the partition coefficient for a chemical between octanol and water.

Moreover, the theoretical basis for such relationships is explained, and some of the published correlations for these data are discussed. This section also briefly discusses the calculation of the octanol-water partition coefficient data used to calculate many of the other partitioning constants.

As discussed in Section 2, the partitioning of a chemical is given by the equation

$$K_{p} = C_{p}/C_{w} \tag{4.1}$$

where C_p and C_w are the concentrations on a particulate material (sediment or biota) and in water, respectively, and K_p is the partitioning constant (or coefficient) whose units are determined by those of C_p and C_w (see section 2). In practice, C_p is usually defined as the amount of chemical per dry weight of sediment (or organisms) to correct for the variability of the particulate water content. The partition coefficient between

This section was taken in whole from W. R. Mabey, J. H. Smith, R. T. Podoll, et al., "Aquatic Fate Process Data for Organic Priority Pollutants," EPA Report No. 440/4-81-014, December 1982.

microorganism and water, K_B , given for individual organic chemicals in Section 3, is in units of micrograms of chemical per gram of microorganism divided by grams of chemical per liter of water. Because the amount of organic chemical sorbed to sediments has been found to depend on the amount of organic carbon in the sediment, it is useful to normalize a measured sediment partition coefficient (K_p) for organic carbon content:

$$K_{oc} = K_{p}/f_{oc} \tag{4.2}$$

where $f_{\rm oc}$ is the fraction of organic carbon and $K_{\rm oc}$ is the normalized (for organic carbon content) partition coefficient. Karickhoff et al. (1979) have also shown that, because $f_{\rm oc}$ varies with sediment particle size, the distribution of sediment particle size will markedly affect measured $K_{\rm oc}$ values.

The octanol-water partion coefficient K_{ow} has commonly been used as a measure of the hydrophobicity of a chemical in medical and toxicological applications as well as in environmental chemistry (Hansch and Leo, 1979; Kenaga and Goring, 1978). A large number of K_{ow} values is therefore available as a result of the number of uses of such data. Most significantly, K_{ow} values can be calculated from molecular structure (see Section 4.4). The K_{ow} data in Section 3 are given to allow calculations of other properties (partitioning coefficients for biota as well as toxicological data) for use in environmental assessments of the organic priority pollutants.

4.2 CALCULATION METHODS

Several correlation equations have been proposed to calculate the water solubility (S_w) , K_{oc} , and K_{B} from K_{ow} values and to calculate K_{oc} values from water solubility. The more widely used of these equations are discussed and analyzed in Section 4.3. Although we recognize that better equations are evolving as more experimental data are obtained, the following equations are recommended for use in environmental fate assessments.

4.2.1 Correlation Equations

In the following equations, all partition coefficients (K_{oc} , K_{ow} and K_{g}) are unitless, and water solubility (S_{w}) is in units of parts per million (ppm). As discussed in Section 4.2.2, however, the solubility units of molarity (moles per liter) or mole fraction are preferred.

 K_{oc} and K_{ow} are correlated by the following equation (Karickhoff, 1979):

$$\log K_{OC} = 1.00 \log K_{OW} - 0.21$$
 (4.3)

Correlation of S_w and K_{ow} was reported by Yalkowsky and Valvani (1980). For organic pollutants that are liquid in their pure state at 25°C:

$$\log S_{w} = -1.08 \log K_{ow} + 3.70 + \log MW$$
 (4.4)

where MW is the molecular weight of the pollutant (g mole $^{-1}$). For organic pollutants that are solid in their pure state at 25°C:

$$\log S_{w} = -1.08 \log K_{ow} + 3.70 + \log MW - \left(\frac{\Delta S_{F}}{1360}\right) \pmod{-25}$$
 (4.5)

where mp is the melting point of the pollutant (°C) and ΔS_f is the entropy of fusion of the pollutant (cal mol⁻¹ deg⁻¹). If ΔS_f is not known, it may be approximated by (Yalkowsky and Valvani, 1980):

$$\Delta S_f \sim 13.6 + 2.5 (n - 5)$$
 (4.6)

where n is the number flexible atoms (i.e., atoms not involved in double bonds, triple bonds, or part of a ring structure) in the pollutant molecule, other than hydrogen. If n is less than 5, (n - 5) is set equal to zero.

The original equations in the literature are different if they were reported in different solubility units. Refer to Section 4.2.2 for the appropriate solubility units conversion factors.

Correlation of K_{OC} and S_{W} is provided by (Kenaga and Goring, 1978):

$$\log K_{QC} = -0.55 \log S_{W} + 3.64$$
 (4.7)

 K_{B} can be correlated with K_{OW} by

$$K_{B} = 0.16 K_{ow}$$
 (4.8)

4.2.2 Units and Conversion Factors

Three commonly used units of aqueous solubility are defined below:

(1) Mole fraction, x, the unitless ratio of the number of moles of solute to the total number of moles of solute plus water. In symbols, for a binary solution of n moles of solute in n moles of water

$$x = n/(n + n_w)$$

$$\sim n/n_w \text{ for } n_w >> n$$
(4.9)

(2) Molarity, S, expressed in moles of solute per liter of solution (M):

$$S(M) = n(mol)/liter of solution$$
 (4.10)

(3) Weight fraction, expressed in milligrams of solute per liter of water, or parts per million, ppm

$$S_{w} (ppm) = \frac{n (mol) MW (g mol^{-1}) 1000 (mg g^{-1})}{1 iter of water}$$
 (4.11)

where MW is the molecular weight of the solute.

For solutions with S < 1 M, one liter of aqueous solution contains approximately 55.5 moles of water. Thus

$$x = \frac{S}{55.5 + S} \sim \frac{S}{55.5}$$
 for S < 1 M (4.12)

or

$$S = \frac{55.5 \text{ x}}{(1-x)} \sim 55.5 \text{ x for } x < 10^{-2}$$
 (4.13)

To convert from molarity to ppm is straightforward by substituting Equation (4.10) into equation (4.11)

$$ppm \sim S(MW)$$
 (1000) for S < 1 M (4.14)

Thus to convert from mole fractions to ppm follows from equations (4.11) and (4.13)

$$ppm = \frac{55.5 \text{ x}}{(1 - \text{x})} \text{ (MW)} \text{ (1000)}$$

$$\sim 55.5(\text{x}) \text{ (MW)} \text{ (1000) for } \text{x} < 10^{-2}$$
(4.15)

These conversion factors are summarized in Table 4.1.

Table 4.1

CONVERSION FACTORS FOR COMPOSITION UNITS

FROM	O ppm	x (mole fraction)	M (Molarity)
ррш		$\frac{1.80 \times 10^{-5}}{MW}$	10 ⁻³ MW
x (mole fraction)	5.55 x 10 ⁴ (MW)		55.5
M (Molarity)	(MW) (10 ³)	<u>1</u> 55.5	

Concentration in aqueous solution is preferably given in mole fraction or molarity units since these units are measures of the amount of solute per amount of solution. The weight fraction or ppm, on the other hand, expresses the weight of solute per weight of solution and is thus a function of the molecular weight of the molecule, which is not relevant to environmental or toxicological effects.

4.3 CALCULATION OF K and S FROM K ow

The sediment partition coefficient, normalized for organic carbon content (K_{oc}) , and aqueous solubility (S_{w}) of an organic pollutant are critical to its environmental fate. Because K_{oc} and S_{w} values may be unmeasured or unreliable, it is important to be able to correlate these environmental parameters with other experimental quantitites, namely, to predict unmeasured values and appraise the reliability of measured values.

It is useful to correlate these parameters with octanol/water partition coefficients (K_{ow}) for practical as well as theoretical reasons. Practically, K_{ow} values are easier to measure and, where K_{ow} measurements have not been made, calculated values may be used with confidence. The theoretical basis for expecting correlations of K_{oc} and S_{w} with K_{ow} is described below. The correlation of K_{B} with other partitioning constants is not discussed in this section since a recent review of the subject is available.

4.3.1 Partitioning Thermodynamics

This discussion first considers the partitioning of a chemical between octanol and water, with octanol being a representative organic phase. If a small amount of a chemical is added to a closed vessel containing n-octanol and water, the vessel is shaken, and the octanol and water are allowed to separate, the chemical will partition between the two phases (see Figure 4.1). By convention, the small amount of chemical in each phase is called the solute. The partitioning of the solute molecules between the two phases can be understood in terms of a simple lattice model. If we assume that every molecule (water, octanol,

C)	0	\$)	0
0		0	()	0
	S	0	()	0
W	w	w	w	W	w
W	W	W	S	W	W
w	W	W	W	W	w
w	w	W	w	w	w
	w w w	s W W W W	0 0 s 0 w w w w w w w w w	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	O O O S O O W W W W W W W S W W W W W W

SA-6729-8

FIGURE 4.1 LATTICE MODEL OF A SOLUTE (S)
PARTITIONING BETWEEN OCTANOL
(O) AND WATER (W) PHASES $K_{ow} = C_o/C_w = 2$

SA-6729-9

FIGURE 4.2 LATTICE MODEL OF A HIGHER MOLE FRACTION OF SOLUTE (S) PARTITIONING BETWEEN OCTANOL (O) AND WATER (W) PHASES

Because the environment of each solute molecule is the same, $K_{ow} = C_o/C_w = 2$ as in Figure 4.1.

and solute) in both phases occupies a particular site on a three-dimensional lattice, with uniform spacing between sites, then the fraction of sites in each phase occupied by the chemical is the mole fraction x. A two-dimensional cross section of this lattice is shown in Figures 4.1 and 4.2.

The tendency for a solute molecule to leave either phase is proportional to the solute mole fraction in that phase and to the forces acting on the solute in that phase. The forces acting on a solute molecule will depend on which molecules occupy neighboring sites on the lattice. Figures 4.1 and 4.2 show that, over the mole fraction range of $x_w = 1/28$ to $x_w = 1/14$, solute molecules in the water phase are surrounded by water molecules. Thus, the forces acting on the solute in the water phase are independent of the solute mole fraction. Consequently, the tendency (f) of a solute molecule to leave the water phase is directly proportional to its mole fraction:

$$f = Hx \tag{4.16}$$

where H is a constant representing the forces exerted on the solute by the solvent. At higher solute mole fractions, where solute-solute interactions become important (that is, where the solute is concentrated enough that solute molecules occupy neighboring lattice sites), H becomes a function [H(x)] of the solute mole fraction, and thus f is no longer directly proportional to x:

$$f = H(x) x \tag{4.17}$$

The partitioning of the chemical between the octanol and water phases depends on this relative tendency of the chemical to leave each phase (f), which is conveniently viewed as a force per unit area. In thermodynamics, f is called the fugacity and, as explained above, is proportional to the relative amount of the solute in the phase, x, and the forces acting on the solute within each phase; explicitly,

See, for example, G. L. Lewis and M. Randall, <u>Thermodynamics</u>, revised by K. S. Pitzer and L. Brewer (McGraw-Hill, NY, 1961).

$$f_{w} = (f^{R}\gamma_{w}) \times_{w}$$
 (4.18)

$$f_o = (f^R \gamma_o) x_o (4.19)$$

where subscripts w and o refer, respectively, to the water and octanol phases, and f^R and γ_i are, respectively, the reference fugacity and activity coefficient, which together represent the forces acting on the solute in the i^{th} phase. At equilibrium

$$f_{w} = f_{0} \tag{4.20}$$

so that

$$x_{o}/x_{w} = \frac{f^{R}\gamma_{w}}{f^{R}\gamma_{o}} = \frac{\gamma_{w}}{\gamma_{o}}$$
 (4.21)

In general, at constant pressure, f^R depends only on the temperature and γ_i depends on the composition as well as the temperature of the i^{th} phase. In sufficiently dilute solutions, however, the forces acting on a solute molecule will be independent of x_i because, as explained above, the environment of a solute molecule will remain constant. Thus $(f^R\gamma_i)$ will be a function only of temperature

$$(f^{R}\gamma_{i}) = H_{i}$$
 (4.22)

where \mathbf{H}_{i} is the Henry's constant for a very dilute solution of the solute in phase i. Thus

$$x_{o}/x_{w} = H_{w}/H_{o}$$
 (4.23)

is a function only of temperature. However, if x_0 or x_w is large enough that γ_0 or γ_w is not constant, then K_{ow} will also no longer be constant.

Because composition is commonly measured in moles liter $^{-1}(M)$, it is convenient to define:

$$K_{ow} = C_{o}/C_{w} = r_{wo}(x_{o}/x_{w}) = r_{wo}(H_{w}/H_{o})$$
 (4.25)

where r_{wo} is a constant equal to the ratio of the molar volume of water

$$r_{wo} = v_{w}/v_{o}$$
 (= 0.115) (4.26)

to that of octanol. (In terms of the lattice mode, r_{wo} is equal to the ratio of the number of sites per unit volume of octanol to that of water.)

Numerous workers have correlated the partitioning of chemicals between sediment and water and between biota and water with octanol/water partition coefficients. Before discussing these specific correlations in detail, it is useful to understand the conditions that must be met for these correlations to be successful.

Partitioning of a solute between water and any other water immiscible phase p (i.e., biota, sediment) may be described by

$$K_{pw} = r_{wp}(H_w/H_p) \tag{4.27}$$

From equation (4.25) for partitioning between octanol and water

$$H_{w} = K_{OW} H_{O}/r_{WO}$$
 (4.28)

thus

$$K_{pw} = (r_{wp}/r_{wo})(H_o/H_p)K_{ow} = r_{op}(H_o/H_p)K_{ow}$$
 (4.29)

where r is the ratio of the molar volume of octanol to that of phase p. Thus, taking the logarithm of both sides of equation (4.29)

$$\log K_{pw} = \log K_{ow} + \log (r_{op}H_{o}/H_{p})$$
 (4.30)

Thus, for the second term on the right-hand side of equation (4.30) to remain constant for a set of chemicals partitioning between water-octanol and water-phase p, phase p must be chemically similar to octanol and both $K_{\mbox{ow}}$ and $K_{\mbox{pw}}$ must be measured at low enough solute concentrations that solute-solute interactions are absent.

The success of K_{ow} - K_{ow} correlations (to be discussed in detail below), for example, may thus be understood. First, by normalizing adsorption for organic carbon content, we ensure the chemical similarity of phase p (that is, the organic content) and octanol. Second, the partitioning of the chemical between the water and sediment phases is usually measured at very low surface coverage (in the linear region of the adsorption isotherm) where adsorbate-adsorbate interactions are minimal.

Octanol/water partition coefficients have been used not only to correlate other partitioning data, but also to predict aqueous solubilities. The assumptions implicit in these predictions become apparent on close examination of the octanol/water partition experiment.

If it is assumed that the ratio of the number of solute molecules in each phase remains constant up to the limit of solubility, then

$$K_{ow} = (C_o/C_w)_{dilute} = (C_o/C_w)_{saturated}$$
 (4.31)

From equation (4.21), this means that the ratio of activity coefficients $\gamma_{\rm w}/\gamma_{\rm o}$ remains constant up to saturation. As explained above, however, the ratio $\gamma_{\rm x}/\gamma_{\rm o}$ will depend on solute concentration, particularly if $C_{\rm w}$ (saturated) is large enough that solute-solute interactions become

^{*}Because of the chemical similarity of a neutral organic solute with n-octanol, it is expected that γ_{o} will not vary significantly with C_{o} .

important. Furthermore, if we assume that the solubility of the chemical in pure water equals its solubility in the octanol-saturated water phase of the partition measurement, then

$$K_{OW} = S_O/S_W \tag{4.32}$$

where S_0 and S_w are solubilities in moles liter $^{-1}(M)$ in pure octanol and pure water, respectively.

To correlate aqueous solubility with $K_{\mbox{ow}}$, many authors have proposed an equation of the form:

$$\log S_w = -(1/a) \log K_{ow} + c$$
 (4.33)

where a and c are constants. Equation (4.33) may be derived by modifying equation (4.32) to account for deviations of real systems from model behavior:

$$K_{ow} = (S_o/S_w)^a$$
 (4.34)

This equation is clearly identical to equation (4.32) for a = 1. Taking the logarithm of both sides of equation (4.34) and rearranging terms:

$$\log S_w = - (1/a) \log K_{ow} + (1/a) \log S_o$$
 (4.35)

If S $_{\rm o}$ is assumed constant for a set of solutes in octanol, equation (4.35)becomes

$$\log S_{w} = - (1/a) \log K_{ow} + c$$
 (4.36)

and the correlation coefficients a and c may be calculated from a plot of known values of log S $_{\!\!\!W}$ versus known values of log K $_{\!\!\!\!OW}$ for the given

set of solutes. Clearly, if the assumptions implicit in equation (4.32) are reasonable, the calculated value of <u>a</u> should be close to one.

The variability of S_O for a set of solutes is difficult to quantify except by comparing liquid and solid solutes. If two solutes are identical except that one is a liquid and the other is a solid in its pure state at temperature T, the solid will be less soluble than the liquid because of the additional energy required to remove solute molecules from the solid phase. Thus, if we assume that all liquid solutes have the same solubilities in n-octanol, and we use this pure liquid solute as the reference state, calculated solid solubilities must be corrected for the energy necessary to transform the solid to the liquid state. This energy is called the enthalpy of fusion, and from simple thermodynamic arguments, we can modify equation (4.35) for solid solutes:

$$\log S_{w} = - (1/a) \log K_{ow} + c - (1/a) \frac{\Delta H_{f}}{2.303 \text{ RT}_{f}} \frac{T_{f} - T}{T}$$
 (4.37)

where $\Delta H_{\hat{f}}$ is the enthalpy of fusion, R is the gas constant, and $T_{\hat{f}}$ is the melting temperature of the solute. At the melting point,

$$\Delta H_{f} = T_{f} \Delta S_{f} \tag{4.38}$$

Therefore at 25°C, equation (4.38) becomes

$$\log S_w = - (1/a) \log K_{ow} + c - \frac{\Delta S_f}{a(1360)}$$
 (mp-25) (4.39)

where mp is the melting point (in $^{\circ}$ C) and $^{\circ}$ C is the entropy of fusion (in cal deg $^{-1}$ mole $^{-1}$). This correction is zero for solutes that are liquid at $^{\circ}$ C, but substantial for solutes with high melting points. Assuming that the theory is approximately correct and the correlation coefficient \underline{a} is approximately equal to one, Table 4.2 and Figure 4.3 illustrate the magnitude of this correction as a function of melting point for a hypothetical solute with an uncorrected solubility of 100 ppm and a typical entropy of fusion of 13.6 entropy units (cal deg $^{-1}$ mol $^{-1}$).

Table 4.2

EFFECT OF MELTING POINT CORRECTION
ON WATER SOLUBILITY VALUES

Solubility (uncorrected) (ppm)	Melting Point	Solubility* (corrected) (ppm)
100	25	. 100
100	50	56
100	100	18
100	200	2
100	300	0.2

^{*}log S_w (corrected) = log S_w (uncorrected) - 0.01 (mp-25) at 25°C, where ΔS_f = 13.6 and a = 1 are assumed in equation (4.39) and S_w is the water solubility in ppm.

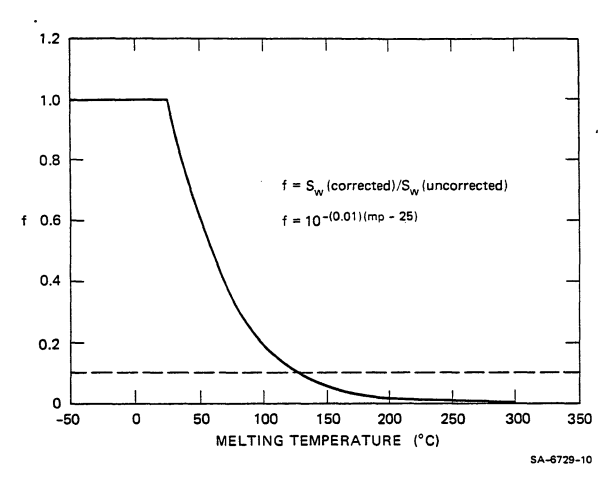


FIGURE 4.3 ENTHALPY OF FUSION CORRECTION FACTOR FOR AQUEOUS SOLUBILITY AT 25°C AS A FUNCTION OF MELTING TEMPERATURE

4.3.2 Comparison of Reported Correlations

Table 4.3 lists a representative sample of recently published correlations among K_{ow} , K_{oc} , S_{w} . This section examines these correlations in detail.

 $\frac{K_{\text{oc}}-K_{\text{ow}}}{\text{ow}}$. As discussed earlier, the sorption constant K_{oc} is the amount of chemical adsorbed per unit weight of organic carbon in the sediment divided by the equilibrium concentration of the chemical in the water phase. This constant is useful because, once K_{oc} has been determined for a chemical, the sorption partition coefficient may be calculated if the fraction organic content (f_{oc}) is known:

$$K_p = K_{oc}(f_{oc}) = C_s/C_w$$
 (4.40)

where

K_D = Sorption partition coefficient

 f_{OC} = Fraction of organic content in the sediment (0 < 0C <1)

 C_s = Concentration of the adsorbed chemical

 $C_{_{\mathbf{W}}}$ = Equilibrium solution concentration.

Furthermore, it is useful to be able to predict $K_{\rm oc}$ values from the more easily measured $K_{\rm ow}$ values. The theoretical basis for expecting good $K_{\rm oc}^{\rm -}K_{\rm ow}$ correlations has been discussed above. Two recent $K_{\rm oc}^{\rm -}K_{\rm ow}$ correlations that have appeared in the literature are listed in Table 4.3. The significantly different correlation equations of Kenaga and Goring (1978) and Karickhoff et al. (1979) probably reflect the different data bases used to correlate $K_{\rm oc}^{\rm with}$ $K_{\rm ow}^{\rm ow}$.

Table 4.3 $\label{eq:correlations} \mbox{REPORTED CORRELATIONS OF } \mbox{K}_{\mbox{ov}}, \mbox{ } \mbox{K}_{\mbox{oc}} \mbox{ AND } \mbox{S}_{\mbox{w}}$

Correlation	Equation	Eq.	Data Base*	Authors
K _{oc} - K _{ow}	log K _{oc} = 0.544 log K _{ow} + 1.377	(4.41)	Pollutants	Kenaga and Goring (1978)
		•	(Aromatic hydrocarbons (8) Carboxylic acids and esters (5) Phosphorus containing insecticions (5) Ureas and uracils (7) Symmetrical triazines (6) Miscellaneous (14)	
			Adsorbents	
			Variety of soils	
Koc - Kow	log K = 1.00 log K - 0.21	(4.3)	Pollutants	Karickhoff et al. (1979)
	· · · · · · · · · · · · · · · · · · ·		Polycyclic aromatics (8) Chlorinated hydrocarbons (2)	
S _w - K _{ow}	log S _w = - 0.922 log K _{ow} + 4.184 S _w in ppm	(4.42)	Substituted benzenes and halobenzenes (12) Halogenated biphenyls and diphenyl oxides (11) Aromatic hydrocarbons (9) Phosphorus containing insecticides (16) Carboxylic acids and esters (9) Ureas and uracils (7) Miscellaneous (24)	Kenaga and Goring (1978)
S _w - K _{ow}	log x _s = - 1.08 log K _{ow} - 1.04	(4.43)	Simple aliphatics and aromatics in the following groups $(n = 114)$	Yalkowsky (1980)
	$- \left \frac{\overline{\Delta}S_{f}}{1360} (mp - 25) \right $		Alcohols Halogens Amines Carboxylic acids and esters Aldehydes and ketones	
	x is the mole fraction solubility at 25°C		Ethers Nitro compounds	
	ΔS_{f} is the entropy of fusion in caldeg $^{-1}$ mol $^{-1}$			
	mp is the melting point in ${}^{\rm O}$ C (if mp \leq 25	4		
	then the term in brackets is zero)			

 $[\]ensuremath{^{\bigstar}}$ Number in parentheses refer to the number of pollutants in the data base.

Correlation	Equation	Eq.	Data Base	Authors	
K _{oc} - S _w	$\log K_{oc} = -0.55 \log S_{w} + 3.64$	(4.7)	Similar to data base for equation (4.41)	Kenaga and Goring (1978)	
	in ppm				
K _{oc} - S	$\log K_{om} = -0.56 \log S_w + 0.70$	(4.44)	Pollutants	Chiou et al. (1979)	
	log K _{oc} = - 0.56 log S _u + 0.93	(4.45)	Polychlorinated biphenyls (3) Pesticides (4) Halogenated ethanes and propanes (6) Tetrachloroethene 1,2-Dichlorobenzene		
			Adsorbents		
			Willamette silt loam Miscellaneous other soils		
K _{oc} - S _w	$\log K_{oc} = -0.54 \log x_{g} + 0.44$	(4.46)	Similar to data base for equation (4.3)	Karickhoff (1979)	
	x in the mole fraction solubility		•		

 K_{om} is the sorption partition coefficient normalized for organic matter reported by Chiou et al. (1979). Assuming $K_a = 1.7 K_{om}$, equation (4.45) is derived.

The theoretical equation of Table 4.4,

$$\log K_{oc} = 1.00 \log K_{ow} + constant \qquad (4.47)$$

follows from assuming that the second term on the right-hand side of equation (4.30) is constant; the data base required for a good fit with equation (4.47) follows from the assumptions used in the derivation of equation (4.30). It is clear from Table 4.4 that the data base and correlation equation of Karickhoff et al. (1979) closely conform with the theoretical model; however, the data base and correlation equation of Kenaga and Goring (1978) do not.

The advantages and disadvantages of using these alternative equations are not as well defined, however. Although the equation of Karickhoff et al. (1979) conforms to a simple model and accurately predicts sorption coefficients from K $_{\rm ow}$ data for a limited class of organic chemicals, it has not been widely tested and may be highly inaccurate for a more universal set of pollutants and soil/sediments. The equation of Kenaga and Goring (1978), however, is strictly empirical and only roughly predicts $K_{\rm oc}$ values from $K_{\rm ow}$ data, but it is applicable to a more universal set of pollutant/adsorbent systems because of the data base used. When more precise $K_{\rm oc}$ and $K_{\rm ow}$ data are available, it will be of interest to assess the predictive value of both of these correlations for both the universal set and individual classes of pollutant/adsorbent systems. It may become apparent that several correlation equations may be required to adequately predict $K_{\rm oc}$ values from $K_{\rm ow}$ values for the variety of systems of interest.

 $\frac{S_w - K_{ow}}{ow}$. Several comparisons of the equations of Kenaga and Goring (1978) and Yalkowsky (1980) can be made. For reasons discussed earlier, the mole fraction units of solubility used by Yalkowsky are to be preferred to the ppm units used by Kenaga and Goring. In fact, to compare equation (4.42) of Kenaga and Goring with equation (4.43) of Yalkowsky, we must assume an average molecular weight for the chemicals in the data

Table 4.4 DATA BASES FOR $K_{OC}^{-}K_{OW}^{-}$ CORRELATIONS

		Kenaga and Goring (1978)	Karickhoff et al. (1979)	Theoretical
	$log K_{oc} =$	$0.54 \log K_{ow} + 1.38$	1.00 log K _{ow} - 0.21	1.00 log K _{ow} + constant
	K ow	Measured and calculated values compiled from literature	Measured by Karickhoff et al.	Measured for very dilute solution
133	K _{oc}	Calculated average values for each chemical from adsorption coefficients for widely differing soils	Measured values for the silt (high organic content) fractions of two natural sediments	Uniform organic content of soil/sediment. Mea-sured for adsorption from very dilute solutions
	Chemicals	Very wide range of organic classes	Nonpolar or slightly polar organics	Nonpolar organics

base of Kenaga and Goring. Converting equation (4.42) from ppm to mole fractions units

$$\log x_s = -0.922 \log K_{ow} - 0.56 - \log MW$$
 (4.48)

where \mathbf{x}_{s} is the mole fraction solubility and MW is the average molecular weight.

The variation of equation (4.48) with MW is shown in Figure 4.4 and compared with Yalkowsky's equation for liquid solutes. Two observations can be made about Figure 4.4. First, the molecular weight dependence of equation (4.48) is not very great for chemicals in the molecular weight range of 100-400. Second, because the average molecular weight of chemicals in the data base used to determine equation (4.48) is in the range of 100-400, it is clear that solubilities predicted by equation (4.48) will be approximately an order of magnitude lower than those predicted by equation (4.43).

A comparison of measured solubilities (in molarity units, M) with those predicted by the equations of Kenaga and Goring and of Yalkowsky is shown in Table 4.5 for a series of chlorinated methanes and ethanes. Note that all the chemicals listed in Table 4.5 (except hexachloroethane, which sublimes) are liquid at 25°C. Furthermore, is is clear from Table 4.5 that equation (4.43) of Yalkowsky predicts the aqueous solubility of chlorinated methanes and ethanes very accurately, whereas the corresponding prediction of equation (4.42) is an order of magnitude lower. Table 4.6, which compares calculated and measured solubilities for some low melting point aromatics, further supports these conclusions.

The cause of this discrepancy becomes clear when we examine the contrasting methods and data bases used by Kenaga and Goring and by Yalkowsky to develop their correlations. Kenaga and Goring empirically correlated Kow with the solubility of a set of chemicals, most of which are solid at 25°C. In other words, Kenaga and Goring implicitly used a solid solute reference state; consequently, their correlation equation cannot accurately predict the solubility of a chemical that is liquid at 25°C.

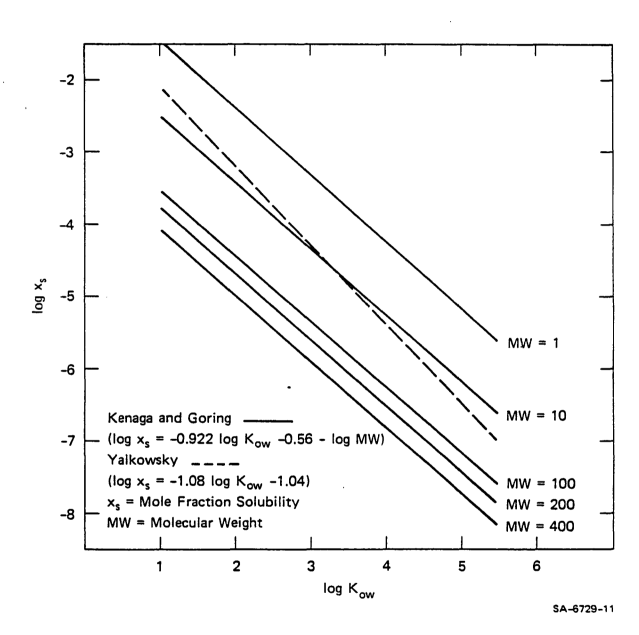


FIGURE 4.4 COMPARISON OF SOLUBILITY - Kow EQUATIONS FOR LIQUID SOLUTES

Table 4.5

CALCULATED VERSUS MEASURED SOLUBILITIES FOR CHLORINATED METHANES AND ETHANES

			log S _w		
	log K	mp (°C)	(M)		
			Kenaga and Goring	Yalkowsky	Measured
Chloromethane	0.95	-98	-1.4	-0.32	-0.89
Dichloromethane	1.26	- 95	-1.87	-0.66	-0.80
Chloroethane	1.49	-136	-2.03	-0.91	-1.05
1,1-Dichloroethane	1.80	-97	-2.45	-1.24	-1.25
Trichloromethane	r. 96	-64	-2.67	-1.41	-1.16
1,1.2-Trichloroethane	2.07	-37	-2.84	-1.53	-1.47
l,l,l-Trichloroethane	2.50	-30	-3.25	-2.00	-2.27
1,1,2,2-Tetrachloroethane	2.66	-36	-3.48	-2.17	-1.76
Tetrachloromethane	2.96	-23	-3.70	-2.49	-2.29
Hexachloroethane	4.62	Sublimes	-5.45	-4.29	-3.68

Yalkowsky, on the other hand, explicitly used a liquid solute reference state. To calculate the solubilities of chemicals that are solid at 25° C, Yalkowsky included an entropy of melting correction term. Thus the equation of Yalkowsky, assuming accurate known values of the entropy of fusion (ΔS_f) and melting point (T_f), is equally valid for liquid and solid solutes.

As discussed earlier, if two solutes are identical except that one is a liquid and the other is a solid in its pure state at 25° C, then the solid will be less soluble than the liquid by a factor of

$$\exp \left[-2.303(\Delta S_f/1360)(mp-25)\right]$$
 (4.49)

where ΔS_f is the entropy of fusion and mp is the melting point ($^{\circ}C$). If ΔS_f is constant, then it is clear from equation (4.46) that solubility decreases as the melting point increases. Assuming $\Delta S_f = 13.6$ entropy units and converting mole fraction solubilities to molarity units, Figure 4.5 illustrates that equation (4.43) of Yalkowsky, in contrast with equation (4.42) of Kenaga and Goring, successfully predicts the decrease in solubility with increase in melting point for α -, β -, δ -, and γ -BHC.

Figure 4.5 also indicates that implicit in equation (4.42) of Kenaga and Goring is an empirical average of the solid solute correction term. Because the solubilities of liquid solutes predicted by equation (4.42) are approximately an order of magnitude lower than measured values, we can assume that this average correction term is approximately equal to 0.10, which is the dashed line in Figure 4.3. Thus, the predicted solubilities of equation (4.42) should approximate those of Yalkowsky and measured values for solutes with melting points in the 100° to 200°C temperature range. Figure 4.6 illustrates, in fact, that for solutes with an approximate molecular weight of 150, an entropy of fusion of 13.6 and a melting point of 125°C, the correlation equations of Yalkowsky and of Kenaga and Goring are similar. Moreover, Table 4.7 illustrates

Table 4.6

CALCULATED VERSUS MEASURED SOLUBILITIES FOR LOW MELTING POINT AROMATICS

	_	mp	log S _w (M)		
	log K	<u>(°c)</u>	Kenaga and Goring	Yalkowsky	Measured
Nîtrobenzene	1.87	5.6	-2.63	-1.32	-1.82
Benzene	2.13	5.5	-2.63	-1.60	-1.64
Toluene	2.79	~9 5	-3.35	-2.31	-2.24
Chlorobenzene	2.84	-45	-3.48	-2.37	-2.37
Ethylbenzene	3.34	-94.9	-3.92	-2.90	-2.85
1,2-Dichlorobenzene	3.56	-17	-4.26	-3.14	-3.00

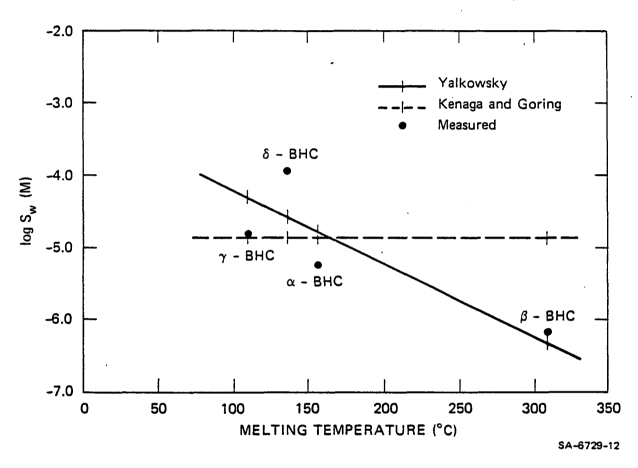


FIGURE 4.5 SOLUBILITIES OF HEXACHLOROCYCLOHEXANES (α -, β -, δ -, γ -BHC) AS A FUNCTION OF MELTING TEMPERATURE

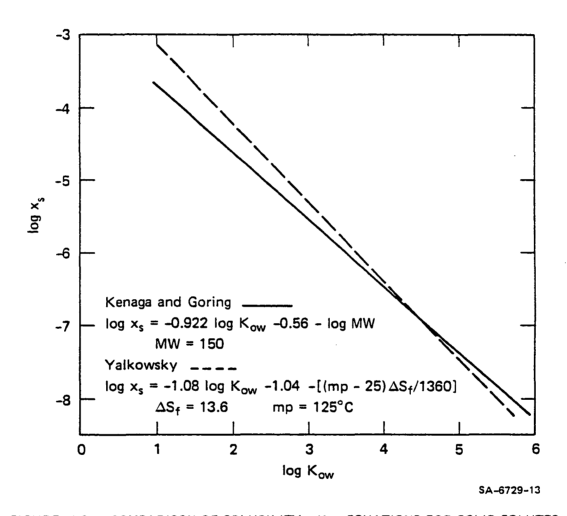


FIGURE 4.6 COMPARISON OF SOLUBILITY - Kow EQUATIONS FOR SOLID SOLUTES

Table 4.7

CALCULATED VERSUS MEASURED SOLUBILITIES FOR SELECTED PESTICIDES

		MP	log	S (M)	
	log K	(°c)	Kenaga and Goring	Yalkowsky	Measured
Lindane	3.89	113	-4.85	-4.38	-4.40 to -5.15
Aldrin	5.30	104	-6.24	-5.80	-6.30 to -7.35
Chlordane	5.48	108	-6.46	-6.04	-5.30 to -6.85
DDD	6.20	112	-7.04	-6.85	-6.5 to -7.2
DDT	6.91	109	-7.74	-7.59	-6.6 to -8.5

that for selected pesticides with melting points around 110°C the correlations of Yalkowsky and of Kenaga and Goring compare equally well with measured values.

Figure 4.5 also suggests that solubilities predicted from equation (4.42) of Kenaga and Goring will become progressively higher relative to measured values as the melting temperature increases above 200°C. Table 4.8 indicates that, indeed, measured solubilities of chemicals with melting points above 200°C systematically fall below those predicted by Kenaga and Goring.

In summary, equation (4.42) of Kenaga and Goring should be restricted to chemicals with melting points in the 100° to 200° C range, but equation (4.43) of Yalkowsky, because it includes a melting point correction factor is not limited by melting point restrictions.

 $\frac{K_{oc} - S_{w}}{w}$. To compare equation (4.7) with equations (4.45) and (4.46), it is again necessary to assume an average molecular weight for the correlation equation of Kenaga and Goring. If an average molecular weight of 200 is assumed, converting equations (4.7) and (4.45) to mole fraction solubility units gives

$$\log K_{\text{oc}} = -0.55 \log x_{\text{s}} - 0.23$$
 (Kenaga and Goring, 1978) (4.50)

$$\log K_{oc} = -0.56 \log x_{s} - 0.04$$
 (Chiou et al., 1979) (4.51)

$$\log K_{oc} = -0.54 \log x_{s} + 0.44$$
 (Karickhoff et al., 1979) (4.46)

Several observations can be made about these equations. First, the similarity of equations (4.50) and (4.51) is remarkable, considering the contrasting data bases used by Kenaga and Goring and by Chiou et al. to determine their correlation coefficients. In fact, equations (4.50), (4.51), and (4.46) may all be written in the form

$$K_{oc} = (constant) x_s^{-0.55(\pm 0.01)}$$
 (4.52)

Table 4.8

AQUEOUS SOLUBILITIES OF HIGH MELTING POINT CHEMICALS

	M 1nd D ton	Solubilities (ppm)	
Chemical Name	Melting Point	Measured	Predicted by Equation (4.42)
Benzo[k]fluoranthene	217	5.6×10^{-4}	0.04
Anthracene	219	0.045	1.2
Benzo[g,h,i]perylene	222	2.6×10^{-4}	0.015
Chrysene	258	1.8×10^{-3}	0.1
Dibenz[a,h]anthracene	270	5×10^{-4}	9×10^{-3}
TCDD	303	2×10^{-4}	7.5×10^{-3}
В-ВНС	309	0.24	4.0

^a Kenaga and Goring (1978)

It is not clear why the solubility coefficient of $-0.55(\pm0.01)$ should appear in each of these correlations. If as expected from the above discussions [see equations (4.3), (4.42), and (4.43)],

$$\log K_{\text{oc}} = a \log K_{\text{ow}} + \text{constant}$$
 (4.53)

and

$$log K_{ow} = - a log x_s + constant$$
 (4.54)

where a \sim 1, then by substituting equation (4.54) into equation (4.53)

$$\log K_{oc} = -a^2 \log x_s + constant$$
 (4.55)

$$\sim$$
 - 1.0 log x_s + constant

It is also apparent that none of these three equations accounts for the variation in solubility and hence variation in $K_{\rm oc}$ value with the melting point of the adsorbed chemical. The difference in adsorption behavior between solid and liquid solutes, in general, has been well documented in the literature (see, for example, Kipling, 1965). In fact, Roe (1975) has accounted for this difference in terms of the solid solute correction factor discussed earlier in this report. Karickhoff et al. (1979), in discussing their relatively poor correlation of $K_{\rm oc}$ with $K_{\rm ow}$, mention that a correction term is probably needed in equation (4.40) to account for the enthalpy of fusion of the chemicals they studied.

 $K_{\rm B}-K_{\rm ow}$. The partitioning of organic chemicals has recently been reviewed by Baughman and Paris (1981), who noted the paucity of reliable data available for correlating $K_{\rm B}$ with other partitioning parameters. For the chemicals in Section 3, the following equation was used to calculate $K_{\rm B}$

$$K_{R} = 0.16 K_{OM}$$
 (4.8)

which is the simplified version of the equation given by Baughman and Paris (1981),

$$\log K_{B} = 0.907 \log K_{OW} - 0.21$$
 (4.56)

The reader is referred to the above review for an excellent exposition on the problems of reliably measuring K_B and the use of correlation equations to calculate K_B from S_W or from K_{QC} or K_{QW} data.

4.4 CALCULATION OF K FROM STRUCTURAL PARAMETERS

The thermodynamics of partitioning of a chemical solution between octanol and water phases was discussed in 4.3.1, and the use of the octanol/water partition coefficient, K_{ow} , for calculating S_{w} , K_{oc} and K_{B} was described in Section 4.3.2. Although K_{ow} is the symbol used by many scientists for this partition coefficient, earlier literature and some current medical toxicology literature has commonly referred to the logarithm value of K_{ow} as "log P" (Hansch and Leo, 1979). For discussion in this section only, the log P nomenclature will be used instead of log K_{ow} , although the K_{ow} term will be used.

The $K_{\rm ow}$ data on the data sheets in Section 3 were calculated using a computer program developed at SRI; it uses the FRAGMENT method for calculating log P values (Hansch and Leo, 1979). The theory and procedures for these calculations are discussed in detail in that reference. Briefly, the method assumes that select groups of atoms in a molecule can be considered fragments, each of which contributes to the total log P value in an additive manner

$$\log P = \sum_{1}^{n} a_{n} f_{n}$$
 (4.57)

where a is the number of occurrences of fragment f of structural type n. Values of f have been empirically derived from the vast body of log P data available in the literature. Since the calculation of log P values

for complex molecules can be time-consuming and subject to numerous calculation errors, the FRAGMENT calculation method and the data base for fragment values have been incorporated into a computer program using the PROPHET computer network. The log P data are generated by first entering the structure on a graphic tablet. The log P program then uses an ordered substructure search routine to obtain fragment values for fragments of the molecular structure. Fragments are used, rather than atoms, because atomic contributions to log P vary with certain structural environments. The program then adds the fragment values to obtain log P values. It also identifies where the log P calculation may be incomplete because of the absence of values for particular fragments or because polar interactions must be accommodated by manual calculations. The log P program is under continuing development and evaluation at SRI and other laboratories.

The manual calculation of log P values using the FRAGMENT method is already established as a valid method for obtaining these data (Hansch and Leo, 1979). The calculations are, of course, subject to errors arising from subtle structural differences that are not recognized or cannot be accounted for when obtaining empirical values for the molecular fragments. In fact, the primary source of error is the original data on which the fragment values are based. The lack of reliable data is also a dilemma for verification of calculated log P values.

As an indicator of the accuracy of the log P calculation program Table 4.9 compares the K values recently published by Hassett et al. (1980) with the K values calculated by the log P program. Although the chemicals are not among the organic priority pollutants, they do represent some of the best K data currently available. The calculated and measured K values agree within the factor of two for 8 of the 14

^{*}PROPHET is a NIH resource available to biological and chemical scientists on a time-share basis. Information on the log P/PROPHET system can be obtained from Dr. Howard L. Johnson at SRI.

Table 4.9 CORRELATION OF MEASURED AND CALCULATED VALUES OF $\boldsymbol{K}_{\mbox{\scriptsize ow}}$

	2	Computer-Calculated	
Compound	Measured K ± S.D. ^a	K ow	r ^b
Pyrene	124,000 ± 11,000	79,400	1.6
7,12-Dimethylbenz[a]anthracene	953,000 ± 59,000	871,000	1.1
Dibenz[a,h]anthracene	$3,170,000 \pm 883,000$	5,890,000	0.54
3-Methylcholanthrene	2,632,000 ± 701,000	9,330,000	0.28
Dibenzothiophene	24,000 ± 2,200	33,900	0.71
Acridine	$4,200 \pm 940$	2,570	1.6
13H-Dibenzo[a,i]carbazole	2,514,000 ± 761,000	692,000	3.6
Acetophenone	38.6 ± 1.2	38.9	0.99
1-Napthol	700 ± 62	417	1.7
Benzidine	46.0 ± 2.2	35.5	1.3
2-Aminoanthracene	$13,400 \pm 930$	1,660	8.1
6-Aminochrysene	96,600 ± 4,200	24,000	4.0
Anthracene-9-carboxylic acid	$1,300 \pm 130$	15,500	0.08

 $^{^{\}rm a}$ Hassett et al. (1980). b Ratio of measured K $_{\rm ow}$ to calculated K $_{\rm ow}$

compounds listed and agree within a factor of five for 12 of the 14 compounds. It is also significant to note that the last three compounds in Table 4.9 show the most disagreement between calculated and measured $K_{\mbox{ow}}$ values, and these compounds are large molecules containing groups that may participate in H-bonding interactions.

In general, the accuracy of log P calculations by this method closely approaches the accuracy of experimental determinations performed over the last ten or twenty years because the fragment values were derived largely from those experimental data (by regression analysis) and incorporate the same experimental errors. It is not uncommon for measured log P values for a given compound in the literature to vary by 1 to 2 units; this corresponds to a factor of 10 to 100 in measured K_{OW} variation.

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15. SUPPLEMENTARY NOTES

16. ABSTRACT

Physical properties, equilibrium, and kinetic constants for evaluating the transformation and transport in aquatic systems for organic chemicals of interest to the U.S. Environmental Protection Agency's Office of Emergency and Remedial Response have been obtained from the literature and calculated from theoretical or empirical relations. Values for selected physical properties such as melting point, boiling point, vapor pressure, water solubility, and octanol/water partitioning, and for rate constants such as hydrolysis, microbial degradation, photolysis, and oxidation are listed for each chemical along with the source of the data. Values are reported in units suitable for use in a current aquatic fate model. A discussion of the empirical relationships between water solubility, octanol/water partition coefficients, and partition coefficients for sediment and biota is presented.

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