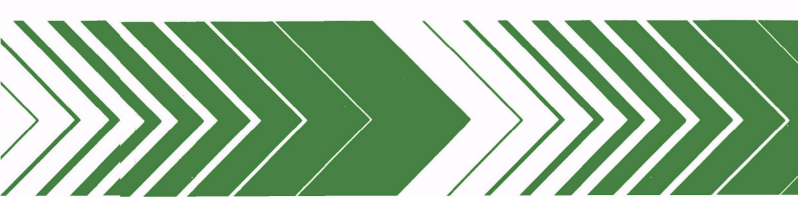
Research and Development



Adsorption of Energy-Related Organic Pollutants

A Literature Review



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ADSORPTION OF ENERGY-RELATED ORGANIC POLLUTANTS: A LITERATURE REVIEW

by

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FOREWORD

Environmental protection efforts are increasingly directed towards prevention of adverse health and ecological effects associated with specific compounds of natural or human origin. As part of this Laboratory's research on the occurrence, movement, transformation, impact, and control of environmental contaminants, the Environmental Processes Branch studies the microbiological, chemical, and physico-chemical processes that control the transport, transformation, and impact of pollutants in soil and water.

Efforts to achieve our national goal of energy independence will require increasing use of our country's vast domestic coal reserves. The combustion of coal or its conversion to a gaseous or liquid fuel, however, can release numerous organic compounds that are potentially toxic, carcinogenic, or mutagenic. This report reviews the literature on the adsorption of energy-related organic pollutants and other compounds on sediments and soils. Information on the adsorption of these pollutants onto sediments is needed to predict their movement and fate in aquatic systems so that potential environmental problems can be anticipated.

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ABSTRACT

This report is a literature review which was completed as the first phase of a research project on sorption properties of sediments and energy-related organic pollutants. Adsorption of organic compounds in general is discussed, and analytical methodology in soil thin-layer chromatography and chemical analysis as applicable to measurement of sorption properties is summarized. The literature on the adsorption of energy-related organic pollutants is reviewed. Reported constants for the adsorption of organic compounds on several adsorbents are tabulated, and factors which influence the adsorption are discussed.

This report was submitted in partial fulfillment of Contract No. 68-03-2555 by the University of Illinois Institute for Environmental Studies in cooperation with the Department of Agronomy under the sponsorship of the U.S. Environmental Protection Agency. This report covers the period of July 1977 to April 1978, and work was completed as of June 1979.

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ACKNOWLEDGMENTS

The authors gratefully acknowledge the assistance and coordination provided by Dr. David Brown, Project Officer, of the EPA Environmental Research Laboratory in Athens, Georgia. Co-principal investigators for the project under which this review was done were Dr. John J. Hassett, associate professor of soils, and Dr. Jay C. Means, assistant research chemist.

The literature review presented in this report was conducted under the direction of Dr. Keturah A. Reinbold, associate research biologist, with the assistance of Ms. Dee Condon and Ms. Carol Wells, research assistants, in conducting the literature search. Information on soil thin-layer chromatography was summarized by Dr. Wayne L. Banwart, assistant professor of soils. Appreciation is expressed to Thomas Knecht, publications director of the Institute for Environmental Studies, and Cindy Bohde, student editorial assistant, for the technical editing of this report and to Jean Clarke, Judith Jones, Trace Black, and Sharon Sparling for typing the report.

1 INTRODUCTION

Fossil fuels are a major source of anthropogenic organic compounds in the environment. Because supplies of petroleum and natural gas are dwindling, it will be necessary to utilize increasing amounts of coal to meet our energy needs in the near future. As Figure 1 illustrates, coal is a complex organic chemical. Either the combustion of coal or its conversion to a gaseous or liquid fuel breaks the coal into numerous simpler organic compounds, which then appear in process waste streams and may be released into the environment. These coal fragments include a great variety of polycyclic aromatics, heterocyclic- and carbonyl-polycyclics, and aromatic amines, groups which all contain known or suspected carcinogens, as well as phenolics and long-chain aliphatic hydrocarbons (TRW Systems and Energy, 1976; Sharkey et al., 1976).

Coal conversion processes produce large volumes of gaseous and aqueous effluents (Magee, Bertrand, and Jahnig, 1976). Hundreds of thousands of tons of gases containing volatile organics, particulates, and combustion gases are released each day. These emissions may disperse over hundreds of miles, distributing the effluent materials in both terrestrial and aquatic systems. With such large volumes of effluents, even trace components are distributed in significant amounts—up to hundreds of pounds per year. Some of these materials are potentially toxic, carcinogenic, or mutagenic.

Aqueous effluents may be released at a rate of up to 5 million gallons each day. Also, large volumes of aqueous leachate from stockpiles of coal or solid wastes, such as ash and char, are produced. The wastes from the various steps of the coal conversion process contain a variety of organic materials.

A number of organic products have been detected in the wastes from coal conversion pilot plants. Forney et al. (1974) identified some of the major organic compounds in tars produced by the Synthane coal gasification process (Table 1), and Schmidt, Sharkey, and Friedel (1974) analyzed the process water (Table 2). TRW Systems and Energy (1976) listed several organic compounds which are associated with the processing and utilization of coal and which are known or suspected carcinogenic or cocarcinogenic compounds (Table 3).

Clearly, large quantities of organic compounds are introduced into aquatic systems, either directly in aqueous effluents or indirectly from gaseous effluents. The behavior of these compounds in aquatic systems depends largely upon the extent to which they are adsorbed on suspended or settled sediments.

Figure 1. Functional group model of bituminous coal (Wiser).

Source: Wewerka, Williams, and Vanderborgh, 1976.

TABLE 1. ORGANIC CONTAMINANTS PRESENT IN COAL TAR FROM THE SYNTHANE COAL CONVERSION PROCESS

		Volu		
~	Illinois		Montana	Pittsburg
Structural Type	No. 6	Lignite	Subbituminous	Seam
Benzenes	2.1	4.1	3.9	1.9
Indenes	8.6	1.5	2.6	6.1
Indanes	1.9	3.5	4.9	2.1
Naphthalenes	11.6	19.0	15.3	16.5
Fluorenes	9.6	7.2	9.7	10.7
Acenaphthenes	13.5	12.0	11.1	15.8
3-ring aromatics	13.8	10.5	9.0	14.8
Phenyl naphthalenes	9.8	3.5	6.4	7.6
4-ring aromatics - peri	7.2	3.5	4.9	7.6
4-ring aromatics - cata	4.0	1.4	3.0	4.1
Phenols	2.8	13.7	5.5	3.0
Naphthols	+	9.7	9.6	+
Indanols	0.9	1.7	1.5	0.7
Acenaphthols	-	2.5	4.6	2.0
Phenanthrols	2.7	-	0.9	-
Dibenzofurans	6.3	5.2	5.6	4.7
Dibenzothiophenes	3.5	1.0	1.5	2.4
Benzonaphthothiophenes	1.7	-	-	-
N-heterocycles	10.8	3.8	5.3	8.8

Source: Forney et al., 1974

TABLE 2. CONTAMINANTS PRESENT IN PRODUCT WATER FROM THE GASIFICATION OF ILLINOIS NO. 6 COAL

Compound	Concentration (ppm)
Phenols	2,660 to 3,400
Cresols	2,610 to 2,840
C ₂ -phenols	560 to 1,170
C ₃ -phenols	70 to 150
Dihydric phenols	60 to 300
Benzofuranols	70 to 120
<pre>Indanols }</pre>	
Acetophenones	60 to 210
Benzoic Acids	
Hydroxybenzaldehyde	40 to 210
Naphthols	110 to 160
Indenols	90
Benzofurans	10 to 30
Dibenzofurans	10
Biphenols	20 to 40
Benzothiophenols	60 to 110
Pyridines	60 to 580
Quinolines	10 to 20
Indoles	20 to 70

Source: Schmidt, Sharkey, and Friedel, 1974.

OBJECTIVES

To examine the adsorption of energy-related organic compounds in the environment, a research project was initiated in the Institute for Environmental Studies at the University of Illinois at Urbana-Champaign. The project is supported by the U.S. Environmental Protection Agency under contract number 68-03-2555. This report presents the results of the first phase of the project, the objective of which was to review published literature on (1) the adsorption of energy-related organic pollutants on sediments, (2) the theory of adsorption, and (3) analytical techniques pertinent to adsorption measurements. Because the literature survey produced only a limited amount

of information on the adsorption of energy-related organic compounds, a variety of organic compounds were included in this review to provide a background of information.

APPROACH

The literature was searched by a combination of computer and manual methods. The 1970-77 volumes of *Chemical Abstracts* were searched using-the computerized Bibliographic Retrieval Service System. The key words used were terms such as *adsorption*, *desorption*, and *sediments* accompanied by the names of compound groups or of specific compounds which may be energy-related organic pollutants, including all those listed in Table 4. Earlier volumes of *Chemical Abstracts* were searched manually, as were pertinent books, series of *Residue Reviews*, current journals, and the 1977 Weekly Government Abstracts from the National Technical Information Service.

The computerized search produced more than 900 citations. Copies of approximately one-third of these references were obtained for review. With the addition of those acquired by manual searching, a total of 670 references were obtained for review. A bibliography is included at the end of this report.

TABLE 3. CLASSES OF KNOWN OR SUSPECTED CARCINOGENIC OR COCARCINOGENIC COMPOUNDS ASSOCIATED WITH PROCESSING AND UTILIZATION OF COAL

Compound Class	Representative Compound	Structure
Polynuclear Aromatic Hy	drocarbons	СН ₃ ,
Anthracenes	9-,10-dimethylanthracene	CH ₃
Chrysenes	chrysene	
Benzanthracenes	benzo(a)anthracene	
Fluoranthenes	benzo(j)fluoranthene	
Cholanthrenes	3 -methylcholanthrene	сн ₃

TABLE 3, continued

Compound Class	Representative Compound	Structure		
Benzopyrenes	benzo(a)pyrene			
Dibenzpyrenes	dibenzo(a,h)pyrene			
Nitrogen-, Sulfur-, an	d Oxygen-Containing Polycyclic C	Compounds		
Mono- and dibenzacridines	dibenz(a,h)acridine			
Benzocarbazoles	7H-benzo(c,g)carbazole			
B enza thrones	7H-benz(d,e)anthracen-7-one			
Aromatic Amines		O		
Aminoazobenzenes	4-dimethylaminœzobenzene	(CH ₃) ₂ N \ \ \ \ \ \ N=N \ \ \ \ \ \		
Naphthylamines	α -naphthylamine	NH ₂		
Cocarcinogens and Promoting Agents				
Phenols/naphthols	α-naphthol			
Long-chain aliphatic hydrocarbons	n-dodecane	CH ₃ -(CH ₂) ₁₀ -CH ₃		

Source: TRW Systems and Energy, 1976.

TABLE 4. SELECTED ENERGY-RELATED ORGANIC COMPOUNDS

Polynuclear Aromatics	S-Heterocyclics	Phenolics	
Anthracene Phenanthrene Acenaphthene Fluorene	2,3-Benzothiophene Dibenzothiophene	l-Naphthol 2-Naphthol Acenaphthol 4-Indanol	
Naphthacene Chrysene Pyrene Triphenylene	N-Heterocyclics Carbazole Indole Acridine Pyridines Pyrroles Benzocarbazole Dibenzocarbazole Benzoquinoline	4-Indahol 4-Benzofuranol 4-Hydroxybenzothiophene 2,3,4-Trimethyl Phenol	
Perylene 1,2-Benzopyrene 3,4-Benzopyrene 1,2-Benzanthracene 1,2,7,8-Dibenzanthracene 1,2,5,6-Dibenzanthracene 1,2,3,4-Dibenzanthracene 7,12-Dimethylbenzanthracene 3-Methylcholanthrene		Miscellaneous Acetophenone Anthraquinone Benzidine Benzophenone	

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2 SUMMARY AND CONCLUSIONS

We have reviewed the literature on the adsorption of energy-related organic pollutants and other organic compounds on sediments, soils, and other selected adsorbents. Adsorption constants reported in the literature for organic compounds are compiled in Tables 6 and 7, and factors which influence adsorption are discussed.

Among the many factors which influence adsorption are several molecular properties of compounds. It has been shown that as chain length, molecular volume, molecular weight, and carbon number increase, and as polarity decreases, adsorption of hydrophobic compounds increases. These properties are related to water solubility of the compound. The adsorption of hydrophobic compounds increases with decreasing water solubility, but for more polar compounds adsorption increases with decreasing water solubility only within a family of compounds. Soil organic matter content influences adsorption and has been shown to correlate with the partitioning of nonpolar organic compounds between octanol and water. This relationship makes possible the calculation of Koc and Kp values when Kow is known.

Little of the published information pertains specifically to the adsorption of energy-related organic pollutants onto sediments or soils. Data are available on the adsorption of a few such compounds onto carbon, but those results cannot be directly extrapolated to sediments or soils. Investigation of adsorption of these compounds on sediments is beginning, as in the laboratory phase of this project. Information on the adsorption of these pollutants onto sediments is needed to predict their movement and fate in aquatic systems. To obtain sufficient information, further research is necessary.

3 ADSORPTION OF ORGANIC COMPOUNDS

J. J. Hassett

THE SOLID-SOLUTION INTERFACE

The adsorption or concentration of organic materials at the solid-solution interface is of particular importance in natural waters. Streams, lakes, and rivers contain a variety of colloidal materials both in their bottom sediments and dispersed throughout the aqueous phase. These colloids can actively sorb organic materials, removing them from solution or suspension, and hence can have a marked effect on the chemical and physiological properties of the organic pollutants.

The sediments in a natural body of water consist predominantly of organic colloids (such as clay minerals and metal oxides, hydroxides, and carbonates), of organic colloids, and of living organisms. The metal oxides and hydroxides and the organic colloids may exist as discrete particles or as coatings on other colloids such as the clay minerals. A major source of these inorganic colloids (and, to a certain extent, of the organic colloids) is the erosion of watershed soils. The properties of the sediments can be quite similar to those of the soils from which they are derived, or they may be substantially altered from the original soil, since the sediments may be subject to physical sorting by the water and often to different chemical environments (e.g., lower redox potentials).

Adsorption at the solid-liquid interface results when the forces of attraction between the surface (adsorbent) and the solute (adsorbate) overcome the forces of attraction between the solvent and the solute. Hence, the degree of adsorption depends on the relative strengths of the adsorbate-adsorbent interactions and the solute-solvent interactions. If the adsorbate-adsorbent interactions dominate due to a strong adsorbate-adsorbent interaction or due to a weak solvent-solute interaction, adsorption will take place and the adsorbing species will be primarily associated with the solid phase. The net interaction of the surface and the adsorbate may result from a variety of chemical and electrical interactions (Stumm and Morgan, 1970; Adamson, 1967).

ADSORPTIVE FORCES

1. Coulombic attraction. This force of attraction results when a charged surface such as a clay mineral attracts an oppositely charged ion to maintain electrical neutrality. The force of attraction is given by Coulomb's law:

$$F = q_1 q_2 / DX^2$$
 (Eq. 1)

where q_1 is the charge density of the surface, q_2 is the charge of the adsorbing species, D is the dielectric constant of the solvent and is a measure of the shielding of q_1 from q_2 by the solvent, and X is the distance between the charges. Sorption of inorganic cations by negatively charged soil colloids in the process of cation exchange is one example of coulombic attraction. The sorption of the organic cations paraquat and diquat by montmorillonite and kaolinite (Weber $et\ al.$, 1965) is also an example of coulombic attraction. In the case of paraquat and diquat sorption within the interlayer of montmorillonite, other sorptive forces add to the coulombic attraction.

- 2. London-van der Waals dispersion forces result from the oscillating electron cloud of one atom rotating in phase with a nonoverlapping oscillating electron cloud of another atom, producing a dipole-like attraction. The differential heats of adsorption are of the order of 1 to 2 kcal mole for small molecules and atoms. For large molecules the heats of adsorption may be much larger. Corkill et al. (1966) gave the heats of adsorption of methane, ethane, pentane and hexane on carbon black as being 3, 4.3, 9.2 and $11.4 \text{ kcal mole}^{-1}$.
- 3. Orientation energy results from the attraction of a permanent dipole for another permanent dipole. The resulting energy of attraction is less than 2 kcal mole $^{-1}$.
- 4. Induction forces result from the attraction of an induced dipole for the inducing species which can be either a permanent dipole or a charged site or species. This force often adds to the adsorptive forces present in cases of coulombic or orientation energy attraction. The energy of attraction is less than 2 kcal mole^{-1} .
- 5. Hydrogen bonding occurs in compounds such as water where electrons are unequally shared between the more electronegative oxygen and hydrogen. This arrangement results in a slight negative charge on the oxygen atom and a slight positive charge on the hydrogen atoms, producing (in the case of water) a dipole moment of 1.84 Debye units (Douglas and McDaniel, 1965). Hence, attraction is possible between dissimilarly charged sites of the adsorbate and the adsorbent. The energy of attraction ranges from 2 to 10 kcal mole⁻¹.
- 6. Chemical forces result when the adsorbate-adsorbent bond approaches an ordinary chemical bond in strength (>>10 kcal mole⁻¹). Chemical forces extend over only very short ranges and often result in the nature of the adsorbate being significantly different in the adsorbed state. Such adsorption is often termed chemisorption to distinguish it from the less specific lowenergy physical sorption. It is often difficult to distinguish between chemical and physical sorption because a chemisorbed layer may have physically sorbed layers upon it.

For some solutes the attractive force of the adsorbate for the solid surface can play a subordinate role to the hydrophilic-hydrophobic balance of the solute with the solvent (Hance, 1967; Stumm and Morgan, 1970). This type

of adsorption is of particular interest when considering the adsorption of organic molecules that are concentrated at the solid-solution interface because of the hydrophobic nature of their hydrocarbon parts. For this type of adsorption the effect of the adsorbent on the interfacial tension or surface free energy would appear to be an important consideration in explaining the observed phenomena. Conversely, some ions that show a strong affinity for the solvent (for example, ones that are highly hydrated) may stay in solution even if they are specifically attracted to the adsorbent.

Traube observed a regularity in the lowering of surface tension by members of three homologous series of esters, alcohols, and fatty acids (Kipling, 1965). The rule derived from that study states, in essence, that the tendency to adsorb organic compounds from aqueous solution increases with increasing molecular weight for members of a homologous series. Hence, adsorptive energy increases systematically for each additional CH₂ group.

Other general rules (Stumm and Morgan, 1970) concerning the adsorption of organics state that a polar adsorbent adsorbs the most polar constituent of a nonpolar solution in preference to the least polar constituent(s). In contrast, a nonpolar surface adsorbs the nonpolar component preferentially from a polar solution.

ADSORPTION ISOTHERMS

Several mathematical expressions—some with a theoretical basis and others entirely empirical in nature—have been employed to describe the relationship between the amount adsorbed and the equilibrium solution concentration. Those equations that have a theoretical basis can (if the assumptions each is based upon are met) provide valuable information about bonding or adsorption energies (affinities), adsorption maxima (capacities), and interfacial free energies. The empirical equations provide a framework for predicting the distribution of adsorbate between the solid and aqueous phases.

Langmuir Adsorption Isotherm

The Langmuir equation (1918), originally developed to describe the adsorption of a gas by a clean solid surface, has been used by numerous investigators to describe adsorption at the solid-liquid interface. The equation usually takes the following form (Veith and Sposito, 1977):

$$\frac{x}{m} = \frac{KbC}{(1 + KC)}$$
 (Eq. 2)

where

x/m = amount of adsorbate adsorbed per unit mass of adsorbent

K = a constant related to the bonding energy of the adsorbate to the adsorbent

b = adsorption maximum or capacity factor

Correct use of the equation requires that two assumptions be met:

- 1. That the adsorbed ions be bound in a monolayer on a homogeneous surface with localized sites.
- 2. That the energy of adsorption be the same for each molecule of adsorbate regardless of the degree to which the monolayer is completed.

Veith and Sposito (1977) have shown that it is necessary not only to meet the basic assumptions of the Langmuir equation but also to have an independent means of determining that the only process taking place is adsorption, since the Langmuir equation will also fit data obtained in situations where secondary precipitation is taking place. Other investigators (Stumm and Morgan, 1970; Kipling, 1965) have also cautioned that although a set of data may fit the equation, that fact is not of itself sufficient evidence that the assumptions have been met.

The normal application of the Langmuir equation to adsorption data involves a least-squares fitting of the data to a linear form of equation 2. At least three linear forms of the Langmuir equation have been used in adsorption studies:

$$C/(x/m) = 1/Kb=C/b$$
 (Eq. 3)

$$1/(x/m) = 1/b+1/KbC$$
 (Eq. 4)

$$(x/m) = b-(x/m)/KC$$
 (Eq. 5)

Equation 3 has been used extensively in adsorption studies involving soils. A plot of C/(x/m) against C for this equation should yield a straight line having a slope of 1/b and an intercept at 1/Kb. For equation 4 the plot would be 1/(x/m) against 1/C, yielding a slope of 1/Kb and an intercept at 1/b. This type of plot is very similar to the double-reciprocal or Lineweaver-Burk plot used in enzyme studies employing Michaelis-Menten kinetics. For equation 5 (x/m) is plotted against (x/m)/C, yielding a slope equal to 1/K and an intercept at b. This plot is of the same form as the Eadie-Hofstee plot (Hofstee, 1952; 1960) also employed in Michaelis-Menten kinetics.

Dowd and Riggs (1965) compared the statistical fit and the predictability of the three linear forms of the Michaelis-Menten equation, which have the same form as the Langmuir adsorption isotherm and its linear equations:

$$v = \frac{V_{\text{max}} C_{\text{s}}}{K + C_{\text{s}}}$$
 (Eq. 6)

$$C_{s}/v = K_{m}/V_{max} + C_{s}/V_{max}$$
 (Eq. 7)

$$1/v = 1/V + K_{\text{max}} + K_{\text{max}} V_{\text{max}}^{C}$$
 (Eq. 8)

$$v = V - K v/C$$
max m s (Eq. 9)

where

v = the initial velocity of the reaction

C = the concentration of the substrate

 V_{max} = the maximum initial velocity

 K_{m} = the Michaelis constant

They found that equation 7, which has the same form as equation 3, and equation 9, which has the same form as equation 5, gave comparable predictions of the two constants when the error in determining the dependent variable (i.e., v or \mathbf{x}/\mathbf{m}) was small, although equation 9 would better show deviations from linearity. Equation 9 gave the best results when the error associated with determining the dependent variable was large. These results are supported by Syers et al. (1973), who in a soil adsorption study compared the form of the Langmuir equation given in equation 3 with that of equation 5. Dowd and Riggs (1965) concluded that the Lineweaver-Burk or double reciprocal plot--equation 8, which has the same form as equation 4--should not be used even if it fits the data well.

Several nonlinear forms of the Langmuir equation are used in adsorption studies:

$$\theta = KC/(1+KC)$$
 (Eq. 10)

$$(x/m) = bC/(K'+C)$$
 (Eq. 11)

Equation 10 is the same as equation 2 except that adsorption is expressed in terms of the percentage of the monolayer that is occupied by adsorbate molecules. This results in b, the monolayer capacity, having a value of 1 unit of adsorption sites and disappearing from the adsorption equation. Equation 11 reduces to equation 2 if K' is replaced with 1/K; hence, K' = 1/K.

Freundlich Adsorption Isotherm

To describe adsorption from dilute solutions, Freundlich (1922) applied the adsorption isotherm:

$$a = \alpha c^{1/n}$$
 (Eq. 12)

or in its more commonly used form:

$$(x/m) = KC^{1/n}$$
 (Eq. 13)

where

C = the equilibrium concentration in the solution after adsorption

a or x/m = the adsorption value

$$\alpha$$
, K, and $1/n = constants$

In studies of the adsorption of a variety of organic compounds on charcoals Freundlich found the adsorption exponent 1/n to vary between 0.1 and 0.5.

The Freundlich adsorption equation is normally considered an empirical equation relating the amount adsorbed to the equilibrium adsorbate concentration. Kipling (1965) cites a derivation of the Freundlich equation by Henry (1922) based on combining an expression for the free energy of a surface with the Gibbs equation. The resulting equation defines the adsorption exponent in terms of the monolayer capacity and surface free energy:

$$(x/m) = KC^{(RT(x/m)m/(\sigma_0-\sigma_1))}$$
 (Eq. 14)

hence

$$1/n = (RT(x/m)m/(\sigma_0 - \sigma_1))$$
 (Eq. 15)

where

(x/m)m =the monolayer capacity

 σ_0 = the surface free energy in contact with pure solvent

 σ_1 = the surface free energy of the surface covered with a monolayer of solute

R = the gas constant

T = temperature

This equation is applicable to dilute solutions where Gibb's surface excess equals (x/m), the amount adsorbed.

When $1/n\simeq 1$ then the Freundlich isotherm reduces to a partition equation:

$$K = \frac{(x/m)}{C} = \frac{C_s}{C_w}$$
 (Eq. 16)

where

 $C_{_{_{\mathbf{S}}}}$ = the concentration of adsorbate in the solid phase

 $C_{\overline{W}}$ = the concentration of adsorbate in equilibrium with the adsorbed phase

Partition equations have been written for both adsorption and desorption

(LaFleur, 1976):
$$C$$

$$K_{f} = \frac{S}{C_{w}} \text{ adsorption}$$
 (Eq. 17)

$$K_{b} = \frac{C_{s}}{C_{w}}$$
 desorption (Eq. 18)

Usually $K_{\rm b}$ > $K_{\rm f}$; that is, a given surface coverage will be in equilibrium with a greater concentration of the solute for an adsorption process than for the desorption process.

The Freundlich equation is normally statistically fit to adsorption data in its linear form:

$$\log(x/m) = \log K + 1/n \log C$$
 (Eq. 19)

A plot of $\log(x/m)$ against $\log C$ yields a slope of 1/n and an intercept equal to $\log K$.

In many studies the difference in adsorption of an organic material by several soils has been correlated with the organic carbon content of the adsorbate. When the adsorption constants are put on an organic carbon basis, the differences in adsorption are often removed.

$$K_{OC} = \frac{K \times 100}{\%OC}$$
 (Eq. 20)

Gibbs Adsorption Equation

The Gibbs equation, originally derived for the adsorption at the liquidliquid interface (Kipling, 1965) has been applied to the adsorption of solutes from dilute solutions by solid surfaces (Stumm and Morgan, 1970):

$$\int_{i}^{1} = \frac{\partial \gamma}{\partial \mu_{i}}$$
 (Eq. 21)

$$\mu_{i} = \mu_{i} + RT \ln a_{i}$$
 (Eq. 22)

hence,

where

$$\sum_{i}$$
 = the adsorption density of component i

 γ = the interfacial tension

 μ_2 = the chemical potential

 a_{i} = the activity of component i

The Gibbs equation defines adsorption in terms of the effect of a solute in either increasing or decreasing interfacial tension. The equation illustrates that solutes which lower the interfacial tension tend to be concentrated at the interface. Many organic substances tend to lower the interfacial tension and hence are accumulated at the interface.

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4 ANALYTICAL METHODS: SOIL THIN-LAYER CHROMATOGRAPHY

W. L. Banwart

The principle of thin-layer chromatography (TLC) and evidence of its widespread use are presented by Stahl (1965). The basis of all chromatographic separations is similar in that a mobile phase passes over a stationary phase and thereby transports different substances with varying speeds in the direction of the flow. Conventional TLC is a form of elution chromatography in which molecules of the test compounds are exchanged between the mobile and stationary phase as a solvent carrying the compounds migrates across a uniform thin layer of stationary phase fixed to a glass plate. The rate or relative rate of movement of the test compounds is dependent on their physical properties and on experimental parameters. The solvent employed may be water, an organic solvent, or a mixture of solvents allowing movement of the test compounds.

The mobility or migration distances of substances on the TLC chromatograms can be expressed in terms of their relative mobility, Rf (Stahl, 1965),

where $R_f = \frac{\text{distance of spot center from starting point}}{\text{distance of solvent from starting point}}$

The use of relative mobilities provides reproducible data, whereas absolute mobilities, which are defined simply as the distances of the spot centers from the starting point, may vary considerably with experimental conditions.

Using TLC to study the mobility of compounds requires a method for documenting the substances' movement. The final position of the compounds on the chromatogram can be made visible with dye indicators, fluorescence under ultraviolet light (Pullan, Howard, and Perry, 1966), a bioassay (Helling, Kaufman, and Dieter, 1971), a spark-chamber apparatus (Pullan et al, 1966), a beta camera (Snyder, 1970), and autoradiography using X-ray film (Mangold, Kammereck, and Malins, 1962). In the last of these methods, chromatograms are developed using 14 C labeled test compounds; X-ray film is then exposed to the chromatogram for several days before development. The developed films make it possible to observe visually the movement of the test compounds and can be used for easy calculations of Rf values. For quantitative measurements, radioactive compounds on TLC plates can be counted directly by commercially available instruments such as strip scanners (Ravenhill and James. 1967) or by zonal analysis where small successive segments of the chromatogram are scraped from the plate and radio-assayed by liquid scintillation spectrometry (Brown and Johnston, 1962). Thin-layer chromatography has been used as a standard method for separating and identifying many synthetic and natural organic compounds (Maier and Mangold, 1964; Stahl and Mangold, 1975), including polynuclear aromatic hydrocarbons (Zoccolillo and Liberti, 1976; Candeli et al., 1975).

Soil thin-layer chromatography was introduced by Helling and Turner (1968). In soil TLC a uniform but relatively thin layer (often less than 1 mm) of a soil-water slurry is spread on TLC plates and allowed to dry. The soil then serves as the stationary phase interacting with compounds carried by the mobile phase. Soil TLC has provided a much simpler and faster way of determining compound mobilities in soil than the traditional leaching columns. Helling and Turner (1968) found that pesticide mobilities determined by soil TLC correlated well with published data. The mobility of substances on soil TLC plates have been reported by Helling and Turner (1968) as frontal Rf values where

$R_f = \frac{\text{distance of frontal edge of spot or streak from starting point}}{\text{distance of solvent edge from starting point}}$

Standard or reference compounds can be spotted on each plate to improve reproducibility.

Data obtained by soil TLC provide information of a nature somewhat different from that obtained with sorption isotherms or partition coefficients. A compound on a soil TLC chromatogram may move as a compact band or as a diffuse streak. From the type of movement it is possible to draw conclusions about the homogeneity of the soil material or of the compound itself. It is also possible to obtain a relative measure of the distance compounds might leach in a soil system when finite amounts of water are applied. $R_{\rm b}$ values have been used (Rhodes, Belasco, and Pease, 1970), where

$R_{b} = \frac{\text{distance moved by bottom of spot}}{\text{distance traveled by solvent}}$

to measure the relative soil depth through which essentially all of a given organic compound (pesticide) applied to a soil has leached. Thus, soil TLC provides some kinds of information that cannot be obtained strictly from adsorption constants or partition coefficients.

Data recalculated from Rhodes, Belasco, and Pease (1970) showed simple correlation coefficients of r = 0.95 between Freundlich K and frontal R_f values for four agricultural chemicals applied to two soils. Helling (1971c) reported highly significant negative correlation coefficients for soil adsorption of nonionic herbicides and Rf values of the same or chemically similar herbicides. As determined by leaching experiments, the mobilities of six organophosphorus insecticides (McCarty and King, 1966), three acidic herbicides (Hamaker, Goring and Youngson, 1966) and their five s-triazine herbicides (Harris, 1966) were inversely related to their adsorption by the soil. Data by Hance (1967), using two soils and 29 organic compounds, show correlation coefficients ranging from 0.85 to 0.91 for the amount of compound adsorbed by the soil and the Rf values for TLC using 40% aqueous ethanol as a solvent. Other workers (Martin and Synge, 1941; Stahl, 1965) have also discussed the relationship between K and Rf values. If a correlation between Rf (mobility) and K (adsorption isotherm) values for a given set of organic compounds can be established, it should be possible to predict K values for additional test compounds on a particular soil.

The greatest use of soil TLC to date has been the study of pesticide mobility in soils (Helling 1971a,b,c; Hance, 1967), but applying the technique to other mobile organic compounds should provide useful data.

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5 ANALYTICAL PROCEDURES: CHEMICAL ANALYSIS

J. C. Means

The analysis of trace organic compounds in water or bound to soil and sediment samples is a complicated task because of a number of factors. First, the water and soil or sediment may contain a wide variety of organic compounds of different polarities, molecular weights, and structures. Second, the affinities of different types of organic compounds for different soil-sediment types span a large range. Third, the solubilities of organic compounds in different solvents vary considerably. Fourth, interactions between soil components and many types of organic compounds are poorly understood.

In general, the process of analyzing trace organic components in water and soil or sediment samples may be divided into these tasks:

- 1. Quantitative recovery of the organic compound of interest
- 2. Separation of that compound from other types of organic compounds
- 3. Quantitation of the organic compound using spectroscopy, chromatography, and radioactivity or a combination of techniques.

RECOVERY OF ORGANIC COMPOUNDS FROM ENVIRONMENTAL SAMPLES

To accurately determine the amount of a given organic compound in a water sample or bound to a soil or sediment sample, the compound must be quantitatively recovered from the gross sample. In the case of a soil or sediment sample, recovery is routinely accomplished by extracting the compound from the soil in a Soxhlet apparatus. Depending upon the variety of compounds which require analysis, a single solvent, a series of solvents, or a mixture of solvents may be required. For example, a cyclohexane or benzene extraction (Sawicki, 1964; Hermann, 1974) is typically used to recover aliphatic and aromatic hydrocarbons from sediments, whereas a benzene and methanol mixture may be used to remove a broad spectrum of nonpolar and polar organics (Giger and Blumer, 1974; Blumer and Youngblood, 1976). Acetone has been used to extract polar organics and humic substances.* Regardless of the extraction procedure used, it is always necessary to perform recovery studies using a reference compound to verify that the expected extraction efficiencies are achieved.

Organic compounds may be quantatively recovered from water samples using one of three techniques. Compounds having low boiling points and low or intermediate polarities may be stripped from the water by passing a pure

^{*}F. J. Stevenson, 1977: personal communication.

inert gas through the sample and collecting it in a trap containing a gas chromatographic column packing (e.g., Tenax GC) (Bellar and Lichtenberg, 1974; Kuo et al., 1977). Soluble organic compounds may be recovered from water by a series of extractions using single or mixed solvents. In some analytical schemes, the pH of the water may be altered (i.e., made basic, neutral, and acidic) to achieve partial separation of compounds having ionizable functional groups. (U.S. EPA, 1977; Acheson et al., 1976; Chang, 1976; Webb et al., 1973). A third technique, one which has been used very successfully, is to collect organic compounds by sorption on purified activated carbon (Keith et al., 1976) or on purified sorbant resins (e.g., XAD-2, 4, and 8) (Malcolm, Thurman, and Aiken, 1977; Junk et al., 1974; Adams, Menzies, and Levins, 1977). These techniques have the advantage that the organics contained in a very large sample volume may be collected on a relatively small amount of resin. However, the recovery efficiencies for different types of organics adsorbed from the water and subsequently desorbed from the resin vary significantly. Therefore, the recovery of each compound of interest should be evaluated. Another advantage of the resin sorption technique is that both polar and nonpolar organics may be sorbed to the resin and eluted for analysis.

Once an extract has been prepared from either a solid or a water sample, it is usually necessary to reduce the volume so as to bring the concentrations of the extracted components into a detectable (ppm) analytical range. The best general technique available for this process is the use of a Kuderna-Danish evaporator (Webb et al., 1973; U.S. EPA, 1977). In some cases air evaporation or evaporation under a stream of nitrogen may be sufficient, but several studies have shown that significant losses of extracted organics may occur (Chiba and Mosley, 1968; Goldberg, Delong, and Sinclair, 1973). Similarly, rotary vacuum evaporation may be appropriate for certain compounds, but losses of many extracted components may occur. Extracts are typically concentrated by a factor of 500 to 10,000, depending upon the origin of the extract and the sensitivity of the analytical systems being used. Here again, it is advisable to test the recovery efficiency for the compound of interest using the concentration technique being considered.

FRACTIONATION, CLEANUP, AND SEPARATION OF ORGANIC EXTRACT COMPONENTS

Organic extracts of soil or sediments and of water, particularly those from industrial areas, may contain hundreds of components. These multiple components tend to complicate the accurate quantitation of individual constituents. Therefore, some steps may be needed to fractionate the extract prior to analysis. To some extent, fractionation begins with the selection of an extracting solvent or sorption resin. However, solvent selectivities are rarely sufficient, especially when concentration factors are high. Solvent-solvent partitioning of the organics in an extract may be useful in crude fractionations (e.g., separating polar compounds from nonpolar compounds).

Liquid chromatography and thin-layer chromatography (TLC) provide the best selectivities in fractionating complex mixtures of organics. For relatively large extracts, separation on liquid chromatographic columns containing such adsorbents as silica gel, alumina, or Florisil is the method of choice. Excellent separations of complex mixtures from acidic, basic, and neutral fractions of coal wastes and cigarette-smoke condensates have been achieved

using these three adsorbents (Swain, Cooper, and Stedman, 1969; Bell, Ireland, and Spears, 1969; Severson et al., 1976).

In the last few years, a number of advances in liquid chromatographic column packings have provided investigators with a number of gel-permeation and adsorption materials which can be extremely useful in fractionating and separating complex mixtures into subsamples which can be analyzed and quantitated.

For relatively small samples, thin-layer chromatography can be used successfully to fractionate a complex mixture. The same basic substrates mentioned above for use in column chromatography are used for TLC separations. Careful selection of solvents and the use of two-dimensional development can yield excellent purifications of individual components prior to quantitation. A large number of investigators have employed TLC to purify compounds for study or to fractionate extracts prior to quantitative analysis (Treiber, 1976; Grant and Meiris, 1977; Bender, 1968; White and Howard, 1967; Pierce and Katz, 1975; Stanley, Bender, and Elbert, 1973; additional references are given in the analytical techniques section of the bibliography at the end of this report).

QUANTITATION OF TRACE ORGANIC COMPOUNDS IN SOLVENT EXTRACTS

The quantitation of individual components of a complex mixture after partial cleanup or fractionation may generally be achieved using the techniques of spectroscopy, gas chromatography, liquid chromatography, or, in appropriate cases, liquid scintillation counting or combinations of the above techniques.

Spectroscopic techniques, which are based on some characteristic absorption wavelength (in the UV, visible, or infrared spectrum) of the compound or a derivative, vary in their selectivity and sensitivity. In cases where the extracts contain relatively few components, spectroscopic methods may be successfully used to quantitate a component without additional cleanup or separation. In most cases, however, the spectroscopic techniques must be used in combination with compatible separation systems (e.g., liquid chromatography or thin-layer chromatography). A large body of literature exists on separations and quantitations of organic pollutants using the above techniques. Representative reports are those of Caton, Matthews, and Walters (1976); Kelly (1967); Willis (1973); Freudenthal et al. (1975); Jenkins and Baird (1975); Brocco, Cantuti, and Cartoni (1970); and IUPAC Applied Chemistry Division (1974). Additional references are listed in the bibliography.

One spectroscopic technique which offers a relatively high degree of both selectivity and sensitivity is the use of fluorescence spectra of various compounds or their fluorescent derivatives (Woo et al., 1978). These techniques are particularly useful when combined with liquid chromatography for the analysis of many coal-derived substances (Stroupe et al., 1977) and pesticides (Mallet, Belliveau, and Frei, 1975). All of these spectroscopic techniques have the advantage that they are not destructive to the sample being analyzed and that the column packing materials and thin-layer supports used

for separations prior to spectral analysis can cover an almost unlimited range of molecular weights.

Gas chromatography (GC) is perhaps the most widely used technique for the separation and quantitation of organic compounds. Methods are available for almost every class of organic pollutant including many energy-related compounds. Column packing materials are available for the separation of compounds representing a broad spectrum of polarities and functional groups. Although fewer liquid phases are readily available, capillary columns have demonstrated increasing utility for the separation of the highly complex mixtures of organics typically found in environmental extracts.

Gas chromatographic separations in general are better than those offered by liquid chromatography, but the compounds which can be analyzed by GC are limited to those having significant vapor pressures at temperatures below 400°C. In most cases, gas chromatographic analysis is destructive of the sample. A wide variety of GC detector systems are available. The flame ionization detector is the most common universal type. A number of other types of detectors demonstrate selectivities for specific types of compounds (e.g., the electron capture detector for halogens, the thermionic detector for nitrogen and phosphorus, and the flame photometric detector for sulfur and phosphorus). All of these detection systems may be used to quantitate organic compounds with the use of appropriate internal standardization techniques. The literature contains a large amount of information on this general topic. Pertinent references are included in the bibliography.

In the last decade, gas chromatography has been combined with mass spectrometry, providing investigators with a very powerful analytical system which can be used to both identify and quantitate organic compounds (John and Nickless, 1977; Janini et al., 1976; Alford, 1977; Oswald, Albro, and McKinney, 1974; O'Reilly and Murrmann, 1974; Lao, Thomas, and Monkman, 1975; and McGuire, Alford, and Carter, 1973).

In certain types of experiments, radiolabeled organic compounds may be used successfully to follow the movement of trace organics in environmental samples. When labeled compounds are used, many types of samples may be analyzed directly without the need for extraction, concentration, or cleanup procedures. Care must be taken, however, to insure that the labeled compound introduced into the experimental system is not degraded in such a way that the radioactivity is lost or transferred to other compounds. The sample is typically separated by (1) thin-layer chromatography followed by radioautography or liquid scintillation counting or (2) by liquid chromatography followed by liquid scintillation spectrometry. Carbon -14 or tritium -3 labeled compounds can be detected in liquid samples by liquid scintillation spectrometry on aliquots of the samples. If solid samples are to be analyzed, the radiolabeled compound may be extracted and then analyzed as a liquid sample. Solids may also be analyzed directly by pyrolytic combustion of the solid and recovery of the radioactivity as ¹⁴CO₂ or ³H₂O.

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6 REVIEW AND INTERPRETATION OF ADSORPTION DATA

K. A. Reinbold and J. J. Hassett

A review of the literature revealed relatively few data on the adsorption of energy-related organic pollutants by sediments or soils. The summary presented here, therefore, covers literature on a variety of types of organic compounds, including pesticide data from references not covered by Farmer (1976).

Some of the published reports included constants for the sorption of organic compounds on various adsorbents. These constants are tabulated at the end of this chapter. However, many sources containing sorption data did not report sorption constants. The latter publications are included in the bibliography.

Except in the case of pesticides, most of the adsorption constants reported for organics were derived using adsorbents other than sediments, soils, and clay minerals. The most commonly used adsorbents were activated carbon or ion exchange resins, which are often used for the removal of organics in water treatment processes. Nylon and cellulose triacetate have also been used. The pesticide data, however, do pertain to sorption on sediments, soils, or clay minerals. The adsorption constants were determined in most cases by fitting experimentally derived data to various forms of the Freundlich or Langmuir adsorption equations.

FACTORS INFLUENCING ADSORPTION

Adsorption is a process in which a solution component is concentrated at the solid-solution interface. Adsorption results when the forces of attraction between the solution component and the surface, that is, the adsorbate-adsorbent interaction, overcomes the forces of attraction between the solution component and the solvent, that is, the solute-solvent interaction. There are two general cases where the adsorbate-adsorbent interaction is greater than the solute-solvent interaction and adsorption results.

In the first case there is a strong positive interaction between the surface and the adsorbate, and this interaction is strong enough to overcome even a fairly strong solute-solvent interaction. In this case adsorption is primarily related to the nature of the bonding between the adsorbate and the adsorbent. The adsorption of organic cations such as paraquat by clay minerals or polar organic molecules within the montmorillonitic interlayer are examples of this type of adsorption (Weber et al., 1965). Reviews emphasizing the effect of adsorbate and adsorbent properties on adsorption have been written by Mortland (1970), Adams(1973), Bailey and White (1970), and Weber (1972).

In the second case, adsorption takes place not because of a strong adsorbate-adsorbent interaction, but rather due to a weak solute-solvent interaction. In this case even a small positive adsorbate-adsorbent interaction can overcome the solute-solvent interaction and result in adsorption. The degree of adsorption or partitioning of the adsorbing material between the solid and aqueous phases is primarily determined by the suitability of the aqueous phase as a solvent for the material. The poorer the aqueous phase as a solvent for the adsorbing species, the weaker the solute-solvent interaction and the greater the adsorption.

The adsorption of nonpolar aromatic hydrocarbons of low water solubility by organic surfaces is an example of this type of adsorption (Karickhoff $et \ al.$, 1979; Means $et \ al.$, 1979). This type of adsorption has been called "hydrophobic adsorption" because of the emphasis on the weak solute-solvent interaction in determining the degree of adsorption in aqueous systems (Horvath and Melander, 1978)

Hydrophobic or nonpolar adsorption can be considered an example of a nonpolar organic compound (adsorbate) partitioning between a polar aqueous phase and a stationary organic phase (adsorbent). In a soil or sediment system the aqueous phase would be the soil solution or the interstitial water in the sediment, while the organic phase would be the naturally occurring humic materials. Factors which either increase the affinity of the adsorbate for the humic surfaces or decrease the affinity of the adsorbate (solute) for the solvent (water) would result in greater adsorption.

Van der Waals forces have been identified as the main source of adsorbate-adsorbent interactions between nonpolar compounds and nonpolar organic surfaces (Horvath and Melander, 1978). The differential heats of adsorption for van der Waals forces are of the order of 1 to 2 kcal mole⁻¹ for small molecules; these forces may be much greater for larger molecules, especially with an increase in the number of double and triple bonds (Bailey and White, 1970). This is illustrated by the adsorption of methane, ethane, pentane and hexane by carbon black. These hydrocarbons gave differential heats of adsorption of 3, 4.3, 9.2 and 11.4 kcal mole⁻¹, respectively (Corkill et al., 1966). The differential heats of adsorption do not increase indefinitely with increasing molecular size; eventually a point is reached where increasing molecular size does not increase adsorption and may even decrease adsorption due to steric hindrance.

Many factors influence the solute-solvent interaction and hence influence adsorption. Molecular properties such as chain length, molecular volume, molecular weight, carbon number, and polarity have all been shown to influence adsorption (Bailey and White, 1970; Gustafson and Paleos, 1971; Lailach et al., 1968; Cummings et al., 1959; Bartell and Miller, 1924; Kipling, 1965; Hansen and Craig, 1954; Parkash, 1974). It has been demonstrated that as chain length, molecular volume, molecular weight, and carbon number increase, and as polarity decreases, the solute-solvent interaction weakens and hydrophobic adsorption increases. The effect of all of these properties on the solute-solvent interaction is integrated into the water solubility of the compound.

The adsorption of hydrophobic compounds has been shown to increase with decreasing water solubility of the compound (Karickhoff $et\ al.$, 1979; Means $et\ al.$, 1979). The adsorption of more polar compounds has also been shown to increase with decreasing water solubility, but only within a family of compounds (Bailey and White, 1970). Aqueous solubilities of organic compounds, while useful in predicting adsorption, are often difficult to determine. Some of the difficulties encountered include the approach of the water solubilities to analytical detection limits, the formation of stable suspensions and the long equilibration times required to establish equilibrium.

Lambert (1968) discussed the similarity between the role of soil organic matter in the sorption of organic compounds and the role of an organic solvent in a liquid-liquid extraction. He observed that the partitioning of a nonpolar organic compound between the soil solution and soil organic matter was highly correlated with the partitioning of the compound between water and an organic solvent. Karickhoff $et\ al$. (1979) reported, for sorption placed on an organic carbon basis (Koc), a significant correlation between the sorption of several aromatic hydrocarbons and the partitioning of the compounds between octanol and water (Kow).

$$\log \text{ Koc} = 1.00 \log \text{ Kow} - 0.21$$
 (Eq. 24)

The linear partition coefficients (Kp) for many compounds may be calculated from the following equation:

$$Kp = Cs/Cw$$
 (Eq. 25)

where

Cs = the concentration of the compound in the solid phase at
 equilibrium

Cw = equilibrium solution concentration of the compound

When the individual linear partition coefficients for the sorption of a hydrophobic organic compound by a variety of different sediments and soils are divided by the respective sediment or soil organic carbon contents, a unique constant Koc is produced.

$$Koc = \frac{Kp \times 100}{\$OC}$$
 (Eq. 26)

This constant is independent of soil or sediment (adsorbent) properties and is only dependent on the nature of the adsorbing species (Karickhoff $et\ al.$, 1979; Means $et\ al.$, 1979).

The relationship between Koc values and Kow values for hydrophobic compounds has several distinct merits. First, Kow determinations are more reliable than water solubility determinations, particularly for the highly hydrophobic compounds. Second, once a compound's Kow value has been measured or calculated, its Koc can be determined from equation 24. Third, if the organic carbon contents of the individual soils or sediments are known, then their respective Kp values for the adsorption of the compound can be calculated.

Table 5 gives an example of linear Kp values, measured Koc values and Koc values calculated from Kow values for the sorption of three hydrophobic compounds, pyrene, dibenzothiophene and acetophenone by a variety of sediments.

TABLE 5. Kow, CALCULATED Koc, LINEAR Kp AND MEASURED Koc VALUES FOR SORPTION OF PYRENE, DIBENZOTHIOPHENE AND ACETOPHENONE BY SOILS AND SEDIMENTS

		Koc	Sam	ple		Koc	Average
Compound	Kow	(calc'd)	No.	%OC	Кр	(meas'd)	Кос
Pyrene	124,000	76 , 400	В2	1.21	774	63,991	63,400
-		•	4	2.07	1098	53,019	•
			5	2.28	1191	52,250	
			6	0.72	633	87,847	
			8	0.15	125	83,333	
			9	0.11	79	71,818	
			14	0.48	285	59 , 271	
			15	0.95	783	82,453	
			18	0.66	509	77,182	
			20	1.30	747	57,469	
			21	1.88	1159	61,628	
			22	1.67	811	48,557	
			23	2.38	1130	47,487	
			26	1.48	1023	69,108	
Dibenzo-	24,000	14,700	B2	1.21	117.5	9711	11,230
thiophene			4	2.07	180.6	8725	
			5	2.28	167.1	7329	
			6	0.72	60.8	8444	
			8	0.15	9.4	6267	
			9	0.11	5.8	5273	
			14	0.48	49.7	10,354	
			15	0.95	179.9	18,937	
			18	0.66	65.1	9864	
			20	1.30	101.4	7800	
			21	1.88	276.0	14,681	
			22	1.67	176.3	10,557	
			23	2.38	388.6	16,328	
			26	1.48	134.5	9088	
Aceto-	38.6	23.8	B2	1.21	0.44	36	38.6
phenone			4	2.07	0.89	43	
			5	2.28	0.56	24	
			6	0.72	0.68	95	
			8	0.15	0.07	48	
			9	0.11	0.09	82	
			14	0.48	0.12	25	
			15	0.95	0.27	28	
			18	0.66	0.30	46	
			20	1.30	0.29	22	
			21	1.88	0.85	45	
			22	1.67	0.53	31	
			23	2.38	0.68	29	
			26	1.48	0.66	45	

The range in Kp values for the adsorption of hydrophobic compounds depends on the compound being adsorbed and the range in organic carbon contents found in the soils or sediments studied. The upper limit for Koc values depends on the compound, but appears to be around 2,000,000 due to present analytical chemistry limitations. The lower limit for the validity of the Koc-Kow relationship has not yet been defined. This limit will be reached when case 1 adsorption, i.e., a specific strong adsorbate-adsorbent interaction, is encountered.

A relationship has been demonstrated for nonlinear Freundlich isotherms when the data are expressed on a molar basis instead of a mass basis (Osgerby, 1970). Molar Kd values may be calculated from mass Kd values by the following equation:

$$Kd(Molar) = \frac{Kd(Mass) \times Molecular \ weight^{1/n}}{Molecular \ weight}$$
(Eq. 27)

Where Kd(Mass) and 1/n are Freundlich constants and the molecular weight is that of the adsorbate.

INTERPRETATION OF TABULATED DATA

Various combinations of the influencing factors discussed above are necessary to explain the adsorption data obtained from the literature and tabulated at the end of this chapter. For the nonpesticide organics, the adsorbents used in most of the studies reviewed were montmorillonite clay or carbon rather than natural sediments or soils. Thus, rather than a variety of adsorbents with a range of characteristics, two distinctly different types are represented: an inorganic clay mineral and an organic.

The chemical characteristics of the adsorbate account for much of the variation in adsorption behavior shown in the tabulated data. These effects were best shown in studies of groups of related compounds. Linner and Gortner (1935) studied the adsorption of 31 organic acids on carbon; the results are summarized in Table 6. Traube's rule that adsorption from aqueous solutions increases with molecular weight for a homologous series was demonstrated for fatty acids but not for other acids. The branched chain had little effect on the maximum adsorption of the acids, but the double bond showed a tendency to decrease adsorption. The introduction of polar groups—carboxyl, hydroxyl, or keto—caused decreased adsorption. The decrease was more pronounced as the number of carboxyl groups increased or as a second hydroxyl group was introduced. The decrease in adsorption caused by the keto group was dependent on the length of the chain and the position of the polar group in the chain.

For 52 structurally related N-phenylcarbamates, acetanilides, and anilines from aqueous 2% ethanol solutions, the inverse relationship between solubility and adsorption accounted for 60 percent of the total variation in adsorption, shown by the data in Table 6 (Ward and Upchurch, 1965). All of the compounds were adsorbed on nylon and cellulose triacetate. Other than solubility, the principal variable factor affecting adsorption was the difference in the molecular structures of the compounds. An investigation of

compounds having systematic variations in molecular structure shows which sites in the molecule can be involved in adsorptive processes and how various substituents may influence the extent of adsorption. For those compounds, differences in molecular structure caused differences in adsorption as a result of steric hindrance, tautomerism, chelation, and induction. The results suggest that the preferred adsorption mechanism of the amido compounds from aqueous solution is via the adsorbate's imino hydrogen and the adsorbent's carbonyl oxygen. If neither of these is available, however, alternative binding sites are utilized.

As a means of establishing a quantitative relationship between soil sorption equilibria and chemical structure, Lambert (1967) proposed the following relationship between parachor of uncharged organic chemicals for which no appreciable hydrogen bonding occurs and the soil sorption of those chemicals:

$$K_{\Delta} = aP\Delta U$$
 (Eq. 25)

The relationship is based upon extrathermodynamic linear free energy approximations and uses of parachor as an approximate measure of the molar volume of the chemical. Distribution equilibria between soil and water for a number of chemical homologs of two chemical classes, including anilines (see Table 6) were used to establish the relationship. The relationship emphasizes the importance of using the partition or distribution coefficient, defined with respect to organic matter, as the most representative index of soil sorption equilibria.

Hance (1969) extended this relationship. A factor given by (parachor-45N), where N is the number of sites in a molecule which can participate in the formation of a hydrogen bond, was correlated with the logarithm of the Freundlich K value for the adsorption of 29 aromatic herbicides. Again, the relationship is valid for soils in which organic matter is the dominant adsorbing constituent.

In the case of a few compounds, adsorption constants were determined for more than one adsorbent (Table 6). Where data were obtained on both a clay mineral and a sediment, the adsorption was generally greater on the sediment. Some data on sediments and soils, including those for benz(a)pyrene, pyrene, methoxychlor, and carbaryl, show a correlation between adsorption and organic matter content (Table 6) (Smith, Mabey, Bohonos, Holt, Lee, Chou, Mill, and Bomberger, 1976a; Karickhoff, Brown, and Scott, 1978; La Fleur, 1976a).

In addition to the adsorption constants for specific compounds and adsorbents, additional pertinent data are tabulated here. Adsorption constants for a number of pesticides averaged over several soils are shown in Table 7. Also, Tables 8 and 9 show the inverse relationship between $\mathbf{R}_{\mathbf{f}}$ values obtained by soil thin-layer chromatographs and adsorption or adsorbate characteristics.

(Note: the references cited in this chapter are listed beginning on p. 110.)

	TABLE 6: ADSORPTION COM	NSTANTS FOR ORGANIC COMPOUNDS	
Compound Name			Ref.
	COMPOUNDS OTH	ER THAN PESTICIDES	
Acids, Aliphatic			
acetic acid	1 January 1	αβ С	Linner and Gortner,
C2H402	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	1935
Compound properties ^a	(decolorizing carbon)	αβ α β	1933
m.w. 60.05	· ,	0.462 0.266 1.736	
m.p. 16.604°C		1/n	
b.p. 117.9°C		Adsorption equation: $a=\alpha C^{1/n}$	
density 1.0492 4°		α 1/n	
water sol ∞		2.46 0.351	

Experimental Conditions: 1 gm of adsorbent and concentrations or reagents varying from 0.01 to 0.25 molar.

^aSee notes at end of table

TABLE 6: Continued

mpound Name			Ref.
adipic acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
C6H10O4	nordite		1935
Compound properties	(decolorizing carbon)	αβ α β	
m.w. 146.14 ^a		2.347 1.886 1.245	
m.p. 151°C ^a		Adsorption equation: $a=\alpha C^{1/n}$	
b.p. 265° C at 100 mm a			
density 1.360 2_4 5a		a 1/n 1.79 0.163	
		1.75 0.103	
water sol 1.5^{15b}			
water sol 1.5^{15b}	Experimental Condition	ons: same as for acetic acid.	
	-		Linner
outyric acid	Adsorbent	ons: same as for acetic acid. $ \text{Adsorption equation:} \text{a=} \ \frac{\alpha\beta C}{1+\alpha C} $	and Gortner
outyric acid C ₄ H ₈ O ₂	-	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	
utyric acid C4 ^H 8 ⁰ 2 Compound properties ^a	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\frac{\alpha \beta \qquad \alpha \qquad \beta}{\alpha \beta \qquad \alpha \qquad \beta}$	and Gortner
outyric acid ^C 4 ^H 8 ^O 2	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortner
outyric acid C4 ^H 8 ⁰ 2 Compound properties	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $1.689 0.863 \qquad 1.957$	and Gortner
Compound properties m.w. 88.12°C	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\frac{\alpha \beta}{\alpha \beta} \qquad \alpha \qquad \beta$	and Gortner
outyric acid C ₄ H ₈ O ₂ Compound properties ^a m.w. 88.12°C m.p4.26°C	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $1.689 0.863 \qquad 1.957$	and Gortner

TABLE	6 •	Continued

Compound Name			Ref.
butyric acid C4H802	Adsorbent activitated	Adsorption equation: y= a.x ^b	Spahn et al.,
m.w. 88.12 m.p4.26 °C b.p. 163.53 °C density 0.9577 % ° water sol ∞	charcoal B10	x - range I (mmol/dm ³) a b 0.08 - 7.18 0.878 0.4	
n-butyric acid	Adsorbent XAD-2 (amberlite resin)	Adsorption equation: $\frac{c}{q} = \frac{1}{Kb} + \frac{c}{b}$ (Langmuir) K(25°C) b(25°C) 23.7 liter/mole 1.46X10 ⁻³ mole/g	Gustafson et al., 1968
		Enthalpy of adsorption: \[\Delta H^\circ = 2.303 \text{ RT}_1 T_2 \text{ (logK}_2' - logK}_1') \\ \text{ (T}_2 - T_1) \\ \text{ q} \qquad - \Delta H^\circ * \\ 7\text{ 7X10}^{-4} \text{ mole } \qquad \text{ 4.1 kcal/mole } \\ \text{ 9X10}^{-4} \text{ mole } \qquad \text{ 2.4 kcal/mole } \\ \text{ 1.2X10}^{-3} \text{ mole } \qquad \text{ 2.1 kcal/mole } \\ \text{ q} \qquad \qquad \text{ 1.2X10}^{-3} \text{ mole } \qquad \qquad \text{ 2.1 kcal/mole } \\ \end{array}	

^{*}AH decreases with increasing surface coverage, i.e. energetically preferred sites are utilized first.

TABLE 6: Continued

ompound Name			Ref.
caproic acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner,
$CH_3(CH_2)_4CO_2H$	nordite	- 1+αC	
Compound properties	(decolorizing carbon)	αβ α β	1935
m.w. 116.16^{α}	Carbony	8.772 4.636 1.892	
m.p. -2 to -1.5 °C $^{\alpha}$		a1/n	
b.p. 205°Ca		Adsorption equation: $a=\alpha C^{1/n}$	
density 0.9274 20		α 1/n	
water sol 0.4 gm/l00 \mathtt{ml}^b		3.03 0.175	
citraconic acid		a 8 C	Linner
C5H604	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortner,
Compound properties	(decolorizing	αβ α β	1935
m.w. 130.10 ^a	carbon)	1.356 1.014 1.337	
m.p. 91°C ^b		Adsorption equation: $a = \alpha C^{1/n}$	
density 1.617^a		α 1/n	
water sol 238 $cold^b$			
1001 001 100 COIG		1.69 0.167	

TABLE 6: Continued

apound Name			Ref.
itric acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
C6 ^H 8 ^O 7	nordite		1935
Compound properties	(decolorizing carbon)	αβ α β	
m.w. 192.14 $^{\alpha}$	carbony	1.444 2.757 0.524	
m.p. 153°C (anhydrous) a		Adsorption equation: $a= \alpha C^{1/n}$	
b.p. $decomp^a$			
density 1.54240		0.73 0.203	
water sol 133 cold gm/100 ml b	Experimental Conditi	ons: same as for acetic acid.	
libromosuccinic acid	Experimental Conditi Adsorbent		Linner and Gortne
libromosuccinic acid ^C 4 ^H 4 ^O 2 ^{Br} 2	Adsorbent nordite	ons: same as for acetic acid. $ \label{eq:acetic} \text{Adsorption equation:} \text{a=} \ \frac{\alpha\beta C}{1+\alpha C} $	
libromosuccinic acid	Adsorbent	Adsorption equation: $a = \frac{\alpha\beta C}{1+\alpha C}$ $\alpha\beta \qquad \alpha \qquad \beta$	and Gortne
dibromosuccinic acid C4 ^H 4 ⁰ 2 ^{Br} 2	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha\beta C}{1+\alpha C}$	and Gortner
dibromosuccinic acid C4 ^H 4 ⁰ 2 ^{Br} 2 Compound properties ^b	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha\beta C}{1+\alpha C}$ $\alpha\beta \qquad \alpha \qquad \beta$	and Gortner
dibromosuccinic acid C4H402Br2 Compound properties m.w. 275.90	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $1.397 1.119 \qquad 1.248$	and Gortner

TABLE 6: Continued

mpound Name			Ref.
formic acid HCO ₂ H	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortne
Compound properties ^a m.w. 46.03	(decolorizing carbon)	αβ α β 0.273 0.159 1.710	1333
m.p. 8.4°C		Adsorption equation: $a=\alpha C^{1/n}$	
b.p. 100.7°C density 1.220 ²⁰		α 1/n	
water sol ∞		2.47 0.435	
fumaric acid			
umaile acid			Linnor
$C_4H_4O_4$	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	
Compound properties		Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $7.097 5.798 \qquad 1.224$	
m.w. 116.07 ^b m.p. 287°C ^b	nordite (decolorizing	αβ α β 7.097 5.798 1.224	and Gortne
m.w. 116.07 ^b	nordite (decolorizing	αβ α β	and Gortne
m.w. 116.07 ^b m.p. 287°C ^b	nordite (decolorizing	Adsorption equation: $a = \alpha C^{1/n}$	and Gortne

TABLE 6: Continued

mpound Name			
glutaric acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortne
C5H8O4	nordite (decolorizing	Adsorption equation: d= 1+αC	1935
Compound properties	carbon)	αβ α β	
m.w. 132.11^{α}		3.697 3.104 1.192	
m.p. 99°C ^a		Adsorption equation: $a = \alpha C^{1/n}$	
b.p. 304 $decomp.b$		Adsorption equation: a= ac	
density 1.424 $^{25a}_{4}$		$\frac{1}{n}$	
water sol 64^{20}		1.96 0.201	
	Experimental Conditi	lons: same as for acetic acid.	
	Experimental Conditi	lons: same as for acetic acid.	
		-	Linner and Gortne
	Adsorbent nordite	Lons: same as for acetic acid. $ \text{Adsorption equation:} a=\frac{\alpha\beta C}{1+\alpha C} $	and Gortne
glyceric acid ^C 3 ^H 6 ⁰ 4	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	
glyceric acid	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortne
glyceric acid C3 ^H 6 ⁰ 4 Compound properties ^a	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $0.668 0.812 \qquad 0.823$	and Gortne
glyceric acid C3 ^H 6 ⁰ 4 Compound properties ^a m.w. 106.08	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $0.668 0.812 \qquad 0.823$ Adsorption equation: $a = \alpha C^{1/n}$	and Gortne
glyceric acid C3H604 Compound properties m.w. 106.08 b.p. disintegrates	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $0.668 0.812 \qquad 0.823$	and Gortne

TABLE 6: Continued

mpound Name			Ref.
glycolic acid HOCH ₂ CO ₂ H	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortne 1935
Compound properties	carbon)	αβ α β	
m.w. 76.05		0.239 0.249 0.958	
m.p. 80°C			
b.p. decomposes		Adsorption equation: $a=\alpha C^{1/n}$	
		α 1/n	
		1.54 0.390	
	Experimental Conditi	ons: same as for acetic acid.	
nlyoxylic acid	Experimental Conditi	ons: same as for acetic acid.	Linner
glyoxylic acid ^C 2 ^H 2 ^O 3	Adsorbent nordite		
•	Adsorbent	ons: same as for acetic acid.	Linner and Gortner 1935
C ₂ H ₂ O ₃	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortne
Compound properties	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortner
$C_2H_2O_3$ Compound properties m.w. 74.04^a m.p. $70-5^\circ C^a$ $(+1/2w)$	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha\beta C}{1+\alpha C}$ $\alpha\beta \qquad \alpha \qquad \beta$ $0.508 0.223 \qquad 2.275$	and Gortner

TABLE 6: Continued

mpound Name			Ref.
isobutyric acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
(Сн ₃) ₂ СнСо ₂ н	nordite	1+αC	
Compound properties	(decolorizing carbon)	αβ α β	1935
m.w. 88.1f		0.883 0.497 1.776	
m.p46.1°C°			
b.p. 153.7°C ^a		Adsorption equation: $a=\alpha C^{1/n}$	
density 0.96815 $^{20}_{4}$; .949 $^{20}_{4}$		α 1/n	
water sol 20 20 gm/l00 ml b		2.36 0.273	
			Linner
isovaleric acid	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortner
^{C5^H10⁰2}	nordite	Title	1935
Compound properties	(decolorizing carbon)	αβ α β	1933
m.w. 102.13 ^a		1.630 0.902 1.807	
m.p29.3°C ^a			
b.p. 176.7°C ^a			
		Adsorption equation: $a=\alpha C^{1/n}$	
density 0.92864°a		Adsorption equation: $a = \alpha C^{1/n}$ ${\alpha} \frac{1/n}$	
density 0.9286^{20a} water sol 4.2^{20b}			

TABLE 6: Continued

ompound Name			Ref.
itaconic acid	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner, 1935
Compound properties	carbon)	αβ ο β	
m.w. 130.10^a		1.167 0.904 1.291	
m.p. 175°C ^a			
b.p. decomposes		Adsorption equation: $a = \alpha c^{1/n}$	
density 1.632^b		α 1/n	
water sol 8.33^{20}		1.54 0.148	
######################################	Experimental Conditi	ons: same as for acetic acid.	
	Experimental Conditi		Linner
lactic acid (DL)	Adsorbent nordite		Linner and Gortner, 1935
lactic acid (DL)	Adsorbent nordite (decolorizing	ons: same as for acetic acid.	and Gortner,
lactic acid (DL)	Adsorbent nordite	ons: same as for acetic acid. $ \label{eq:acetic} \text{Adsorption equation:} a = \frac{\alpha \beta C}{1 + \alpha C} $	and Gortner,
lactic acid (DL) C3H603 Compound properties m.w. 90.08a m.p. 18°Ca	Adsorbent nordite (decolorizing	ons: same as for acetic acid. Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $0.437 0.415 \qquad 1.054$	and Gortner,
lactic acid (DL) C ₃ H ₆ O ₃ Compound properties m.w. 90.08 ^a m.p. 18°C ^a b.p. 122°C ^a	Adsorbent nordite (decolorizing	ons: same as for acetic acid. Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	and Gortner,
lactic acid (DL) C3H603 Compound properties m.w. 90.08a m.p. 18°Ca	Adsorbent nordite (decolorizing	ons: same as for acetic acid. Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$ $0.437 0.415 \qquad 1.054$	and Gortner,

TABLE 6: Continued

ompound Name			Ref
levulinic acid C ₅ H ₈ O ₃	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
m.w. 116.13 ^a m.p. 37.2°C ^a	nordite (de∞lorizing carbon)	αβ α β 2.990 2.289 1.307	1935
b.p. 246°C slight decomp. ^a density 1.1335 $^{20a}_{4}$; 1.1395 $^{20b}_{4}$ water sol very sol ^b	Experimental conditions:	Adsorption equation: $a=\alpha C^{1/n}$ $\alpha \qquad 1/n$ $1.83 \qquad 0.183$ same as for acetic acid	
maleic acid $C_4^H{}_4^O{}_4$ Compound properties m.w. 116.07^a m.p. $139-140^{\circ}\text{C}^a$	Adsorbent nordite (decolorizing (carbon)	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\frac{\alpha \beta}{\alpha \beta} \qquad \alpha \qquad \beta$ 1.233 0.884 1.395	Linner and Gortne 1935
b.p. 135 decomp. ^b density 1.590 20ba water sol 78.825; 392.697.5b		Adsorption equation: $a=\alpha C^{1/n}$ $\alpha \qquad 1/n$ $1.90 \qquad 0.203$	

TABLE 6: Continued

mpound Name			Ref
malic acid (1) C4 ^H 6 ^O 5	Adsorbent	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
Compound properties	nordite (decolorizing carbon)	αβ α β	1935
m.w. 134.09 m.p. 100°C	· · · · · · · · · · · · · · · · · · ·	0.531 0.574 0.927	
b.p. 140°C decomp.		Adsorption equation: $a=\alpha C^{1/n}$	
density 1.595		a 1/n	
water sol very sol		1.28 0.252	
	Experimental conditions:	same as for acetic acid	
malonic acid C ₃ H ₄ O ₄	Adsorbent	Adsorption equation: $a \approx \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner
Compound properties	nordite (decolorizing carbon)	αβ α β	1935
m.w. 104.06 ^b		1.897 1.540 1.232	
m.p. 135.6 °C subl. ^{t} b.p. decomp. at 140 °C ^{a}		Adsorption equation: $a=\alpha C^{1/n}$	
1 _		1 /	
density 1.631, 15b ; 1.619 water sol 61.10 gm/100 mlb 73.5 gm/100 mlb 92.65 gm/100 mlb		α 1/n	

TABLE 6: Continued

mpound Name						Ref
mesaconic acid	Adsorbent	Adapy		ntion	$a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortne
^С 5 ^Н 6 ^О 4		Adsorptio	n equa	ation:	$a = \frac{1+\alpha C}{1+\alpha C}$	
Compound properties	nordite (decolorizing carbon)	αβ	α		β	1935
m.w. 130.10 ^a	calbon)	2.7	6 1	.886	1.435	-
m.p. 204.5					1/n	
b.p. 250°C decomp.		Adsorption	n equa	ation:	$a=\alpha C^{1/n}$	
density 1.466^{20}_{4} g/m 1^{b}		 α		./n		_
water sol 2.7^{18}^{b} ; 118^{100}^{b}		1.8		.133		
	Experimental conditions					-
methylsuccinic acid	-	: same as for	cetic	acid	$a = \frac{\alpha \beta C}{1}$	Linner and Gortne
nethylsuccinic acid ^{C5^H8^O4}	Adsorbent	: same as for	cetic	acid	$a = \frac{\alpha\beta C}{1 + \alpha C}$	and Gortne
_	Adsorbent nordite (decolorizing	: same as for α	on equ	acid	β	Linner and Gortne
С ₅ ^Н 8 ^О 4	Adsorbent nordite	: same as for a	on equ	acid		and Gortne
C ₅ H ₈ O ₄ Compound properties ^b	Adsorbent nordite (decolorizing	: same as for α	on equa	acid	β 1.092	and Gortne
C ₅ H ₈ O ₄ Compound properties ^b m.w. 132.11	Adsorbent nordite (decolorizing	: same as for α	on equa	acid	β	and Gortne
m.w. 132.11 m.p. 111°C	Adsorbent nordite (decolorizing	: same as for α	on equal	acid	β 1.092	and Gortne

TABLE 6: Continued

mpound Name			Ref
nonobromosuccinic acid	Adsorbent	Adsorption equation: $a = \frac{\alpha RC}{1+\alpha C}$	Linner and Gortner
C ₄ H ₅ O ₄ Br	nordite		1935
Compound properties	(decolorizing	αβ α β	4,500
m.w. 197.00^{t}	carbon)	0.934 0.643 1.451	
m.p. 159 °C b 161 °C a			
density 2.073^a		Adsorption equation: $a=\alpha C^{1/n}$	
water sol 19^{15}			
		α 1/n	
	Experimental conditions	1.82 0.195 same as for acetic acid	
xalic acid	Experimental conditions	same as for acetic acid	Linner
	Experimental conditions Adsorbent		
^C 2 ^H 2 ^O 4	Adsorbent nordite	same as for acetic acid	
^C 2 ^H 2 ^O 4	Adsorbent nordite (decolorizing	same as for acetic acid	and Gortner
	Adsorbent nordite	same as for acetic acid	and Gortner
Compound properties	Adsorbent nordite (decolorizing	same as for acetic acid	and Gortner
C ₂ H ₂ O ₄ Compound properties m.w. 90.04 ^a	Adsorbent nordite (decolorizing	same as for acetic acid	and Gortner
C ₂ H ₂ O ₄ Compound properties m.w. 90.04 ^a m.p. 189.5°C ^a	Adsorbent nordite (decolorizing	same as for acetic acid $Adsorption \ equation: \ a = \frac{\alpha\beta C}{1+\alpha C}$ $\frac{\alpha\beta}{\alpha\beta} \alpha \beta 0.440 0.332 1.325$ $Adsorption \ equation: \ a = \alpha C^{1/n}$	and Gortner
m.w. 90.04 ^a m.p. 189.5°C ^a b.p. 157°C subl. a	Adsorbent nordite (decolorizing carbon)	same as for acetic acid	and Gortner

TABLE 6: Continued

mpound Name			Ref.
propionic acid C3H602	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner 1935
Compound properties ^a m.w. 74.08	carbon)	αβ α β 0.925 0.491 1.885	
m.p20.8°C b.p. 140.99°C		Adsorption equation: $a = \alpha C^{1/n}$	
density 0.9930^{20} water sol ∞		α 1/n 2.46 0.236	
	Experimental condici	ons: same as for acetic acid.	
	Experimental conditi	ons: same as for acetic acid.	Chahn
propionic acid C3 ^H 6 ^O 2	Adsorbents	Adsorption equation: $y=a \cdot x^b$	Spahn et al.,
C ₃ H ₆ O ₂ Compound properties	·	Adsorption equation: y= a·x ^b Adsorption constants: x-range I	
C ₃ H ₆ O ₂ Compound properties m.w. 74.08	<u>Adsorbents</u> Activated	Adsorption equation: $y=a \cdot x^b$ Adsorption constants:	et al.,
Compound properties	<u>Adsorbents</u> Activated	Adsorption equation: y= a·x ^b Adsorption constants: x-range I (mmol/dm ³) a b	et al.,

TABLE 6: Continued

ompound Name			Ref.
pyruvic acid C3H4O3	Adsorbent nordite (decolorizing	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner 1935
Compound properties	carbon)	αβ α β	
m.w. 88.06 ^a		0.979 0.585 1.674	
m.p. $13.6^{\circ}C^{a}$ b.p. $165^{\circ}C$ slight decomp.		Adsorption equation: $a = \alpha C^{1/n}$	
density 1.2272^{20}_{4}		1/-	
water sol ∞^{b}		α 1/n 2.44 0.273	
	Experimental Conditi	ons: same as for acetic acid.	
succinic acid			
C4H6O4	Adsorbent nordite	Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$	Linner and Gortner, 1935
C ₄ H ₆ O ₄ Compound properties		Adsorption equation: $a = \frac{\alpha \beta C}{1 + \alpha C}$ $\alpha \beta \qquad \alpha \qquad \beta$	and Gortner
•	nordite (decolorizing		and Gortner
Compound properties	nordite (decolorizing	αβ α β 0.865 0.467 1.854	and Gortner
Compound properties ^b m.w. 118.09	nordite (decolorizing	αβ α β	and Gortner
Compound properties ^b m.w. 118.09 m.p. 185°C	nordite (decolorizing	αβ α β 0.865 0.467 1.854	and Gortner

TABLE 6: Continued

mpound Name					Ref
tartaric acid ${^C_4}^{\rm H_6}{^O_6}$ Compound properties m.w. 150.09^b	Adsorbent nordite (decolorizing carbon)	Adsorption αβ 0.322	equation: α 0.468	$a = \frac{\alpha\beta C}{1+\alpha C}$ β 0.687	Linner and Gortner 1935
m.p. 100°C ^b density 1.697 ^b ; 1.788 ^a	h	Adsorption	ecuation	$: a=\alpha C^{1/n}$	
water sol 20.6 ²⁰ b; 9.23 ⁰ b;	185 1000	α	1/n 0.275	-	
		0.94	0.275		
	Experimental conditions:				
valeric acid CH ₃ (CH ₂) ₃ CO ₂ H Compound properties	Experimental conditions: Adsorbent nordite (decolorizing carbon)	Same as for ace Adsorption α	tic acid	: a= αC ^{1/n}	Linner and Gortne 1935
сн ₃ (сн ₂) ₃ со ₂ н	Adsorbent nordite (decolorizing carbon)	same as for ace Adsorption	equation 1/n 0.182		and Gortne

TABLE 6: Continued

Compound Name								Ref
Acids, Aromatic								
benzoic acid	Adsorbents	Adsorption equa	tion:	$\frac{x}{m} = \frac{abC}{(1+aC)}$)			Ward and Getzen,
C ₇ H ₆ O ₂ Compound properties	Type BL - activated carbon (Pittsburgh	Adsorption cons		(I 1 d c)	,			1970
m.w. 122.13 ^a	Chemical Co.)			pH of				
m.p. 122.4°C	Surface area $1000-1100 \mu^2/g$	Compound	<u>pka</u>	soln.	a*_	<u>b</u>	x/m	
b.p. 249°C	1000-1100 μ ⁻ /g	benzoic acid	4.20	3 7	0.238 0.108	510 124	489 113	
density 1.2659^{15b}_{4}				11	0.081	75	67	
b		-2,4-dichloro	2.76	3	0.259	676	651	
water sol 0.18		,		7	0.123	159	147	
water sol 0.184b 0.2718b 2.275b				11	0.108	73	69	
		-3-amino-2,5-	3.40	3	0.928	515	510	
		dichloro		7	0.283	131	127	
		(amiben)		11	0.025	72	51	
		-3-nitro-2,5-	3.23	3	0.360	505	491	
		dichloro		7	0.118	130	120	
				11	0.035	93	72	
		-2-methoxy-3,6-	1.94	3	0.181	394	313	
		dichloro		7	0.317	154	149	
		(dicamba)		11	0.068	68	59	

Experimental conditions: 100 ml aqueous acid solution: 10mg carbon

*Units: a = liters/μ mole

b = μ moles/g c = μ moles/liter

 $\frac{x}{m} = \mu \text{ moles/g}$

TABLE 6: Continued

ompound Name							Ref.
benzoic acid	Adsorbent	Adsorpt	ion equat:	ion: y= a·	· x ^b		Spahn et al.,
^C 7 ^H 6 ^O 2 Compound properties	activated charcoal B10	Adsorpt	ion consta	ant:			1974
m.w. 122.13 ^a		x-ran					
m.p. 122.4°C°		(mmol	$/dm^3$	<u>a</u> <u>b</u>			
b.p. 249°C ^b		0.1 -	6	3.06 0.1	81		
density 1.2659_4^{15}							
water sol 0.18*b 0.27 ¹⁸ 2.2 ⁷⁵							
benzoic acid and substituted benzoic acids ${^{\rm C}7}^{\rm H}6^{ m 0}2$	Adsorbent charcoal, activated, 200 mesh	-	ion equat: g x/m = lo	ion: og k + l/n	log c		Hartman, Kern, Bobalek,
		1/:	n and log	k Values			
	- 11	1 /	20°	log 30°	k 40°	50°	
	Acids	1/n					
	Benzoic	0.3680	0.0613	0.0265	-0.1091	-0.2551	
	o-Chlorobenzoic	0.4060	0.2208	0.1834	0.1303	0.0936	
	o-Aminobenzoic	0.3693 0.3840	0.3472 0.2700	0.3031 0.2385	0.2678 0.2091	0.2218 0.1844	
	o-Hydroxybenzoic	0.3840	-0.0215	-0.0719	-0.2510	-0.5232	
	o-Toluic			0.0/13	0.2310	0.0232	
	o-Toluic m-Toluic	0.3581	-0.0092	-0.0582	-0.1791	-0.3732	

Adsorption from benzene solutions.

TABLE 6. Continued

	TA	BLE 6: Continu	<u>ed</u>					
ompound Name								Ref.
phenyl acetic acid	Adsorbent	Adsorption	eguatio	n. v- a	.vb			Spahn et al.
с ₆ н ₅ сн ₂ со ₂ н	Activated	Adsorption	equacio	11. y- a	- X			
Compound properties	charcoal							1974
m.w. 136.16								
m.p. 77°C		Adsorption	constan	ts:				
b.p. 265.5°C		x-range I			x-range II			
density 1.091 ⁷⁷ ; 1.228 ⁴		(mmol/dm ³)	<u>a</u>	<u>b</u>	$(mmo1/dm^3)$	<u>a</u>	<u>b</u>	
water sol slightly sol.	B10	0.1 - 10	2.51	0.16	0.001 - 0.1	2.6	0.32	
	TM	0.05 - 12	1	0.165				
	Lev 634	0.1 - 8	0.84	0.38	0.01 - 0.11	1.76	0.56	
	BD	0.1 - 22	1.14	0.15				
	Dl	0.05 - 7	1.93	0.162				
	D2	0.1 - 8	1.57	0.245				
	B2	0.1 - 11	1	0.145				
	LS-supra	0.1 - 7	1.75	0.18	0.001 - 0.31	1.53	0.23	
	Decaka 9	0.1 - 8	1.82	0.03				
	106/427/1	0.1 - 7	1.53	0.11	0.001 - 0.12	2.05	0.33	
	106/427/5	0.1 - 6.5	1.87	0.144	0.001 - 0.11	1.85	0.3	
	106/427/3	0.1 - 8	2.08	0.244	0.001 - 0.12	2.6	0.44	

TABLE 6: Continued

	TABLE 0:	continuea						
Compound Name								Ref.
Carboxylic acids, aromatic								
Phenoxyacetic acid				x ah	oC.			Ward and
с ₈ н ₈ о ₃	Adsorbents	Adsorption equa	tion:	$\frac{1}{m} = \frac{3}{(1+1)}$	aC)			Getzen,
	Type BL - activated carbon							1970
Compound properties	(Pittsburgh	Adsorption cons	tants:					
m.w. 152.14	Chemical Co.)			pH of			x	
m.p. 99°C	Surface Area $\frac{2}{1000-1100} \mu^2/g$	Compound	<u>pka</u>	soln.	_ <u>a*_</u>	<u>b</u>	<u>x</u>	
b.p. 285°C slight decomp.	1000 1100 μ / g	Phenoxyacetic		_			426	
•		Acid	3.03	3 7	0.423	446 105	4 36 86	
water sol 1.2 10				ıí	0.209	64	43	
		-4-chloro-	2.36	3	0.362	575	560	
				7	0.101	198		
				11	0.115	145	133	
		-2,4-dichloro-	3.31	3 7	0.279	629	607	
					0.112	223		
				11	0.070	154	135	
		-2,4,6-						
		trichloro-	3.35	3 7	0.672 0.095	676 239	666	
				11	0.073	144	127	
		2,4-dichloro-	3.92	3 7	0.431 0.125	645 235		
				11	0.123	235 59	54	

Experimental Conditions: 100 ml aqueous acid solution: 10 mg carbon.

Units: a = liters/µ mole b = 1 moles/g c = 1 moles/liter

 $\frac{x}{m} = \mu \text{ moles/g}$

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Compound Name							Re f.
2,4-dichloro- phenylacetic aci	d <u>Adsorbents</u>	Adsorp	tion equat	tion: $\frac{x}{m} =$	abC (1+aC)		Ward and Getzen,
	Type BL - activated carbon (Pittsburgh Chemical Co.)	Adsorp	tion const	tants:			1970
	Surface area 1000-1100 µ ² /g	pka	pH of soln.	_a*_	<u>b</u>	x/m	
		3.92	3 7 11	0.431 0.125 0.100	645 235 59	630 218 54	
	Experimental conditions:	100 ml aqu	eous acid	solution:	10 mg	carbon	
*Units: $a = liters/b = \mu \text{ moles}$ $c = \mu \text{ moles}$ $\frac{x}{m} = \mu \text{ moles}$	/g /liter						

TABLE 6: Continued

TABLE 6: Continued

	14011	so. contr	nuea			
Compound Name						Ref.
lcohol						
n-butyl alcohol	Adsorbent	Adsor	ption eq	uation: $\frac{x'}{m} = \frac{1}{2}$	$\frac{x}{m}(1+N_{b}\overline{v}_{b}/N_{a}\overline{v}_{a})$	Bartell, Thomas,
С ₄ н ₉ он	Graphite	N =	mole fr	action		and Fu,
Compound properties	Blood Char.	$\overline{v} =$	1951			
m.w. 74.12 ^c		a =				
m.p. −90°C°		b =	solute			
b.p. 117-118°C°; 117.71°C		<u>t°C</u>	<u>K*</u>	ΔF°** kcal/mole	ΔH°*** kcal/mole	
density $0.810^{20^{\mathcal{C}}}_{4}$		0	63.3	-2.24	-0.16	
water sol 7.9 gm/l00 ml^b		25 4 5	61.6 52.7	-2.43 -2.50	-1.47	
		0	477	-3.34	-0.94	
		25 45	412 390	-3.54 -3.76	-0.52	

*Equilibrium constant K obtained from the limiting slope of the adsorption isotherms at zero concentration

$$\star\star_{\Delta}F^{\circ} = -RT \ln K$$

$$\Delta H^{\circ} = R\left(\frac{T_2T_1}{T_2-T_1}\right) \ln \frac{K_2}{K_1}$$

TABLE 6: Continued

mpound Name					Ref.	
ides acetanilide and derivatives, adsorption from 2% ethanol		Adsorption equation: solubility		$(\frac{x}{m}) = KC^{1/n}$ $1/n = 1$	Ward and Upchure	
C _a h ₉ on					1965	
Compound properties b	Compound	(x 10 ⁻⁴ M) 2% ethanol	<u>k</u> *	k _A **		
m.w. 135.16	N-Ethylacetanilide	2320	0.05	0.20		
m.p. 114°C	4-Hydroxyacetanilide	1580	0.25	0.13		
_	Acetanilide	624	0.21	0.32		
b.p. 305°C	2-Chloroacetanilide	375	0.25	0.47		
density 1.214	4-Aminoacetanilide	352	0.06	0.05		
water sol 0.563 ²⁵ : 3.5 ⁸⁰	<i>N-n-</i> Butylacetanilide	167	0.23	0.89		
Water 301 0.303 , 3.3	2-Nitroacetanilide	138	0.16	0.72		
	2-Hydroxyacetanilide	32	0.26	0.15		
	N-Phenylacetanilide	31	0.26	1.59		
	4-Chloroacetanilide	16	1.19	2.64		
	3,4-Dichloroacetanilide	e 12	6.05	6.83		
	4-Nitroacetanilide	11	1.02	2.18		
	4-Bromoacetanilide	11	1.73	3.30		
	2,5-Dichloroacetanilide	10	0.92	1.59		

Method of analysis: UV spectrophotometry with Beckman DU spectrophotometer

 $[*]k_N$ = adsorption on nylon

^{**} k_A = adsorption on cellulose triacetate

TABLE 6: Continued

mpound Name					Ref.
niline and derivatives, adsorption from 2% ethanol		Adsorption 6	equation	$: (\frac{x}{m}) = KC^{1/n};$	Ward and Upchurch,
C ₆ H ₅ NH ₂ Compound properties	<u>Compound</u>	solubility (x 10 ⁴ M) 2% ethanol	k _N *	$1/n = 1$ k_{A}^{**}	1965
m.w. 93.13 ^a	Aniline	4450	0.07	0.27	
m.p6.3°C ^a	2-Methylaniline	1670	0.10	0.44	
b.p. 184.13°C ^a	<pre>4-Methylaniline N-Methylaniline</pre>	1240 560	0.05 0.24	0.50 0.92	
density 1.02173 20	4-Chloroaniline	520	0.68	3.31	
water sol 3.4^{20} ; 6.4^{90}	2-Chloroaniline	450	0.55	1.94	
water sor 3.4 / 0.4	${\it N-}$ Ethylaniline	250	0.32	1.10	
	2-Nitroaniline	103	0.91	2.84	
	N, N -Dimethylaniline	95	0.52	2.12	
	2,3-Dichloroaniline	85	3.45	7.20	
	3,4-Dichloroaniline	52	4.32	10.20	
	2,5-Dichloroaniline	49	3.78	11.20	
	4-Nitroaniline	38	1.10	3.40	
	N-n-Butylaniline	16	1.42	6.55	
	2-Nitro-4-chloroaniline	9	3.70	7.80	
	2,4-Dinitroaniline	5	1.94	5.65	
	N-Benzylaniline	4.	3.46	19.48	
	N-Phenylaniline	3	16.50	24.00	

Method of analysis: UV spectrophotometry with Beckman DU spectrophotometer.

 $^{{}^{\}star}k_{\stackrel{\cdot}{N}}=$ adsorption on nylon ${}^{\star\star}k_{\stackrel{\cdot}{A}}=$ adsorption on cellulose triacetate

TABLE 6: Continued

mpound Name					-	Ref.
4-(R ₂ sulfonyl)-2,6-dinitro-N,N- (di-R ₁) aniline				Adsorption equat	ion: ln K _e = aPΔU*	Lambert,
(1967			
	SD	R ₁	R ₂	Ke va Ripperdan soil 1% 0.m	Sacramento soil 5% 0.m	
	11830	CH ₃	CH ₃	125	145	
	12639	^С 2 ^Н 5	CH ₃	230	193	
	11831	с ₃ н ₇	CH ₃	500	520	
	13207	^С 2 ^Н 5	с ₂ н ₅	320	269	
	12030	с ₃ н ₇	с ₂ н ₅	750	702	
	12346	с ₃ н ₇	с ₃ н ₇	1170	-	
	12400	CH ₃	iso-C ₃ H ₇	222		

^{*}K_e = equilibrium constant, estimated from adsorption data $P = \frac{1/4}{\dot{\rho}}, P = \text{parachor}, \rho = \text{density of the liquid, } \gamma = \text{surface tension, } M = \text{molecular weight}$ $\Delta U = \text{difference in internal pressures of the solvent phases}$

TABLE 6: Continued

								Ref.
Adsorbent activated		sorption equ	ation:	у= a·x	b			Spahn et al. 1974
charcoal E	•							257.
	Ads	sorption con	stants:					
	x-1							
	<u>(mr</u>	mol/dm ³	<u>a</u>	<u>b</u>				
	0.3	1 - 2	0.84	0.09				
Adsorbent	Ads	Adsorption equation: $M = \frac{abC}{1+bC}$					Nogami, Nagai, and Uchida,	
carbon black								1968
Urea	a	b	ΔF	ΔН	ΔS	۵S _t	ΔS _s	
Concentration M	10 ⁻³ liter mole	10 ⁻³ liter mole	$\frac{\texttt{Kcal}}{\texttt{mole}}$	Kcal mole	e.u.	e.u.	e.u.	
0	1.63	1.70	-4.45	0.22	15.4	0	15.4	
0.11	1.60	1.70	-4.45		15.4	0.05	15.4	
1.01	1.54	1.08	-4.18		14.5	0.38	14.9	
3.05	1.28	0.615	-3.87		13.5	1.15	14.7	
5.03	1.18	0.472	-3.70		12.9	1.68	14.6	
7.07	0.92	0.397	-3.57		12.5	2.08	14.6	
	Adsorbent carbon black Urea Concentration M 0 0.11 1.01 3.05 5.03	Adsorbent Adsorbent a Carbon black Urea Concentration M 1.63 0.11 1.60 1.01 1.54 3.05 1.28 5.03 1.18	Adsorption con x-range I (mmol/dm ³) 0.1 - 2 Adsorbent carbon black Urea carbon black Oncentration M nole mole 0 1.63 1.70 0.11 1.60 1.70 1.01 1.54 1.08 3.05 1.28 0.615 5.03 1.18 0.472	Adsorption constants: X-range I (mmol/dm ³ a 0.1 - 2 0.84	Adsorption constants: x-range I (mmol/dm ³ a b 0.1 - 2 0.84 0.09 Adsorbent carbon black Urea Concentration M = 10 ⁻³ liter mole M = 10 ⁻³ liter M	Adsorption constants: x-range I	activated charcoal B10 Adsorption constants: x-range I (mmol/dm ³ a b 0.1 - 2 0.84 0.09 Adsorbent carbon black Urea Concentration Mole Mole Mole Mole Mole Mole Mole Mole	activated charcoal B10 Adsorption constants: x-range I

 ΔF = free energy change

 ΔS = entropy change

 $\Delta S_s = \Delta S + \Delta S_t$

 ΔS_{t} = entropy change of the transfer of tryptophan from aqueous solution to aqueous urea solution

e.u. = entropy units

TABLE 6: Continued

	TABLE 6:	Contr	пиец					
Compound Name					Ref.			
Benzene		Adaar	ption equatio					
<pre>benzene - adsorption from binary mixture</pre>			-		Zettlemoyer and Micale,			
с ₆ н ₆		$\frac{x_1 x_2}{n^{\circ} \Delta x/m} = \frac{1}{n^s} \left[x_1 + \frac{1}{K-1} \right]$						
Compound properties								
m.w. 78.11 ^d								
density $.87901^{b}$								
m.p. 5.5°C ^e								
b.p. 80.1°C ^e								
water sol 820 ppm at 22°C f OCT/water part coeff 130 g			Area of adsorbent	(ΔH° - ΔH°*)				
Solvent	Adsorbent	_K_	m^2/g	cal/g				
Ethylene dichloride	Graphon	18.5	119	0.54				
	Spheron 6	5	57	-				
Cyclohexane	Coconut shell charcoal	19.9	677	12.0				
	Decolorizing charcoal	16.4	596	7.1				
Carbon tetrachloride	Decolorizing charcoal	16.1	374	4.0				
	Coconut shell charcoal	18.2	350	3.8				

$$\star_{\Delta H} = X_1^{s} \Delta H_1^{o} + X_2^{s} \Delta H_2^{o}$$

$$K = exp - (\frac{\Delta Ha}{RT} - \frac{\Delta Sa}{R})$$

TABLE 6: Continued

	TABLE 6: Continue	e d			
Compound Name					Ref.
Carbamates					
carbamate derivatives, adsorption from 2% ethanol	Adsorpt	ion equation:	$(\frac{\mathbf{x}}{m}) =$	KC ^{1/n}	Ward and Upchurch,
R ₁ NHCOOR ₂			l/n =	= 1	1965
	<u>Compound</u>	solubility 2% ethanol	k _N *	<u>k_A**</u>	
	Ethyl-N-methyl-N-phenyl	172	0.22	1.42	
	Ethyl-N-benzyl	121	0.49	2.03	
	Ethyl-N-phenyl	96	1.03	3.68	
	Ethyl-N-ethyl-N-phenyl	78	0.25	1.83	
	Ethyl-N-(2-nitrophenyl)	71	0.90	3.25	
	N-Phenylglycine ethyl ester	64	0.16	0.83	
	Phenyl	28	0.40	0.58	
	Isopropyl-N-phenyl	16	1.16	4.95	
	Ethyl-N-(4-chlorophenyl)	9.6	5.70	14.70	
	Ethyl-N-butyl-N-phenyl	8.5	0.80	6.75	
	$n extsf{-} extsf{Butyl-} extsf{N} extsf{-} extsf{phenyl}$	8.5	2.20	9.62	
	<pre>Ethyl-N-(4-nitrophenyl)</pre>	5.7	3.65	8.75	
	Ethyl-N-benzyl-N-phenyl	2.9	1.85	12.65	
	Ethyl-N-(2,5-dichlorophenyl)	2.6	4.00	19.40	
	Ethyl-N-(2,3-dichlorophenyl)	2.5	3.65	27.90	
	Ethyl- N , N -diphenyl	1.7	1.02	9.10	
	<pre>Isopropyl-N-(2-methyl-5- chlorophenyl)</pre>	1.6	3.37	14.00	
	Methyl-N-(2,4-dichlorophenyl)	0.72	15.10		
	<pre>Isopropyl-N-(3-chlorophenyl)</pre>	0.50	9.00	8.45	
	Isopropyl-N-(3,4-dichloropheny	1) 0.32	23.00		

Method of analysis: UV spectrophotometry with Beckman DU spectrophotometer

 $^{{}^{\}star}k_{N}$ = (amount adsorbed/weight of nylon)/(C^{1/n})

^{**} $_{A}$ = (amount adsorbed/weight of cellulose triacetate)/($C^{1/n}$)

TABLE 6: Continued

Compound Name				Ref.
carbamate derivatives, adsorption from 2% ethanol	Adsorption equation:	$\frac{\mathbf{x}}{\mathbf{m}} = \mathbf{KC}^{1/n}$	(1/n = 1)	Ward and Upchurch,
				1965

			^K Ν	
Adsorbent	Compound	10°C	26.5°C	50°C
Nylon	Ethyl-N-phenyl	1.03	1.03	1.03
(Zytel-101) (80 mesh)	<pre>Isopropyl-N-phenyl</pre>	1.16	1.16	1.16
(00	Ethyl-W-methyl-W-phenyl	0.22	0.22	0.22
	Ethyl-N-ethyl-N-benzyl	0.25	0.25	0.25
	Ethyl-N-benzyl-N-phenyl	1.33	1.85	1.85
	Ethyl-N, N-diphenyl	0.66	1.02	1.02
	<pre>Isopropyl-N-(3-chlorophenyl)</pre>	9.00	9.00	9.00
	Ethyl-N-(2-nitrophenyl)	0.75	0.90	0.90
	Ethyl-N-(4-nitrophenyl)	3.65	3.70	3.70
	Ethyl-N-(2,3-dichlorophenyl)	3.70	3.70	3.70

Method of analysis: UV spectrophotometry with Beckman DU spectrophotometer.

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Compound Name					Ref.
Carbonyls					
acetophenone	Adsorbent	Adaomni	ion oguat	ion: y≈ a·x ^b	Spahn et al.,
C6H5COCH3	activated	Adsorpt	.10n equat.	ion: y- a-x	1974
Compound properties	charcoal B 10				±21.
m.w. 120.16 ^h		Adsorpt	cion const	ants:	
m.p. $20.5^{\circ}C^{h}$		u-range	、 T		
b.p. 202.0°C ^h		x-range (mmol/d	_	a_ b_	
density 1.0281 at 20°C/4°C h		0.1 - 4	_	.46 0.155	
water sol $5.6 \times 10^{-3} \text{M}$ at $25 ^{\circ} \text{C}^i$					
benzo[f]quinoline C _{1.2} H ₀ N		Adsorpt	ion equat	ion: $S_s = K_p S_w$, $n=1$	Lee, Chou,
benzo[f]quinoline Cl3H9N Compound properties	Adsorbent	Adsorpt	cion equat	ion: $S_s = K_p S_w$, $n=1$	Bohonos, Holt, Lee, Chou, Bomberger, Mil
C ₁₃ H ₉ N	Coyote Creek			-	Bohonos, Holt, Lee, Chou,
C ₁₃ H ₉ N Compound properties		TOC*	CEC**	<u>кр</u>	Bohonos, Holt, Lee, Chou, Bomberger, Mil
C ₁₃ H ₉ N Compound properties m.w. 179.22	Coyote Creek	TOC*	CEC**	<u>кр</u>	Bohonos, Holt, Lee, Chou, Bomberger, Mil
C ₁₃ H ₉ N Compound properties m.w. 179.22 m.p. 93.5°C b.p. 350°C (721 torr)	Coyote Creek	TOC*	CEC**	<u>кр</u>	Bohonos, Holt, Lee, Chou, Bomberger, Mil
C ₁₃ H ₉ N Compound properties m.w. 179.22 m.p. 93.5°C b.p. 350°C (721 torr) 202-5°C (8 torr)	Coyote Creek	TOC*	CEC**	<u>кр</u>	Bohonos, Holt, Lee, Chou, Bomberger, Mil
C ₁₃ H ₉ N Compound properties m.w. 179.22 m.p. 93.5°C b.p. 350°C (721 torr) 202-5°C (8 torr) vapor pressure at 20°C (torr) water sol (µg ml ⁻¹)	Coyote Creek	TOC*	CEC**	<u>кр</u>	Bohonos, Holt, Lee, Chou, Bomberger, Mil

TABLE 6: Continued

npound Name					Ref.
PH-carbazole C ₆ H ₄ NHC ₆ H ₄		Adsorp	tion equat	ion: $C_s = K_p C_w$	Smith, Mabey, Bohonos, Holt, Lee, Chou,
Compound properties	Adsorbent	<u>Kp</u>			Mill, Bomberge
m.w. 167.21	Ca-montmorillonite	3.20±1	.06		1977ь
m.p. 247-248°C	Coyote creek sediment	175 ±2	0.9		
water sol (1.0 μ g/ml)					
масст 501 (1.0 дд/мг/					
7H-dibenzocarbazole (DBC)					
7H-dibenzocarbazole (DBC)		Adsorp	tion equat	ion: $C_s = K_p C_w$	Smith, Mabey, Bohonos, Holt,
	Adsorbent	Adsorp	tion equat <u>CEC**</u>	ion: $C_s = K_p C_w$ $K_{\underline{p} * * * *}$	Bohonos, Holt, Lee, Chou,
7H-dibenzocarbazole (DBC) C10 ^H 6 ^{NHC} 10 ^H 6	Des Moines River				Bohonos, Holt, Lee, Chou,
7H-dibenzocarbazole (DBC) C10 ^H 6 ^{NHC} 10 ^H 6 Compound properties	Des Moines River sediment	TOC*	CEC**	K _{p ***} 32,600	Bohonos, Holt, Lee, Chou, Mill, Bomberge
7H-dibenzocarbazole (DBC) C10 ^H 6 ^{NHC} 10 ^H 6 Compound properties m.w. 267.31	Des Moines River	TOC*	CEC**	K _{p ***}	Bohonos, Holt, Lee, Chou, Mill, Bomberge

^{*}TOC = total organic carbon

^{**}CEC = cation exchange capacity

^{***}Based on analysis of the supernatant at equilibrium

ompound Name						Ref.
pyridine					n: y= a·x ^b	Spahn
с ₅ н ₅ и		Adsorp	tion	equation	n: y= a·x	et al.,
Compound properties j						1974
m.w. 79.10	Adsorbent	x-rang (mmol/	~	_a_	_b_	
m.p. 42.0°C	activated	0.14 -	7.4	1.2	2 0.2	
b.p. 115.3°C	charcoal B 10					
density .982 at 20°C						
water sol ∞						
pyridine		Adsorp	tion (equation	$n: X = KCe^n, X = mg/g;$	Baker and Luh,
					Ce = mg/1	1971
	Adsorbent	Temp.	рН	<u>K</u>	<u>n</u>	
	Na-kaolinite	1°	2	0.03	1.01	
		24°	2	0.01	1.03	
	Na-montmorillonite	1°	2	0.12	1.04*	
		24°	2	0.06	1.02	

TABLE 6: Continued

Method of analysis: liquid scintillation with Packard Tri-Carb spectrometer

Note: clay: pyridine ratio was varied from 12.15 to 6.25x10⁴ for ratio (12.15) effect of pH was studied:

max adsorption for Na-kaolinite was found at pH 5.5; pka pyridine = 5.25 max adsorption for Na-montmorillonite was found at pH 4.0

^{*}discrepancy noted between data given Table 2 p.842 and regression equation p.843, believe this correct value.

TABLE	6.	Continued

mpound Name					Ref.
quinoline			, _		Smith,
С ₉ н ₇ N		-	ion equa		Mabey,Bohonos Holt, Lee,
Compound properties		K	$=\frac{ug\ ad}{g}$	sorbed/µg in solution	Chou, Mill, Bomberger,
m.w. 129.15	Adsorbent	TOC	CEC	<u>K</u> p	1976b
water sol 6.11 μ g/ml	Ca-montmorillonite	0.05	69	7.28±0.52	
$pR_b = 9.5*$	Coyote Creek sediment	1.4	13.5	10.9±0.4	
m.p14.5°C					
b.p. 161.9°C					

Experimental conditions: quinoline concentrations 4 and 8 μ g/ml⁻¹; sediment concentrations 1000 to 3000 times that of quinoline.

Method of analysis: UV spectrophotometry with Cary Model 11 spectrophotometer.

*Quinoline is calculated to be 97%, 24%, and 0.32% protonated at pH 3, 5, and 7, respectively.

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	TABLE 6: Cont	inued				
Compound Name						Ref.
S-Heterocyclics						
$\mathtt{benz}_o[b]$ thiophene						Smith, Mabey,
с ₈ н ₆ s		Adsor	ption eq	uation:	$S_s = K_p S_w, n=1$	Bohonos, Holt, Lee, Chou,
Compound properties	Adsorbent	TOC	CEC	K _p		Mill, Bomberge
m.p. 31.3°C	Coyote Creek sediment	1.4	13.5	50±5		1976c
b.p. 212.9°C						
water sol 127.3±2.5 at 20°C	!					
	Experimental conditions: 5000:1 sediment: BT by		loading	s of 200	0:1 and	
dibenzothiophene		Adeor	ption eq	nation:	S = K S . n=1	Smith, Mabey,
с ₁₂ н ₈ s		Ausor	brion ed	uu cion.	s = K S , $n=1$	Bohonos, Holt, Lee, Chou,
Compound properties	Adsorbent	Toc	CEC	<u>ъ</u>		Bomberger, Mil
m.w. 184.27	Coyote Creek sediment	1.4	13.5	1380±1	.30	1977
m.p. 99-100°C						
b.p. 332-333°C						
water sol 1.11±0.09						

TABLE 6: Continued

Compound Name			Ref.
Phenols		Adsorption equation: y=a·x ^b	Spahn et al.,
phenol $C_6^{H}{}_5^{OH}$ Compound properties m.w. 9411^d m.p. 40.9°C^e $\text{b.p. 180°C/740mm}^k$ $\text{density 1.072 g/m1}^b$ $\text{water sol 84.12 mg/m1}^k$	Adsorbent activated charcoal B 10	X-range I $\frac{(mmol/dm^3)}{0.1-8}$ a b $\frac{b}{0.23}$ X-range II $\frac{(mmol/dm^3)}{0.001-0.06}$ a b $\frac{b}{0.39}$	1974
Polynuclear aromatics benz[a]anthracene C18H12 Compound properties m.w. 228.28 m.p. 155-7°C b.p. (at 760 torr) 435°C water sol at 27°C	Adsorbent Coyote Creek sediment	Adsorption equation: $s_s = K_p S_w$, $n=1$ TOC CEC $\frac{K_p}{1.4}$ 13.5 26,200±1700	Smith, Mabey, Bohonos, Holt, Lee, Chou, Bomber ger, Mill,

ompound Name					Ref
benzo[a]pyrene		Adsorption equa	ition: x=K _p C		Smith, Mabey Bohonos,
Compound properties m.w. 252.32 ^a m.p. 178°C ¹ b.p. ~500°C 311°C at 10 torr water sol 1.2±0.1 ng/ml	Adsorbent Ca-montmorillonite Des Moines River Coyote Creek Searsville Pond	Total organic carbon (%) 0.06 0.6 1.4 3.8	CEC meg. 100g ⁻¹ 69.0 10.5 13.5 34.5	$ \frac{(\times 10^{-4})}{1.7 \pm 0.5} $ 3.5\pm 2.7 7.6\pm 2.4 15.0\pm 2.2	Holt, Lee, Chou, Mill, Bomberger, 1976a
pyrene C16 ^H 10 Compound properties m.w. 202 ^I m.p. 150°C T		Adsorption equation: q = heat evolved when q° = total heat evolv Equilibrium constants	the [cyclohex		Groszek, 1975
b.p. 393°C e density 1.1271 a water sol .135±.005 mg/1 Oct water part coeff: g KOW=150,000	Adsorbent graphon oleophilic graphite "polar" graphite	-	st. from heat	18.7	H /mole 23 18 26

Method of analysis: UV spectrophotometry with Unicam SP 500 spectrophotometer.

^{*}at 21°

	TABLE 6:	Continued		
Compound Name				Ref.
pyrene Compound properties		Adsorption equation:	x=k _p C; koc=k _p /oc	Karickhoff, Brown, and Scott,
<pre>water sol (mole fractionX10⁹)=12</pre>				1979
Koctanol water=150,000				

Adsorbent	% 0.C.	K _p	$KOC(X10^{-5})$
Hickory Hill			
sand coarse silt medium silt fine silt clay	0.13 3.27 1.98 1.34 1.20	42 3000 2500 1500 1400	0.32 0.92 1.3 1.1
Doe Run			
sand coarse silt medium silt fine silt clay	0.086 2.78 2.34 2.89 3.29	9.4 2100 3000 3600 3800	0.11 0.76 1.3 1.2
Oconee River			
sand coarse silt medium silt fine silt clay	0.57 2.92 1.99 2.26	68 3200 2300 2500	0.12 1.1 1.2 1.1

Experimental conditions: sorbent concentrations 400 mg/ml of suspension for sand, 20 mg/ml for coarse and medium silt, 10 mg/ml for fine silt and 1 mg/ml for clay.

Method of analysis: UV spectrophotometry with Perkin Elmer 356 spectrophotometer.

· · · · · · · · · · · · · · · · · · ·			Ref.
Quinone	Adsorbent		
alizarin	activated Adsorption ed	uation: y= a·x ^b	Spahn et al.,
C ₁₄ H ₈ O ₄	В 10	,	·
Compound properties ^a	Adsorption co	onstants:	1974
m.w. 240.23	x-range I		
m.p. 289-90°C (cor)	$(mmol/dm^3)$	<u>a</u> <u>b</u>	
b.p. 430°C (sub)	0.1 - 8	1.39 0.095	
water sol slightly sol			
16			
ulfonate sodium naphthalenesulfonate	Advertion of	$C = \frac{1}{2} + \frac{C}{2}$	
		quation: $\frac{C}{q} = \frac{1}{Kb} + \frac{C}{b}$	et al.
	Adsorbent Adsorption eq	quation: $\frac{C}{q} = \frac{1}{Kb} + \frac{C}{b}$	
		ΔFu°** ΔSu°***	et al.
	Amberlite XAD-2 t K b ΔH°*	ΔFu°** ΔSu°*** e Kcal Kcal	Gustafso et al. 1968
Sulfonate sodium naphthalenesulfonate	Amberlite XAD-2 t K b AH°*	ΔFu°** ΔSu°*** <u>Kcal</u> <u>Kcal</u> -5.7 +4.4	et al.
	t K b AH°* 1/mole mole/g Kcal/mol 25°C 319 4.58x10 ⁻⁵ -4.4 Method of analysis: UV spectrophoto	ΔFu°** ΔSu°*** <u>Kcal</u> <u>Kcal</u> -5.7 +4.4	et al.
sodium naphthalenesulfonate $^{\star}\Delta \text{H}^{\circ} = 2.303 \frac{\text{RT}_{1}\text{T}_{2}}{(\overline{\text{T}}_{2}^{-}\text{T}_{1}^{-})} (\text{LogK}_{2}^{\prime} - \text{logK}_{2}^{\prime})$	t K b AH°* 1/mole mole/g Kcal/mol 25°C 319 4.58x10 ⁻⁵ -4.4 Method of analysis: UV spectrophoto	ΔFu°** ΔSu°*** <u>Kcal</u> <u>Kcal</u> -5.7 +4.4	et al.

TABLE 6: Continued

mpound Name									Ref.
		PESTICIDES							
rbamates									
carbaryl									LaFleur
N-methyl-1-naphthylcarbamate		Desor	Desorption Equation: K_{b}		$K_b = 1$	P _s /P _w			1976a
C ₁₂ H ₁₁ NO ₂		Adsor	Adsorption Equation:			$K_f = P_s/P_w$			
Compound properties ^p	Adsorbent	<u>рН</u>	go.m.	1/0.5	1/1	1/2	1/4	1/1	
m.w. 201	Lakeland s	5.3	0.22	0.44	0.21	0.16	0.13		
water sol 350 µmol/l	Norfolk ls	6.0	0.57	1.4	0.36	0.31	0.25		
, , , , , , , , , , , , , , , , , , ,	Norfolk scl	5.4	0.15	0.72	0.12	0.10	0.08		
	Cecil sl	6.3	1.77	5 .2	1.8	1.6	1.3	0.79	
	Cecil c	5.7	0.53	0.88	0.27	0.22	0.19		
	Okenee sl	4 65	5.16	7.6	3.7	2.6	2.3		

Chlorinated hydrocarbons		Adsorption	equation:	$x/m = KC^{1/n}$	Huang and Liao,
1,1'-(2,2,2-trichloroethylidene)-	Adsorbent	K	<u>1/n</u>		1970
bis[4-chlorobenzene]	illite	2.72x10 ⁻³	3.28		
C ₁₄ H ₉ C1 ₅	kaolinite	7.37×10^{-6}	5.08		
Compound properties	montmorillonite	1.10x10 ⁻⁵	5.97		
m.w. 354.49	Method of analysis	· GC - elect	tron cantur	- 6	

Method of analysis: GC - electron capture

TABLE 6: Continued

mpound Name				Ref.
dieldrin			x - val/n	Huang and
1,2,3,4,10,10-hexachloro- 6,7-epoxy-1,4,4a,5,6,7,8,8a- octahydro-endo-1,4-exo-5,8- dimethanonaphthalene		Adsorption equa	tion: $\frac{x}{m} = KC^{1/n}$	Liao, 1970
C ₁₂ H ₈ C1 ₆ 0				
Compound properties	Adsorbent	<u>K</u>	<u>1/n</u>	
m.w. 380.91	illite	9.45×10^{-16}	8.82	
	kaolinite	1.46×10^{-21}	11.63	
	montmorillonite	1.05x10 ⁻¹⁶	9.24	
	Method of analysis:	GC-electron captu	ure.	
otachlor		Adcorption ogua	tion: $\frac{\kappa}{m} = \kappa c^{1/n}$	Huang and Liao,
1,4,5,6,7,8,8-heptachloro- 3a,4,7,7a-tetrahydro-4,7- methano-1#-indene		Adsorption equa	m - Ke	1970
C ₁₀ H ₅ Cl ₇				
	Adsorbent	<u>K</u>	<u>1/n</u>	
	illite	1.09x10 ⁻⁹	6.07	
	kaolinite	5.00x10 ⁻⁶	4.51	
		1.48X10 ⁻⁴		

Method of analysis: GC-electron capture.

TABLE 6: Continued

•	TABLE	6: Continue	: a		
mpound Name					Ref.
methoxychlor					Karickhoff,
^C 16 ^H 15 ^{Cl} 3 ⁰ 2		Adsorpti	on equat	zion: $X = K_pC$; $KOC = K_p/OC$	Brown, and Scott,
Compound properties c	Adsorbent	% O.C.	<mark>к</mark> р	$KOC(X10^{-5})$	1979
m.w. 345.65	Hickory Hill				
m.p. 78-78.2°C or	sand	0.13	53	0.41	
86-88°C	coarse silt	3.27	2600	0.80	
	medium silt	1.98	1800	0.91	
insol. in water	fine silt	1.34	1400	1.0	
	clay	1.20	1100	0.92	
	Doe Run				
	sand	0.086	8.3	0.097	
	coarse silt	2.78	2200	0.80	
	medium silt	2.34	1700	0.73	
	fine silt	2.89	2300	0.80	
	clay	3.29	2400	0.73	
	Oconee River				
	sand	0.57	95	0.17	
	coarse silt	2.92	2500	0.86	
	medium silt	1.99	2000	1.0	
	fine sand	2.26	2100	0.93	
	clay				

Experimental conditions: sorbent concentrations 400 mg/ml of suspension for sand, 20 mg/ml for coarse and medium silt, 10 mg/ml for fine silt and 1 mg/ml for clay.

Method of analysis: UV spectrophotometry with Perkin Elmer 356 spectrophotometer.

	TABLE 6: Co	ntinue	d			
Compound Name						Ref
mirex					, , , , , , , , , , , , , , , , , , ,	Crith Mahou
c ₁₀ c1 ₁₂			Adsorpti	on equa	tion: $S = KS$, $n=1$	Smith, Mabey, Bohonos, Holt, Lee,
Compound properties	Adsorbent	TOC	CE	EC.	К	Chou, Bomber ger, Mill,
m.w. 546.0	Coyote Creek	===	<u> </u>	==	K _P	1977
m.p. 485 (decomp)	sediment	1.4	13	3.5	460,000 ± 110,000	
<pre>vapor pressure at 50° torr 6 x 10⁻⁶</pre>						
water sol at 22°C						
(pg/ml) 70±20						
Organophosphates						
aminoparathion			Adsorpti	ion egua	tion: $x/m = KC_e^{1/n}$	Bowman and Sans,
C ₁₀ H ₁₆ NO ₃ PS						1977
0, 0 -diethyl 0 - p -aminophenyl phosphorothioate	Adsorbents		<u>1/n</u>	<u>K</u>		
Compound properties	Na-montmorillonite		0.954	46.3		
	Ca-montmorillonite	*	1.151	43.3		
m.w. 261.3	Fe-montmorillonite	*	>99.9%			

Experimental conditions: $450\ \mathrm{mg}$ clay in 30 ml aqueous insecticide solution

Method of analysis: GC - alkali flame ionization detector

adsorption

water sol 390.0 µg/ml

^{*}Prepared from Wyoming bentonite, < 2 µm fraction

TABLE 6: Continued

### Recall	ompound Name				Ref.	
0,0-dimethyl 0-(3-methyl-4-nitrophenyl) phosphorothioate 1977	fenitrothion	Adsorption equation: $\frac{x}{n} = KC_{o}^{1/n}$				
Compound properties m.w. 277.2 Na-montmorillonite* 1.163 71.1 b.p. 118°C Ca-montmorillonite* 1.952 64.0 Fe-montmorillonite* 1.773 740.1 Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector.			-	- m e	1977	
m.w. 277.2 Na-montmorillonite* 1.163 71.1 b.p. 118°C Ca-montmorillonite* 1.952 64.0 density 1.322745 water sol 25.4 µg/ml Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector. Method of anal	C ₉ H ₁₂ NO ₅ PS					
b.p. 118°C density 1.3227% Fe-montmorillonite* 1.952 64.0 fee-montmorillonite* 1.773 740.1 Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector. Method of analysis: GC-alkali flame ionization. Method of analysis: GC-alkali flame ionization. Me	Compound properties	Adsorbent	<u>1/n</u>	<u>K</u>		
density 1.3227 ² / ₄ 5 water sol 25.4 μg/ml Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector. Method of analysis: GC-alkali f	m.w. 277.2	Na-montmorillonite*	1.163	71.1		
density 1.3227 $_{4}^{25}$ water sol 25.4 μ g/ml Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector. Method of analysis: $\frac{x}{m} = KC_{e}^{1/n}$ Bowman and Sa 1977 $C_{8}H_{10}NO_{5}PS$ Compound properties M. W. 263.2 M. Ma-Montmorillonite* 1.032 Ma-Montmorillonite* 1.032 Ma-Montmorillonite* 1.663 Ma-Montmorillonite* 1.663 Mensity 1.358 $_{4}^{2}$ 0° Water sol 45.0 μ g/ml Experimental conditions: same as for aminoparathion.	b.p. 118°C					
Experimental conditions: same as for aminoparathion. Method of analysis: GC-alkali flame ionization detector. Bowman and Salling an	density 1.3227 ²⁵	Fe-montmorillonite*	1.773	740.1		
Method of analysis: GC-alkali flame ionization detector. Bowman and Sa 1977 C ₈ H ₁₀ NO ₅ PS Compound properties Adsorbent 1/n \underline{K} m.w. 263.2 Na-Montmorillonite* 1.032 65.4 m.p. 37-38°C° Ca-Montmorillonite* 1.663 56.3 density 1.358 $_4^2$ 0° Fe-Montmorillonite* 1.463 147.3 water sol 45.0 μ g/ml Experimental conditions: same as for aminoparathion.	water sol 25.4 μg/ml					
methyl parathion O,O-dimethyl O-p- nitrophenyl phosphorothioate C ₈ H ₁₀ NO ₅ PS Compound properties m.w. 263.2 Ma-Montmorillonite* 1.032 Ma-Montmorillonite* 1.663 M.p. 37-38°C° density 1.358 2_4 °° water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.		Experimental condition	ns: same as f	for aminoparathion.		
Adsorption equation: $\frac{x}{m} = KC_e^{1/n}$ and Sand Sand Sand Sand Sand Sand Sand		Method of analysis:	GC-alkali flam	ne ionization detector.		
0,0-dimethyl 0-p- nitrophenyl phosphorothioate $C_8H_{10}NO_5PS$ Compound properties Madsorbent 1/n M.w. 263.2 Na-Montmorillonite* 1.032 M.p. 37-38°C° Ca-Montmorillonite* 1.663 Mensity 1.358 $_4^2$ 0° water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.	nethyl parathion		Adsorption e	equation: $\frac{x}{x} = KC^{1/n}$	Bowman and Sans,	
Compound properties m.w. 263.2 Na-Montmorillonite* 1.032 65.4 m.p. $37-38^{\circ}C^{\circ}$ Ca-Montmorillonite* 1.663 fe-Montmorillonite* 1.463 water sol $45.0 \mu g/ml$ Experimental conditions: same as for aminoparathion.				т пе		
m.w. 263.2 Na-Montmorillonite* 1.032 65.4 m.p. 37-38°C° Ca-Montmorillonite* 1.663 fe-Montmorillonite* 1.463 147.3 water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.	C ₈ H ₁₀ NO ₅ PS					
m.p. 37-38°C° Ca-Montmorillonite* 1.663 56.3 density 1.35840° Fe-Montmorillonite* 1.463 147.3 water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.	Compound properties	Adsorbent	<u>l/n</u>	<u>K</u>		
m.p. 37-38°C density 1.3584°C water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.	m.w. 263.2	Na-Montmorillonite*	1.032	65.4		
density 1.35840° Fe-Montmorillonite* 1.463 147.3 water sol 45.0 µg/ml Experimental conditions: same as for aminoparathion.	m.p. 37-38°C°		1.663	56.3		
50 ppm ^c		Fe-Montmorillonite*	1.463	147.3		
50 $ exttt{ppm}^{\mathcal{C}}$ Method of analysis: GC-alkali flame ionization detector.		Experimental condition	ons: same as i	for aminoparathion.		
	50 ppm^c	Method of analysis:	GC-alkali flam	ne ionization detector.		

^{*}Prepared from Wyoming bentonite <2 μm fraction.

TABLE 6: Continued

Ref.
Smith, Mabey, Bohonos, Holt Lee, Mill,
Bomberger,
1976
ז

Experimental conditions: methyl parathion concentrations at 4 and 8 µg/ml. Sediment concentrations were 1000 to 2000 times as concentrated as methyl parathion at 8 µg/ml and 1000 to 3000 times as concentrated as MP at 4 µg/ml.

TABLE 6: Continued Compound Name Ref parathion Adsorption equation: $x/m = KC^{1/n}$ or $\log x/m = \log K + 1/n \log C$ Wahid and Sethunathan, 1978 Adsorbent Soil organic Oxidized Soils Natural Soils Soil No.* рН matter, % CEC K 1/n K <u>1/n</u> 10 6.20 0.75 18.6 7.67 1.04 8 6.25 1.62 26.6 12.30 1.05 3.16 1.33

2.88

5.52

8.21

24.62

Experimental conditions: 1 g soil in 10 ml aqueous parathion solution

Method of analysis: liquid scintillation

6.30

5.20

3.50

3.30

11

13

15

14

amitrole

3-amino-1,2,4-triazole

C2H4N4

 $pKa = 4.14 (3-ATH^{+}=H^{+}+3-AT)$

Compound properties

water sol 28 g/100 ml (23°C)

Adsorbent H sat. organic matter

Al sat. organic matter

. X_m 63 mM/100g

42.8 38.02

19.2 125.90

21.2 213.80

28.9 457.10

1.11

1.05

1.03

1.02

Adsorption equation: $X = X_mbC/(1+bC)$

b**

0.015

0.010

10.72

1.33

Nearpass,

1969

49 mM/100g

Experimental conditions: 20:1 water to organic matter

Method of analysis: liquid scintillation

*all soils from India **calculated from figure 2

∞
1.1
•••

	TABLE 6:	Cont	inued					
Compound Name								Ref.
s-Triazine								
cyanazine		Adso	orption eq	uation	: <u>×</u> =	= KC ^{l∕n}		Majka and Lavy,
2-[[4-chloro-6-(ethylamino)- s-triazin-2-yl]amino]-2- methylpropionitrile			Chromato					1977
^C 9 ^H 13 ^{C1N} 6	Adsorbents	<u>рН</u>	% o.m.	CEC	<u>K</u>	<u>1/n</u>	Rf	
	Monona silty clay loam (Typic Hapludol1)	6.5	2.9	21.2	4.6	0.96	0.39	
	Valentine loamy fine sand (Typic Ustipsamment)	6.6	1.4	10.1	3.4	0.86	0.74	
	Experimental condition 0.2 ml methanol:			th cya	nazine	e added	in	
	Method of analysis: 1 spectrometer.	iquid	scintill	ation	with F	Packard	3320	

TABLE 6: Continued

Compound Name							-			Ref
prometryne				_	eption I	1	s			Lafleur,
$c_{10}^{H}_{19}^{N}_{5}^{S}$				Desoi	rption I	K _b = 7	W			1976b
2,4-bis(isopropylamino)- 6-methylthio-1,3,5- triazine	Adsorbent	рн	%O.M	<u>ccc</u> *	1/0.5	1/1	1/2	1/4	ΔG _{sp} ** (Kcal/mol)	K _o (calc)***
Compound properties	Lakeland s	5.3	0.22	3.0	1.5	0.86	0.74	0.67	-4.1	1060
m.w. 241	Norfolk ls	6.0	0.57	6.5	2.1	1.5	1.2	1.0	-4.1	990
water sol 200 µM/l at	Norfolk scl	5.4	0.15	28	5.7	3.4	2.6	2.3	-3.6	420
20°C	Cecil sl	6.3	1.77	15	7.9	4.0	3.2	2.8	-4.1	1110
pKa = 4.05	Cecil c	5.7	0.53	18	3.7	2.3	1.7	1.6	-3.3	280
	Okenee sl	4.65	5.16	19	19	12	9.0	7.0	-4.6	2340

Method of analysis: UV spectrophotometry

^{*}Cation combining capacity - methylene blue adsorption at pH5. ** $\Delta G_{\rm sp} = -RTlnk_{\rm o}$ *** $K_{\rm o} = P_{\rm a}/P_{\rm w}P_{\rm a}$ is $P_{\rm s}$ adjusted to soil weight

compound Name								Ref.
'rea								
diuron		Adsor	otion equ	uation:	$\frac{\mathbf{x}}{\mathbf{x}} = \mathbf{K}$	C ^{l/n}		Majka and Lavy,
3-(3,4-dichlorophenyl)-1,1-dimethylurea			Chromato				il)	1977
$C_9^{H}_{10}^{C1}_{2}^{N}_{2}^{0}$	Adsorbents	DĦ	% o.m.	CEC	ĸ	l/n	$^{\mathrm{R}}f$	2377
Compound properties c		рH						
m.w. 233.10	Monona silty clay loam (Typic Hapludoll)	6.5	2.9	21.2	14.3	0.77	0.18	
m.p. 158-159°C	Valentine loamy fine sand	6.6	1.4	10.1	6.5	0.74	0.39	
water sol 42 ppm at 25°C	(Typic Ustipsamment)							
	Experimental condition	s: sam	ne as fo	r cyana	zine.			
	Method of analysis: 1	iania s	cintill:	ation w	ith Pac	kard 33	20	
	spectrometer.	rquru s	CHICLL				20	
fluometuron		iquiu s	,cincili				20	_
1,1-dimethy1-3-(α,α,α-								_
1,1-dimethy1-3-(α , α , α -trifluoro- m -toly1)urea			Chromato					Stritzke
1,1-dimethyl-3-(a,a,a-trifluoro-m-tolyl)urea C10H11F3N20	spectrometer.	(Chromato	graphy	thin-la			Stritzke
l,l-dimethyl-3-(a,a,a-trifluoro-m-tolyl)urea	spectrometer.	€ O.m.	Chromato	graphy <u>p</u> H	thin-la			Stritzke
1,1-dimethyl-3-(a,a,a-trifluoro-m-tolyl)urea CloHl1F3N20	spectrometer.	(Chromato	graphy	thin-la			Chang and Stritzke 1977
1,1-dimethy1-3-(\alpha,\alpha,\alpha-ctifluoro-m-toly1)urea C10H11F3N20 Compound properties	spectrometer.	€ O.m.	Chromato	graphy <u>pH</u> 6.3	thin-la			Stritzke
l,l-dimethyl-3-(a,a,a-trifluoro-m-tolyl)urea CloHl1F3N20 Compound properties m.w. 232.21	Adsorbent Bethony silt loam	% o.m.	Chromato <u>CEC</u> 12.4	graphy <u>pH</u> 6.3	thin-la R <u>f</u> 0.57			Stritzke
1,1-dimethyl-3-(a,a,a-trifluoro-m-tolyl)urea C10H11F3N20 Compound properties m.w. 232.21 m.p. 163-164°C	Adsorbent Bethony silt loam Hector loam	% o.m. 4.4 4.8 0.3 s: 10	Chromato . <u>CEC</u> 12.4 9.2 0.6 g soil:	graphy <u>pH</u> 6.3 6.3 6.7	thin-la Rf 0.57 0.65 0.98 of 0.0	yer (so	il)	Stritzke

TARLE	6 •	Continued

mpound Name	TABLE 6	: Continu	<u> </u>			Ref.
tebuthiuron N-[5-(1,1-dimethyl)ethyl]- thiadiazol-2-yl-N,N'-dime						Chang and Stritzke 1977
C ₉ H ₁₆ N ₄ OS	_	Chromat	ography	thin-	layer (soil)	
Compound properties		Water s	olvent			
water sol 2500 μg/ml	Adsorbent	% O.m.	CEC	рН	$\frac{R_{f}}{}$	
	Bethony silt loam	4.4	12.4	6.3	0.58	
	Hector loam	4.8	9.2	6.3	0.66	
	Eufaula sand	0.3	0.6	6.7	0.98	

Experimental conditions: 10 gm of soil: 10 ml of 0.01 N CaCl₂ solutions containing 1,2,4, or 8 ppm tebuthiuron.

Method of analysis: liquid scintillation.

TABLE 6: Continued

Compound Name				Ref.
paraoxon 0,0-diethyl 0-p-		Adsorption equati	on: $\frac{x}{m} = KC_e^{1/n}$	Bowman and Sans,
nitrophenyl phosphate				1977
^C 10 ^H 14 ^{NO} 6 ^P				
Compound properties	Adsorbent	<u>1/n</u>	K	
m.w. 275.2	Na-Montmorillonite*	1.157	53.7	
water sol 3,250.0 μg/ml	Ca-Montmorillonite*	1.449	42.9	
	Fe-Montmorillonite*	1.376	81.7	
	Experimental condition	ons: same as for am	ninoparathion.	
	Method of analysis:	GC-alkali flame ion	ization detector.	
parathion		Adsorption equati	on: $\frac{x}{m} = KC_0^{1/n}$	Bowman and Sans,
0,0-diethyl 0-p- nitrophenyl phosphorothioate			ш е	
• • •				1977
^C 10 ^H 14 ^{NO} 5 ^{PS}				
Compound properties	Adsorbent	<u>1/n</u>	<u>K</u>	
m.w. 291.3	Na-Montmorillonite*	1.042	119.8	
m.p. 6.1^{α}	Ca-Montmorillonite*	1.513	120.8	
b.p. 375°C ^a	Fe-Montmorillonite*	1.194	605.8	
-	Experimental condition	ons: same as for am	inoparathion.	
density $1.2704\frac{2}{2}8^{a}$	Method of analysis:	GC-alkali flame ion	nization detector.	
water sol 12.9 μ g/ml	-			

^{*}Prepared from Wyoming bentonite, <2 µm fraction.

NOTES, TABLE 6

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 $[^]m {\it Mackay}$, D., and W. Y. Shiu. 1977. Aqueous solubility of polynuclear aromatic hydrocarbons. J. Chem. Eng. Data 22:399-402.

TABLE 7: AVERAGE VALUES OF ADSORPTION CONSTANTS FOR ORGANIC PESTICIDES

Compound Name

S-triazines

	Average*	Average* K±S%	Soils (No.)	Average** Kd±S%	Average*** K tS%	Soils (No.)
simazine	0.820±3.48		6	1.93±6.69	135±6.33	174
6-chloro-N, N'-diethyl-1,3,5-triazine-2, 4-diamine				16.3±30	159±5.66	2
^C 7 ^H 12 ^{C1N} 5 m.w. 201.66 ^a						
propazine	0.95±1.7	2.68±33.0	4	2.37±7.06	160±11.6	54
2-chloro-4,6-bis (isopropylamino)-8-triazine				26.7±9.71	152±3.7	17
^C 9 ^H 16 ^{ClN} 5 m.w. 229.71 ^a						

$$*K = \frac{C_s}{C_w^{1/n}}$$

**
$$K_d = \frac{C_s}{C_w}$$

$$K_{OC} = \frac{X/m(\mu g/g \text{ of organic carbon})}{C_{eq}} = \frac{K_{d(\mu g/g \text{ of soil})}}{% \text{ organic carbon}}$$

Value given is average of K and $K_{\overline{d}}$ adjusted for organic carbon

TABLE 7: Continued

	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soil (No.
atrazine	0.772±8.65	Kiba	19	2.94±10.8	172±6.42	79
6-chloro-N-ethyl-N'- (1-methylethyl)-1,3,5- triazine-2,4-diamine	0.709±10.7		6	25.5±20.4	102±5.80	_24
C8H14ClN5						
m.w. 215.69 ^a						
prometone	0.81±2.1	3.65±26	4	3.31±16.5	300±19.5	49
2,4-bis(isopropylamino)- 6-methoxy-s-triazine				13.1±11.5	73.8±5.1	<u>15</u>
^C 10 ^H 19 ^N 5 ⁰						
m.w. 225.30 ^a						
metryne				6.17±11.1	380±10.0	33
<pre>N-ethyl-N'-(l-methylethyl) -6-(methylthio)-1,3,5-triazine- 2,4-diamine</pre>				167.4	802	1
с ₉ н ₁₇ n ₅ s						
m.w. 227.33 ^a						
prometryne	0.855±1.1	7.83±31	4	7.03±8.78	513±13.0	53
2,4-bis(isopropylamino)-6-(methylthio)- s -triazine	1.76	50.3	<u>_1</u>	55.8±8.46	311±5.12	_18
с ₁₀ н ₁₉ N ₅ s						
m.w. 241.36 ^a						

TABLE 7: Continued

	TA	BLE 7: Contin	иеа			
ompound Name						
	Average*	Average* K±S%	Soils (No.)	Average** K + S%	Average*** K _{oc} +S%	Soils (No.)
ipazine				47.7	2571	1
6-chloro-N, N-diethyl- N'-(1-methylethyl)-1,3,5- triazine-2,4-diamine						
^C 10 ^H 18 ^{C1N} 5						
m.w. 243.74 ^a						
hydroxyatrazine				51.5	888	1
G30026	0.702	<u>37.4</u>			81.8	
Norazine						
C7H12ClN5						
m.w. 201.66 ^a						
eas and Uracils						
Urea	0.717	5.22			14.3	_1
CH ₄ N ₂ 0						
Compound properties b						
density 1.32^{18}_4						
m.w. 60.06						
m.p. 132.7°C						

m.p. 157.5-160°C

TABLE 7: Continued

	Average*	Average*	Soils (No.)	Average** Kd ±S%	Average*** Koc ^{±S%}	Soi:
fenuron	0.953±0.95	0.554±20.6	3	0.781±17.0	0.554±20.6	13
N, N-dimethyl-N'-phenylurea; 1,1-dimethyl-3-phenylurea	1.01±3.20	6.22±35.3	_3_	1.01±3.2	0.622±35.3	1
$^{\rm C}_{9}^{\rm H}_{12}^{\rm N}_{2}^{\rm O}$						
m.w. 164.21 ^a						
methylurea	0.604±2.38	1.73±26.3	3		70.2±7.59	
	0.832±4.71	12.5±48.3	_3_		57.6±8.43	
phenylurea	0.744±1.34	1.80±14.8	3		76.7±8.83	
с ₇ н ₈ № ₂ 0	0.733±2.90	21.7±52.9	_3_		98.3±12.9	
Compound properties b						
m.w. 136.15						
density 1.302						
m.p. 147°C						
b.p. 238°C						
promacil	0.58, 0.85	0.08, 1.5	2	19, 123		
5-bromo-6-methyl-3-(l-methylpropuracil	py1) -					
$^{\mathrm{C_9^H_{13}BrN_2^0_2}}$						
Compound properties b						

TABLE 7: Continued

	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soils (No.)
terbacil	0.50, 0.96	0.15, 1.7	2		37, 140	
5-chloro-3-(1,1-dimethylethyl)-6-methyl-2,4($1H$, $3H$)-pyrimidinedione						
$c_9^{H_{13}^{ClN_2O_2}}$						
m.w. 216.67 ²						
monuron	0.74	5.98	1	2.17±18.9	83.1±22.9	31
N' - (4-chloropheny1) - N, N-dimethylurea	0.83	23.6	1	33.3±22.3	163±16.7	6
C9H11ClN20						
Compound properties b						
m.w. 198.65						
m.p. 170.5-171.5°C						
water sol 230 ppm at 25°C						
diuron	0.818±6.45		10	6.29±9.33	351±8.95	79
3-(3,4-dichlorophenyl)- 1,1-dimethylurea	0.707±5.10		_3_	196±39.2	902±31.9	4
$c_9H_{10}cl_2N_20$						
Compound properties b						
m.w. 233.10						
water sol 42 ppm at 25°C						

TABLE 7: Continued

	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** Kd ^{±S%}	Average*** Koc ±S%	Soils (No.)
metobromuron					60±20	4
3-(p-bromophenyl)-1- methoxy-1-methylurea						
$^{\mathrm{C}_{9^{\mathrm{H}}_{11}^{\mathrm{BrN}}_{2}^{\mathrm{O}}_{2}}$						
m.w. 259.11 ^a						
chlorobromuron					217±20	4
monolinuron	0.810±0.302	9.24±9.75	3	9.24±9.75	195±38.9	7
3-(4-chloropheny1)-1- methoxy-1-methylurea	0.832±2.33	43.0±48.5	_3_	43.0±48.5	198±8.67	3
C9H11C1N2O2						
m.w. 214.65a						
neburon	0.956±2.51	27.9±53.4			787±6.89	3
1-buty1-3-(3,4-dichlorophenyl)- 1-methylurea						
$c_{12}H_{16}Cl_2N_2O$						
m.w. 275.18						
inuron	0.75, 0.75	45.2, 67.8	2	18.1±18.7	840±20	26
N'-(3,4-dichlorophenyl)- N-methoxy-N-methylurea				147±77.2	653±13.4	3_
Compound properties						
m.w. 249.10						
m.p. 93-94°C						
water sol 75 ppm						

TABLE 7: Continued

	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K OC C OC	Soils (No.)
logenated Hydrocarbons						
ethylene dibromide	0.967, 0.970	0.408, 0.803	2	0.537±24.8	32.4±23.7	3
1,2-dibromoethane	1.04	2.16	1	2.13±1.41	14.4±19.4	_2_
C2H4Br2						
m.w. 187.87a						
beta-BHC	0.861	148	1		4254	
1,2,3,4,5,6-hexachlorocyclo-hexane						
C6H6C16	0.950	456_	1_		3573	
m.w. 290.85 ^a						
lindane	0.841,080	45.7, 7.91	2	24.7±27.8	1342±15.3	6
$1\alpha, 2\alpha, 3\beta, 4\alpha, 5\alpha, 6\beta$ - hexachlorocyclohexane	0.981	331	1	321±6.10	1943±20.4	_4

Compound properties

m.w. 290.85

m.p. 112.5°C

insol. in water

TABLE 7: Continue	ıued
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	Average*	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soils (No.)
N-serve				6.80±30.1	271±11.5	9
2-chloro-6-(trichloromethyl)- pyridine				44.5	238	1_
C6H3Cl4N						
Compound properties b						
m.w. 230.93						
m.p. 62.5-62.9°C						
b.p. 136-137.5°C						
DDT				1.3x10 ⁵ , 1.37x10 ⁴	1.31x10 ⁵ 3.55x10 ⁵	2
				1.063X10 ⁵	$2.29x10^{5}$	_1_
rbamates						
chloropropham	1.00	21	1	9.28±11.8	590±8.35	65
isopropyl-m-chlorocarbanilate	1.00	<u>132</u>	_1_			
^C 10 ^H 12 ^{C1NO} 2						
m.w. 213.67 ^a						
propham					51	4
isopropyl carbanilate						
$^{\rm C}_{10}^{\rm H}_{13}^{\rm NO}_{\rm 2}$						
m.w. 179.22 ^a						

l	1	
8	•	•
		•

m.w. 203.35^a

	Average* 1/n±S%	Average* <pre>K±S%</pre>	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soils (No.)
EPTC				5.96±24.6	283±18.9	3
$\underline{\mathtt{S-ethyl}}$ dipropylcarbamothioate				52.6	109	_1_
C ₉ H ₁₉ NOS						
m.w. 189.32 ^a						
cycloate				7.32±25	345±17.4	3
S-ethyl cyclohexylethyl- carbamothicate				66.6	222	1
C ₁₁ H ₂₁ NOS						
m.w. 215.36 ^a						
pebulate				14.5±20.1	719±21.4	3
S-propyl butylethylthio-carbamate				109	363	1
C ₁₀ H ₂₁ Nos						

TABLE 7: Continued

TABLE	7•	Continued

	TA	BLE 7: Contin	ued			
Compound Name						
	Average*	Average*	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soils (No.)
Phosphates						
nellite				0.525±72.5	30.89±25.9	8
$^{\mathrm{C}}{_{8}^{\mathrm{H}}}_{13}^{\mathrm{N}}{_{2}}^{\mathrm{0}}{_{2}}^{\mathrm{P}}$				6.30, 6.02	43.5, 33.2	
m.w. 200.18 a						
crotoxyphos				6.09±15	173±34	3
(E)-1-phenylethyl- 3-[(dimethoxyphosphinyl)oxy]- 2-butenoate						
^C 14 ^H 19 ⁰ 6 ^P						
m.w. 314.28 ^a						
phorate	0.97±2.1	8.76±39	4		3199±25	
<pre>0,0-diethyl S-[(ethylthio) methyl] phosphorodithioate</pre>						
^C 7 ^H 17 ⁰ 2 ^{PS} 3						
m.w. 260.38 ^a						

	Average* l/n±S%	Average* K±S%	Soils (No.)	Average** Kd ±S%	Average*** Koc±S%	Soils (No.)
disulfoton	0.930±2.38	20.1±18.9	10		2132±25	16
0,0-diethyl S-[2- (ethylthio)ethyl] phosphorodithioate						
^С 8 ^Н 19 ⁰ 2 ^{PS} 3						
Compound properties ^b						
m.w. 274.41						
b.p. 108°C						
density 1.144_{4}^{20}						
insol. in water						
methyl parathion	1.04	18.43±20	4		9799± 4 1	
θ , θ -dimethyl θ -(4-nitrophenyl) phosphorothioate						
C ₈ H ₁₀ NO ₅ PS						
Compound properties L						
m.w. 263.23						
m.p. 37-38°C						
density 1.358^{20}						
water sol 50 ppm						
parathion	1.03±1.5	2.19±32	4		10454±38	
θ , θ -diethyl θ -(4-nitrophenyl) phosphorothioate						
^C 10 ^H 14 ^{NO} 5 ^{PS}						
m.w. 291.26						

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T	ABLE	7:	Continued

	TAI	BLE 7: Continu	ıed			
ompound Name						
	Average*	Average* K±S%	Soils	Average** K _d ±S%	Average*** K ± S%	Soils (No.)
ethion	0.945±2.5	28.5±15.9	4		15435±39.3	
heta,0,0',0'-tetraethyl <u>S,S</u> '-methylenebisphosphorodithioate						
Compound properties						
m.w. 384.48						
m.p12 to -13°C						
density 1.220^{20}_{4}						
slightly sol in water						
carbophenothion	0.940±3.7	74.5±13	4		45368±40.6	
S-[[(4-chlorophenyl)thio]methyl] 0,0-diethyl phosphorodithioate						
с ₁₁ н ₁₆ с10 ₂ рѕ ₃						
Compound properties b						
m.w. 342.85						
b.p. 82°C						
density 1.2712 ⁵						
insol. in water						

Compound	Name
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	Average*	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K _{OC} ±S%	Soils (No.)
Amino, Nitrophenyl Sulfones						
$R_2 SO_2 \longrightarrow N(R_1)_2$ NO_2						
nitralin				50, 26	500, 520	2
$(R_1 = CH_3H_7; R_2 = CH_3)$						
4-(methylsulfonyl)-2,6-dinitro- \underline{N} , \underline{N} -dipropylbenzenamine						
$^{\mathrm{C}_{13^{\mathrm{H}}_{19}\mathrm{N}_{3}\mathrm{O}_{6}\mathrm{S}}$						
m.w. 345.38						
5D11830				12.5, 7.25	125, 145	2
$(R_1 = CH_3; R_2 = CH_3)$					143, 143	2
SD12639				23, 9.65	230, 193	2
$(R_1 = C_2H_5; R_2 = CH_3)$						
SD13207				32, 13.5	320, 269	2
$(R_1 + C_2H_5; R_2 = C_2H_5)$						
SD12030				75, 35.1	750, 702	2
$(R_1 = C_3H_7; R_2 = C_2H_5)$						
D12346				117	1170	2
$(R_1 = C_3H_7; R_2 = C_3H_7)$						
D12400				22.2	222	2
$(R_1 = CH_3; R_2 = C_3H_7)$						

	TA	BLE 7: Contin	ued			
Compound Name						
	Average*	Average* K±S%	Soils	Average** K _d ±S%	Average*** K _{OC} ±S%	Soils (No.)
Carboxylic Acids						
dicamba				0.9	3.8	1
3,6-dichloro-2- methoxybenzoic acid						
^С 8 ^н 6 ^{С1} 2 ⁰ 3						
m.w. 221.04^{α}						
chloramben NH ⁺ ₄ salt				0.320±30.4	15.4±37	9
chloramben: 3-amino-2,5-dichlorobenzoic acid				4.1	17.4	1_
^C 7 ^H 5 ^{C1} 2 ^{N0} 2						
m.w. 206.03 ^a						
chloramben methyl ester				3.5±35	507±24	4
MCPA				0.420±37.4		
(4-chloro-2-methylphenoxy)-acetic acid						
_{С9} н ₉ с10 ₃						
m.w. 200.62						

TABLE 7: Continued

### Dictoram 4-amino-3,5,6-trichloro-2-pyridinecarboxylic acid C6H302Cl3N2	14
2-pyridinecarboxylic acid $ {}^{C}_{6}{}^{H}_{3}{}^{0}{}_{2}{}^{C1}{}_{3}{}^{N}{}_{2} $ m.w. 241.46 a	7
m.w. 241.46 ^a ,4-D (2,4-dichlorophenoxy) - acetic acid $C_8^H {}_6^{Cl} {}_2^0 {}_3$ m.w. 221.04 ^a ,4,5-T (2,4,5-trichlorophenoxy) - acetic acid	7
$(2,4-\text{dichlorophenoxy}) - \\ \text{acetic acid} \\ C_8^{\text{H}}{}_6^{\text{Cl}}{}_2^{\text{O}}{}_3 \\ \text{m.w. } 221.04^{\alpha} \\ (2,4,5-\text{trichlorophenoxy}) - \\ \text{acetic acid} \\ (2,4,5-\text{trichlorophenoxy}) - \\ \text{acetic acid} \\$	7
(2,4-dichlorophenoxy) - acetic acid $C_8^{H_6}C^{1}_2O_3$ m.w. 221.04 ^a 1.05±42.3 (2,4,5-trichlorophenoxy) - acetic acid	7
acetic acid $ {^{\text{C}}8}^{\text{H}}6^{\text{Cl}}2^{\text{O}}3 $ m.w. 221.04 $^{\text{A}}$ 1.05±42.3 (2,4,5-trichlorophenoxy) - acetic acid	•
m.w. 221.04 ^a ,4,5-T (2,4,5-trichlorophenoxy) - acetic acid	
,4,5-T (2,4,5-trichlorophenoxy) - acetic acid	
(2,4,5-trichlorophenoxy) - acetic acid	
acetic acid	7
8*5**3*3	
m.w. 255.49 ^a	
ichlebenil 167 4.19±11.4 164±3.57	

freely sol. in water

	TAI	BLE 7: Contin	ued			
Compound Name						
	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** Kd±S%	Average*** K _{oc} ±S%	Soils (No.)
Acids						
linear alkyl				34.2±57.4	1222±10.2	30(K _d ,) 20(K _{oc}
sulfonates				55.3	170	1
pentachlorophenol				8.96±20.38		10
С ₆ нС1 ₅ 0						
Compound properties b						
m.w. 266.35						
m.p. 190-191°C						
b.p. 309-310°C						
density 1.978 ²²						
water sol. 8 mg/100 ml						
methane arsonate				12.4±38.9	770±19.1	4
CH ₅ As0 ₃						
Compound properties b						
m.w. 139.96						
m.p. 161°C						

TABLE 7: Continued

	Average* 1/n±S%	Average* K±S%	Soils (No.)	Average** K _d ±S%	Average*** K _{oc} ±S%	Soils (No.)
Miscellaneous						
silvex	0.639, 0.987	42.1, 34.2	2		2786, 4682	
2-(2,4,5-trichlorophenoxy)-propionic acid	1.05	162	1_		440	
с ₉ н ₇ с1 ₃ 0 ₃						
Compound properties b						
m.w. 269.53						
m.p. 181.6°C						
0.014% sol. in water at 25°C						
chlorthiamid	0.868±1.28	4.72±8.13	5		107.2±6.42	
2,6-dichlorobenzene carbothioamide						
C7H5Cl2NS						
chloroneb	1.3	20	1		1159	
<pre>1,4-dichloro-2,5-dimethoxy- benzene</pre>						
$^{\mathrm{C_{8}H_{8}Cl_{2}O_{2}}}$ m.w. 207.06 a						
paraquat	0.566±19.6	353±97			20152±65	
<pre>l,l'-dimethyl-4,4'- bipyridinium</pre>	0.360±13.3	5501±16.6				
^C 12 ^H 14 ^N 2						
m.w. 186.26 ^a						

NOTES, TABLE 7

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TABLE 8: RELATIONSHIP BETWEEN OCTANOL/WATER PARTITION COEFFICIENT AND R $_{\mathbf{f}}$ VALUES OF PESTICIDES ON A SOIL

		*	**	
Mobility Class	$\frac{R}{f}$	log P	2	
Immobile	0 - 0.09	>3.78	>398	
Low	0.10 - 0.34	3.78-2.39	398-74	
Intermediate	0.35 - 0.64	2.39-1.36	74-29	
Mobile	0.65 - 0.89	1.36-0.08	29-4.5	
Very Mobile	0.90 - 1.00	<0.08	<4.5	

^{*} P = Octanol/water partition coefficient

^{**} Q = Soil organic matter/water partition coefficient

TABLE 9: LEACHING OF PESTICIDES FROM A SOIL

Compound Name Ref.

Adsorbent = Hagerstown silty clay loam \$0.M = 2.41, \$0.C = \$0.M/1.724 = 1.40

Hamaker,

1974

Pesticide	R _f (soil TLC)	Mobility class	K _{oc} *
chloramben	0.96	5	12.8
2,4-D	0.69	4	32
propham	0.51	3	51
bromacil	0.69	4	71
monuron	0.48	3	83
simazine	0.45	3	135
propazine	0.41	3	152
dichlobenil	0.22	2	164
atrazine	0.47	3	172
chlorpropham	0.18	2	245
prometone	0.60	3	300
ametryne	0.44	3	380
diuron	0.24	2	485
prometryne	0.25	2	513
chloroxuron	0.09	1	4,986
paraquat	0.00	1	20,000
DDT	0.00	1	243,000

^{*}K = adsorption coefficient on the basis of organic carbon

TABLE 10: SORPTION DEPENDENCE ON SORBATE PROPERTIES

ound Name	Compo	Ref.		
Compound	Water solubility (mole fraction x 109	K _{oc} (x10 ⁻³)*	K _{OW} (X10 ⁻³) **	Karickhoff, Brown and Scott,
pyrene	12	84	150	1979
methoxychlor	6.3	80	120	
naphthalene	4460	1.3	2.3	
2-methylnaphthalene	3220	8.5	13	
anthracene	7.57	26	35	
9-methylanthracene	24.4	65	117	
phenanthrene	130	23	37	
tetracene	0.037	650	800	
hexachlorobiphenyl	0.048	1200	2200	
benzene	189,000	0.083	0.13	

^{*}K = partition coefficient based on organic carbon oc

^{**}K = octanol/water distribution coefficient

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APPENDIX: FORMULAS OF ORGANIC COUMPOUNDS

COMPOUNDS OTHER THAN PESTICIDES

Acids,	Alipha	atic

acetic acid formic acid

CH₃COOH НСООН

adipic acid fumaric acid HOOCCH2CH2CH2COOH

butyric acid

glutaric acid $\mathrm{CH_{3}CH_{2}CH_{2}COOH}$ ${\tt HOOCCH_2CH_2CH_2COOH}$

caproic acid glyceric acid $\mathrm{CH_3CH_2CH_2CH_2CH_2COOH}$

HOCH₂CHCOOH citraconic

 $\text{CH}_{3\parallel}\text{CCOOH}$ glycolic acid

HCCOOH HOCH₂COOH citric acid

СН₂СООН НОССООН СН₂СООН glyoxylic acid НСОСООН

isobutyric acid

dibromosuccinic CH₃ CHCOOH BrCHCOOH BrCHCOOH

isovaleric acid oxalic $^{\mathrm{CH}}_{3}$ CHCH $_{2}$ COOH нооссоон propionic acid CH₃CH₂COOH itaconic acid CH_{2} =CCOOH CH_{2} COOH pyruvic acid сн зсосоон lactic acid succinic acid ноосси, си, соон tartaric acid levulinic acid CH3COCH2CH2COOH CHOH COOH maleic acid valeric acid **HCCOOH** $\mathrm{CH_{3}CH_{2}CH_{2}CH_{2}COOH}$ malic acid Acids, Aromatic CH₂COOH benzoic acid malonic acid ноосси, соон mesaconic acid CH₃CCOOH phenylacetic acid HOOCCH methylsuccinic acid CH₃OOCCH₂CH₂COCH₃ phenoxyacetic acid monobromosuccinic

HOOCCH₂CHBrCOOH

Alcohol

n-butyl alcohol ${\rm CH_3CH_2CH_2CH_2OH}$

Amines and derivatives

acetanilide

aniline

4-(R₂ sulfonyl)-2,6-dinitro-N.N.-(di-R₁)aniline

$$R_z - S \longrightarrow NO_z$$
 NO_z
 NO_z

crystal violet

$$(\operatorname{CH}_3)_2 \operatorname{N} - \operatorname{N}(\operatorname{CH}_3)_2$$

Amino Acid

tryptophan

Benzene



 $\frac{\text{Carbamate derivatives}}{\text{R}_1\text{NHCOOR}_2}$

Carbonyl compounds acetophenone



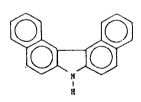
N-Heterocyclics

benzo[f]quinoline



9H carbazole

7-H dibenzocarbazole



pyridine



quinoline

S-Heterocyclics

benzo[b]thiophene

dibenzothiophene



Phenols

phenol



Polynuclear aromatics

benz[a]anthracene

benz[a]pyrene

pyrene

Quinone



alizarin

Sulfonate

sodium napthalenesulfonate

PESTICIDES

Carbamates

carbaryl

cycloate

$$\begin{array}{c} \text{EPTC} \\ \text{CH}_3\text{CH}_2 \\ \text{SCN} \\ \end{array} \begin{array}{c} \text{CH}_2\text{CH}_2\text{CH}_3 \\ \text{CH}_2\text{CH}_2\text{CH}_3 \end{array}$$

pebulate

$$\begin{smallmatrix} \operatorname{CH_3CH_2CH_2SCNCH_2CH_2CH_2CH_3} \\ \operatorname{OCH_2CH_3} \end{smallmatrix}$$

propham

Carboxylic Acids

chloramben

2,4-D

dicamba

dichlobenil

MCPA

picloram

2,4,5-T

Halogenated Hydrocarbons

внс

DDT

dieldrin

ethylene dibromide BrCH₂CH₂Br

heptachlor

lindane

methoxychlor

mirex

n-serve

Organophosphates

carbophenothion

crotoxyphos

disulfoton

$$\begin{array}{c} \operatorname{CH_3CH_2O} \\ \operatorname{CH_3CH_2O} \\ \end{array} \\ \operatorname{PSCH_2CH_2SCH_2CH_3} \\ \end{array}$$

ethion

fenitrothion

methyl parathion

nellite
$$\bigcirc$$
 -OP $<$ NHCH $_3$

paraoxon

parathion

phorate

Amino, Nitrophenyl Sulfones
Nitralin

S-Triazines

ametryne

amitrole

atrazine

cyanazine

ipazine

norazine

prometone

prometryne

propazine

simazine

Ureas & Uracils

bromacil

diuron

fenuron

fluometuron

linuron

methylurea

metrobromuron

monolinuron

monuron

neburon

phenylurea

Miscellaneous

chloroneb

tebuthiuron

chlorthiamid

terbacil

paraquat

urea

silvex

TECHNICAL REPORT DATA (Please read Instructions on the reverse before completing)			
1 REPORT NO.	2.	3. RECIPIENT'S ACCESSION NO.	
EPA-600/3-79-086			
4. TITLE AND SUBTITLE Adsorption of Energy-Related Organic Pollutants: A Literature Review		5. REPORT DATE August 1979 issuing date	
		6. PERFORMING ORGANIZATION CODE	
7. AUTHOR(S)		8. PERFORMING ORGANIZATION REPORT NO.	
K.A. Reinbold, J.J. Hassett, J.C. Means, and			
W.L. Banwart			
9. PERFORMING ORGANIZATION NAME AND ADDRESS		10. PROGRAM ELEMENT NO.	
Institute for Environmental Studies		1BB770	
University of Illinois at Urbana-Champaign		11. CONTRACT/GRANT NO.	
Urbana, Illinois 61801		68-03-2555	
1			
12. SPONSORING AGENCY NAME AND ADDRESS Environmental Research LaboratoryAthens, Ga.		13. TYPE OF REPORT AND PERIOD COVERED	
		Final, 7/77 to 4/78	
Office of Research and Development		14. SPONSORING AGENCY CODE	
U.S. Environmental Protection Agency		EPA/600/01	
Athens, Georgia 30605			
15. SUPPLEMENTARY NOTES			

16. ABSTRACT

This report is a literature review on sorption properties of sediments and energy-related organic pollutants. Adsorption of organic compounds in general is discussed, and analytical methodology in soil thin-layer chromatography and chemical analysis as applicable to measurement of sorption properties is summarized. The literature on the adsorption of energy-related organic pollutants is reviewed. Reported constants for the adsorption of organic compounds on several adsorbents are tabulated, and factors that influence adsorption are discussed.

7. KEY WORDS AND DOCUMENT ANALYSIS			
a. DESCRIPTORS	b.IDENTIFIERS/OPEN ENDED TERMS	c. COSATI Field/Group	
Adsorption Chemical analysis Coal Coal gasification Energy Organic compounds Sediments		68C 68D 99A 99D	
13. DISTRIBUTION STATEMENT RELEASE TO PUBLIC	19. SECURITY CLASS (This Report) UNCLASSIFIED 20. SECURITY CLASS (This page) UNCLASSIFIED	21. NO. OF PAGES 178 22. PRICE	