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AN EQUILIBRIUM MODEL FOR THE PARTITIONING
OF SYNTHETIC ORGANIC COMPOUNDS:
FORMULATION AND CALIBRATION*

Brian J. Eadie

An equilibrium toxic organic distribution model has been
designed. This simple model, needing only information on the con-
taminants, water solubility, and vapor pressure, yields useful
information on the distribution of environmentally persistent orga-
nic contaminants.

The model was calibrated for total DDT in three ecosystems: a
representative coastal regime, Lake Michigan, and a global system.
There are some discrepancies between model output and available
data; while the model calibrated well for the coastal regime and
the Lake Michigan ecosystem, it failed for the global ecosystem.
This is presumably because of the uneven application of DDT and the
large biomass of terrestrial plants, which are relatively uncon-
taminated. Owing to its low vapor pressure, DDT has not and will
not come to a global equilibrium.

The Lake Michigan model was also run for four other organic
contaminants, which span several orders of magnitude in solubility
and vapor pressure. These will be discussed, as will sensitivity
of the model to input parameters.

1. INTRODUCTION

In August 1979, the Great Lakes Environmental Research Laboratory
(GLERL) initiated a research program entitled “The Cycling of Toxic Organic
Substances in the Great Lakes Ecosystem.” This research was partially sup-
ported by the Office of Marine Pollution Assessment (OMPA) under Section 202
of PL 92-532, which states that NOAA should initiate “a comprehensive and
continuing program of research with respect to the possible long-range
effects of pollution, overfishing, and man-induced changes of ocean
ecosystems:”

GLERL's approach consists of a series of models designed to simulate
ecosystems at different scales of time and space and to improve these models
through research on various important processes. This report describes one
of these models.

2. MODEL DESCRIPTION

Conceptually, this is a simple model that assumes that the ecosystem
under consideration is in equilibrium with the toxic organic contaminant.

*GLERL Contribution No. xxx.
It is recognized that the equilibrium assumption is naive, but for persistent organic contaminants in well-mixed environments, the approach does yield worthwhile information.

The environment or ecosystem under consideration is divided into compartments, such as atmosphere, water, sediments, biota, etc. At equilibrium, the toxic organic contaminants' "escaping tendency" or fugacity from each compartment is equal. Mackay (1979) clarifies the concept through the following analogy:

Fugacity is to mass diffusion as temperature is to heat diffusion. Mass (or heat) always diffuses from high to low fugacity (or temperature). Diffusion directions are not obvious from concentrations such as g mol m$^{-3}$ (or cal m$^{-3}$), but they are obvious when expressed in atm (or °C). The insights into heat diffusion provided by temperature can be obtained for mass by using fugacity.

Fugacity has units of pressure and, at the very low concentrations encountered with trace organic contaminants, it is linearly proportional to concentration. In Mackay's (1979) terms,

$$C = Zf, \quad (1)$$

where $C$ = concentration (g mol m$^{-3}$), $Z$ = fugacity capacity (g mol m$^{-3}$ atm$^{-1}$), and $f$ = fugacity (atm).

The fugacity capacity ($Z$) for each ecosystem compartment must be estimated. In this work each was calculated in the following manner:

Vapor phase (atmosphere)--for an ideal gas, fugacity is exactly equal to partial pressure ($P$). At the concentrations with which this model deals, the vapor phase will be close to ideal. Thus, from

$$PV = nRT$$

$$fV = nRT,$$

from equation (1)

$$\frac{CV}{Z} = nRT$$

and $CV$ = n (number of g moles)

$$Z = CV (nRT)^{-1} = RT^{-1} (\approx 40-45)$$

where $R = 82 \times 10^{-6}$ and

$$T = \, ^\circ R.$$
The fugacity capacity in the vapor phase is independent of compound characteristics.

Liquid phase (water)—the ratio of contaminant vapor pressure (P) to solubility (S) is the Henry's Law constant (H). From

\[ H = \frac{P S}{1} = C_{\text{vapor}} (C_{\text{dissolved}})^{-1} = fC_d^{-1}, \]

and combining with equation (1), it follows that \( Z = 1 \text{ H}^{-1} \).

Sorbed phases (particulates, sediments, etc.)—if the equilibrium partition coefficient \( (K_p) \) is defined as the ratio of sorbed concentration (\( \mu g \text{ g}^{-1} \)) to water concentration (\( g \text{ m}^{-3} \)), then

\[ C_{\text{sorbed}} = K_p C_{\text{dissolved}} \]

since

\[ f = HC_d = \frac{HC_s}{K_p} ; \]

then substituting equation (1)

\[ Z = K_p H^{-1}. \]

The \( K_p \) is calculated from the contaminant's octanol-water partition coefficient \( (K_{ow}) \) as discussed below.

Biotic phases (plankton, fish, benthos, etc.)—there have been several attempts at correlating concentration in fish based on the contaminant's \( K_{ow} \) (Vieth et al., 1979; Chou et al., 1977; Thomann, 1979). Mackay (1979) assumes that the biotic phases act as passive substrates for sorption. Although there is considerable evidence for bioaccumulation and biomagnification up the food chain, this equilibrium approach (which is based on mass, not surface area) appears to be reasonable.

Thus

\[ Z = K_p H^{-1}. \]

2.1 Estimation of the Equilibrium Partition Coefficient

\( K_p \), the equilibrium partition coefficient is defined as follows:
Contaminant concentration in the sorbed phase (ppm)

\[ K_p = \frac{\text{Contaminant concentration in the dissolved phase (ppm)}}{\text{Contaminant concentration in the sorbed phase (ppm)}} \] (2)

Karickhoff et al. (1979) and others have shown that \( K_p \) is primarily a function of the octanol-water partition coefficient of the contaminant (\( K_{ow} \)) and the organic carbon content of the substrate. For neutral, hydrophobic organic contaminants, this result agrees with intuition; the contaminant preferentially dissolves (partitions) more favorably into those substrates that are highest in organic carbon content. \( K_p \) can be described as

\[ K_p = K_{oc} \times \frac{\text{percent substrate organic carbon}}{100} \]

where \( K_{oc} \) is the partition coefficient normalized to organic carbon content (Means et al., 1979).

Life is never quite so simple; there are several other variables that appear to affect \( K_p \); among them are substrate surface area (Karickhoff et al., 1979), concentration of substrate (O'Connor and Connolly, 1980), and the nature of the substrate organic matter. These complications appear to be second order, and the approximation based on \( K_{oc} \) (described below) will be used in this preliminary model.

\( K_{oc} \) has been shown to be well correlated to the octanol-water partition coefficient (\( K_{ow} \)) of the contaminant. Figure 1 illustrates two of the most recent of these correlations (which therefore employ the most data). The Smith and Bomberger (1979)/Chiou et al. (1977) line was derived from their individual correlations of \( K_{oc} \) with solubility and \( K_{ow} \) with solubility, respectively. These will be described in more detail below. The dashed line was used in this model. (\( \log K_{oc} = 1.05; \log K_{ow} = 0.300 \).)

\( K_{ow} \) is a relatively simple laboratory measurement, and values of \( K_{ow} \) are available in the literature (Leo et al., 1971). \( K_{ow} \) can also be approximated directly from the molecular structure of the compound by the "Hansch method," described in Hansch (1980) and supported by numerous references. That article, along with Tulip and Hutzinger (1978), discusses the limits of this approach, generically termed structure-activity relationships, and the extension of the technique into the estimation of toxicity. Although not included in the current version of our model, this approach is certainly promising and will be pursued by our laboratory in future modeling efforts.

A third technique for estimating \( K_{ow} \) is the relationship developed between \( K_{ow} \) and solubility. Chiou et al. (1977) derived the relationship

\[ \log K_{ow} = 5.00 - 0.670 \log S, \]
Figure 1.—The relationship between $K_{OC}$ and the octanol-water partition coefficient, $K_{ow}$. For Karickhoff et al., $\log K_{OC} = \log K_{ow} - 0.21 (r^2 = 1.00, n = 10)$. The Smith and Bomberger/Chou et al. line was derived from their correlations of $K_{OC}$ and $K_{ow}$ versus solubility, respectively ($n > 33$). Sample compounds are listed on the abscissa and illustrate the properties of a range of contaminants. The dashed line was used in the model.
where

\[ S = \text{aqueous solubility in micromols per liter} \]

\[ r^2 = 0.970, N = 33, \log K_{ow} \text{ range } = 1.26-6.72. \]

This relation is shown in figure 2, along with a relation generated by Smith and Bomberger (1979), which relates \( K_{oc} \) directly with solubility. Although statistics are not provided, the scatter of data indicates a poorer correlation than was found by Chiou et al. (1977), which is to be expected since the nature of the substrate is now part of the variability. Recall that the subject models parameterization of \( K_{oc} \) is

![Figure 2](image-url)
In our model, $K_{oc}$ is calculated from Chiou's equation. Our calculations can also be presented in terms of $\log K_{oc} = f(S)$ as follows:

$$\log K_{oc} = 1.05 \log K_{ow} - 0.500.$$ 

$$\log K_{ow} = 5.00 - 0.670 \log S$$

and

$$\log K_{oc} = 1.05 \log K_{ow} - 0.500.$$ 

Substituting $1\% K_{oc} = 4.75 - 0.704 \log S$, plotted as the dashed line in figure 2, $K_p$ is then calculated from $K_{oc}$ as

$$K_p = K_{oc} \times \frac{\text{percent substrate organic carbon}}{100}$$

2.2 Biotic Phases

As mentioned earlier, $Z = K_{H-1}$ is used for all sorbed phases including biotic compartments. Figure 3 illustrates the problem with this approach. The above formulation, using 50 percent organic carbon (dry weight) for fish, yields a bioconcentration function that differs from the results of four other syntheses of real data. The bioconcentration factor (BCF) correlations presented in figure 3 were considered, and an information item was included in the model output indicating the concentration of contaminant in large fish based on

$$\log BCF = \log K_p = 0.80 \log K_{ow} - 0.50,$$

which is approximately midway between the values obtained by Vieth et al. and Chiou et al.

2.3 Benthic Organisms

Difficulty was encountered when the passive sorption concept was applied to benthic organisms. The sparse environmental data indicate that contaminant concentrations in benthic organisms (dry weight) are approximately twice those of dry sediments. Since the organic carbon concentration of a benthic organism is 10-40 times as high as that of the sediments, and the model is based on $K_p$, which is linearly dependent on organic carbon content, the benthos values predicted by the model are too high by an order of magnitude. At this level of model development, the benthic organisms have been left out, although they are considered a prime target for further research since benthos are exposed to long-term high concentrations of contaminants in the surficial sediments.
2.4 Calculating the Distribution of Contaminants Within the Ecosystem

Once the $Z$ values for each environmental compartment have been calculated, the concentration in each compartment is calculated in the following manner:

At equilibrium $f_1 = f_2 = f_i$, $i = \text{compartment}$, and the total mass of contaminant ($M$) equals

$$M = \sum f_i Z_i V_i,$$

where

$V_i = \text{volume of the } i\text{th compartment}$

and from $C = fZ$ [equation (1)],

$$M = \sum f Z V_i,$$
\[ M = f_1 E V_i Z_i, \text{ and} \]

\[ f_1 = M (\Sigma V_i Z_i)^{-1}. \]

The terms on the right are either input \((M, V_i)\) or calculated from inputs \([Z_i = f(\text{solubility, vapor pressure, K octanol-water})]\). The mass in each compartment is

\[ M_i = f_1 Z_i V_i \]

and the concentration is

\[ C_i = f_1 Z_i. \]

The total mass of a contaminant in the ecosystem \((M)\) is rarely known; therefore, for an initial run, an arbitrary value may be used (e.g., 1.0). Model output will then give relative concentrations in each compartment, and if data exist for any compartment, the value of \(M\) can be scaled up by

\[ \frac{C_i(\text{data})}{C_i(\text{model output})}, \]

which will yield approximate concentrations in all compartments.

2.5 Description of an Ecosystem

As developed in the previous section, the volume \((V_i)\) of each environmental compartment needs to be input. This is not too difficult for the large, physically distinct compartments of atmosphere, water, and sediments (except for estimating the mixing depths of the atmosphere and sediments), but how does one estimate the dry weight volume of any biotic compartments? It turns out that, although concentrations of a contaminant are high in biota, the mass of the contaminant in these phases is bound to be small for large lakes and marine systems, and the model is insensitive to order-of-magnitude errors in these compartment volumes.

Three model ecosystems were run; they are described below. They are (ECOL), a 1-m² horizontal area, 100-m-deep water body; (LKH), an approximation of Lake Michigan; and (ECOALL), a representation of a global ecosystem.

All volumes are calculated on a dry-weight basis, and comparative data should be carefully examined and corrected to a dry-weight basis.
2.6 Compartmentalization of ECOL

Atmosphere (10-km mixed depth)

\[ V = 1 \text{ m}^2 \times 10,000 \text{ m} = 10^4 \text{ m}^3. \]

Water

\[ V = 1 \text{ m}^2 \times 100 \text{ m} = 10^2 \text{ m}^3. \]

Sediment (2-percent organic carbon, 5-cm mixed depth)

\[ V = 1 \text{ m}^2 \times 5 \text{ cm} \times 0.5 \text{ (porosity)} = 2.5 \times 10^{-2} \text{ m}^3. \]

Suspended matter (1.5 mg l\(^{-1}\), \( \rho = 1.5 \text{ g m}^{-3}\), 15-percent organic carbon)

\[ V = \frac{1.5 \text{ g}}{\text{m}^3} \times 10^2 \text{ m}^3 \times \frac{1 \text{ cm}^3}{1.5 \text{ g}} \times \frac{\text{m}^3}{10^6 \text{ cm}^3} = 1 \times 10^{-4} \text{ m}^3. \]

Biota (1 \( \mu \text{g l}^{-1} \) chlorophyll \( a = 50 \mu \text{g C l}^{-1} \) total plankton, \( \rho = 1.0 \text{ g/cm}^3 \)).

\[ V = 50 \frac{\text{mg C}}{\text{m}^3} \times 10^2 \text{ m}^3 \times \frac{2.5 \text{ mg org. matter}}{\text{mg org. carbon}} \times \frac{1 \text{ cm}^3}{10^{-3} \text{ mg}} \times \frac{\text{m}^3}{10^6 \text{ cm}^3} = 1.25 \times 10^{-5} \text{ m}^3. \]

The above values are representative of Lake Michigan or a coastal marine ecosystem.
2.7 Compartmentalization of \( LKM_i \)

Lake Michigan

Surface area \((5.8 \times 10^{10} \text{ m}^2)\).

Average depth \((86 \text{ m})\).

Atmosphere \((10\text{-km mixed depth})\)

\[
V = 5.8 \times 10^{14} \text{ m}^3
\]

Water

\[
V = 5.8 \times 10^{10} \times 86 = 5 \times 10^{12} \text{ m}^3
\]

Suspended matter \((1.5 \text{ mg l}^{-1}; \rho = 1.5 \text{ g cm}^{-3}, 10\text{-percent organic carbon})\)

\[
V = 1.5 \frac{\text{g}}{\text{m}^3} \times 5 \times 10^{12} \text{ m}^3 \times \frac{1 \text{ cm}^3}{1.5 \text{ g}} \times \frac{m^3}{10^6 \text{ cm}^3} = 5 \times 10^6 \text{ m}^3
\]

Phytoplankton cell density \(= 1500 \text{ cells ml}^{-1}\) (Tarapchak and Stoermer, 1976)

\(= 100 \text{ pg cell}^{-1}\) (Stoermer and Ladewski, 1978)

\[
V = 1500 \frac{\text{cells}}{\text{cm}^3} \times 5 \times 10^{18} \text{ cm}^3 \times \frac{100 \times 10^{12} \text{ g}}{\text{cell}} \times 1 \text{ cm}^3 \times \frac{m^3}{10^6 \text{ cm}^3} = 7.5 \times 10^5 \text{ m}^3.
\]

Zooplankton \((30 \text{ mg m}^{-3}, \rho = 1)\)

\[
V = \frac{30 \text{ mg}}{\text{m}^3} \times 5 \times 10^{12} \text{ m}^3 \times \frac{1 \text{ cm}^3}{\text{g}} \times \frac{\text{g}}{1000 \text{ mg}} \times \frac{m^3}{10^6 \text{ cm}^3} = 1.5 \times 10^5 \text{ m}^3.
\]
Forage fish ($10^5$ MT, $\rho = 1$ g cm$^{-3}$, 40-percent organic carbon)

\[
V = 10^5 \text{MT} \times \frac{10^6 \text{g}}{\text{MT}} \times \frac{1 \text{ cm}^3}{\text{g}} \times \frac{1 \text{ g day}}{5 \text{ g wet}} \times \frac{m^3}{10^6 \text{ cm}^3} = 2 \times 10^4 \text{ m}^3. 
\]

Top predators; salmonids ($1.6 \times 10^3$ MT, $\rho = 1$ g cm$^{-3}$)

\[
V = 1.6 \times 10^3 \text{MT} \times \frac{10^6 \text{g}}{\text{MT}} \times \frac{1 \text{ cm}^3}{\text{g}} \times \frac{1 \text{ g dry}}{5 \text{ g wet}} \times \frac{m^3}{10^6 \text{ cm}^3} \times 3.2 \times 10^2 \text{ m}^3. 
\]

Sediment (5-cm mixed depth, 2-percent organic carbon)

\[
V = 5.8 \times 10^{10} \text{ m}^2 \times 5 \text{ cm} \times \frac{m}{10^2 \text{ cm}} \times \frac{1 \text{ g dry}}{2 \text{ g wet}} \approx 1.5 \times 10^9 \text{ m}^3. 
\]

2.8 Compartmentalization of ECOALL

Earth (McLellan, 1968)

Surface area (land) ($1.5 \times 10^{14}$ m$^2$).

Surface area (water) ($3.6 \times 10^{14}$ m$^2$).

Augmented depth ($3.8 \times 10^3$ m).

Atmosphere (10-km depth)

\[
V = 5.1 \times 10^{14} \text{ m}^2 \times 10^4 \text{ m} = 5.1 \times 10^{18} \text{ m}^3. 
\]

Water

\[
V = 3.6 \times 10^{14} \text{ m}^2 \times 3.8 \times 10^3 \text{ m} \approx 1.4 \times 10^{18} \text{ m}^3. 
\]

Suspended matter (2 $\mu$g l$^{-1}$, 10-percent organic carbon)

\[
V = \frac{2 \text{ mg}}{\text{m}^3} \times \frac{cm^3}{g} \times \frac{m^3 g}{10^3 \text{mg}} \times \frac{1.4 \times 10^{18} \text{ m}^3}{10^6 \text{ cm}^3} = 2.8 \times 10^9 \text{ m}^3. 
\]
Plant biomass

Aquatic ($1.8 \times 10^{15}$ g C; Whittaker and Likens, 1973).

$$V = 1.8 \times 10^{15} \text{ g C} \times \frac{2.5 \text{ g biomass}}{8 \text{ g}} \times \frac{1 \text{ cm}^3}{8 \text{ g}} \times \frac{m^3}{10^6 \text{ cm}^3} = 4.5 \times 10^9 \text{ m}^3.$$

Terrestrial ($1.9 \times 10^{12}$ MT; Woodwell et al., 1971)

$$V = 1.9 \times 10^{18} \text{ g} \times \frac{1 \text{ cm}^3}{8 \text{ g}} \times \frac{m^3}{10^6 \text{ cm}^3} = 1.9 \times 10^{12} \text{ m}^3.$$

Animal biomass

Aquatic ($0.45 \times 10^{15}$ g C; Whittaker and Likens, 1973).

$$V = 0.45 \times 10^{15} \text{ g C} \times \frac{2.5 \text{ g biomass}}{8 \text{ g}} \times \frac{1 \text{ cm}^3}{8 \text{ g}} \times \frac{m^3}{10^6 \text{ cm}^3} = 1.1 \times 10^9 \text{ m}^3.$$

Terrestrial ($0.5 \times 10^9$ MT; Woodwell et al., 1971).

$$V = 0.5 \times 10^{15} \text{ g} \times \frac{1 \text{ cm}^3}{8 \text{ g}} \times \frac{m^3}{10^6 \text{ cm}^3} = 5 \times 10^8 \text{ m}^3.$$

Sediments

Aquatic (1-cm mixed depth, 50-percent dry, 1-percent organic carbon).

$$V = 3.6 \times 10^{14} \text{ m}^2 \times 0.01 \text{ m} \times 0.5 = 1.8 \times 10^{12} \text{ m}^3.$$

Terrestrial (1-cm mixed depth, 2-percent organic carbon).

$$V = 1.5 \times 10^{14} \text{ m}^2 \times 0.01 \text{ m} = 1.5 \times 10^{12} \text{ m}^3.$$

3. MODEL CALIBRATION

The minimum input that the model requires is a description of the ecosystem in terms of volume and the fugacity capacity values ($Z$), which are
calculated from solubility, vapor pressure, and temperature. For the synthetic organics that are of initial interest, the solubilities are so low (microgram per liter range) that measured values often range over an order of magnitude. The same range exists for vapor pressures. This allows for a small amount of "tuning" if we have reliable environmental data for one or more of our ecosystem compartments.

Another tuning factor is the percent organic carbon of sorption substrates since partitioning is linearly related to this value. Reliable numbers are usually available for percent organic carbon of sediments or suspended matter, but most ecosystems are heterogeneous and an average value needs to be estimated.

The final input variable is the total amount of contaminant in the ecosystem, a number rarely available. As mentioned earlier, this can be scaled by first running the model with an input of 1.0 and then multiplying this input (for the second run) by the ratio of the contaminant concentration (data) in any compartment to the model output for that compartment.

These calibration knobs are straightforward and are constrained within generally acceptable limits. Sensitivity to these calibration terms will be discussed in the following section, which describes model output for several representative compounds.

4. MODEL ANALYSIS

The most extensive model analysis has been done with total DDT (DDT plus decomposition products, mostly DDE and DDD), for which a reasonable amount of environmental data is available. The first run was with the global system, ECOAll. Accepted values for solubility (1.2 x 10^-3 ppm) and vapor pressure (1.6 x 10^-4 mm Hg) were input, along with the estimated value for total world DDT production of ~ 2.5 x 10^12 g (Woodwell et al., 1971). Model output concentrations, shown in table 1, were close to those summarized in Woodwell, with the notable exception of land plant data. Results from this run indicate that DDT is stored primarily in terrestrial plants (60 percent) and water (36 percent). The mass of DDT stored in land plants is in excess of that estimated by Woodwell et al. (1971) by a factor of 1000. The sediment (land) values output from the model (4.0 g m^-2) are in the middle of their agricultural soils range data (0.15-11), but are much higher than for unsprayed forests (0.0004-0.004), which accounts for most of the land plant biomass. The global ecosystem has not been uniformly exposed to DDT, and the equilibrium assumption leads to an inaccurate distribution pattern. Presumably, this is because of DDT usage patterns and the compound's low vapor pressure. The model's atmospheric concentration (0.17 x 10^-6 g m^-3) is within the data range of 0.1-10 x 10^-6 g m^-3, but represents less than 0.1 percent of atmospheric saturation (~ 3 x 10^-3 g m^-3).

The second model ecosystem considered was the simple 1-m^2 coastal representation. A single run for DDT was made simply by scaling the DDT mass in this system to that in the global system by the volume ratio of the atmospheres (0.10 x 10^9 / 0.51^-1 x 10^-19 = 1.96 x 10^15), yielding 1.4 x 10^-5 g
TABLE 1.—Comparison of model output and summarized data for the global system. DDT concentrations are in ppm.

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<th>Compartment</th>
<th>Data\textsuperscript{1}</th>
<th>Model</th>
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<tbody>
<tr>
<td>Atmosphere</td>
<td>$1-100 \times 10^{-9}$</td>
<td>$1.7 \times 10^{-9}$</td>
</tr>
<tr>
<td>Water</td>
<td>$10^{-4}-10^{-1}$</td>
<td>$0.81$</td>
</tr>
<tr>
<td>Land Plants</td>
<td>$0.1-1$</td>
<td>$0.81$</td>
</tr>
<tr>
<td>Aquatic Plants</td>
<td>$1$</td>
<td>$1$</td>
</tr>
<tr>
<td>Land Animals</td>
<td>$1$</td>
<td>$1$</td>
</tr>
<tr>
<td>Aquatic Animals</td>
<td>$0.1-10$</td>
<td>$4.0$</td>
</tr>
<tr>
<td>Land Sediment</td>
<td>$2 \times 10^{-2}$</td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{1}Data synthesized in Woodwell et al., 1971 order of magnitude.
\textsuperscript{2}Model output DDT Mass = $2.5 \times 10^{12}$ g; appendix 1.
\textsuperscript{3}Units of g m$^{-2}$; model assumes 1-cm thickness.

in the system. In this case, DDT more realistically resides in the sediments (91 percent), for the same approximate concentration in the biota (1-3 ppm). This single output is listed below; other model outputs discussed in this report are listed in appendix 1.

The final ecosystem calibrations and sensitivity testing were performed on the Lake Michigan ecosystem. Data were summarized for Lakes Michigan and Ontario from the International Joint Commission (International Joint Commission, 1977, 1978, 1980) and Sonzogni et al. (1981), and ranges are compared to 10 model outputs in figure 4. The variations in the 10 outputs are described in table 2 below; for clarity only those variables that were altered are listed.

In the initial runs, the concentrations in the sediments were too high. The discrepancies were eventually reduced by lowering the percent organic carbon in the sediments from 2 percent to 1 percent. The solubility was increased by a factor of 10 and kept at that level for the last 5 runs because the octanol-water partition coefficient (calculated from the solubility) for this solubility was closer to measured $K_{ow}$ (5.5-6.2).

These 10 runs also show that a two-order-of-magnitude change in solubility (runs 1-4) or vapor pressure (runs 8-10) has very little impact on the concentrations of contaminant in sediments and biota. The insensitivity to these 2 input variables is important because reported values for contaminants often range over a factor of 10 or more.
Figure 4.--Model output versus DDT data for Lake Michigan. See table 2 for input variables, appendix I for total outputs. Dot with error bars connecting to dashed lines represent data mean + standard deviation. Individual dots represent model output for each run.
### TABLE 2: Input variables for the 10 calibration and sensitivity runs. Only changes are listed; subsequent runs used the last listed value. Outputs are in Appendix 1.

<table>
<thead>
<tr>
<th>Model Run</th>
<th>Solubility</th>
<th>Vapor present</th>
<th>DDT Mass (g)</th>
<th>Sed. org. C (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.2 x 10^{-3}</td>
<td>1.6 x 10^{-7}</td>
<td>1 x 10^6</td>
<td>2.0</td>
</tr>
<tr>
<td>2</td>
<td>6.0 x 10^{-3}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>1.2 x 10^{-2}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>1.2 x 10^{-1}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>1.2 x 10^{-3}</td>
<td></td>
<td></td>
<td>1.5</td>
</tr>
<tr>
<td>6</td>
<td>1.2 x 10^{-2}</td>
<td></td>
<td></td>
<td>1.0</td>
</tr>
<tr>
<td>7</td>
<td></td>
<td>5 x 10^5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td></td>
<td>1.6 x 10^{-8}</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td></td>
<td>1.6 x 10^{-6}</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The final model exercise was to take the calibrated Lake Michigan ecosystem (the reduction of sediment organic carbon from 2 percent to 1 percent) and to examine other organic contaminants. Unfortunately, data for comparison are sparse.

This exercise yields two important pieces of information (shown in Table 3). First, the model appears to work well for contaminants other than DDT (runs 11 and 12), and second, the model yields useful information on the relative distribution of contaminants even when environmental data are not available (runs 13 and 14).

The mirex run (#12) was a particularly interesting exercise. Mirex is not known to be a problem in Lake Michigan, but is known to be a contaminant in Lake Ontario. Not having a Lake Ontario ecosystem (which is similar in relative volume distribution to Lake Michigan), I scaled the estimated mass of mirex in Ontario (668 kg; Holdrinen et al., 1978) to Michigan as follows:

\[ g_{\text{mirex}} = 6.88 \times 10^5 \times A \times S \times \frac{100}{p} \times c, \]
TABLE 3.—Model outputs for selected contaminants.

<table>
<thead>
<tr>
<th>Model run</th>
<th>Compound</th>
<th>Fish Concentrations (PPM)</th>
<th>Water</th>
</tr>
</thead>
<tbody>
<tr>
<td>11*</td>
<td>Dieldrin</td>
<td>0.12 0.095 2.4 x 10^{-3} 3.1 x 10^{-6}</td>
<td></td>
</tr>
<tr>
<td>Data</td>
<td></td>
<td>0.2±0.2 0.14±0.1 1.8±1.2 x 10^{-3} 3.7±2.9 x 10^{-6}</td>
<td></td>
</tr>
<tr>
<td>12†</td>
<td>Mirex</td>
<td>0.35 0.28 7 x 10^{-3} 2.3 x 18</td>
<td></td>
</tr>
<tr>
<td>Data</td>
<td></td>
<td>1.0±1.0 -- 7.5±8.3 x 10^{-3} --</td>
<td></td>
</tr>
<tr>
<td>13†</td>
<td>Naphthalene</td>
<td>552. 447. 11. 1.0</td>
<td></td>
</tr>
<tr>
<td>14†</td>
<td>Benzene</td>
<td>24. 19. 0.5 1.0</td>
<td></td>
</tr>
</tbody>
</table>

*Contaminant mass scaled to known approximate concentrations.
†Compartment concentrations are reported as a ratio to the water concentration.

where

\[ A = \text{area ratio of Lake Michigan to Ontario} = 3.03, \]
\[ S = 5/3; \text{mixed sediment thickness in model} \]
\[ P = \text{percent of mirex in sediment (62 percent from model),} \]
\[ C = 3; \text{mirex only in 1/3 of Lake Ontario sediment, model assumes equilibrium,} \]

\[ g \text{ mirex} = 1.7 \times 10^7 \text{ g.} \]

This mass gave concentrations in fish and sediments in agreement with reported data.
5. RUNNING THE MODEL

The information necessary to run this model consists of the contaminants' solubility in water, its vapor pressure, and a description of the ecosystem. The program is interactive and will prompt the user to supply information. We have created three procedure files for the three ecosystems described in this report. They are shown below. Ecosystem files are LKI, ECOALL, and ECOL (also listed below). To generate a new procedure file, simply create a new file, then change the LKIIs to the new file name in the procedure file (e.g., RUNMI).

The model is initiated by typing in (e.g.) CALL, RUNMI.
COPY=RUN1.  
SET FUGMOD1.  
GET=SUB1.  
GET TAPE=ECO1.  
EDIT=FUGMOD1;I=SUB1.  
REWRIND;LGO.  
REWRIND TAPE=ECO1.  
LGO.  
REPLACE=TAPE=ECO1.  
REWRITE=TAPE=ECO1.  
COPY=TAPE=OUTPUT.  
ED1 ENCOUNTERED.

COPY=RUN1.  
SET FUGMOD1.  
GET=SUB1.  
GET TAPE=ECO1.  
EDIT=FUGMOD1;I=SUB1.  
REWRIND;LGO.  
REWRIND TAPE=ECO1.  
LGO.  
REPLACE=TAPE=ECO1.  
REWRITE=TAPE=ECO1.  
COPY=TAPE=OUTPUT.  
ED1 ENCOUNTERED.

COPY=RUN1.  
SET FUGMOD1.  
GET=SUB1.  
GET TAPE=LKMI.  
EDIT=FUGMOD1;I=SUB1.  
REWRIND;LGO.  
REWRIND TAPE=LKMI.  
LGO.  
REPLACE=TAPE=LKMI.  
REWRITE=TAPE=LKMI.  
COPY=TAPE=OUTPUT.  
ED1 ENCOUNTERED.

COPY=RUN1.  
SET FUGMOD1.  
GET=SUB1.  
GET TAPE=ECD2.  
EDIT=FUGMOD1;I=SUB1.  
REWRIND;LGO.  
REWRIND TAPE=ECD2.  
LGO.  
REPLACE=TAPE=ECD2.  
REWRITE=TAPE=ECD2.  
COPY=TAPE=OUTPUT.  
ED1 ENCOUNTERED.

COPY=ECD2.  
GET=LKMI.  
ATMOSPHERE(10 K) .430E+15-.100E+01  
WATER <100 M> .100E+03-.200E+01  
DTRITUS 1.50PPM:5X .100E-03:.500E+01  
SEDIMENT 1PPM CHL. .125E-04:.400E+02  
ED1 ENCOUNTERED.

COPY=ECD2.  
GET=LKMI.  
ATMOSPHERE(10 K) .540E+15-.100E+01  
WATER <100 M> .490E+13-.200E+01  
DTRITUS 1.50PPM:5X .500E+07-.100E+02  
SEDIMENT 1PPM CHL. .150E+00:.500E+02  
ED1 ENCOUNTERED.

COPY=ECD2.  
GET=LKMI.  
ATMOSPHERE(10 K) .510E+19-.100E+01  
WATER <3800 M> .140E+19-.200E+01  
DTRITUS 2PPM:100C .280E+10-.100E+02  
PLANT HUQTIC .450E+10-.400E+02  
PLANTS LAND .900E+13-.400E+02  
ANIMALS HUQTIC .110E+10-.500E+02  
ANIMALS LAND .500E+09-.500E+02  
SEDIMENT HUQTIC .180E+13-.100E+01  
SEDIMENT LAND .150E+13-.200E+01  
ED1 ENCOUNTERED.
If there is an ecosystem file, the model is run as follows:

CALL,RUNMCI
XEDIT,3,1.00
END

If there is an ecosystem file, the model is run as follows:

MIREX

ENTER THE COMPOUNDMOLECULARWEIGHT

546

ENTER TOTAL MASS OF COMPOUND IN SYSTEM (G) OR 1.0

2.1E5

IS THERE A FILE DESCRIBING THE ECOSYSTEM? I YES=1, NO=2

ENTER ECOSYSTEM FILE NUMBER, NUMBER OF COMPARTMENTS

7.8

ENTER A TITLE FOR THIS RUN (80 CHARACTERS)

TEST

ENTER THE SYSTEM TEMPERATURE (C)

15

ENTER THE COMPOUND WATER SOLUBILITY (G/M3)

7.0E-04

ENTER THE COMPOUNDS VAPOUR PRESSURE (MM HG)

1E+0

MIREX

MOLECULAR WEIGHT = 546

SYSTEM TEMPERATURE = 15.0 C

WATER SOLUBILITY = 7.00E-04 (G/M3)

VAPOUR PRESSURE = 1.00E-05 (MM HG)

LOG(k10) (OCTANOL-WATER COEFFICIENT) = 7.608

TOTAL MASS OF COMPOUND IN SYSTEM = 0.385E+04 MOL

6.2E+10 = L x G x M x 3

EQUILIBRIUM FUGACITY = 0.530E-13 (ATM)

HENRY'S CONSTANT = 0.103E-01 (ATM M3/MOL)

:: COMPARTMENT VOL(M3) CONC(G/M3) MASS(MOL)

====================================================================

1 ATMOSPHERE(10 KM) .55E+15 .42E+02 .34E+02 .12E+05 .13E+04
2 WATER (86 M) .49E+13 .97E+08 .69E+00 .38E-08 .25E+02
3 DETRITUS 1.5PPM 0C .50E+07 .30E+06 .21E+01 .87E+02 .76E+02
4 SEDIMENT 3CM .15E+10 .30E+08 .62E+02 .37E-03 .24E+04
5 PHYTO 15000CEL/ML .75E+06 .12E+10 .12E+01 .35E+01 .49E+02
6 ZOOPLANK (0.5M) .5E+06 .12E+10 .25E+00 .35E-01 .95E+01
7 FORG FISH (165 MT) .20E+05 .15E+10 .41E-01 .43E-01 .16E+01
8 SALMONID (1.6E3 MT) .32E+03 .15E+10 .66E-03 .43E-01 .25E-01
9 TOP AQUATIC PREDATOR .54E-02
If there is no ecosystem file, the model is run as follows:

```plaintext
CALL RUNME
XEDIT3,1.00
END
FUGMODI$ A LOCAL FILE
ENTER THE COMPOUND NAME (20 CHARACTERS)
? TOTAL DDT
ENTER THE COMPOUND'S MOLEULARWEIGHT?
? 356
ENTER TOTAL MASS OF COMPOUND IN SYSTEM (G) ? R 1.0
? 1
IS THERE A FILE DESCRIBING THE ECOSYSTEM? YES=1 NO=2
? 2

Z'S ARE CALCULATED AS FOLLOWS
VAPOR PHASE: P = 1/P
LIQUID PHASE: P = 1/H
SORBED: R BIOTIC PHASE: Z = KP/H
KP = KC * ORGANIC CARBON / 100
KDC = 1.05 * MLKD = 0.50

FOR Z CLASSIFICATION, ENTER
- 1. FOR VAPOR PHASE (ATM)
- 2. FOR LIQUID PHASE
- SUBSTRATE ORGANIC CARBON FOR SORBED OR BIOTIC PHASE

ENTER A TITLE FOR THIS RUN (80 CHARACTERS)
? 155
ENTER THE SYSTEM TEMPERATURE (C) ? 155
ENTER THE COMPOUND'S WATER SOLUBILITY (G/M3)
? 1.2E-2
ENTER THE COMPOUNDS VAPOR PRESSURE (MM Hg)
? 1.6E-06
ENTER THE NUMBER OF ECOSYSTEM COMPARTMENTS
NAME (IN QUOTES) , INFO (IN QUOTES) ARE 10 CHARACTERS MAX

ENTER COMPARTMENT 1 NAME , INFO , VOL (M3) , Z CLASS (-1 , -2 , , KDC)
? "ATMOSPHERE" "(7.555 KM) " , 1E6 + 1
ENTER COMPARTMENT 2 NAME , INFO , VOL (M3) , Z CLASS (-1 , -2 , , KDC)
? "WATER" " (170 M) " 6.3E3 + 2
ENTER COMPARTMENT 3 NAME , INFO , VOL (M3) , Z CLASS (-1 , -2 , , KDC)
? "SEDIMENTS" "(10 Kg/C)" , 4E1 , 4

EXAMPLE

TOTAL DDT

MOLECULAR WEIGHT = 356
SYSTEM TEMPERATURE = 15.0 C
WATER SOLUBILITY = 1.2E-02 (G/M3)
VAPOR PRESSURE = 1.6E-06 (MM Hg)
LOG (10) (OCTANOL-WATER COEFFICIENT) = 5.656
```
TOTAL MASS OF COMPOUND IN SYSTEM = 2.0E-02 MOL
EQUILIBRIUM FUGACITY = 3.55E-13 (ATM)
HENRY'S CONSTANT = 6.25E-04 (ATM M3/MOL)

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>VOL (M3)</th>
<th>I</th>
<th>%</th>
<th>CONC (PPM)</th>
<th>MASS (MOL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ATMOSPHERE (7.55Km)</td>
<td>1.0E+07</td>
<td>4.2E+02</td>
<td>5.3E-01</td>
<td>5.3E-09</td>
<td>1.5E-05</td>
</tr>
<tr>
<td>2 WATER (170 M)</td>
<td>6.3E+04</td>
<td>1.6E+05</td>
<td>1.3E+00</td>
<td>2.0E-06</td>
<td>3.6E-05</td>
</tr>
<tr>
<td>3 SEDIMENTS (1CM 4% C)</td>
<td>1.40E+02</td>
<td>1.0E+00</td>
<td>1.0E+03</td>
<td>2.5E-01</td>
<td>2.8E-02</td>
</tr>
</tbody>
</table>

TOP AQUATIC PREDATOR
ED ENCOUNTERED.

The input ecosystem file is in temporary local storage and can be retained by copying to a permanent file.

GET-LKMI
.COPY-LKMI
ATMOSPHERE (7.55KM) 1.00E+07 -1.00E+01
WATER (.70 M) .820E+04 -.200E+01
SEDIMENTS (1CM 4% C) .400E+02 -.400E+01
EXAMPLE

6. ACKNOWLEDGMENTS

Partial funding for this work was provided by the Office of Marine Pollution Assessment (OMPA).


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Appendix 1 – Model Runs

The following section lists the model inputs and outputs for the runs discussed in this report.

Global Ecosystem Run

**TOTAL DDT**

<table>
<thead>
<tr>
<th>MOLECULAR WEIGHT = 356</th>
</tr>
</thead>
<tbody>
<tr>
<td>SYSTEM TEMPERATURE = 15.10°C</td>
</tr>
<tr>
<td>WATER SOLUBILITY = .120E-02 (g/m³)</td>
</tr>
<tr>
<td>VAPOR PRESSURE = .160E-06 (mm Hg)</td>
</tr>
<tr>
<td>( \log(10) ) (OCTANOL-WATER COEFFICIENT) = 6.656</td>
</tr>
<tr>
<td>TOTAL MASS OF COMPOUND IN SYSTEM = .720E+10 g/mol</td>
</tr>
<tr>
<td>EQUILIBRIUM FUGACITY = .115E-12 (atm)</td>
</tr>
<tr>
<td>HENRY'S CONSTANT = .625E-04 (atm M3/mol)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>VOL(M³)</th>
<th>Z</th>
<th>%</th>
<th>CONC(PPM)</th>
<th>MASS(MOL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ATMOSPHERE (10 KM)</td>
<td>.51E+19</td>
<td>.42E+02</td>
<td>.24E+00</td>
<td>.17E-08</td>
<td>.25E+08</td>
</tr>
<tr>
<td>2 WATER &lt;3000 M)</td>
<td>.14E+13</td>
<td>.16E+05</td>
<td>.36E+02</td>
<td>.65E+06</td>
<td>.26E+10</td>
</tr>
<tr>
<td>3 DETRITUS 2PBB 1 %C</td>
<td>.28E+10</td>
<td>.49E+10</td>
<td>.22E-01</td>
<td>.20E-00</td>
<td>.16E+07</td>
</tr>
<tr>
<td>4 PLANTS AQUATIC</td>
<td>.45E+10</td>
<td>.20E+11</td>
<td>.14E+00</td>
<td>.81E+00</td>
<td>.10E+08</td>
</tr>
<tr>
<td>5 PLANTS LAND</td>
<td>.15E+13</td>
<td>.20E+11</td>
<td>.60E+02</td>
<td>.81E+00</td>
<td>.43E+10</td>
</tr>
<tr>
<td>6 ANIMALS AQUATIC</td>
<td>.11E+10</td>
<td>.25E+11</td>
<td>.43E-01</td>
<td>.10E+01</td>
<td>.31E+07</td>
</tr>
<tr>
<td>7 ANIMALS LAND</td>
<td>.50E+09</td>
<td>.25E+11</td>
<td>.20E-01</td>
<td>.10E+01</td>
<td>.14E+07</td>
</tr>
<tr>
<td>8 SEDIMENTS AQUATIC</td>
<td>.10E+13</td>
<td>.49E+09</td>
<td>.14E+01</td>
<td>.20E-01</td>
<td>.10E+09</td>
</tr>
<tr>
<td>9 SEDIMENT LAND</td>
<td>.15E+13</td>
<td>.99E+09</td>
<td>.24E+01</td>
<td>.40E-01</td>
<td>.17E+09</td>
</tr>
<tr>
<td>TOP AQUATIC PREDATOR</td>
<td></td>
<td></td>
<td></td>
<td>.22E+00</td>
<td></td>
</tr>
</tbody>
</table>
DDT Calibration for the Lake Michigan Ecosystem (See Table 2).

Run 1

CALIBRATION ⬇️ FLAKE MICHIGAN

TOTAL DDT

- MOLECULAR WEIGHT = 356
- SYSTEM TEMPERATURE = 15.0°C
- WATER SOLUBILITY = 1.20E-02 (g/M³)
- VAPOR PRESSURE = 1.60E-06 (MM Hg)
- LOG10 (OCTANOL-WATER COEFFICIENT) = 6.656
- TOTAL MHSS OF COMPOUND IN SYSTEM = 1.00E+07 GMOL
- EQUILIBRIUM FUGACITY = 6.14E-12 (ATM)
- HENRYS CONSTANT = 6.25E-04 (ATM M³/MOL)

<table>
<thead>
<tr>
<th>#</th>
<th>COMPARTMENT</th>
<th>VOL(M³)</th>
<th>%</th>
<th>CONC(PPM)</th>
<th>MASS(MOL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ATMOSPHERE (10 KM)</td>
<td>5.8E+15</td>
<td>42E+02</td>
<td>1.5E+01</td>
<td>9.3E-08</td>
</tr>
<tr>
<td>2</td>
<td>WATER (86 M)</td>
<td>4.9E+13</td>
<td>16E+05</td>
<td>4.8E+01</td>
<td>3.5E-05</td>
</tr>
<tr>
<td>3</td>
<td>DETRITUS (1.5PPM)</td>
<td>5.0E+17</td>
<td>4.9E+10</td>
<td>1.5E+01</td>
<td>1.1E+01</td>
</tr>
<tr>
<td>4</td>
<td>SEDIMENTS 5CM(L)</td>
<td>1.5E+10</td>
<td>9.9E+09</td>
<td>9.1E+02</td>
<td>2.2E+00</td>
</tr>
<tr>
<td>5</td>
<td>PHYTO 1500CEL/ML</td>
<td>7.5E+06</td>
<td>2.0E+11</td>
<td>9.1E+01</td>
<td>4.3E+01</td>
</tr>
<tr>
<td>6</td>
<td>ZOOPLANK (306/M³)</td>
<td>1.5E+06</td>
<td>2.0E+11</td>
<td>1.8E+00</td>
<td>4.3E+01</td>
</tr>
<tr>
<td>7</td>
<td>FISH FISH (1.5 M)</td>
<td>2.5E+05</td>
<td>2.0E+11</td>
<td>3.0E-01</td>
<td>5.4E-01</td>
</tr>
<tr>
<td>8</td>
<td>SALMONIDS (1.6E3 M)</td>
<td>3.2E+03</td>
<td>2.5E+11</td>
<td>4.9E-03</td>
<td>5.4E+01</td>
</tr>
</tbody>
</table>

TOP AQUATIC PREDATOR = 1.2E+01
### Run 2

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>VOL (M3)</th>
<th>Z</th>
<th>%</th>
<th>CONC (PPM)</th>
<th>MASS (MOL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATMOSPHERE (10 KM)</td>
<td>.58E+15</td>
<td>.42E+02</td>
<td>.64E+00</td>
<td>.39E-08</td>
<td>.64E+04</td>
</tr>
<tr>
<td>WATER (86 M)</td>
<td>.46E+13</td>
<td>.16E+06</td>
<td>.21E+02</td>
<td>.15E-04</td>
<td>.21E+06</td>
</tr>
<tr>
<td>DETRITUS (1.5PPM/10C)</td>
<td>.50E+07</td>
<td>.98E+10</td>
<td>.13E+01</td>
<td>.91E+00</td>
<td>.13E+05</td>
</tr>
<tr>
<td>SEDIMENTS (CHL:0.5DC)</td>
<td>.15E+10</td>
<td>.20E+10</td>
<td>.27E+02</td>
<td>.18E+00</td>
<td>.27E+06</td>
</tr>
<tr>
<td>PHYTO 1500CEL/ML</td>
<td>.75E+06</td>
<td>.32E+11</td>
<td>.77E+00</td>
<td>.36E+01</td>
<td>.77E+04</td>
</tr>
<tr>
<td>ZOOPLANK (306/M3)</td>
<td>.15E+06</td>
<td>.39E+11</td>
<td>.15E+00</td>
<td>.36E+01</td>
<td>.15E+04</td>
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<tr>
<td>FORPS FISH (1.5ES MT)</td>
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<td>.26E+01</td>
<td>.45E+01</td>
<td>.26E+03</td>
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<tr>
<td>SALMONIDS (1.6ES MT)</td>
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<td>.41E+03</td>
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**TOP AQUATIC PREDATOR**

### Run 3

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<tr>
<th>COMPARTMENT</th>
<th>VOL (M3)</th>
<th>Z</th>
<th>%</th>
<th>CONC (PPM)</th>
<th>MASS (MOL)</th>
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<tbody>
<tr>
<td>ATMOSPHERE (10 KM)</td>
<td>.58E+15</td>
<td>.42E+02</td>
<td>.64E+00</td>
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<td>.64E+04</td>
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<tr>
<td>WATER (86 M)</td>
<td>.46E+13</td>
<td>.16E+06</td>
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<td>.21E+06</td>
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<td>DETRITUS (1.5PPM/10C)</td>
<td>.50E+07</td>
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<td>PHYTO 1500CEL/ML</td>
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<td>ZOOPLANK (306/M3)</td>
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<td>.26E+03</td>
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<td>SALMONIDS (1.6ES MT)</td>
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**TOP AQUATIC PREDATOR**

### Run 4

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<th>%</th>
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<td>WATER (86 M)</td>
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<td>SEDIMENTS (CHL:0.5DC)</td>
<td>.15E+10</td>
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<td>.42E+06</td>
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<tr>
<td>PHYTO 1500CEL/ML</td>
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**TOP AQUATIC PREDATOR**

.12E+01
Run 5

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<th>%</th>
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<tr>
<td>ATMOSPHERE (10 KM)</td>
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<td>WATER (86 M)</td>
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<td>.16E+06</td>
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<td>DETRITUS 1.5PPM10C</td>
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<td>SEDIMENTS SC1: 1.5%DC</td>
<td>.15E+10</td>
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<td>.21E+00</td>
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<td>PHYTO 15000CEL/ML</td>
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<td>FORAG FISH (155 MT)</td>
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<tr>
<td>SALMONIODS (1.6E3 MT)</td>
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TOP AQUATIC PREDATOR: .15E+01

Run 6

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<td>.25E+02</td>
<td>.16E-04</td>
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<td>DETRITUS 1.5PPM10C</td>
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<td>FORAG FISH (155 MT)</td>
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TOP AQUATIC PREDATOR: .18E+01

Run 7

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<tr>
<td>DETRITUS 1.5PPM10C</td>
<td>.50E+07</td>
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<tr>
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<tr>
<td>ZOOPLANK (300G/M3)</td>
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<tr>
<td>FORAG FISH (155MT)</td>
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<td>.49E+11</td>
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<td>.74E+01</td>
</tr>
<tr>
<td>SALMONIODS (1.6E3 MT)</td>
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TOP AQUATIC PREDATOR: .23E+01
### Run 8

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<th>Z</th>
<th>CONC (PPM)</th>
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<td><strong>ATMOSPHERE (10 KM)</strong></td>
<td>.58E+15</td>
<td>.42E+02</td>
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<td><strong>WATER (86 M)</strong></td>
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<tr>
<td><strong>DETRITUS (5 PPM 10°C)</strong></td>
<td>.50E+07</td>
<td>.90E+10</td>
<td>.21E+01</td>
<td>.74E+00</td>
</tr>
<tr>
<td><strong>SEDIMENTS 5CM1 0°C</strong></td>
<td>.15E+10</td>
<td>.90E+09</td>
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<tr>
<td><strong>ZOOPLANK (3006/3)</strong></td>
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<td>.29E+01</td>
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<tr>
<td><strong>FORAG FISH (15E MT)</strong></td>
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<tr>
<td><strong>SALMONOID (1.6E3 MT)</strong></td>
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**TOP AQUATIC PREDATOR** .12E+01

### Run 9

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<td>.16E+01</td>
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<td><strong>WATER (86 M)</strong></td>
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<td>.16E+06</td>
<td>.33E+02</td>
<td>.12E-04</td>
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<tr>
<td><strong>DETRITUS 1 5 PPM 10°C</strong></td>
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<td>.90E+10</td>
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<td>.74E+00</td>
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<tr>
<td><strong>SEDIMENTS 5CM1 0°C</strong></td>
<td>.15E+10</td>
<td>.90E+09</td>
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<td>.74E-01</td>
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<tr>
<td><strong>PHYTO 1500000 ML</strong></td>
<td>.75E+06</td>
<td>.39E+11</td>
<td>.12E+01</td>
<td>.29E+01</td>
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<tr>
<td><strong>ZOOPLANK (3006/3)</strong></td>
<td>.15E+06</td>
<td>.39E+11</td>
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<td>.29E+01</td>
</tr>
<tr>
<td><strong>FORAG FISH (15E MT)</strong></td>
<td>.20E+05</td>
<td>.49E+11</td>
<td>.41E+01</td>
<td>.37E+01</td>
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<tr>
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<td>.32E+03</td>
<td>.49E+11</td>
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**TOP AQUATIC PREDATOR** .11E+00

### Run 10

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<th>Z</th>
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<td><strong>WATER (86 M)</strong></td>
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<td>.16E+07</td>
<td>.34E+02</td>
<td>.12E-04</td>
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<tr>
<td><strong>DETRITUS 1 5 PPM 10°C</strong></td>
<td>.50E+07</td>
<td>.90E+11</td>
<td>.21E+01</td>
<td>.74E+00</td>
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<tr>
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<td>.74E-01</td>
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<tr>
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<td>.15E+06</td>
<td>.39E+12</td>
<td>.23E+00</td>
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<tr>
<td><strong>FORAG FISH (15E MT)</strong></td>
<td>.20E+05</td>
<td>.49E+12</td>
<td>.42E+01</td>
<td>.37E+01</td>
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<tr>
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<td>.32E+03</td>
<td>.49E+12</td>
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</table>

**TOP AQUATIC PREDATOR** .12E+01
Run 11

CALIBRATION: NEW COMPOUND

DIELDRIN

MOLECULAR WEIGHT = 377
SYSTEM TEMPERATURE = 15.0 C
WATER SOLUBILITY = .240E+00 (G/M3)
VAPOR PRESSURE = .100E-06 (MMHG)

LOG(10)(OCTAHOL-WATER COEFFICIENT) = 5.131
TOTAL MASS OF COMPOUND IN SYSTEM = .500E+05 (G MOI)

EQUILIBRIUM FUGACITY = .169E-14 (ATM)
HENRY'S CONSTANT = .207E-06 (ATM M3/MOL)

<table>
<thead>
<tr>
<th>#</th>
<th>COMPARTMENT</th>
<th>VOL(M3)</th>
<th>CONC(PPM)</th>
<th>MASS(MOL)</th>
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<tbody>
<tr>
<td>1</td>
<td>ATMOSPHERE (10KM)</td>
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<td>.23E+01</td>
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<tr>
<td>2</td>
<td>WATER (86 M)</td>
<td>.49E+13</td>
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</tr>
<tr>
<td>3</td>
<td>DETRITUS L.5PPM10;C</td>
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<tr>
<td>5</td>
<td>PHYTO 1500CEL-ML</td>
<td>.75E+06</td>
<td>.15E+12</td>
<td>.38E+00</td>
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<tr>
<td>6</td>
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<tr>
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<tr>
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</table>

TOP AQUATIC/PREDATOR

.62E-01
CALIBRATION

MIREX

MOLECULAR WEIGHT = 546
SYSTEM TEMPERATURE = 15.0 C
WATER SOLUBILITY = 7.00E-04 (G/M3)
VAPOR PRESSURE = 1.00E-05 (MM HG)
LOG (10)(OCTANOL-WATER COEFFICIENT) = 7.608
TOTAL ASS OF COMPOUND IN SYSTEM = .31E+05 MOL
EQUILIBRIUM FUGACITY = .429E-12 (ATM)
HENRYS CONSTANT = .103E-01 (ATM M3/MOL)

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>VOL (M3)</th>
<th>Z</th>
<th>%</th>
<th>CONC (PPM)</th>
<th>MASS (MOL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ATMOSPHERE (10 KM)</td>
<td>.56E+15</td>
<td>.42E+02</td>
<td>.34E+02</td>
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<tr>
<td>2 WATER (86 M)</td>
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<td>.97E+02</td>
<td>.66E+00</td>
<td>.25E-07</td>
<td>.20E+03</td>
</tr>
<tr>
<td>3 DETRITUS 1.5PPM</td>
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<td>.30E+09</td>
<td>.21E+01</td>
<td>.70E-01</td>
<td>.64E+03</td>
</tr>
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<td>.30E+08</td>
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<td>.19E+05</td>
</tr>
<tr>
<td>5 PHYTO 1500CEL/ML</td>
<td>.79E+06</td>
<td>.12E+10</td>
<td>.12E+01</td>
<td>.28E-00</td>
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<td>7 FORGS FISH 1E5 MT</td>
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<td>.35E+00</td>
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<tr>
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<td>.66E+03</td>
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</tr>
</tbody>
</table>

TOP AQUATIC PREDITOR
EDI ENCOUNTERED. .44E-01
Run 13

ALIBRATION: NEW COMPOUND

NHPHTALEN

CLEUCULAR WEIGHT = 128
SYSTEM TEMPERATURE = 15.0 C
WATER SOLUBILITY = 330E+02 (G/M3)
VAPOR PRESSURE = 230E+00 (MM Hg)
LOG10(OCTANOL-WATER COEFFICIENT) = 3.384
TOTAL MASS OF COMPOUND IN SYSTEM = 100E+01 G MOLE
EQUILIBRIUM FUGACITY = 346E-16 (ATM)
HENRYS CONSTANT = 117E-02 (ATM M3/MOLE)

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>VOL (M3)</th>
<th>2</th>
<th>%</th>
<th>CONC (PPM)</th>
<th>MASS (MOLE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ATMOSPHERE (10 KM)</td>
<td>5.9E+15</td>
<td>4.2E+02</td>
<td>8.5E+02</td>
<td>1.9E-12</td>
<td>8.5E+00</td>
</tr>
<tr>
<td>2 WATER (86 M)</td>
<td>4.9E+13</td>
<td>8.5E+03</td>
<td>1.5E+02</td>
<td>3.8E-11</td>
<td>1.5E+00</td>
</tr>
<tr>
<td>3 DETRITUS 1.5PPM100C</td>
<td>5.0E+07</td>
<td>9.6E+05</td>
<td>1.7E-02</td>
<td>4.3E-09</td>
<td>1.7E-04</td>
</tr>
<tr>
<td>4 SEDIMENTS SCM: 1.0PPM</td>
<td>1.5E+10</td>
<td>9.6E+04</td>
<td>5.6E-01</td>
<td>4.3E-10</td>
<td>5.6E-03</td>
</tr>
<tr>
<td>5 PHYTO 1 5 00CEL/ML</td>
<td>7.5E+06</td>
<td>3.9E+06</td>
<td>1.0E-02</td>
<td>1.7E-08</td>
<td>1.0E-04</td>
</tr>
<tr>
<td>6 COOPLANK (306/M3)</td>
<td>1.5E+06</td>
<td>3.9E+06</td>
<td>2.0E-03</td>
<td>1.7E-08</td>
<td>2.0E-05</td>
</tr>
<tr>
<td>7 FORAGFISH (1E5 M3)</td>
<td>2.0E+05</td>
<td>4.8E+06</td>
<td>3.4E-04</td>
<td>2.1E-08</td>
<td>3.4E-06</td>
</tr>
<tr>
<td>8 SALMONOIDS (1.6E3 M3)</td>
<td>3.2E+03</td>
<td>4.8E+06</td>
<td>5.4E-06</td>
<td>2.1E-08</td>
<td>5.4E-08</td>
</tr>
<tr>
<td>TOP AQUATIC PREDATOR</td>
<td>.31E-08</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

33
Run 14

CALIBRATION NEW COMPOUND

BENZENE

MOLECULAR WEIGHT = 78
SYSTEM TEMPERATURE = 15.10 °C
WATER SOLUBILITY = 1.75E+04 (G/M³)
VAPOR PRESSURE = 988E+02 (MM HG)

LOG(10)(OCTANOL-WATER COEFFICIENT) = 2.080

TOTAL MASS OF COMPOUND IN SYSTEM = 1.10E+01 G MOL
EQUILIBRIUM FUGACITY = 3.93E-16 (ATM)
HENRY'S CONSTANT = 5.70E-02 (ATM M³/MOL)

# COMPARTMENT          VOL(M³)      Z         %        CONC (PPM)    MASS (MOL)
-----------------------------------------------------------------------------------------------
1 ATMOSPHERE (10 KM)   .58E+15      .42E+02   .97E+02   .13E-12   .97E+00
2 WATER (86 M)         .49E+13      .18E+03   .34E+01   .54E-12   .34E-01
3 DETRITUS 1.5PPM  (102 C)  .50E+07      .85E+03   .17E-04   .26E-11   .17E-06
4 SEDIMENTS (1.0°C)    .15E+10      .85E+02   .54E-03   .26E-12   .50E-05
5 PHYTO 1500CEL/M³     .75E+06      .34E+04   .13E-06   .85E-10   .13E-08
6 ZOOPLANK (306/M³)    .15E+06      .24E+04   .10E-05   .10E-10   .20E-07
7 FORAG FISH (1E5 MT)  .20E+05      .42E+04   .33E-06   .13E-10   .33E-08
8 SALMONID (1.6E3 MT)  .38E+03      .42E+04   .53E-08   .13E-10   .53E-10

TOP AQUATIC PREDATOR   .39E-10
Appendix 2 - Programs

Procedure File

GET+RUNMI
./COPY+RUNMI
GET+FUGMODI.
GET+SUBSI.
GET+TAPE7=KMK.
EDIT+FUGMODI+I=SUBSI.
REWIND+LGO.
REWIND+TAPE6.
MAP+OFF.
FIN. I=FUGMODI+L=0.
LGO.
REPLACE+TAPE7=KMK.
REWIND+TAPE6.
COPY+TAPE6+OUTPUT.
EOI ENCONTERED.

GET+SUBSI
./COPY+SUBSI
B
READP SFUGI SFUGO ZIN
EOI ENCONTERED.
Main Program

//GET,FUGMOD
//COPY,FUGMOD
PROGRAM MOD (INPUT,OUTPUT,TAPE6,TAPE5=INPUT,TAPE7)

MAIN PROGRAM FOR FUGACITYMODEL 1 (NODECOMPOSITION)

FOR INFORMATION SEE MACKAY (1979) EST & T 13 OCT 1979 1218-1223

COMMON /DATA/MW,TC,S6,VPW,XML,DW,C,M,N,CNAME,F,W
COMMFW/ID,JV(20),Z(20),IDC(20),CM(20),PCM(20),CC(20),NAME(20)

NO 10 I = 1,NC

V(I) = 0.
Z(I) = 0.
ID(I) = 0.
CM(I) = 0.
PCM(I) = 0.
CC(I) = 0.
10 NAME(I) = 0

CALL FUG

CALCULATE THE FUGACITY

SUMF = 0.
DO 40 I = 1,NC

40 SUMF = SUMF + V(I)*Z(I)

F = C/SUMF

CALCULATE THE COMPOUNDS DISTRIBUTION AT EQUILIBRIUM

SUMM = 0.
DO 50 I = 1,NC

50 CM(I) = F*V(I)*Z(I)

SUM = SUM + CM(I)

CM(I) = MASS (MOL) IN EACH COMPARTMENT SUM = C

CALCULATE PERCENTAGE DISTRIBUTION

DO 60 I = 1,NC

60 PCM(I) = 100*CM(I)/SUMM

CALCULATE COMPARTMENT CONCENTRATIONS

NO 70 I = 1,NC

70 CC(I) = CM(I)*MW/V(I)

CALL FUG

STOP

END ENCOUNTERED.
SUBROUTINES

SUBROUTINE FUGI

COMMON /DATA/ MM, TC, VM, YL, HK, NC, CNAM, FH
COMMON /ID/ V(20), X(20), ID(20), CH(20), PCM(20), CC(20), NAME(20)
COMMON /CK/ ICHK, CD(120)

SUBROUTINE CONTROLS INTERACTIVE INPUT TO FUGACITY MODELS

C

PRINT *,'ENTER THE COMPOUND NAME (20 CHARACTERS)'
READ (5, 11) NAME

11 FORMAT (2810)
PRINT *,'ENTER THE COMPOUND'S MOLECULAR WEIGHT'
READ *, MW
PRINT *,'ENTER TOTAL MASS OF COMPOUND IN SYSTEM (G) OR 1.0'
READ *, C
C = C / MW
PRINT *,'IS THERE A FILE DESCRIBING THE ECOSYSTEM? YES=1, NO=2'
READ *, ICHK
IF (ICHK EQ. 2) GO TO 100
PRINT *,'ENTER ECOSYSTEM FILE NUMBER, NUMBER OF COMPARTMENTS'
READ * NNC

C

REWIND
DO 10 I = 1, NNC
10 FORMAT (2810, 2E9.3)
READ (H, 15) NAME(I), ID(I), V(I), CD(I)

C

100 CALL CREHTZ
C
101 CONTINUE
C
THIS HAS DEFINED THE ECOSYSTEM AND COMPARTMENT PROPERTIES
C
RETURN
END

EDIT ENCOUNTERED.
SUBROUTINE CREHTZ
COMMON/DATA/ MW, TC, SG, VPM, XLDW, MC, NC, NAME, F, H
COMMON/ID/ V(20), Z(20), ID(20), CM(20), PC(20), CC(20), NAME(20)
COMMON/XICL, CD(20)
DIMENSION (8)

ROUTINE CALCULATES Z'S FUGACITY CAPACITIES

IF (ICHK .EQ. 1) GO TO 16
WRITE(5, 22)
2 FORMAT('/" )
1"Z'S ARE CALCULATED ASFOLLOWS"
2"VAPOR PHASE : Z = 1/RT"
3"LIQUID PHASE : Z = 1/H"
4"SORBED OR BIOTIC PHASE : Z = KP/H"
5"KP = KOC * % ORGANIC CARBON / 100"
6"KOC = 1.05 * XLDW - 0.50"

WRITE(5, 52)
5 2 FORMAT('/" )
1"FOR CLASSIFICATION" ENTER"
1"1 FOR VAPOR PHASE (ATM)"
2"2 FOR LIQUID PHASE"
3"3 SUBSTRATEog ORGANIC CARBON FOR SORBED OR BIOTIC PHASE"
4"4 FOR SOLVENT"

16 CONTINUE
PRINT *, "ENTER A TITLE FOR THIS RUN (80 CHARACTERS)"
READ(5, 31) (SYSTEM(J), J = 1, 8)
31 FORMAT(80)

PRINT .. "ENTER THE SYSTEM TEMPERATURE (C)"
READ **. TC
PRINT *, "ENTER THE COMPOUNDS WATER SOLUBILITY (G/M3)"
READ **, SG
PRINT * "ENTER THE COMPOUNDS VAPOR PRESSURE (MMHG)"
READ **, VPM

RG = 82.0 . IE-06
TK = TC + 273.
RT = RG . TK
VPM = VPM * 760.
SM = SG / RW
H = VPM / SM
XLDW = 5.00 - 0.670 * LG(10 SM + 1000)

IF (ICHK .EQ. 1) GO TO 77

PRINT .. "ENTER THE NUMBER OF ECOSYSTEM COMPARTMENTS"
READ **, NC
REWIND 7
PRINT "8"
90 FORMAT('/" NAME < IN QUOTES ) INFO (IN QUOTES) HERE 10 CHARACTERS MAX
1"
DO 41 I = 1, NC
PRINT 99, I
99 FORMAT('/" ENTER COMPARTMENT"I3,1X, "N H M E I N F O " VOL (M3) Z CLASS"
1 "(-1 , 1) Vol(%)"
READ .. NAME(I), ID(I), V(I), CD(I)
IF (CD(I), .GT. 0.) Z(I) = 10**(1.05 * XLDW - 0.50) * 0.01 * CD(I)/H
IF (CD(I), .EQ. 1.) Z(I) = 1.0 / RT
IF (CD(I), .EQ. -2.) Z(I) = 1.0 / H
11 FORMAT(29, 2) 41 WRITE(7, 11) NAME(I), ID(I), V(I), CD(I)
SUBROUTINE FUGO
COMMON /DATA/ ML, SG, VM, XLMKDW, WC, NAME(20)
COMMON /ID/ ID(20), CM(20), PC(20), CC(20)
C SUBROUTINE FOR DATA OUTPUT FROM FUGACITY MODELS
C
WRITE(6,200) NAME
WRITE(6,201) MH
200 FORMAT(3X,2A10)
201 FORMAT(*" MOLECULAR WEIGHT ="/I4,"/
WRITE(6,202) TC
202 FORMAT(*" SYSTEM TEMPERATURE ="/F6.1,"/
WRITE(6,203) SG
203 FORMAT(*" WATER SOLUBILITY="/E10.3," (G/M3)="/)
WRITE(6,204) VM
204 FORMAT(*" VAPOR PRESSURE="/E10.3," (MMHG)="/)
XLMKDW = ML*SG*VM/CWCM
WRITE(6,205) XLMKDW
205 FORMAT(*" TOTAL MASS OF COMPOUND IN SYSTEM="/E10.3," MOL="/)
WRITE(6,206) F
206 FORMAT(*" EQUILIBRIUM FUGACITY="/E10.3," (ATM)="/)
WRITE(6,207) H
207 FORMAT(*" EQUILIBRIUM FUGACITY="/E10.3," (ATM)="/)
WRITE(6,208) C
208 FORMAT(*" HENRY'S CONSTANT="/E10.3," (ATM.M3/MOL)="/)
WRITE(6,209) C
209 FORMAT(*" MASS OF COMPOUND VOL(M3)="/)
1 FORMAT(1E18.6) DO 12 I = 1, NC
12 WRITE(6,211) I, NAME(I), ID(I), CM(I), PC(I), CC(I)
11 FORMAT(*"================================================================="/)
111 FORMAT(12X,2A10,5X,E10.2,1X)
C BIOVARIABLES FOR AQUATIC PREDATOR-33X, E10.2)
RETURN
EOI ENCLOSED.
C
60 TO 44
X = 451 - 1, NC
45 IF CO(I).GT.0 Z(I) = 10**((1.05*XLMKDW-0.5)*0.01+CO(I))/H
45 IF CO(I).EQ.-1. Z(I) = 1./FT
45 IF CO(I).EQ.0. Z(I) = 1./H
C
44 PRINT 66
66 FORMAT(*") PRINT 31, I, SYSTEM(I) J=1,8
PRINT 66 XLMKDW = 10 ** XLMKDW
RETURN
EOI ENCLOSED.