The relative importance of water and ingested food as possible avenues of contaminant accumulation was examined. *Mysis relicta* was exposed to two representative non-polar contaminants: benzo(a)pyrene (BaP) and 2,2',4,4',5,5'-hexachlorobiphenyl (HCB). The accumulation was examined kinetically in the presence and absence of two food types, *Daphnia pulex* (water flea) and *Tabellaria flocculosa* (a diatom). The availability of either food enhanced HCB accumulation and reduced BaP accumulation. Feeding did not affect HCB elimination but enhanced BaP elimination. The BaP was eliminated primarily as metabolites. Essentially all of the elimination occurred via the fecal route for both compounds. The fraction of the accumulation via ingestion of contaminated food was greater when mysids fed on *Daphnia* than *Tabellaria* and was greater for HCB than BaP.

**Introduction**

The role of ingested contaminant in bioaccumulation is important to fully understand contaminant transport in aquatic ecosystems and, specifically, the importance of biomagnification. If significant biomagnification occurs, energy transfer to upper trophic levels with the concomitant flow of contaminant influences contaminant fate. Such trophic transfer may augment the potential deleterious health effects on higher trophic levels by increasing the contaminant exposure.

Until recently, evidence for the popular conception of food chain transfer and the importance of biomagnification was the subject of much debate (Biddinger and Gloss 1984). Laboratory data indicated that the extent of xenobiotic accumulation from food is quite variable and may depend on (1) the extent to which contaminants are metabolized and subsequently eliminated (Neff 1979) and (2) the differences between the concentrations in the water and food sources (Opperhuizen 1991). Accumulation of p,p'-DDT from food ranged between 30% and 100% of the p,p'-DDT body burden for fish (Macek and Korn 1970, Macek et al. 1970, Rhead and Perkins 1984) and the extent of kepone accumulation from a dietary source for shrimp was additive to the water source (Fisher and Clark 1990). Further, both feeding rate and food concentration affected the efficiency of accumulation for compounds accumulated from food (Opperhuizen and Schrap 1988, Weston 1990). Thus, food chain transfer is expected to produce concentrations above those that can be achieved through water-borne accumulation alone (Gobas et al. 1988).

Modeling efforts to describe the extent of hydrophobic contaminant accumulation have incorporated food...
chain transfer to bring the models in line with the field data (Thomann and Connolly 1984, Weininger 1978, Connolly and Tonelli 1985, Connolly and Pedersen 1988, Gobas et al. 1988, Connolly 1991). These models have had to assume factors such as contaminant assimilation efficiency and metabolizing capability of the animal. Recent field work has demonstrated that trophic structure, specifically the number of links in the food chain, is important for determining the final contaminant concentrations in the upper trophic levels for polychlorinated biphenyls (PCB) (Rasmussen et al. 1990). This field effort identifies biomagnification and food chain transport as important processes in contaminant accumulation.

In the above models, accumulation by the invertebrates is important for correctly predicting the transfer further up the food chain. However, mathematical models have not been developed to predict the contaminant body burden for invertebrates as a function of the specific kinetics or feeding. Conceptually, the role of feeding on contaminant accumulation by aquatic animals may (1) be additive, (2) negligible or (3) reduce the contaminant body burdens (Biddinger and Gloss 1984). While some experimental work has been performed with warm water species (Biddinger and Gloss 1984, Weston 1990, Fisher and Clark 1990), no research has been conducted to assess the effect of feeding on contaminant accumulation in cold water invertebrates of oligotrophic systems such as those studied by Rasmussen et al. (1990).

Our work examines the effects of food consumption on contaminant accumulation by the invertebrate Mysis relicta, which comprises an integral part of the ecology of the upper Laurentian Great Lakes. M. relicta is a vertically migrating omnivore (Beeton and Bowers 1982), consuming zooplankton (Bowers and Vanderploeg 1982) and algae (Bowers and Grossnickle 1978). Mysis may comprise up to 50% of the diet of juvenile lake trout (Dryer et al. 1965) and bloater (Weils and Beeton 1963). Because of its role as both predator and prey, M. relicta is a potential food chain vector for contaminants. Our study measured the toxicokinetics, from water and food sources, of the polycyclic aromatic hydrocarbon (PAH), benzo(a)pyrene (BaP) and the polychlorinated biphenyl (PCB) congener, 2,2',4,4',5,5'-hexachlorobiphenyl (HCB).

Materials and Methods

Collection of Mysis relicta: Mysids were collected at night from Lake Michigan from a station four miles west of Grand Haven, MI at a maximum depth of 65 m by obliquely towing a nitex net (1.0 m x 3.0 m; 565 μm mesh, fit with a bayonet-mounted weighted PVC cod-end, 2 L capacity) near the lake bottom. After collection, the mysids were transferred to aquaria containing approximately 50 L of hypolimnetic water and kept at approximately 4°C. The animals were transported to the laboratory within 24 h.

In the laboratory, Mysids were gently transferred to rectangular aquaria containing aerated and charcoal-filtered lakewater at 4°C. They were held in the dark because of their sensitivity to light (Smith 1970). Water volume changes of approximately 10% were made weekly to reduce the affects of waste accumulation. The mysids were fed daily rations of fresh trout chow. Approximately every 2 days, unconsumed trout chow was removed to minimize deleterious BOD and COD effects. Mysids were used for experiments within one month of collection to minimize potential deleterious effects due to containment.
**Daphnia pulex Collection, Growth and Maintenance:** *Daphnia pulex* were frequently collected simultaneously with *M. relicta*. Large, presumably gravid adults were gently seined from the mysid collection and placed in 20 L polyethylene carboys filled with Lake Michigan surface water. The daphnids were then transported to the laboratory and maintained on a light:dark cycle of 12h:12h. The daphnids were fed daily with aliquots of (1) uni-algal *Chlamydomonas oblonqa* cultures (obtained from H.A. Vanderploeg, Great Lakes Environmental Research Laboratory, NOAA, Ann Arbor, MI.) collected from Lake Huron, (2) a suspension of a water extract of Cerophyll, and (3) water reconstituted yeast cells. Neonates were removed by screening from the initial culture aquaria, using a 400 μm mesh nitex net. The neonates were then transferred to fresh filtered lakewater and grown to maturity.

**Tabellaria flocculosa Growth and Maintenance:** A uni-algal culture of *T. flocculosa* collected from Lake Huron was obtained from H.A. Vanderploeg (Great Lakes Environmental Research Laboratory, NOAA, Ann Arbor, MI.). A 100 mL inoculum was added to sterile WC medium excluding Tris buffer (Stein 1973) and grown to an optically dense, presumably late log phase culture.

Prior to use in feeding studies, the chlorophyll a (chl a) content of culture samples was determined. Aliquots were removed and centrifuged at 1400 g for 10 min to concentrate the algal cells. The cells were then added to fresh lakewater to give an approximate concentration of 15 mg-chl a m⁻³. In later studies the chl a content was determined along with the mass of *T. flocculosa*, in order to determine the chl a:mass ratio.

**Compounds:** ^3H-Benz(a)pyrene (BaP, specific activity = 23.8 Ci mMol⁻¹) was obtained from Amersham while ^14C-2,2',4,4',5,5'-hexachlorobiphenyl (HCB, specific activity = 13.1 mCi mMol⁻¹) was obtained from Pathfinder Laboratories (currently Sigma Radiochemicals). The compounds were tested for radiopurity with a combination of thin layer chromatography (TLC) and liquid scintillation spectrometry. The compounds were chromatographed on silica gel plates (250 μm, E Merck) in hexane:benzene (8:2 V:V). The silica gel was scraped from the plates at the retention position corresponding to the parent compound and at several other segments, including the origin. Each segment was placed in a separate scintillation vial with 12 mL scintillation cocktail (3a70B, Research Products International). The radioactivity of the samples was determined on a Packard 460C scintillation spectrometer. The radiopurity was determined as the percent parent compound activity divided by the total activity for the particular compound under test. All compounds used for study were at least 98% radiopure. Compounds found to be less than 98% pure were purified by TLC using the same solvent system as above.

**Determination of Xenobiotic Metabolism:** During certain experiments, whole mysids, fecal material, and exposure water were analyzed for degradation products. Wet mysids and dried fecal pellets were weighed and extracted in a 1:1 mixture of acetone:ethyacetate (EtOAc) by grinding in a 10 mm X 100 mm ground glass tissue grinder. The extract slurry was then quantitatively transferred to a conical centrifuge tube, capped with a Teflon-lined screw cap, and centrifuged at 1400 g for 10 min. The supernatant was transferred to a clean centrifuge tube using a Pasteur pipette. The tissue pellet was rinsed and centrifuged three more times, using fresh EtOAc.
The combined extract was concentrated to 1 mL or less with a gentle stream of nitrogen. The tissue residue was air dried, placed directly into 12 mL scintillation cocktail and allowed to soak overnight before being counted.

Aliquots of the concentrated extract were chromatographed on silica gel TLC plates (250 μm thickness- E. Merck). Each TLC plate was developed under a solvent-unsaturated atmosphere using a 4:1 hexane:benzene mixture. The dried plate was cut into 17 one-cm sections that were scraped onto glassine powder paper and placed into vials containing 12 mL of scintillation cocktail. The radioactivity of each section was determined by liquid scintillation counting.

Water from static uptake studies was pooled and extracted using a liquid-liquid dichloromethane (DCM)/cyclohexane system. The water was extracted three times with a 10% volume of DCM and three times with a 10% volume of cyclohexane. The extracts were combined and dried by passing them through granular anhydrous Na2SO4. The dried extract was then concentrated to approximately 10 mL using a rotary evaporator, followed by further concentration to 1 mL using nitrogen. The extracts were analyzed using TLC and scintillation counting, as described earlier.

**Toxicokinetic Experiments:** Experimentally, a three-compartment model was simulated such that mysids accumulated contaminant from both food and water sources. As references, parallel experiments were conducted to determine the uptake rates from water alone. In the three-compartment experiments, the food sources were pre-equilibrated with a concentration of xenobiotic in water which was the same as that used for the mysid exposure. Methanol was used as a carrier solvent for all experiments, because it has no observable effects on toxicokinetic parameters at concentrations used (approximately 50 mg L⁻¹)(Landrum, unpublished data).

**Feeding Experiments Utilizing Daphnia pulex:** Adult D. pulex were placed in fresh filtered lakewater containing radiolabeled ¹⁴C-HCB (approximately 200 DPM mL⁻¹) and ³H-BaP (approximately 1000 DPM mL⁻¹). The D. pulex were allowed to accumulate the radiolabeled compound for 48 h at 4°C in the dark. After 24 h of loading, fresh filtered lakewater was dosed with radiolabeled ¹⁴C-HCB and ³H-BaP at the same concentrations as above, poured into 700 mL serum bottles, and capped with Teflon-lined screw caps. At 48 h after the start of the loading phase, the D. pulex were removed from the loading aquarium and placed in eight, 700 mL serum bottles. An individual mysid was then placed in each serum bottle. For reference, eight mysids were also placed in bottles containing no D. pulex for contaminant uptake from water only. The capped bottles were placed on a wheel rotating at approximately 1 revolution per min (RPM).

At approximately 2, 4, 12, 24, and 48 h, reference and experimental bottles were removed from the rotating wheel. Two mL of whole water were sampled and placed in 12 mL scintillation cocktail for a determination of the total radioactivity present. Mysids and daphnids were recovered by pouring the exposure water through a 400 μm screen, rinsing with cold filtered lakewater to remove droplets of radioactive solution, and blotting to dryness. Mysids were weighed wet and placed in 12 mL scintillation cocktail at least 24 h prior to counting. Daphnids from each bottle were counted, combined in tared aluminum foil boats, dried at approximately 60°C for 48 h, and weighed. The daphnia were also held in 12 mL scintillation cocktail at least 24 h prior to counting.

Fecal material was collected from exposure water by applying suction to a 100 μL pipette. The fecal material
was then rinsed and placed in cold filtered lakewater and pipetted into tared aluminum boats. The fecal material was dried at 60°C for 48 h and weighed.

Elimination studies were conducted independently of the uptake studies. Mysids were "loaded" with $^{14}$C-HCB and $^3$H-BaP via exposure to contaminated water for 48 h. Fresh adult D. pulex were removed from their maintenance aquaria, placed in 700 mL serum bottles containing filtered lakewater at 4°C, and kept in the dark. Individual radiolabeled mysids were then transferred to serum bottles containing D. pulex and also to reference bottles having no food. The bottles were capped and placed on the rotator wheel at 4°C in the dark. Triplicate samples of the reference and the experimental bottles were removed at 2, 4, 8, 24, and 48 h after the start of the elimination phase. Mysids and fecal material were collected and treated as previously discussed. Daphnia were removed using a screen, placed in tared aluminum drying boats, dried at 60°C for 48 h, and weighed to estimate the mass of daphnia consumed during the experiment.

Feeding Experiments Utilizing Tabellaria flocculosa: Tabellaria flocculosa cells were harvested by centrifuging aliquots of culture at 1400 g for 10 min. The cells were resuspended in 50 mL filtered lakewater dosed with radiolabeled $^{14}$C-HCB (approximately 200 DPM mL$^{-1}$) and $^3$H-BaP (approximately 1000 DPM mL$^{-1}$) at 18°C in the dark. At approximately 36 h, fresh filtered lakewater dosed with $^{14}$C-HCB and $^3$H-BaP at the above concentrations was poured into 700 mL serum bottles. Aliquots of the labeled T. flocculosa suspension were pipetted into the serum bottles at concentrations of approximately 15 mg-chl a m$^{-3}$. Reference bottles contained radiotracer-dosed water only. Individual mysids were then placed in the serum bottles and the capped bottles were placed on a rotating wheel at 4°C in the dark. Mysids were removed at 2, 4, 8, 24, and 48 h after the start of the uptake phase. The mysids were sampled and prepared as described for the daphnia uptake experiments.

Initially, T. flocculosa was quantified on a mg-chl a m$^{-3}$ basis. However, to adequately determine the mass feeding rate and compare the kinetics with those from the daphnia experiments, the T. flocculosa mass was determined in subsequent experiments. Additionally, chl a:mass ratios were determined to convert chl a based kinetics data to mass-based kinetics data.

Chl a was determined on 100 mL aliquots of filtered (25 mm Gelman A/E glass fiber filters) exposure water. Each filter was then soaked in cold pH-buffered acetone overnight. The absorbance of the extracted chl a was measured on a Turner fluorometer at 520 nm. The chl a concentration was determined by comparison to a standard curve generated by using a spinach chl a (Sigma Inc.) solution in acetone. The radioactivity of the T. flocculosa was determined by filtering 100 mL of the exposure water through two 25 mm Gelman A/E glass fiber filters. The cells were then rinsed with approximately 25 mL of cold filtered lakewater to remove non-absorbed radioactivity.

The radioactivity in the cells was determined by placing the filters in 12 mL of scintillation cocktail and counting. The second filter was used to estimate the extent of radioactive contaminant sorption on the filters.

In an effort to determine the mass feeding rate of the mysids, the mass of algae associated with each bottle was determined gravimetrically. After sub-samples had been taken for chl a and radioactivity determinations and fecal pellets had been removed (as described above), the entire volume of water in the bottle was filtered through a rinsed, dried, and tared 25 mm Gelman A/E glass fiber filter. The filter was removed, placed on a 47 mm Teflon disc (to prevent adhesion during drying), dried in a desiccator, and weighed.
The radioactivity of filtered aliquots of water was determined by placing them in 12 mL of scintillation cocktail and counting them on a Packard 460 Scintillation Counter.

For elimination experiments utilizing T. flocculosa, mysids were loaded in the same manner as those utilizing D. pulex. T. flocculosa was concentrated from the culture flasks by centrifugation and resuspension in 700 mL cold filtered lakewater at a chl \_a concentration of 15 mg m\(^{-3}\). The mysids were added to the serum bottles that were capped and placed on a rotating wheel (1 RPM) at 4\(^{\circ}\)C and kept in the dark. The mysids were sampled at 4, 8, 16, 24, and 48 h. Mysids and fecal pellets were collected, prepared, and analyzed as described earlier.

**Kinetic Analyses:** The uptake clearance from water was determined for water-only exposures by using a two-compartment mass balance model where the mass of the xenobiotic is conserved (Equation 1).

\[
Q_a = \frac{k_{um} \cdot Q_s \cdot (1 - e^{-(k_{um} + k_e) \cdot t})}{(k_{um} + k_e)} \quad \text{Equation (1)}
\]

where,

- \(Q_a\) = the quantity (mg) of contaminant in the animal,
- \(t\) = time (h),
- \(k_{um}\) = the uptake rate constant from water (h\(^{-1}\)),
- \(Q_s\) = the quantity (mg) of contaminant in the system, and
- \(k_e\) = the elimination rate constant (h\(^{-1}\)) of the contaminant from the animal.

Under mass balance conditions, \(Q_t = Q_w + Q_a\) where \(Q_w\) amount of compound in the water compartment.

Initially, when \(Q_a\) is very small or if \(k_e\) is small relative to \(k_{um}\), the uptake clearance can be calculated from a simplification (Equation 2).

\[
k_{um} = -\frac{\ln (1 - \frac{Q_a}{Q_s})}{t} \quad \text{Equation (2)}
\]

\(k_{um}\) must then be converted to the system independent uptake clearance \(k_s\) by multiplying by the quantity \(V/M\) (Landrum 1983) where,

- \(V\) = the volume (mL) of the static exposure system, and
- \(M\) = the mass (g) of animal in the static exposure system.

Conversion to \(k_s\) is required so that comparisons can be made between experiments.

The elimination rate constants were calculated as the slope of the first order decay model (Equation 3).

\[
\ln C_a = \ln C_a^0 - k_e \cdot t \quad \text{Equation (3)}
\]

where, \(C_a^0\) = the concentration (ng g\(^{-1}\)) of the contaminant in the animal at the start of the elimination phase. Independent elimination rate constants can be added to determine the overall elimination rate constant.
Consequently, in the presence of feeding, \( k_u \) is the sum of elimination due to feeding (\( k_{s1} \)) and elimination due to water exposure (\( k_{s2} \)). The \( k_{s1} \) was determined independently from water controls while \( k_{s2} \) was determined by difference between \( k_u \) and \( k_{s1} \).

Uptake from water and food may be expressed mathematically as a three-compartment model (Equation 4).

\[
\frac{dC_a}{dt} = k_w \cdot C_w + k_f \cdot C_f - k_e \cdot C_a \quad \text{Equation (4)}
\]

where,

- \( C_a \) = the concentration (ng g\(^{-1}\)) of contaminant in the animal,
- \( t \) = time (h),
- \( k_w \) = the uptake clearance from water (ml g\(^{-1}\) h\(^{-1}\)),
- \( C_w \) = the concentration (ng mL\(^{-1}\)) of contaminant in water,
- \( k_e \) = the elimination rate constant (h\(^{-1}\)) of the contaminant from the animal,
- \( k_f \) = the uptake clearance (g g\(^{-1}\) h\(^{-1}\)) from food, and
- \( C_f \) = the concentration (ng g\(^{-1}\)) of contaminant in the food.

Because an exact integral of \( dC_a \) does not exist, Equation 4 must be rearranged to provide a numerical estimate of \( k_f \) (Equation 5).

\[
k_f = \frac{\left[ \frac{dC_a}{dt} - k_w \cdot C_w + k_f \cdot C_f \right]}{C_f} \quad \text{Equation (5)}
\]

For this study, \( k_w \) and \( k_{s1} \) were determined for the water controls while \( k_{s2} \) was determined under feeding conditions. \( k_f \) was then calculated as described in Equation 5.

Statistics: Means and slopes were compared using Student’s t test and were considered significant if \( p<0.05 \). Regressions were performed using the linear and nonlinear regression packages in SAS (1985).

Results

Metabolism of Benzo(a)pyrene: Over the course of a 6-h exposure at 4°C in the dark, approximately 24% of the radioactivity associated with mysids was found in the form of non-parent material. Of the total non-parent material associated with the animal, 5.7% was not extractable from the tissue and was, presumably, bound. The calculated metabolic rate for BaP under these conditions is 2.4 pmol g\(^{-1}\) dry weight h\(^{-1}\). Alternatively, HCB was not metabolized over a 24 h period under similar conditions.

Because both of the above experiments were conducted as single-labeled experiments, a dual-labeled experiment was conducted to determine if metabolic rates were different. As in the single labeled experiments,
HC8 was not metabolized; BaP was metabolized at a rate of approximately 2.2 pmol g\textsuperscript{-1} dry weight h\textsuperscript{-1}.

To crudely assess the mechanism of metabolite elimination, water was extracted from the uptake phase of a kinetics experiment. In the absence of feeding, 86 ± 2% of the tritium label was parent BaP and 98 ± 1% of the \textsuperscript{14}C label was parent HC8 from the water extracts. In the mysid, the majority of BaP was parent compound (68 ± 7%) at the beginning of the experiment; at the end of the experiment, non-parent BaP (70 ± 3%) predominated, suggesting that BaP metabolites accumulate in the absence of feeding.

BaP metabolite elimination appears to be influenced by fecal elimination. A TLC analysis of the fecal material indicated that 66% of the associated radioactivity was non-parent. Unfortunately, the amount of error could not be placed on this estimate because there was not a sufficient mass of fecal material necessary to provide replication.

After a 48 h-exposure to both contaminated water and food, the mysid BaP body burden was 30.3 ± 6.7% parent material. However, 77.7 ± 6.3% of the BaP in \textit{D. pulex} and 94.2 ± 3.3% of the BaP in the water was parent material, indicating that chemical changes in the water and/or metabolism by the food source cannot explain the metabolic rate of BaP observed in the mysid. Again, no HCB metabolism or degradation was observed in \textit{D. pulex} or water using these techniques.

**Feeding Rates**: The feeding rate (FR, grams consumed per gram organism per hour) for mysids exponentially decreased with time for both \textit{D. pulex} and \textit{T. flocculosa} (Figure 1). The average number of \textit{D. pulex} consumed per hour for three experiments was 0.97 ± 0.17 (n=3). The average feeding rates for mysids consuming \textit{D. pulex} and \textit{T. flocculosa} were 0.0008 ± 0.0003 mg dry wt. mg\textsuperscript{-1} h\textsuperscript{-1} (n=8) and 0.0019 ± 0.0006 mg dry wt. mg\textsuperscript{-1} h\textsuperscript{-1} (n=12), respectively. The FR was independent of both the mass of \textit{D. pulex} and \textit{T. flocculosa} available. Further, the FR did not depend on the size of the mysids.

![Figure 1](image1.png)

**D. pulex Feeding Studies**: The concentrations of HCB in the \textit{D. pulex} food source remained relatively constant
while BaP concentrations appeared to increase (Figure 2).

![Figure 2. Benzo(a)pyrene and Hexachlorobiphenyl concentrations in *Daphnia pulex.*](image)

Additionally, the concentrations of HCB and BaP in water decreased over time, as expected for a static system. The accumulation of HCB from a combined food and water source was approximately two times greater than the accumulation from a water source alone; BaP accumulation was approximately 2 times less than from a water source alone (Figure 3). *M. relicta* that consumed *D. pulex* exhibited increased elimination (k_{e} = 0.025 ± 0.001 h^{-1}) of BaP compared to unfed mysids (k_{e} = 0.0054 ± 0.001 h^{-1}).

![Figure 3. The accumulation of Benzo(a)pyrene (A) and hexachlorobiphenyl (B) in the presence and absence of feeding on *D. pulex.*](image)

However, the presence of uncontaminated food did not significantly affect the elimination of HCB by mysids (k_{e})
The amount of BaP and metabolites associated with fecal material was greater than 2 times that for HCB. Additionally, BaP appeared in the fecal material earlier than HCB (Figure 4).

Figure 4. The concentration of benzo(a)pyrene and hexachlorobiphenyl in fecal material eliminated by Mysis relicta while feeding on Daphnia pulex.

Tabellaria flocculosa Feeding Studies: The concentrations of HCB and BaP in T. flocculosa also were relatively constant over the course of the exposure while the concentrations of both decreased in the water. This trend was similar to that observed for the feeding studies with D. pulex.

HCB concentrations in mysids increased when exposed to T. flocculosa and water, relative to animals exposed to water alone (Figure 5). As in the case of the D. pulex experiments, BaP concentrations for mysids exposed to the T. flocculosa and water source were approximately two times less than mysids exposed to water alone (Figure 5).

Mysid elimination of HCB in the presence of T. flocculosa and water \( (k_b = 0.0015 \pm 0.0009 \text{ h}^{-1}) \) was similar to elimination in water alone \( (k_b = 0.0013 \pm 0.0006 \text{ h}^{-1}) \). For mysids exposed to BaP, the elimination was approximately four times greater in the presence of food \( (k_b = 0.042 \pm 0.003 \text{ h}^{-1}) \) relative to those exposed to water alone \( (k_b = 0.0098 \pm 0.0016 \text{ h}^{-1}) \). Further, mysids that produced fecal pellets had significantly lower BaP concentrations than those not producing fecal pellets (Table 1), while HCB concentrations did not differ for the two groups. Elimination for both compounds in the presence of T. flocculosa appeared to be dominated by the fecal route based on the concentration found in the fecal pellets (Figure 6).

As with D. pulex as a food source, elimination of fecal material with T. flocculosa as a food source resulted in greater BaP elimination than HCB elimination. Although the previous discussion suggests that fecal elimination only occurs for BaP, the elimination of HCB also appeared to occur in part via a fecal route. However, BaP was eliminated earlier and achieved maximum fecal elimination earlier than HCB.
Figure 5. The accumulation of benzo(a)pyrene (A) and hexachlorobiphenyl (B) in the presence and absence of *T. flocculosa*.

The calculated biological half-lives of HCB were not statistically different for mysids consuming *T. flocculosa* and *D. pulex* compared to the water controls. However, the half-lives of BaP for mysids consuming food sources were significantly reduced by 2 to 10 times compared to the water controls. Additionally, the half-life of BaP was 6 to 35 times smaller than that for HCB (Table 2).

Figure 6. Concentration of benzo(a)pyrene and hexachlorobiphenyl in fecal material eliminated by *M. relicta* feeding on *T. flocculosa*. 
Table 1. Concentrations of Benzo(a)pyrene and Hexachlorobiphenyl in *M. relicta* related to fecal pellet production

<table>
<thead>
<tr>
<th></th>
<th>Benzo(a)pyrene ( \text{dpm mg}^{-1} \text{ wet weight} )</th>
<th>Hexachlorobiphenyl ( \text{dpm mg}^{-1} \text{ wet weight} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fecal Pellet Production</td>
<td>228 ± 92(^1) (11)(^2)</td>
<td>434 ± 52 (10)</td>
</tr>
<tr>
<td>No Fecal Pellet Production</td>
<td>402 ± 178(^1) (4)</td>
<td>483 ± 96 (5)</td>
</tr>
</tbody>
</table>

1. Significantly different \((\alpha=0.05)\)
2. Numbers in parentheses represent number of replicates

**Discussion**

**Feeding Rates:** Selection of the food sources to test with mysids reflects the general omnivorous feeding habits of the organism. The cladoceran *D. pulex* was chosen as a representative zooplankton food source because in the field it is consumed by *Mysis relicta* at high rates (Bowers and Vanderploeg 1982). As a consequence, it is likely to contribute to the contaminant body burden of foraging mysids. The diatom, *Tabellaria flocculosa*, was chosen as an algal surrogate due to the preference of *M. relicta* for larger diatoms (Bowers and Vanderploeg 1982, Bowers and Grossnickle 1978) and the importance of diatoms in the Great Lakes system.

The mysids feed well on both sources confirming their omnivorous habits. Both food sources strongly affected the toxicokinetics of non-polar contaminants in the mysid.

Table 2. Food and water dependent half-Lives \( (t_{1/2}) \) for *M. relicta* previously exposed to benzo(a)pyrene and hexachlorobiphenyl in water

<table>
<thead>
<tr>
<th>Exposure Regime</th>
<th>Half-life Benzo(a)pyrene (h)</th>
<th>Half-life Hexachlorobiphenyl (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>T. flocculosa</em> and water</td>
<td>17 ± 1 (3)(^1)</td>
<td>576 ± 203 (3)</td>
</tr>
<tr>
<td>Water Controls</td>
<td>72 ± 13 (3)</td>
<td>347 (2)</td>
</tr>
<tr>
<td><em>D. pulex</em> and water</td>
<td>28 ± 1 (3)</td>
<td>463 ± 327 (3)</td>
</tr>
<tr>
<td>Water Controls</td>
<td>132 ± 26 (3)</td>
<td>147 ± 87 (3)</td>
</tr>
</tbody>
</table>

1. Numbers in parentheses equals number of replicates.

The controlling factors for the observed feeding rates for both food types are not well defined. However, an inflection point in the curve occurs for both food types at approximately 10 to 15 hours, suggesting that the
feeding rate may depend on satiation. Such a "gut-fullness" hypothesis is not particularly novel. However, it must be considered because changes in feeding rates can affect the accumulation of contaminant from food and, consequently, confound the uptake clearance coefficient from food ($k_u$).

Within the first 10 h of the experiments, the average number of $D. pulex$ consumed per hour compared favorably with in situ predation rates of 1.84 ± 0.95 cladocerans consumed per hour (Bowers and Vanderploeg 1982). The feeding rates expressed as the percentage of mysid dry weight consumed per hour also agree with those determined by Bowers and Vanderploeg (1982). The feeding rates appear to depend on the food type offered. $M. relicta$ fed on $T. flocculosa$ almost three times as much as they did on $D. pulex$. This difference may be due in part to daphnia's enhanced escape capability. A similar mechanism is attributed to mysid's prey selection of cladocerans over copepods by (Bowers and Vanderploeg 1982). Alternatively, differential feeding rates may be controlled by food preferences that may in turn be driven by chemoreceptive mechanisms. Identification of the mechanisms actually controlling the feeding rate and producing the observed differences requires further research.

Toxicokinetics: The apparent increase in the BaP concentration in the $D. pulex$ food source may be related to a buildup of metabolic products in the animal. This mechanism for BaP accumulation is reasonable, given that BaP metabolites are reactive and bind to macromolecules. Metabolite buildup has been observed in invertebrates including mysids (Gardner et al. 1990) and chironomids (Leversee et al. 1982) with BaP metabolism. The decrease in contaminant concentrations in the water indicates that the mass balance kinetics approach must be used.

The consumption of contaminated food increased the HCB body burden of $M. relicta$ by a factor of almost two. The relatively large increase in HCB from a food source is explained in part because $M. relicta$ is not able to eliminate the compound rapidly, even when fed a clean food supply. These findings suggest HCB mobility within the animal is restricted, and once accumulated, remains associated with the animal. The consumption of $D. pulex$ contaminated with BaP does not increase but rather reduces the BaP body burden for $M. relicta$. Elimination experiments show that BaP concentrations are substantially reduced by consumption of this food source.

Similar changes in mysid toxicokinetics are observed when $T. flocculosa$ is used as the food source in place of $D. pulex$, however, the elimination rate constants are considerably larger with $T. flocculosa$. This difference most likely results from the increased mass feeding rates for $M. relicta$ feeding on the diatom compared to feeding on $D. pulex$. The more rapid elimination rate for animals feeding on $T. flocculosa$ may be partially related to the rate of food movement through the gut, coupled with the increased sorptive capacity of diatom frustules, as compared to cladoceran carapaces. In support of this hypothesis, the more rapid consumption of $T. flocculosa$ appears to be related to more frequent fecal pellet production. Such increases in elimination with feeding have also been found in fish exposed to BaP (Jimenez et al. 1987).

The differential metabolism of these compounds also explains some of the differences in their kinetic behaviors. However, feeding coupled with fecal elimination appears to be necessary to maintain these differences. Overall, the results suggest that during feeding, the HCB body burden is likely to increase while
the BaP/BaP metabolite body burden will decrease. The increase in accumulation of the non-metabolizable chlorinated biphenyls with feeding is supported by the concurrence of a food chain accumulation model for PCBs with field observations (Connolly 1991).

Elimination and Metabolism: *M. relicta* is not able to biotransform HCB to any substantial extent. While biotransformation of PCBs occur in aquatic animals, it is usually restricted to the less chlorinated congeners (Melanchon and Lech 1976). Thus, HCB biotransformation is not expected, and it has been found to be extremely recalcitrant based on a wide range of studies with a variety of organisms. Alternatively, *M. relicta* biotransformed BaP at 2.4 pmol g$^{-1}$ dry wt h$^{-1}$. The mixed function oxidase enzymes required to biotransform xenobiotics have been reported for other aquatic invertebrates (Singer and Lee 1977, Stegeman and Kaplan 1981, and Kreiger and Lee 1973) and the capability to biotransform BaP, in particular, has been reported for *Chironomus riparius*. *C. riparius* can metabolize BaP at a rate of 3.2 nmol g$^{-1}$ h$^{-1}$ (Leversee et al. 1982). The difference in metabolic rate between the chironomids and mysids can be attributed to the exposure, 0.6 - 1 $\mu$g L$^{-1}$ for chironomids and 4 ng L$^{-1}$ for mysids, a difference of a factor of 250 and the differences in temperature 25°C for chironomids and 4°C for mysids. Additionally, the amphipod *Hyalella azteca* metabolizes the PAH anthracene at a rate of 2.4 nmol g$^{-1}$ dry wt. h$^{-1}$ (Landrum and Scavia 1983). Again, these rates compare favorably with the rate for BaP metabolism by *M. relicta* of 2.4 pmol g$^{-1}$ dry wt. h$^{-1}$ if differences in exposure and temperature are considered. Finally, the metabolic rate for mysids in this experiment was approximately ten times that for non-fed mysids exposed at somewhat lower concentrations 1 to 2 ng L$^{-1}$ (Gardner et al. 1990).

The overall effect of such BaP biotransformation by *M. relicta* appears to be the enhanced potential for metabolite elimination. However, such elimination appears to also require coupled fecal material production to realize the full elimination potential. The dependence on fecal elimination may be the result of the absence of a functional kidney, typically used to eliminate polar metabolites in higher trophic level organisms. The loss of xenobiotic compounds via a fecal route may be due to their incorporation in the peritrophic membrane that encases the feces (Maucheline 1980). This mechanism has been postulated in the loss of p,p'-DDT residues from mosquito larvae (Abedi and Brown 1961).

Elimination and Feeding: Enhanced elimination with feeding has been reported by other investigators. Anthracene elimination is accelerated by consumption of a sediment for the amphipod *Hyalella azteca* (Landrum and Scavia 1983). Similarly, the midge, *Chironomus riparius*, exhibited increased elimination of BaP while feeding on a paper towel substrate (Leversee et al. 1982). Although fecal material was not collected for either study, it is likely that mechanisms (i.e., biotransformation and fecal elimination) similar to those operative in *M. relicta* are involved.

In *M. relicta*, BaP accumulation and elimination can be conceptualized as a multi-compartment process. Uptake and elimination of BaP occurs via both food and water routes. The majority of the accumulated material eliminated in the feces appears to be metabolites. The TLC of the water extracts suggests that parent material is eliminated directly to the water. These findings suggest that different mechanisms operate based on the nature of molecule.
Molecular specific elimination mechanisms appear to be primarily responsible for the observed differences in the \( k_e \) estimates of BaP and HCB. Additionally, \( k_e \) measured for BaP appears to be related to the type of food offered and is primarily a function of feeding rate. Consequently, the half-life of BaP in _M. relicta_ feeding on _T. flocculosa_ is reduced by almost a factor of two compared to mysids feeding on _D. pulex_. Further, the HCB half-life is 17 to 33 times that of BaP and does not depend on the food source. These observations are significant for estimating biocencentration factors (BCF) for hydrophobic chemicals from log octanol:water partition coefficients. Indeed, if the increased elimination via feeding is ignored, the mysid BCF may be over-estimated by as much as one order of magnitude.

Our results have important implications for the interpretation of toxicant concentrations for field collected _M. relicta_. The increased elimination of BaP and metabolites associated with a diatom suggests that the compound’s concentration and potentially toxic metabolites are lowest during periods of diatom abundance. Consequently, seasonally dependent toxic responses may occur that are driven primarily by varying internal toxicant doses as a function feeding-mediated elimination. Alternatively, although feeding can presumably increase the concentration of HCB in the animal, feeding-mediated elimination is not important and therefore concentrations and the potential toxicity of HCB are expected to be directly related to concentrations in the food source.

**Feeding and Accumulation:** Initially, the feeding rate constants, \( k_f \), were determined by solving Equation 5 for \( k_e \). However, this approach resulted in estimates of \( k_f \) for both food types that exceeded all the experimentally determined feeding rates. This finding posed a dilemma in that the assimilation efficiency (defined as the ratio of the uptake clearance to the feeding rate) was substantially greater than 100%. Conceptually, these results mean that the amount of assimilated contaminant would exceed the total amount of contaminant consumed by the animal. Because of this contradiction, the numerical calculation of \( k_f \) must not be the appropriate model to use to estimate the real \( k_f \). Additionally, without an estimate of \( k_f \), one is not able to estimate the assimilation efficiency.

An examination of the \( k_f \) expression suggests that \( k_f \) would be overestimated if (1) the rate of accumulation \( (dC_a/dt) \) is overestimated, (2) the product \( k_f C_a \) is underestimated, (3) \( C_a \) is underestimated, or (4) the product \( k_f C_a \) is overestimated. In this experimental design, \( k_a \) is estimated by application of the reference \( k_a \) to the feeding animals. Indeed, the activity of feeding may accelerate the flow of contaminant past the animal and hence elevate \( k_a \) for feeding organisms relative to non-feeding organisms. This scenario may be plausible if contaminant uptake is proportional to increases in respiration rates due to feeding. Earlier studies suggest that respiration rates are partially related to the uptake clearances from water for BaP and HCB in _Mysis relicta_ (Landrum et al. 1992). Therefore, \( k_a \) may differ substantially during feeding due to respiration rate changes. Additionally, if uptake can occur across the carapace and is diffusion limited, then animals passing through a greater water volume during prey search and selection may increase their contaminant body burden and consequently show an increased \( k_a \).

To test the general hypothesis that \( k_a \) with and without feeding are essentially the same, the experimentally determined feeding rates were substituted into the \( k_a \) expression, assuming that the observed feeding rates were
equal to $k_i$ and xenobiotic assimilation efficiencies were 100%. The uptake clearances ($k_{wi}$) were then calculated from the expressions and compared to the measured reference $k_e$ values. The calculated $k_{wi}$ was not statistically different from the experimentally determined $k_e$ (Table 3). However, the variability in the estimates was so large that it was not possible to tell whether the uptake clearance was altered. Consequently, it appears that the substitution of the control $k_{wi}$ into the $k_i$ expression is valid, and an alternative model must be used for estimating $k_i$.

If the experimentally obtained feeding rates are substituted for $k_i$ under the assumption that the assimilation efficiencies are equal to 1.0, then the percentage of contaminant obtained from food is at the most 41% (Table 4). Additionally, the use of the feeding rate for the $k_f$ value provides the most realistic and liberal estimate of the fraction of contaminant obtained from a food source for this animal. As presented, a greater fraction of HCB is derived from both D. pulex and T. flocculosa compared to BaP. These observations may be explained by differential molecular transfer throughout the animal because of substantial differences in the stereochemistry and electronegativity of the molecules.

Table 3. Comparison of derived ($k_{wi}$) and experimental ($k_e$) uptake clearance constants ($k_{wi}$) from water for Mysis relict.a.

<table>
<thead>
<tr>
<th>Food Type</th>
<th>Benzo(a)pyrene</th>
<th>Hexachlorobiphenyl</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$k_{w1}$ (ml g$^{-1}$ h$^{-1}$)</td>
<td>$k_{w2}$ (ml g$^{-1}$ h$^{-1}$)</td>
</tr>
<tr>
<td>T. flocculosa</td>
<td>90 ± 61</td>
<td>113 ± 111</td>
</tr>
<tr>
<td>D. pulex</td>
<td>87 ± 30</td>
<td>56 ± 31</td>
</tr>
</tbody>
</table>

Table 4. Estimated percentages of benzo(a)pyrene and hexachlorobiphenyl obtained from food using the measured feeding rates ($k_f$).

<table>
<thead>
<tr>
<th>Food Type</th>
<th>Percent Obtained From Food</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Benzo(a)pyrene</td>
</tr>
<tr>
<td>D. pulex</td>
<td>22 ± 17</td>
</tr>
<tr>
<td>T. flocculosa</td>
<td>17 ± 10</td>
</tr>
</tbody>
</table>

More HCB and BaP are estimated to be obtained from D. pulex than from T. flocculosa. This difference may be related to higher feeding rates on T. flocculosa. Mechanistically, this hypothesis implies that the gut residence time for T. flocculosa mass is less than that for D. pulex. A reduced residence time of food in the gut may consequently restrict the time necessary for enzymatic processes to degrade gut materials and transfer associated contaminant to the tissues. Conversely, an increase in the gut residence time (as hypothesized for
D. pulex) would permit a more extensive digestive process and subsequent contaminant transfer with an apparent increased accumulation of contaminant from food. It is also possible that the surface areas of the food sources play an important role in determining contaminant bioavailability.

Various estimates of the amount of contaminant derived from the food source have been made. Sediment and associated porewater is estimated to contribute 77% of the anthracene body burden to the amphipod *Hyalella azteca* (Landrum and Scavia 1983). It is likely that material sorbed to sediment is less bioavailable than that associated with porewater. Consequently, the actual percentage of anthracene from the sediment food source is probably less than 77%. Additionally, food for fish species such as lake trout and salmon is estimated to contribute up to 90% of the contaminant load of these animals (Thomann and Connolly 1984, Weininger 1978), while the marine fish *Loiostomus xanthurus*, accumulates only 54% of its PCB body burden from food. In the case of BaP, bluegills are estimated to obtain 40 to 50% of their dose from food (Jimenez et al. 1987). However, fish species (particularly larger individuals) have smaller surface area-to-volume ratios relative to macro-invertebrates. This would reduce the relative contribution from a water source for fish. In studies performed on the smaller fathead minnow *Pimephales promelas*, a daphnia food source accounted for less than 1% of the total acridine body burden (Southworth et al. 1979), thus supporting the hypothesis that food does not contribute substantially to the xenobiotic body burden of aquatic animals.

The results presented in this study suggest that at most, food contributes less than 45% of the contaminant input to *Mysis relicta*. Consequently, mathematical models assuming a 90% contribution from food to the xenobiotic body burden of macro-invertebrates may overestimate the extent of food contribution by a factor of at least a factor of two and may not account for feeding induced changes in compound elimination.

**ACKNOWLEDGEMENTS**

We wish to thank the crew of the R/V Shenehon for assistance in collecting mysids. We also wish to thank Wendy Dupuis for technical assistance. GLERL contribution number 797.

**REFERENCES**


