

Methods for Assessing Bioaccumulation of Sediment-Associated Contaminants with Freshwater Invertebrates

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ver the past 10 years, a variety of methods have been described for evaluating the toxicity of sediment-associated contaminants with freshwater invertebrates (i.e., USEPA, 1994; ASTM, 1997a). However, only a limited number of standard methods are currently available for assessing bioaccumulation of contaminants from field-collected or laboratory-spiked sediments (see page 1-31). Standard guides have recently been published for conducting 28-day bioaccumulation tests with the oligochaete Lumbriculus variegatus including determination of bioaccumulation kinetics for different compound classes (USEPA, 1994; ASTM, 1997b). These methods have been applied to a variety of sediments to address issues ranging from site assessments to bioavailability of organic and inorganic contaminants using field-collected and laboratory-spiked samples (Schuytema et al., 1988; Nebeker et al., 1989; Ankley et al., 1991; Call et al., 1991; Carlson et al., 1991; Ankley et al., 1993; Kukkonen and Landrum, 1994; Brunson et al., 1998; see ASTM, 1997b for a listing of these citations). Results of laboratory bioaccumulation studies with L. variegatus have been confirmed with comparisons to residues (polychlorinated biphenyls, PCBs; polycyclic aromatic hydrocarbons, PAHs) present from field populations of oligochaetes collected from the same sites as sediments used in the laboratory exposures (Ankley et al., 1992; Brunson et al., 1998). Additional method development is under way to evaluate bioaccumulation kinetics and to provide additional data confirming responses observed in laboratory sediment tests with benthic communities in the field.

Selection of Test Organisms

The choice of a test organism has a major influence on the relevance, success, and interpretation of a test. Various organisms have been suggested for use in studies of chemical bioaccumulation from freshwater sediments (Table 1). The following criteria outlined in Table 1 were used to select L. variegatus for bioaccumulation method development by USEPA (1994) and ASTM (1997b): (1) ease of culture and handling, (2) known chemical exposure history, (3) adequate tissue mass for chemical analyses, (4) tolerance to a wide range of sediment physicochemical characteristics, (5) low sensitivity to contaminants associated with sediment, (6) amenability to longterm exposures without feeding, (7) ability to accurately reflect concentrations of contaminants in field-exposed organisms (i.e., exposure is realistic), and (8) data confirming the response of laboratory test organisms with natural benthic populations. Thus far, extensive interlaboratory testing has not been conducted with L. variegatus. Other organisms did not meet many of the selection criteria outlined in Table 1, including mollusks (valve closure), midges (short life cycle), mayflies (difficult to culture), amphipods (i.e., Hyalella azteca: small tissue mass, too sensitive), cladocerans and fish (not in contact with sediment).

Testing Procedures for *Lumbriculus variegatus*

The 28-day bioaccumulation test with *L. variegatus* described in USEPA (1994) and ASTM (1997b) is conducted with adult oligochaetes at 23° C with a 16L:8D photoperiod at an illuminance of about 500 to 1000 lux. Test chamber size ranges from 4 to 6 L, and the chamber contains 1 to 2 L of sediment and 1 to 4 L of overlying water with five replicates recommended for routine testing. To minimize depletion of sediment contaminants, a ratio of 50:1 total organic carbon in sediment to dry weight of organisms is recommended. A minimum of 1 g (wet weight)/replicate, with up to 5 g/replicate should be tested. Organisms are not fed during a bioaccumulation test (see page 1-36).



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Criterion	Lumbriculus variegatus	Mollusks	Midges	Mayflies	Amphipods	Cladocerans	Fish
Laboratory culture	+	-	+	-	+	+	+
Known chemical exposure	+	-	+	+/-	+	+	+
Adequate tissue mass	+/-	+	-	+	-	-	+
Low sensitivity to contaminants	+	+	-	-	-	-	+/-
Feeding not required during testing	+	+	-	+	-	-	+
Realistic exposure	+	+/-	+	+	+	-	-
Sediment physico- chemical tolerance	+	?	+/-	-	+	NA	NA
Response confirmed with benthic populations	+	?	?	?	+	?	-

Table 1. Selection criteria for sediment bioaccumulation test organisms (EPA, 1994; ASTM, 1997b; Ingersoll et al., 1995). A "+" or "-" rating indicates a positive or negative attribute; "NA" is not applicable; and "?" is unknown.

If sediments could be toxic to *L. variegatus*, a 4-day toxicity screening test should be conducted before starting a bioaccumulation test (ASTM, 1997b). Endpoints monitored in the toxicity test are survival and behavior. Test organisms should burrow into test sediment because avoidance of test sediment by *L. variegatus* may reduce bioaccumulation. Survival of *L. variegatus* in the toxicity screening test should not be significantly reduced in the test sediment relative to a control sediment. Additional requirements for test acceptability are outlined in USEPA (1994) and ASTM (1997b).

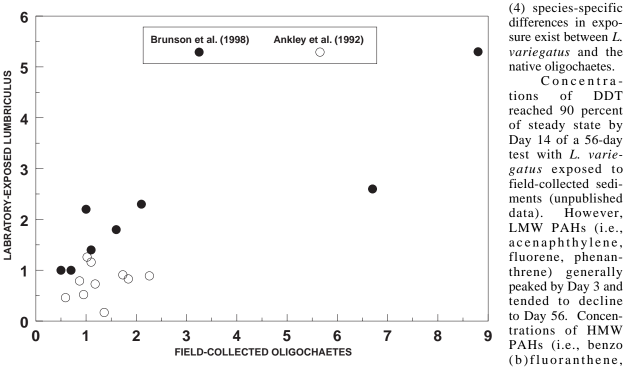
At the end of the bioaccumulation test, live oligochaetes are transferred to a 1-L beaker containing overlying water without sediment for 24 hours to eliminate gut contents (oligochaetes clear more than 90 percent of the gut contents in 24 hours). A correction for the extent of elimination from the body burden may need to be made for compounds with $\log K_{ow}$ less than 5. Oligochaetes are not placed in clean sediment to eliminate gut contents because clean sediment can contribute 15 to 20 percent to the dry weight of the oligochaetes, resulting in a dilution of contaminant concentrations on a dry weight basis. Minimum tissue mass required for various analyses at selected lower limits of detection are listed in USEPA (1994) and ASTM (1997b). Depending on study objectives, total lipids can be measured on a subsample of the total tissue mass of each replicate sample. Dry weight of oligochaetes can be determined on a separate subsample from each replicate.

Because bioaccumulation tests are often used in ecological or human health risk assessments, the procedures are designed to generate estimates of steady-state tissue residues. Eighty percent of steady state is used as the general goal for a test (ASTM, 1997b). An option when conducting a bioaccumulation test is to perform a kinetic study to estimate steady-state concentrations instead of conducting a 28-day bioaccumulation test (e.g., sample on Days 1, 3, 7, 14, 28). A kinetic test can be used when 80 percent of steady state will not be obtained within 28 days or when more precise estimates of steadystate tissue residues are required (see page 1-37).

Case Studies

Methods for conducting bioaccumulation tests with L. variegatus have varied slightly over the years; however, test conditions (e.g., test length, exposure systems) have been consistent enough for evaluation of the robustness of the guidance outlined in USEPA (1994) and ASTM (1997b). In a study with sediments from the lower Fox River in Green Bay, Wisconsin, Ankley et al. (1992) compared the bioaccumulation of PCBs by L. variegatus exposed in the laboratory to PCB residues in collections of oligochaetes from the field. Good agreement was observed between PCB concentrations in the laboratory and field organisms, particularly for those congeners with K_{ow} values <7 (see Figure 1). This indicates that for super-hydrophobic chemicals, laboratory exposures longer than 28 days may be required to reach equilibrium.

Good agreement was also observed in bioaccumulation between *L. variegatus* exposed in the laboratory for 28 days and field-collected oligochaetes from sediments collected from the upper Mississippi River (Brunson et al., 1998). About 90 percent of the corresponding concentrations of PAHs were within a factor of 3 between the laboratory-exposed and fieldcollected oligochaetes (see Figure 1). Concentrations that differed by more than a factor of 3 included



variegatus and the native oligochaetes. Concentrations of DDT reached 90 percent of steady state by Day 14 of a 56-day test with L. variegatus exposed to field-collected sediments (unpublished data). However, LMW PAHs (i.e., acenaphthylene, fluorene, phenanthrene) generally peaked by Day 3 and tended to decline to Day 56. Concentrations of HMW PAHs (i.e., benzo (b)fluoranthene, benzo(e)pyrene, indeno(1,2,3c,d)pyrene) typically either peaked

Figure 1. Biota-sediment accumulation factors (BSAFs) for laboratory-exposed Lumbriculus variegatus and field-collected oligochaetes for PAHs (Brunson et al., 1998) and PCB homologs (Ankley et al., 1992).

naphthalene, 1-methylnaphthalene, 2-methylnaphthalene, 2,6-dimethylnaphthalene, 1,6,7-trimethylnaphthalene, phenanthrene, 1-methylphenanthrene, and benz(a)anthracene. Tissue concentrations of naphthalenes were generally higher in field-collected oligochaetes relative to laboratory-exposed oligochaetes (naphthalenes are low molecular weight (LMW) PAHs with log Kow values less than 4.5). Compounds with similar concentrations in both the laboratory-exposed and field-collected oligochaetes included a similar number of high molecular weight (HMW) and LMW PAHs. These compounds included biphenyl, fluorene, 1-methylphenanthrene, pyrene, fluoranthene, chrysene, and benzo(e)pyrene. Most of these compounds are intermediate in molecular weight and log K_{ow} (except for benzo(e)pyrene, which has the highest molecular weight and $\log K_{ow}$ compared to these other compounds). Compounds with concentrations typically higher in the laboratory-exposed oligochaetes compared to field-collected oligochaetes were primarily HMW PAHs. These compounds included phenanthrene, benz(a)anthracene, benzo(b,k)fluoranthene, and perylene (with log K_{ow} greater than 4.5).

Differences between tissue concentrations in the laboratory-exposed and field-collected oligochaetes may be the result of differential exposure, including the following factors: (1) LMW PAHs may be lost during the sampling of sediments from the field; (2) spatial heterogeneity of contaminants in the field may have resulted in differential accumulation; (3) the route of exposure for oligochaetes in the field is through sediment, food, and overlying water, while the primary route of exposure to oligochaetes in the laboratory is sediment; and by Day 28 or continued to increase during the 56-day exposure. Bioaccumulation of contaminants by indigenous oligochaetes that were recovered on Day 28 from the same chamber with introduced L. variegatus were also evaluated. Peak concentrations of select PAHs and DDT were similar in the indigenous oligochaetes and in L. variegatus exposed in the same chamber (unpublished data). Bioaccumulation of metals from sediments has also been evaluated using L. variegatus. Ankley et al. (1991) reported elevated concentrations of Cd and Ni in worms after 10-day exposures to field-collected sediments where the metal (Cd + Ni):acid-volatile sulfide ratio exceeded 1, but not in samples where the ratio was <1. Ankley et al. (1994) also found that worms did not bioaccumulate metals from three sediments containing elevated concentrations of Cd, Ni, Zn, Cu and Pb, when there was sufficient acid-volatile sulfide to complex metals.

Biota-Sediment Accumulation Factors

Biota-sediment accumulation factors (BSAFs) were calculated for L. variegatus by dividing the lipid-normalized tissue concentrations by the organic carbon-normalized sediment concentrations (Table 2; Brunson et al., 1998). For laboratory-exposed oligochaetes, mean BSAFs ranged from 1.1 for benz(a)anthracene to 5.3 for naphthalene. For field-collected oligochaetes, mean BSAFs ranged from 0.5 for benz(a)anthracene to 8.8 for naphthalene. For individual samples, BSAFs for naphthalene ranged from 1.6 to 10.1 in laboratory-exposed oligochaetes and

Compound	Lee (1992)	Brunson et al. (1998) Lab-exposed oligochaetes	Brunson et al. (1998) Field-collected oligochaetes
Naphthalene	NR	5.3 (1.6-10.1)	8.8 (2.5-26.6)
2-methyl naphthalene	NR	2.6 (0.9-5.1)	6.7 (2.2-12.2)
Pyrene	0.4 (0.18-0.5)	2.3 (0.8-3.9)	2.2 (0.7-5.6)
Fluoranthene	NR	1.8 (0.9-3.9)	1.6 (0.6-4.9)
Chrysene	NR	1.5 (0.7-2.4)	1.1 (0.3-2.0)
Benz(a)anthracene	0.4 (0.2-0.6)	1.1 (0.4-2.5)	0.5 (0.4-0.7)
Benzo(b,k)fluoranthene	0.4 (0.2-1.0)	NR	NR
Perylene	NR	2.24 (0.5-4.7)	1.02 (0.3-1.9)

Table 2. Mean biota-sediment accumulation factors (range in parentheses) reported by Lee (1992) and by Brunson et al. (1998). NR is not reported.

2.5 to 26.6 in field-collected oligochaetes. The BSAFs for pyrene, benz(a)anthracene, and benzo(b,k)fluoranthene were typically greater that BSAFs reported for marine organisms in Lee (1992) for these compounds (Table 2). BSAFs were also calculated using PCB homolog data reported in Ankley et al. (1992) for laboratory-exposed *L. variegatus* and field-collected oligochaetes (Figure 1). BSAFs were similar between laboratory-exposed and field-collected oligochaetes in both Ankley et al. (1992) and Brunson et al. (1998); however, BSAFs reported in Brunson et al. (1998) were typically greater (0.5 to 8.8) than BSAFs from Ankley et al. (1992; 0.17 to 2.26; Figure 1).

A theoretical value of 1.7 for BSAFs has been estimated based on partitioning of nonionic organic compounds between sediment carbon and tissue lipids (ASTM, 1997b). A BSAF of less than 1.7 indicates less partitioning into lipids than predicted, and a value greater than 1.7 indicates more uptake than can be explained by partitioning theory alone (Lee, 1992). The majority of the BSAFs in Figure 1 and Table 2 were within a range of about 0.5 to 2.6, suggesting the theoretical BSAF value of 1.7 could be used to predict these mean BSAFs with a fair amount of certainty. However, mean BSAFs for naphthalene (8.8) and 2-methyl naphthalene (6.7) in the fieldcollected oligochaetes were elevated relative to a theoretical BSAF of 1.7 (Table 2), with BSAFs for individual samples as high as 10.1 for laboratory-exposed oligochaetes and 26.6 for field-collected oligochaetes. The higher BSAFs in the field-collected oligochaetes may be the result of (1) exposure to contaminants in the overlying water; (2) spatial differences in sediment contamination (i.e., sediments were not sampled from a depth representative of the habitat of the oligochaetes); or (3) taxonspecific differences in exposure. BSAFs substantially different from the theoretical value of 1.7 may also result from the system not being at equilibrium (i.e., depletion or release of contaminants in pore water).

In summary, procedures for evaluating the bioaccumulation of contaminants associated with freshwater sediment using the oligochaete *L. variegatus* have been well described. Results of laboratory studies using these procedures are generally similar to the bioaccumulation of contaminants exhibited by oligochaetes in the field. Ongoing research includes further evaluations of bioaccumulation kinetics and field validation of laboratory bioaccumulation methods, use of formulated sediments and sediment spiking, and standardization of micro-lipid analytical methods.

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Objectives:

- Standard methods
- Approaches
- Laboratory exposures
 - -Methods
 - -Lab to Field comparisons
 - -Kinetic studies
- Differences among standard methods
- Future directions

Standard sediment methods					
	ASTM	FPA			

	ASTM	EPA	EC
Toxicity: Fresh water	E1706	1994a	1996a,b
Toxicity: Estuarine & Marine	E1367, E1611	1994b	1992a, 1997a?
Toxicity: Soil	E1676	1986	1994a, 1999?
Bioaccumulation	E1688	1989, 1994a	none
Collection	E1391	1995, 1996?	1994b
Manipulation	E1391	1995, 1996?	1995
Guidance	E1525	1994a,b	1996b, 1997b
Quality Assurance	E1525*	1995	1992b, ISO 9000

Approaches:

- Laboratory-exposed organisms
- Field-collected organisms
- Bioaccumulation factors (BAF)
- Equilibrium partitioning models (BSAF)
- Kinetic models
- Bioenergetic models

Bioaccumulation factor:

BAF = [tissue]/[sediment]

• Equilibrium partitioning models:

Biota-sediment accumulation factor

BSAF = [tissue/lipid]/[sediment/TOC] ~ 1.7 (4.0 USEPA-USCOE; 1991)

Approaches (cont.):

- Assumptions associated with BSAFs:
 - -sediment only source
 - -equilibrium & not kinetically limited
 - -no metabolic degradation
 - -lipid = lipid, TOC = TOC

Selection criteria: Toxicity testing organisms									
	HA	DS	СТ	CR	LV	Π	HS	MO	CL
Sensitivity	+	-	+	-	+	-	-	-	-
Round robin	+	-	+	-	-	-	-	-	-
Contact sed.	+	+	+	+	+	+	+	+	-
Culture	+	-	+	+	+	+	-	-	+
Taxonomy	+	+/-	+/-	+/-	+	+	+	+	+
Ecological	+	+	+	+	+	+	+	+	+
Geographical	+	+/-	+	+	+	+	+	+	+/-
Physico-chem.	+	+	+/-	+	+	+	-	+	NA
Field validation	+	+	+	+	+	+	+	-	+
Peer review	+	+	+	+	+	+	+	-	+/-
Endpoints	SGM	SBA	SGE	BS	BR	SR	SG	В	SGR

Selection Criteria: Freshwater bioaccumulation testing organisms

	LV	Mol	Mdg	Мау	Amp	Cla	Fish
Culture	+	-	+	-	+	+	+
Tissue mass	+/-	+	-	+	-	-	+
Sensitivity	+	+	-	-	-	-	+/-
Feeding	+	+	-	+	-	-	+
Realistic exposure	+	+/-	+	+	+	-	-
Physico-chem.	+	?	+/-	-	+	NA	NA
Field validation	+	?	?	?	+	?	-

LV: Lumbriculus variegatus, Mol: Mollusks, Mdg: Midges, May: Mayflies, Amp: Amphipods, Cla: Cladocerans

Selection Criteria: Recommended freshwater							
bioa	bioaccumulation testing organisms						
Species	Feeding	Biomass	Sensitive	Culture	Data		
Chironomus tentans	FF/ SDF	+	-	++	++		
Chironomus riparius	FF/ SDF	+	-	++	++		
Diporeia spp.*	SSDF	-	+	-	++		
Hexagenia spp.	SDF	+	-	-	++		
Hyalella azteca	SSDF	-	-	++	++		
Lumbriculus variegatus*	SSDF	-	++	++	++		
Earthworms	SSDF	++	?	++	-		

FF = filter feeder; SDF = surface deposit feeder

SSDF = subsurface deposit feeder; Adapted from ASTM E1688

Chemical	Hyalella azteca	Chironomus tentans	Lumbriculus variegatus
Copper	35	54	35
Zinc	73	1125	2984
Nickel	2.8	NT	158
Cadmium	780	NT	12160
Lead	<16	NT	794
p,p'-DDT	0.07	1.23	NT
p,p'-DDD	0.17	0.18	NT
p,p'-DDE	1.39	3.0	>3.3
Dieldrin	7.6	1.1	NT
Chlorpyrifos	0.086	0.07	ΝΤ

Lumbriculus variegatus (oligochaeta):

- Location: North America and Europe
- Habitat: tunnels aerobic sediments lakes, rivers, ponds
- Behavior:
 - buries anterior portion in sediment and undulates posterior end in overlying water for respiration
 - -processes >12 x weight/day

Lumbriculus variegatus (cont.):Adults:

- -40 to 90 mm length
- -1.0 to 1.5 mm diameter
- -5 to 12 mg wet weight
- -about 1% lipid
- Reproduction: asexual (i.e., architomy)
- Culture:
 - -adults of various size
 - population doubles in about 8 to 12 days at 23C

Laboratory exposures:

- Single sampling time:
 - -steady state (i.e., Day 28?)
 - -ANOVA and BSAFs
- Kinetic study:
 - -time course (i.e., Day 1, 3, 7, 14, 28, 56)
 - -regression models (i.e., Ks and K₂)
- Depuration:
 - -experimental (i.e., 24-h gut purge)
 - -regression models (i.e., Ks and K2)

BIOACCUMULATION	<i>Lumbriculus:</i> EPA & ASTM	<i>Macoma</i> : EPA & ASTM	Polychaetes: EPA & ASTM
Temperature (C)	23	NS	10-25
Luminance (lux)	750	750	750
Photoperiod	16:8	16:8 to 12:12	16:8 to 12:12
Chamber (L)	4-6	NS	NS
Sediment (L)	<u>></u> 1.0	NS	NS
Water (L)	<u>></u> 1.0	NS	NS
Water renewal	R	S/R	S/R
Age	adult	adult	juvenile
Loading	>1 g/rep.	1 g/50 g sed.	1 g/200 g sed.
Feeding	No	No	No
Replicates	5	8	8
Duration (days)	28/kinetics	28/kinetics	28/kinetics
Endpoints	В	В	В
Acceptability	T,A,B,R	T,A,B,R	T,A,B,R

Percent of steady state (ASTWIE1000)					
Compound	Day 10	Day 28	Organism		
Phenanthrene	67	95	amphipod		
Benzo(a)pyrene	96	100	mayfly		
Benzo(a)pyrene	43	75	clam		
Benzo(a)pyrene	32	66	amphipod		
Chrysene	43	87	clam		
Hexachlorobenzene	35	70	clam		
Hexachlorobiphenyl	88	100	mayfly		
Aroclor 1242	18	87	polychaete		
Total PCBs	23	87	clam		
Cadmium	17	50	shrimp		

Parcent of steady state (ASTM E1688)

Percent loss during gut purging (ASTM E1688)

Compound	24 h	72 h	Organism
PCB	3	8	shrimp
Hexachlorobenzene	4	12	clam
Benzo(a)pyrene	4	12	amphipod
Phenanthrene	11	33	amphipod
Benzo(a)pyrene	14-26	43-99	mayfly
Phenanthrene	77-100	-	mayfly
HCBP	14-26	43-99	mayfly

Errors associated with gut purging:

- Gut sediment error greatest:
 - -selective ingest high TOC
 - -large gut
 - -early in exposure (low uptake)
 - -cmpds. not bioaccumulated
- Purging error greatest:
 - -rapidly depurated/metabolized cmpds.
 - -dilution by uncontaminated sediment

Performance-based criteria:

- Survival (should; 4-d screening test for LV)
- Avoidance (should)
- Food (should; measure chemicals of concern)
- Water quality (should)
- Culture conditions (should)
- Reference toxicants (must: monthly/start of test)
- Physico-chemical characteristics (should)
- Temperature (must; i.e., consistent life stage)
- Storage sediment (2-8 weeks; no consensus)
- Spiked sediment (1 month holding before testing)

Mean BSAFs for	PCBs	(Lee 1992)
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ORGANISM	BSAF	
Yoldia limatula	10.6	
Nereis virens	10.0	
Macoma nasuta	5.9	
Yoldia limatula	5.7	
Nereis virens	5.2	
Macoma nasuta	3.4	
Nereis virens	3.2	
Nereis virens	1.9	
Macoma nasuta	1.8	
Nereis virens	0.5	
Macoma nasuta	0.4	
Oligochaetes*	0.8/0.9	

Mean BSAFs for other compounds

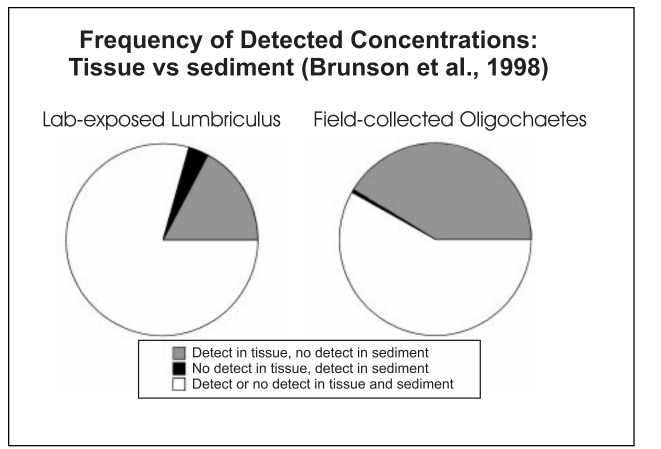
COMPOUND	BSAF1	RANGE	BSAF2	RANGE
Chlordane	4.7	4.0-5.9	-	-
Hexachlorobenzene	3.1	2.1-4.1	-	-
DDD	2.1	0.4-4.8	-	-
DDE	1.3	0.7-2.8	-	-
2,3,7,8-TCDD	0.7	0.5-0.8	-	-
Pyrene	0.4	0.2-0.5	1.1-2.3	0.7-5.6
Benzo(b,k)fluoranthene	0.4	0.2-1.0	0.6-0.8	0.3-1.5
Chrysene	0.4	0.2-0.6	1.0-1.4	0.3-2.4
Benz(a)anthracene	0.4	0.2-0.6	0.5-1.0	0.4-2.5
Benzo(a)pyrene	0.2	0.05-0.9	-	-

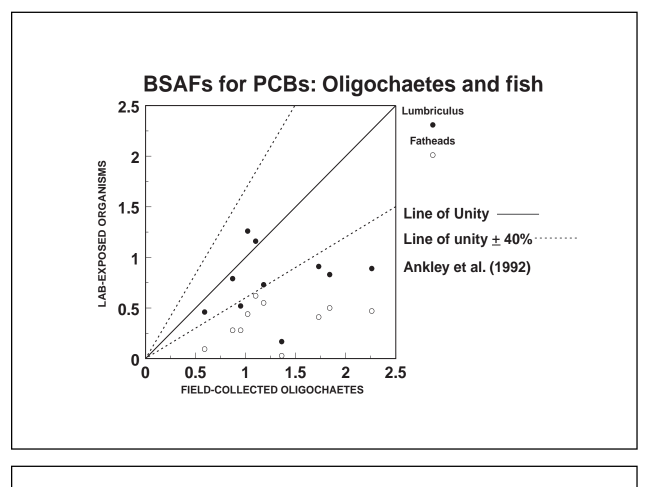
BSAF1 (Lee 1992) and BSAF2 (Brunson et al. 1998)

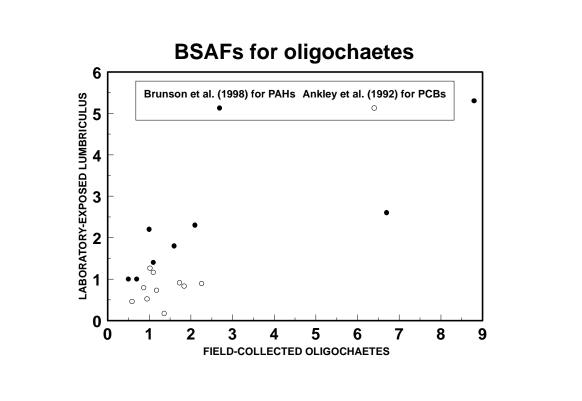
COMPOUND	LAB	RANGE	FIELD	RANGE
2-methylnaphthalene	2.6	0.9-5.1	6.7	2.2-12.2
Benz(a)anthracene	1.0	0.4-2.5	0.5	0.4-0.7
Benzo(b,k)fluoranthene	1.0	0.6-1.5	0.7	0.3-1.5
Chrysene	1.4	0.7-2.4	1.1	0.3-2.0
Fluoranthene	1.8	0.85-3.9	1.6	0.6-4.9
Naphthalene	5.3	1.6-10.1	8.8	2.5-26.6
Perylene	2.2	0.5-4.7	1.0	0.3-1.9
Pyrene	2.3	0.8-3.9	2.1	0.7-5.6

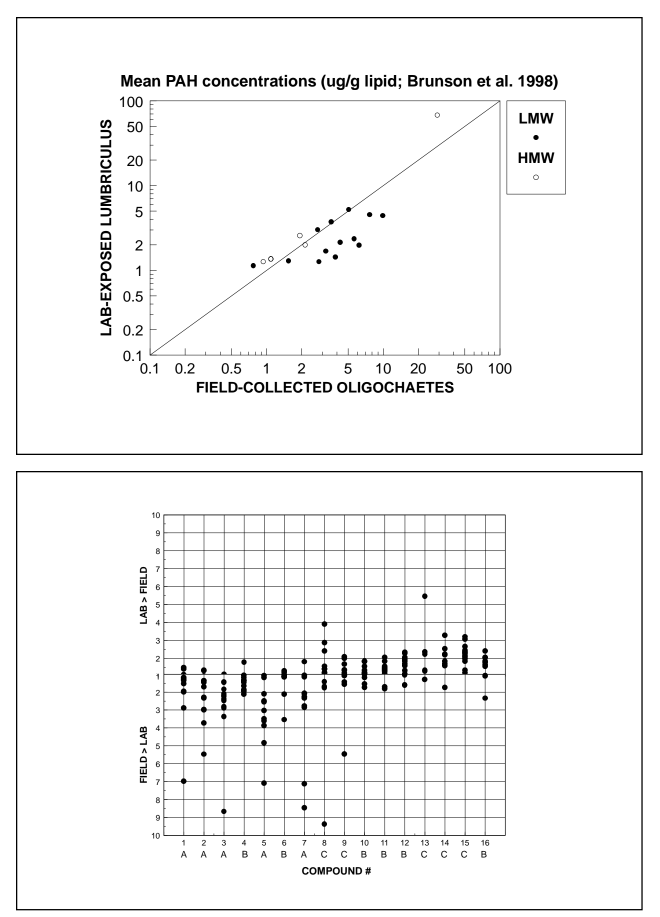
Mean BSAFs for oligochaetes

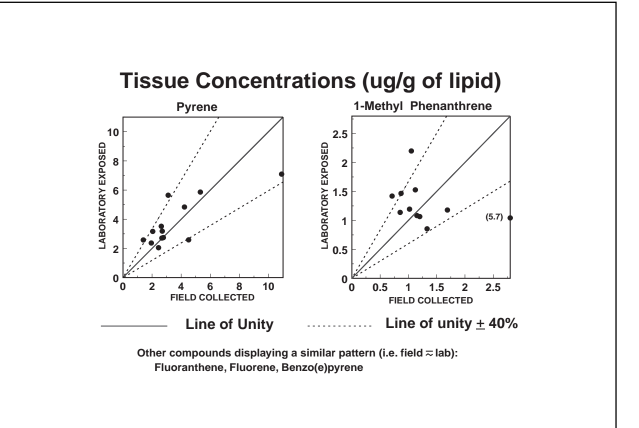
Brunson et al. 1998

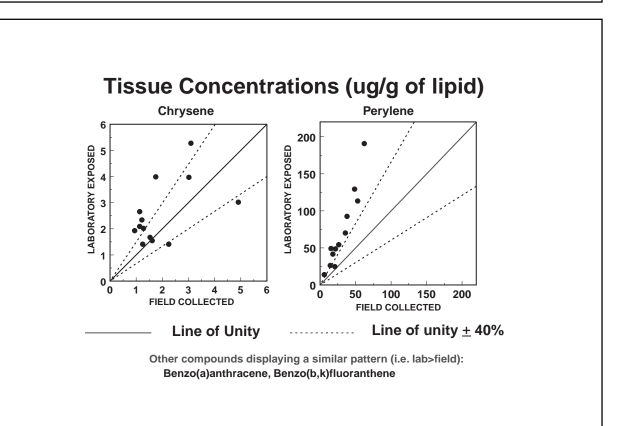


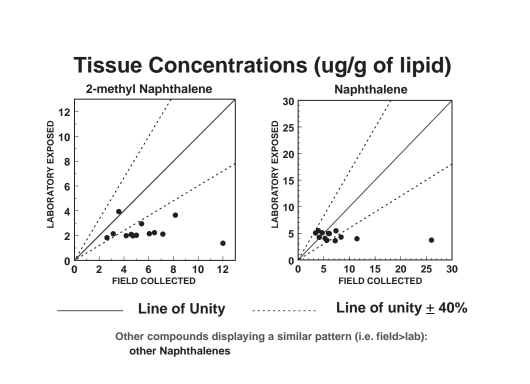












Differences among standard sediment methods:

- Static vs. flow-through (fresh vs. marine)
- Type & quantity of food (toxicity vs. bioaccum.)
- Age (Hyalella and Chironomus: EC vs. EPA)
- Duration & endpoints (Hyalella: EC vs EPA)
- Sieving sediment (EC vs. EPA and ASTM)
- Sediment storage (2 to >8 weeks; consensus?)

Future directions:

- Spiking & formulated sediments
- Quality assurance:
 - -Lab certification (EC)
 - -Reference toxicants
- Standardization of micro-lipid methods
- Kinetics and bioenergetics models
- Field validation

