Prediction of sediment toxicity using consensus-based freshwater sediment quality guidelines

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Christopher G Ingersoll¹, Donald D MacDonald², Ning Wang³, Judy L Crane⁴, L Jay Field⁵, Pam S Haverland¹, Nile E Kemble¹, Rebekka A Lindskoog², Corinne Severn⁶, and Dawn E Smorong²

> ¹Columbia Environmental Research Center, USGS 4200 New Haven Road, Columbia, MO 65201

²MacDonald Environmental Sciences Limited, 2376 Yellow Point Road, Nanaimo, British Columbia, Canada V9X 1W5

> ³Fisheries and Wildlife Sciences, 302 ABRN Building, University of Missouri, Columbia, MO 65211

⁴Minnesota Pollution Control Agency, 520 Lafayette Rd., St. Paul, MN 55155

⁵National Oceanic and Atmospheric Administration (NOAA), 7600 Sand Point Way NE, Seattle, WA 98115

⁶EVS Environment Consultants, 200 West Mercer Street, Seattle, WA 98119

Project Officer: Marc Tuchman USEPA GLNPO 77 West Jackson Chicago, IL 60604

Abstract

The primary objectives of this study were to: (1) evaluate the ability of consensus-based SQGs (sediment quality guidelines) to predict toxicity in a freshwater database for field-collected sediments in the Great Lakes basin; (2) evaluate the ability of SQGs to predict sediment toxicity on a regional geographic basis elsewhere in North America; and (3) compare approaches for evaluating the combined effects of chemical mixtures on the toxicity of field-collected sediments. A database was developed from 92 published reports which included a total of 1657 samples with high-quality matching sediment toxicity and chemistry data. The database was comprised primarily of 10- to 14-day or 28- to 42-day toxicity tests with the amphipod Hyalella azteca (designated as the HA10 or HA28 tests) and 10- to 14-day toxicity tests with the midges Chironomus tentans or C. riparius (designated as the CS10 test). Endpoints reported in these tests were primarily survival or growth. Because field-collected sediments typically contain complex mixtures of contaminants, the predictive ability of a sediment assessment is likely to increase when SQGs are used in combination to classify toxicity of sediments. For this reason, the evaluation of the predictive ability of probable effect concentrations (PECs) was conducted to determine the incidence of effects above and below various mean PEC quotients (mean quotients of 0.1, 0.5, 1.0, and 5.0). The PECs are SOGs that were established as concentrations of individual chemicals above which adverse effects in sediments are expected to frequently occur. A PEC quotient was calculated for each chemical in each sample in the database by dividing the concentration of a chemical by the PEC for that chemical. A mean quotient was calculated for each sample by summing the individual quotient for each chemical and then dividing this sum by the number of PECs evaluated.

When mean quotients were calculated using an approach of equally weighting up to 10 reliable PECs (PECs for metals, total polycyclic aromatic hydrocarbons (PAHs), total polychlorinated biphenyls (PCBs), and sum DDE), there was an overall increase in the incidence of toxicity with an increase in the mean quotient in all three tests. For example in the HA10 test, the toxicity of samples was 20% at mean quotients of <0.1 and increased to 67% at mean quotients of >5.0. Similarly, for the CS10 test there was a 20% incidence of toxicity at mean quotients of <0.1 increasing to a 64% incidence of toxicity at mean quotients of >5.0. In contrast, the incidence of toxicity in the HA28 test was only 8% at mean quotients of <0.1 and increased to 91% at mean quotients of >1.0. In all three tests, there was a consistent increase in the toxicity at mean quotients of >0.5. However, the overall incidence of toxicity was greater in the HA28 test compared to the short-term tests.

The incidence of toxicity at mean quotients of <0.1 was somewhat higher in the HA10 and CS10 tests (20%) compared to the HA28 test (8%). This toxicity at low mean quotients does not appear to be related to total organic carbon in sediment. There was insufficient information in the database to evaluate effects of grain size on toxicity. Unmeasured contaminants in these field-collected sediments or contaminants for which we do not have reliable PECs (i.e., pesticides, herbicides) may have contributed to this toxicity at low mean quotients. Alternatively, the data for HA10 and CS10 tests were obtained from numerous laboratories which may have contributed to variability in the data reported in these studies. In contrast, a limited number of laboratories conducted most of the HA28 tests.

The reason for the higher incidence of toxicity with increasing mean quotients in the HA28 test compared to the short-term tests may be due to the duration of the exposure or the sensitivity of

growth in the longer HA28 test. A 50% incidence of toxicity in the HA28 test corresponds to a mean quotient of 0.63 when survival or growth were used to classify a sample as toxic. By comparison, a 50% incidence of toxicity is expected at a mean quotient of 3.2 when survival alone was used to classify a sample as toxic in the HA28 test. In the CS10 test, a 50% incidence of toxicity is expected at a mean quotient of 9.0 when survival alone was used to classify a sample as toxic or at a mean quotient of 3.5 when survival or growth were used to classify a sample as toxic. In contrast, similar mean quotients resulted in a 50% incidence of toxicity in the HA10 test when survival alone (mean quotient of 4.5) or when survival or growth (mean quotient of 3.4) were used to classify a sample as toxic. Results of these analyses indicate that both the duration of the exposure and the endpoints measured can influence whether a sample is found to be toxic or not. The longer-term tests in which growth and survival are measured tended to be more sensitive than shorter-term tests, with acute to chronic ratios on the order of 6 indicated for *H. azteca*.

We were also interested in determining the predictive ability of PEC quotients for major classes of compounds. Therefore, we evaluated the incidence of toxicity based on a mean quotient for metals, a quotient for total PAHs, or a quotient for total PCBs. Different patterns of toxicity associated with the various procedures for calculating quotients were observed. For example in the HA28 test, a relatively abrupt increase in toxicity was associated with elevated PCBs alone or with elevated PAHs alone, compared to the pattern of a gradual increase in toxicity observed with quotients calculated using a combination of metals, PAHs, and PCBs. These analyses indicate that the different patterns in toxicity may be the result of unique chemical signals associated with individual contaminants. While mean quotients can be used to classify samples as toxic or non-toxic, individual quotients might be useful in helping to identify substances that may be causing or substantially contributing to the observed toxicity.

We chose to make comparisons across geographic areas using mean quotients calculated by equally weighting the contribution of the three major classes of compounds (metals, or PAHs, or PCBs). This approach assumes that these three diverse groups of chemicals exert some form of joint toxic action. Use of this approach also maximized the number of samples that were used to make comparisons across geographic areas. Generally, there was an increase in the incidence of toxicity with increasing mean PEC quotients within most of the regions, basins, and areas for all three toxicity tests. For the HA10 and HA28 tests, the incidence of toxicity for samples from each of the Great Lakes and within the areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database. However, the relationship between the incidence in toxicity and mean quotients in the CS10 test was more variable among geographic areas compared to either the HA10 or HA28 test. The results of these analyses indicate that the consensus-based PECs can be used to reliably predict toxicity of sediments on both a regional and national basis.

This paper presents results of the first analyses completed on the entire freshwater sediment database. Some of the additional analyses planned for the database that are beyond the scope of this paper include: (1) comparing approaches for designating samples as toxic; (2) evaluating logistic-regression models; (3) identifying a list of optimal analytes for broad scale application and testing the relative efficacy of the mean versus the sum PEC quotients; (4) evaluating the influence of grain size and ammonia on the incidence of toxicity; and (5) developing a guidance manual for conducting an integrated assessment of sediment contamination.

Contents

Abstract	2
List of Figures	4
List of Tables	
List of Appendices	4
Acknowledgments	
Introduction	
Methods	7
Results and Discussion	
Summary	
References	
Figures	
Tables	
Appendices	
11	

List of Figures

- Figure 1. Relationship between the geometric mean of the mean PEC quotients and the incidence of toxicity in the three tests, based on survival or growth.
- Figure 2. Relationship between the geometric mean of the mean PEC quotients and the incidence of toxicity in the three tests, based on survival or growth, or based on survival alone.

List of Tables

- Table 1.
 Sediment quality guidelines that reflect probable effect concentrations.
- Table 2.Incidence of sediment toxicity within ranges of PEC quotients (calculated using
various approaches) for freshwater tests based on survival or growth.
- Table 3.Incidence of sediment toxicity within ranges of PEC quotients for the Hyalella
azteca 10- to 14-day test in various geographic areas within the database based on
survival or growth.
- Table 4.Incidence of sediment toxicity within ranges of PEC quotients for the Hyalella
azteca 28- to 42-day test in various geographic areas within the database based on
survival or growth.
- Table 5.Incidence of sediment toxicity within ranges of PEC quotients for the *Chironomus*
spp. 10- to 14-day test in various geographic areas within the database based on
survival or growth.

List of Appendices

Appendix 1. Summary of the data presented in Figures 1 and 2.

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Introduction

Numerical sediment quality guidelines (SQGs) have been developed by a variety of federal, state, and provincial agencies across North America using matching sediment chemistry and biological effects data. These SQGs have been routinely used to interpret historical data, identify potential problem chemicals or areas at a site, design monitoring programs, classify hot spots and rank sites, and make decisions for more detailed studies (Long and MacDonald 1998). Additional suggested uses for SQGs include identifying the need for source controls of problem chemicals before release, linking chemical sources to sediment contamination, triggering regulatory action, and establishing target remediation objectives (USEPA 1997). Numerical SQGs, when used with other tools such as sediment toxicity tests, bioaccumulation, and benthic community surveys, can provide a powerful weight of evidence for assessing the hazards associated with contaminated sediments (Ingersoll et al. 1997).

A critical component in the application of SQGs for assessing sediment quality is a demonstration of the ability of the guidelines to predict the absence or presence of toxicity in field-collected sediments (Ingersoll et al. 1996, Smith et al. 1996, Long et al. 1998a, Swartz 1999, Fairey et al. 2000, MacDonald et al. 2000a,b). This paper is the fourth in a series that is intended to address the ability of various SQGs to predict toxicity in contaminated sediments. The first paper in the series focused on resolving the "mixture paradox" that is associated with the application of empirically-derived SQGs for individual polycyclic aromatic hydrocarbons (PAHs). In this case, the paradox was addressed by developing consensus-based SQGs for total polychlorinated biphenyls (PCBs) to address a similar mixture paradox for that group of contaminants (MacDonald et al. 2000b).

A third paper developed consensus-based SQGs for freshwater sediments (MacDonald et al. 2000a). The published SQGs for 28 chemical substances were assembled and classified into two categories in accordance with their original narrative intent. These published SQGs were then used to develop two consensus-based SQGs for each contaminant, including a threshold effect concentration (TEC; below which adverse effects are not expected to occur) and a probable effect concentration (PEC; above which adverse effects are expected to frequently occur; Table 1; MacDonald et al. 2000a). A preliminary evaluation of the predictive ability of these consensus-based SQGs for freshwater sediment was conducted using a database of 347 samples obtained from 15 separate studies. The results of these three previous investigations demonstrated that the consensus-based SQGs provide a unifying synthesis of the existing guidelines, reflect causal rather than correlative effects, and account for the effects of contaminant mixtures in sediment (Swartz 1999, MacDonald et al. 2000a,b).

The primary objective of this fourth paper is to further evaluate the predictive ability of the consensus-based PECs developed by MacDonald et al. (2000a). The database used by MacDonald et al. (2000a) was expanded to include a total of 1657 samples from 92 published reports with high-quality matching sediment toxicity and chemistry data. The majority of the data from these reports were for 10- to 28-day toxicity tests with the amphipod *Hyalella azteca* or 10- to 14-day toxicity tests with the midges *Chironomus tentans* or *C. riparius*. The endpoints

measured in these toxicity tests primarily included survival or growth of test organisms at the end of the sediment exposures. A second objective of this paper is to evaluate the predictive ability of these PECs on a regional basis within the larger database. We were interested in determining if there are differences in the predictive ability of the PECs across the entire database compared to various geographic areas within the database, such as all of the samples from the Great Lakes or all of the samples from an area within a Great Lake, such as Indiana Harbor or Waukegan Harbor located within Lake Michigan.

Methods

Development of consensus-based sediment quality guidelines

Individual SQGs for freshwater ecosystems have previously been developed using a variety of approaches (Table 1). Each of these approaches has certain advantages and limitations which influence their application in the sediment quality assessment process (Ingersoll et al. 1997). In an effort to focus on the agreement among these various published SQGs, consensus-based SQGs were developed by MacDonald et al. (2000a) for 28 chemicals of concern in freshwater sediments (i.e., metals, PAHs, PCBs, and pesticides). For each contaminant of concern, two consensus-based SQGs were developed from published SQGs, including a threshold effect concentration (TEC) and a probable effect concentration (PEC). The TECs were calculated by determining the geometric mean of the SQGs that were included in this category (MacDonald et al. 2000a). Likewise, consensus-based PECs were calculated by determining the geometric mean of the PEC-type values (Table 1). The geometric mean, rather than the arithmetic mean or median, was calculated because it provides an estimate of central tendency that is not unduly affected by extreme values and because the distributions of the SQGs were not known (MacDonald et al. 2000a). Consensus-based TECs or PECs were calculated only if three of more published SQGs were available for a chemical substance or group of substances. The evaluations of toxicity in the present study were based on the use of PECs because TECs were developed to provide an estimate of conditions where toxicity would not be expected and PECs were developed to provide an estimate of conditions where toxicity would be expected. Evaluations of SQGs in the present study were based on dry-weight concentrations because previous studies have demonstrated that normalization of SQGs for PAHs or PCBs to total organic carbon (Barrick et al. 1988, Long et al. 1995, Ingersoll et al. 1996) or normalization of metals to acidvolatile sulfides (Long et al. 1998b) did not improve the predictions of toxicity in field-collected sediments.

The consensus-based PECs listed in Table 1 were critically evaluated by MacDonald et al. (2000a) to determine if they would provide effective tools for assessing sediment quality conditions in freshwater ecosystems. The criteria for evaluating the reliability of the consensus-based PECs were adapted from Long et al. (1998a). Specifically, the individual TECs were considered to provide a reliable basis for assessing the quality of freshwater sediments if >75% of the sediment samples were correctly predicted to be not toxic. Similarly, the individual PEC for each substance was considered to be reliable if >75% of the sediment samples were correctly predicted to be toxic using the PEC. Therefore, the target levels of both false positives (i.e., samples incorrectly classified as toxic) and false negatives (i.e., samples incorrectly classified as

not toxic) was 25% using the TEC and PEC. To assure that the results of this evaluation were not unduly influenced by the number of sediment samples, the various SQGs were considered to be reliable only if a minimum of 20 samples were included in the evaluation (i.e., 20 samples above at a PEC or 20 samples below a TEC; CCME 1995). The results of this evaluation described in MacDonald et al. (2000a) indicated that most of the TECs (i.e., 21 of 28) provide an accurate basis for predicting the absence of sediment toxicity. Similarly, most of the PECs (i.e., 16 of 28) provide an accurate basis for predicting sediment toxicity (Table 1).

A reliable TEC or PEC was not available for mercury, an important contaminant of concern in sediments (Table 1). This lack of reliability is most likely due to the speciation of mercury in the sediments, as well as the ability of methyl mercury to bioaccumulate in organisms. Sediment quality guidelines developed using a tissue residue approach are needed to establish safe sediment concentrations for human health and piscivorus-wildlife receptors (Ingersoll et al. 1997). For this report, only direct toxic effects on benthic invertebrates are considered in the evaluation of the predictability of the consensus-based SQGs.

Development of the sediment toxicity database

To support the development of the sediment toxicity database, matching sediment chemistry and biological effects data were compiled for various freshwater locations across North America (in addition to the data that were used in the analyses performed by MacDonald et al. 2000a). Candidate data sets were identified by reviewing the published literature and by contacting individuals active in the field of sediment quality assessment. More than 1500 documents were reviewed and evaluated to obtain the data required to evaluate SQGs in the present study. Because these data sets were generated for a wide variety of purposes, each study was critically-evaluated to assure the quality of the data used for evaluating the predictive ability of the SQGs (Long et al. 1998a, Ingersoll and MacDonald 1999). Data from individual studies were considered acceptable for use in the present study if:

- The study was conducted in a freshwater location in North America;
- Appropriate procedures were used to collect, handle, and store sediments (e.g., ASTM 2000, USEPA 2000);
- Matching sediment chemistry and biological effects data were reported and concentrations of contaminants were measured in each sample or treatment group.
- Minimum data quality requirements were reported. For example, analytical detection limits were lower than freshwater probable effect levels (PEL; Smith et al. 1996), accuracy and precision were within acceptable limits, and analytes were not present at detectable levels in method blanks;
- Appropriate analytical methods were used to generate chemistry data. For metals, concentrations of total metals needed to be reported. However, other measures of metal concentrations were used (i.e., simultaneously extracted metals) if sufficient information was available to demonstrate that these measures are comparable to total metal concentrations (Ingersoll et al. 1996, 1998). For organic compounds, the concentrations needed to be measured using gas chromatography-mass spectroscopy, high pressure liquid chromatography, or comparable methods; and,

• Toxicity tests needed to meet test acceptability criteria outlined in ASTM (2000) and USEPA (2000) and endpoints measured needed to be ecologically relevant (likely to influence the viability of the organism in the field) or needed to be indicative of ecologically-relevant endpoints.

Using these selection criteria, a total of 92 freshwater data sets were incorporated into a database that included 1657 individual sediment samples. The toxicity tests in this database primarily include tests with the amphipod, Hyalella azteca; the midges, Chironomus tentans or C. riparius; the mayfly, Hexagenia limbata; the oligochaete, Lumbriculus variegatus; the daphnids, Ceriodaphnia dubia or Daphnia magna; and, the bacterium, Vibrio fisheri (Microtox). We selected a subset of the samples from this database that reported sediment chemistry for at least one of the substances for which reliable SQGs were listed in Table 1. We then selected a subset of these resultant samples with matching toxicity data for the amphipod Hyalella azteca or the midges Chironomus tentans or C. riparius. Tests with H. azteca and Chironomus spp. were selected because the samples that were tested represented a broader geographic area compared to the other tests in the database. The selected studies provided 670 samples for H. azteca 10- to 14day tests (designated as HA10), 160 samples for H. azteca 28- to 42-day tests (designated as HA28) and 632 samples for Chironomus spp. 10- to 14-day tests (CS10; 556 of the samples were for tests with C. tentans and 76 of the samples were for tests with C. riparius). We combined the data for the two midge species due to the limited amount of data for C. riparius. Preliminary analyses of the database indicated similar sensitivity for these two species of midge. The selected studies represented a broad range in both sediment toxicity and contamination. A total of 28% of the samples were toxic in the HA10 test, 35% of the samples were toxic in the HA28 test, and 27% of the samples were toxic in the CS10 test (28% of the samples were toxic in the C. tentans tests and 21% of the samples were toxic in the C. riparius tests). Toxicity of samples was determined as a significant reduction in survival or growth relative to a control or reference sediment (as designated in the original study or determined using appropriate statistical procedures). Sexual maturation and reproduction were periodically reported in the HA28 test; however, these two additional endpoints did not identify any additional samples as toxic relative to effects reported on survival or growth of amphipods.

The total PCB concentration in each sediment sample in the database was calculated by summing dry-weight concentrations of individual congeners. If only aroclors concentrations were reported, total PCBs were calculated as the sum concentration of all individual aroclors. If both congeners and aroclors were reported, the congeners were used to calculate the concentration total PCBs in a sample. If only total PCBs was reported for a sample, then this value was used. The total PAH concentration in each sediment sample was generally calculated by summing the dry-weight concentrations from as many of the following 13 compounds that were reported: acenaphthene, acenaphthylene, anthracene, fluorene, 2-methylnaphthalene, naphthalene, phenanthrene, benz(a)anthracene, dibenz(a,h)anthracene, benzo(a)pyrene, chrysene, fluoranthene, and pyrene. Total PAHs were calculated using eight or less of these individual PAHs for <5% of the samples in the database. In calculating total PCBs or total PAHs, half the detection limit was used for concentrations of metals below the detection limit. For DDTs, the concentrations of p,p'-DDD and o,p'-DDD, p,p'-DDE and o,p'-DDE, and p,p'-DDT and o,p'-DDT were summed to calculate

the concentrations of sum DDD, sum DDE, and, sum DDT, respectively. Total DDT was calculated by summing the concentrations of sum DDD, sum DDE, and, sum DDT. If all individual PCBs, PAHs, DDD, DDE, or DDT were less that the detection limit, the detection limits were summed and reported as a less than value for the sum.

Analysis of data

The initial evaluation of predictive ability by MacDonald et al. (2000a) focused primarily on determining the ability of each SQG, when applied alone, to correctly classify samples as toxic or not toxic. Because field-collected sediments typically contain complex mixtures of contaminants, the predictive ability of these sediment quality assessment tools is likely to increase when the SQGs are used in combination to classify toxicity of sediments. For this reason, the evaluation of the predictive ability of the SQGs in the present study was conducted to determine the incidence of effects above and below various mean PEC quotients (mean quotients of 0.1, 0.5, 1.0, and 5.0; Ingersoll et al. 1998, Long et al. 1998a, Fairey et al. 2000).

A PEC quotient was calculated for each chemical in each sample in the database by dividing the concentration of a chemical by the PEC for that chemical. A mean quotient was then calculated for each sample by summing the individual quotient for each chemical and dividing this sum by the number of reliable PECs evaluated. MacDonald et al. (2000a) found that some PEC values were more reliable predictors of toxicity and that use of only these PECs reduced the variability in the prediction of sediment toxicity compared to using all available PECs. The PEC for total PAHs, instead of the PECs for the individual PAHs, was used in the calculation to avoid double accounting of the PAH data (MacDonald et al. 2000a). This resulted in the use of up to 10 reliable PECs in calculating the mean quotient (arsenic, cadmium, chromium, copper, lead, nickel, zinc, total PAHs, total PCBs, and sum DDE; Table1 and designated "Mean - all" in Table 2). This approach to the calculation of mean quotients weighs each of the chemicals and chemical classes equally (Ingersoll et al. 1998, Long et al. 1998a, MacDonald et al. 2000a).

In the present study, a second approach was also used to calculate mean PEC quotients. We were interested in equally weighting the contribution of metals, PAHs, and PCBs in the evaluation of sediment chemistry and toxicity (assuming these three diverse groups of chemicals exert some form of joint toxic action). For this reason, we first calculated an average PEC quotient for up to seven metals in a sample. A mean quotient was then calculated for each sample by summing the average quotient for metals, the quotient for total PAHs, and the quotient for total PCBs, and then dividing this sum by three $(n = 3 \text{ quotients/sample}; \text{designated "Mean - MPP (and)" in Table$ 2). Another approach for evaluating mean quotients was to calculate the mean of the average quotient for metals, the quotient for total PAHs, or the quotient for total PCBs (n = 1 to 3 quotients/sample; designated "Mean - MPP (or)" in Table 2). Hence, the "Mean - MPP (or)" approach uses any or all of the three classes of contaminants as available. For example, if a sample only had a measure of total PAHs and total PCBs, then the mean quotient would be calculated using just the quotients for these two classes of compounds. In contrast, the "Mean -MPP (and)" approach only uses samples with measures of metals, total PAHs, and total PCBs. Sum DDE was not included in these calculations of the mean quotient because there were a limited number of samples in the database with elevated concentrations of DDE and we were

interested in equally weighting the contributions of the major groups of contaminants in the database (metals, total PAHs, and total PCBs). Therefore, the differences in this "MPP approach" from the approach used by MacDonald et al. (2000a) are: (1) an average quotient for metals was used instead of the individual quotients for metals and (2) sum DDE was not used in the calculation.

Results and Discussion

Evaluation of approaches for calculation of mean PEC quotients

The results of the evaluations of the incidence of sediment toxicity within ranges of mean PEC quotients in the entire database for each of the three tests (HA10, HA28, CS10) are summarized in Table 2. When mean quotients were calculated using the approach of weighting equally up to 10 reliable PECs (designated "Mean - all" in Table 2), there was an increase in the incidence of toxicity with an increase in the mean quotient in all three tests. For the HA10 test, the incidence of toxicity was 20% at mean quotients of <0.1 and increased to 67% at mean quotients of >5.0. Similarly, for the CS10 test there was a 20% incidence of toxicity at mean quotients of <0.1 increasing to a 64% incidence of toxicity at mean quotients of <0.1 and increased to 91% at mean quotients of >1.0 (the incidence of toxicity at a quotient >5.0 was not calculated for the HA28 test due to a limited number of samples above a quotient of 5.0; Table 2).

Long et al. (1998a) conducted a similar analysis of the incidence of toxicity in sediment tests using a database developed for 10-day marine amphipod tests (n=1068). The incidence of toxicity was only 12% at mean quotients of <0.1 (quotients calculated using either marine effect range median (ERM) or probable effect level (PEL) guidelines; Long et al. 1998a). In the present study, the incidence of toxicity at mean quotients of <0.1 was somewhat higher in the HA10 and CS10 tests (20%) compared to the tests with marine amphipods (12%; Long et al. 1998a) or the HA28 test (8%; Table 2).

The reason for this higher incidence of toxicity at mean quotients of <0.1 in the HA10 and CS10 tests is not clear. This toxicity at low mean quotients does not appear to be related to total organic carbon in sediment. There was insufficient information in the database to evaluate effects of grain size on toxicity. USEPA (2000) and ASTM (2000) reported that amphipods and midges are relatively intolerant to effects of sediment grain size. Unmeasured contaminants in these field-collected sediments or contaminants for which we do not have reliable PECs (i.e., pesticides, herbicides) may have contributed to this toxicity at low mean quotients (see discussion of Figures 1 and 2 below). Alternatively, the data for HA10 and CS10 tests were obtained from numerous laboratories which may have contributed to variability in the data reported in these studies. In contrast, a limited number of laboratories conducted most of the toxicity tests for the marine amphipod or HA28 tests.

In all three tests, there was a consistent increase in the toxicity at mean quotients of >0.5 (designated "Mean - all" in Table 2). However, the overall incidence of toxicity was greater in the HA28 test (91% toxicity at mean quotients of >1.0) compared to the short-term tests (57%

toxicity at mean quotients of >1.0 and 67% toxicity at mean quotients of >5.0 in the HA10 test and 51% toxicity at mean quotients of >1.0 and 64% toxicity at mean quotients of >5.0 in the CS10 test). Similarly, Long et al. (1998a) reported a 56 to 71% incidence of toxicity at mean quotients of >1.0 in the 10-day sediment tests with marine amphipods. The reason for the higher incidence of toxicity in the HA28 test compared to the short-term tests may be due to the duration of the exposure or the sensitivity of growth in the longer HA28 test (see discussion of Figures 1 and 2 below). However, comparisons of the sensitivity between these tests needs to be made with some caution. There were very few samples in the database where tests were conducted using splits of the same samples. Therefore, the differences observed in the responses of organisms may also be due to differences in the types of sediments evaluated in the individual databases for each test.

We were also interested in determining the predictive ability of PEC quotients for major classes of compounds. Therefore, we evaluated the incidence of toxicity based on an average quotient for metals, a quotient for total PAHs, or a quotient for total PCBs (second, third, and fourth rows for each toxicity test listed in Table 2). For the HA10 test, the incidence of toxicity across quotients of <0.1 to >1.0, based on metals alone, total PAHs alone, or total PCBs alone were similar to the incidence of toxicity that was calculated for the mean quotient using up to10 PECs (designated "Mean - all" in Table 2). The incidence of toxicity in the HA10 test was somewhat higher at quotients >5.0 for total PAHs (80%) or total PCBs (73%) compared to metals alone (62%).

For the CS10 test, the incidence of toxicity was also similar across quotients of <0.1 to >5.0, calculated based on metals alone compared to a mean quotient calculated using up to 10 PECs (designated "Mean - all" in Table 2). However, the incidence of toxicity at a total PCB quotient <0.1 was 46%, suggesting that other compounds may be contributing to toxicity at low concentrations of PCBs. The incidence of toxicity in the CS10 test was somewhat higher for quotients based solely on total PAHs compared to quotients based on metals alone, total PCBs alone, or mean quotients calculated using up to 10 PECs. These analyses suggest that the CS10 test may be more sensitive to PAHs compared to the other chemical classes.

For the HA28 test, the incidence of toxicity was similar across the quotients of <0.1 to >1.0, calculated based on metals alone compared to a mean quotient calculated using up to 10 PECs (designated "Mean - all" in Table 2). However, the incidence of toxicity at quotient of 0.1 to <0.5 was higher for PAHs (61%) compared to the other three quotients (6 to 20% toxicity for metals alone, total PCBs alone, or mean quotients based on up to 10 PECs). The incidence of toxicity at PCB quotients of <1.0 was lower (4 to 17%), while the incidence of toxicity at a PCB quotient of >1.0 was higher (97%) compared to the other three quotients. Results of these analyses indicate a relatively abrupt increase in toxicity associated with elevated PCBs alone or elevated PAHs alone compared to the pattern of a gradual increase toxicity observed with quotients calculated using the up to 10 PECs (designated "Mean - all" in Table 2). These results suggest that *H. azteca* may be more sensitive to PAHs and PCBs in longer-term tests than it is to metals.

In the next analysis, we were interested evaluating the incidence in toxicity by equally weighting

the combined influence of metals, PAHs, and PCBs in a sample. A mean PEC quotient was calculated for each sample by summing the average quotient for metals, the quotient for total PAHs, and the quotient for total PCBs, and then dividing this sum by three (designated as "Mean - MPP (and)" in Table 2). This calculation was done for only those samples with reported concentrations of metals, total PAHs, and total PCBs (266 of 670 samples in the HA10 test, 109 of 160 samples in the HA28 test, and 177 of 632 samples in the CS10 test). Results of this analysis indicate a higher incidence of toxicity in the HA10 (66%), HA28 (100%), and CS10 (60%) tests at mean quotients >1.0 based on an equal weighting of metals, total PAHs, and total PCBs compared to a mean quotient calculated using up to 10 PECs (57, 91, and 51%, respectively; designated "Mean- all" in Table 2).

The different patterns of toxicity associated with these various procedures for calculating quotients may be the result of unique chemical signals associated with individual contaminants in each sample. For example, there was a higher incidence in toxicity with quotients calculated using total PCBs alone compared to quotients calculated using metals alone in the HA10 test. Alternatively, these different patterns may also be influenced by the total number of samples used to make these comparisons. For example, there were 670 total samples for the HA10 test. Of these 670 samples, 623 had metal chemistry data, 488 had measured concentrations of total PAHs, 326 had measured concentrations of total PCBs, and all three of these classes of compounds were measured in only 266 samples. In order to determine the influence of sample number on the observed incidence of toxicity, we first selected the same samples used in the analysis described in the previous paragraph (samples in which metals, total PAHs, and total PCBs were all measured). Quotients were then calculated for these subsets of samples: (1) using up to 10 PECs (designated "Mean - all (select1)"), (2) using up to 9 PECs (not including DDE; designated "Mean - all (select2)", or (3) using metals alone ("total PCBs (select2)"; Table 2).

Results of these analyses indicate that the incidence of toxicity for all three tests was similar in these subsets of samples using the three procedures to calculate mean quotients ("Mean - MPP (and)" versus "Mean - all (select1)" versus "Mean - all (select2)"). In the HA10 test, the incidence of toxicity in the subset of samples (n=266) was similar based on an average quotient for metals alone, a quotient for total PAHs alone, or a quotient for total PCBs alone compared to the "Mean - all (select2)". In contrast, there were different patterns of toxicity associated with individual classes of compounds in the subsets of samples in the HA28 (n=109) or CS10 (n=177) tests where metals, PAHs, and PCBs were all reported. In the HA28 test, a relatively abrupt increase in toxicity was associated with elevated PCBs alone or with elevated PAHs alone, compared to the pattern of a gradual increase toxicity observed with quotients calculated using the "Mean - all (select2)". Similarly in the CS10 test, a increase in toxicity was observed at lower quotients of PAHs alone or metals alone compared to the pattern of a gradual increase toxicity observed to the pattern of a gradual increase toxicity observed with quotients calculated using the "Mean - all (select2)".

These analyses indicate that the different patterns in toxicity may be the result of unique chemical signals associated with individual contaminants in samples rather than the result of a limited number of samples used to make these comparisons. However, we could only make these comparisons with a limited number of samples where metals, PAHs, and PCBs were all reported.

Fairey et al. (2000) conducted a similar analysis of a larger marine database for amphipods and found that the number and type of SQGs used in the calculation of a mean quotient influenced the predictions of sediment toxicity to amphipods. The incidence of toxicity to amphipods increased with increasing numbers of contaminants included in the derivation of the mean quotient (Fairey et al. 2000). While mean quotients can be used to classify samples as toxic or non-toxic, individual quotients might be useful in helping to identify substances that may be causing or substantially contributing to the observed toxicity (MacDonald et al. 2000b).

To use all samples in the database and equally weight the contribution of metals, PAHs, and PCBs, a final analysis was conducted where the mean quotient was calculated as the average of the three major classes available in a sample. For example, if a sample only had a measure of total PAHs and total PCBs, then the mean quotient would be calculated using just the quotients for these two classes of compounds (designated as "Mean – MPP (or)" in Table 2). Results of these analyses indicate that the incidence of toxicity in all three tests was similar when either "Mean - all" (based on up to 10 PECs) or "Mean - MPP (or)" were used to calculate the mean quotients (Table 2).

Evaluation of exposure duration and endpoints measured in toxicity tests

In Figures 1 and 2, we evaluated the relationship between mean PEC quotients and the incidence of toxicity as a function of the duration of the exposure or of the endpoints measured in the toxicity tests. In this analysis, a mean quotient for each sample was calculated using the "Mean - MPP (or)" approach. The samples within each test were ranked in ascending order by mean quotient. The incidence of toxicity and geometric mean of the mean quotients within groups of 20 samples for the HA10 and CS10 tests or within groups of 10 samples for the HA28 test was then plotted (Figures 1 and 2, Appendix 1). The geometric mean, rather than the arithmetic mean or median of the quotients, was calculated because it provides an estimate of central tendency that is not unduly affected by extreme values and because the distributions of the mean quotients were not known.

In Figure 1, samples were classified as toxic based on an adverse effect on survival or growth in the three tests. Results of these analyses plotted in Figure 1 are consistent with the analyses presented in Table 2. Importantly, the incidence of toxicity increases with increasing level of contamination in all three tests. This increase was particularly pronounced at mean quotients of >0.5 in all three tests. There was a slightly elevated incidence of toxicity at the very lowest mean quotient in all three tests. Long et al. (1998a) also observed an elevated incidence of toxicity with marine amphipods at low mean quotients. Long et al. (1998a) suggested that these samples were sometimes fine-grained sediments with low concentrations of organic carbon and detectable concentrations of butyltins, chlorinated pesticides, alkyl-substituted PAHs, ammonia, or other substances not accounted for with the SQGs. In the present study, the incidence of toxicity at low mean quotients did not appear to be related to total organic carbon in sediment. There was insufficient information in the database to evaluate effects of grain size on toxicity. USEPA (2000) and ASTM (2000) reported that amphipods and midges were relatively intolerant to effects of sediment grain size.

We also evaluated relationships between the toxicity and mean quotients calculated using up to 10 reliable PECs or mean quotients calculated using all of the available PECs in Table 1 regardless of their reliability (plots not included in this paper). An increase in toxicity was observed with increasing contamination using either 10 reliable PECs or all of the PECs to calculate mean quotients. However, the variability was higher when all of the PECs listed in Table 1 were used in this analysis. Therefore, the use of reliable PECs improved the relationship between mean PEC quotients and the incidence of toxicity.

In Figure 2, samples in the three tests were classified as toxic based on an adverse effect on survival alone or based on an adverse effect on survival or growth. The relationship between the incidence of toxicity and the geometric mean of the mean quotients was best described by a three parameter logistic model (SigmaPlot 1997; Figure 2; see Appendix 1 for the equations and coefficients). The best fit of the data was observed in the HA28 test ($r^2 = 0.79$ based on survival or 0.93 based on survival or growth) relative to the HA10 test ($r^2 = 0.73$ based on survival or 0.78 based on survival or CS10 test ($r^2 = 0.56$ based on survival or 0.76 based on survival or growth; Figure 2). In the HA10 test, the relationship between toxicity and mean quotient was similar when either survival alone or survival or growth together were used to classify a sample as toxic. However, in the HA28 and CS10 tests, the relationship between the incidence of toxicity and mean quotient was different when survival or growth were used to classify a sample as toxic compared to survival alone (Figure 2).

The incidence of toxicity in the HA28 and CS10 tests based on survival or growth was often double the incidence of toxicity based on survival alone at mean quotients of >0.3. A 50% incidence of toxicity in the HA28 test corresponds to a mean quotient of 0.63 when survival or growth were used to classify a sample as toxic (Figure 2, Appendix 1). By comparison, a 50% incidence of toxicity was estimated at a mean quotient of 3.2 when survival alone was used to classify a sample as toxic in the HA28 test. In the CS10 test, a 50% incidence of toxicity was estimated at a mean quotient of 9.0 when survival alone was used to classify a sample as toxic or at a mean quotient of 3.5 when survival or growth were used to classify a sample as toxic. In contrast, similar mean quotients resulted in a 50% incidence of toxicity in the HA10 test when survival alone (mean quotient of 4.5) or when survival or growth (mean quotient of 3.4) were used to classify a sample as toxic.

Results of these analyses indicate that both the duration of the exposure and the endpoint measured can influence whether a sample is found to be toxic or not. Again, comparisons of the sensitivity between these tests needs to be made with some caution. There were very few samples in the freshwater database where tests were conducted using splits of the same samples. Therefore, the differences observed in the responses of organisms may also be due to differences in the types of sediments evaluated in the individual databases for each test. Nevertheless, it appears that longer-term tests in which survival and growth are measured tend to be more sensitive than short-term tests, with acute to chronic ratios on the order of 6 indicated for *H. azteca*. Similar differences in sensitivity of *H. azteca* have been observed in 10- and 42-day water-only exposures to cadmium, fluoranthene or DDD (unpublished data).

Evaluation of the predictive ability of mean PEC quotients across various geographical areas with the database

A primary objective of this paper was to determine if there were differences in the predictive ability of PEC quotients within geographic areas compared to the entire database. We chose to make these comparisons of geographic areas using the "MPP (or)" mean quotients calculated as the mean of the average quotient for metals, the quotient for total PAHs, or the quotient for total PCBs (Table 2). Use of this approach maximized the number of samples used in the evaluation and equally weighted the contribution of the three major classes of compounds (metals, or PAHs, or PCBs) to the observed incidence of toxicity. The relationship between the incidence of toxicity and mean quotients is presented in Tables 3 to 5 for the entire database and for various geographical areas represented within the database. Designation of region, basin, and area for each geographic area in Tables 3 to 5 is based on information obtained from the original report or by contacting the authors of the report. Comparisons of toxicity among the entire geographic areas listed in Tables 3 to 5 should be done with caution given the limited number of samples from each area (i.e., only 5 samples for the HA10 test from the entire Buffalo River). Samples were grouped by geographic areas in Table 3 to 5 to determine how well toxicity and mean quotients correspond in the entire database compared to subsets of samples within the database. Control samples were not included within each geographical area. However, the incidence of toxicity with and without control samples was similar for the entire database (first and second rows for each of the three tests listed in Tables 3 to 5).

For the HA10 test, there was typically an increase in the incidence of toxicity with an increase in the mean quotient within most of the regions, basins, and areas (Table 3). The incidence of toxicity for samples from each of the Great Lakes and within the areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database. No one area influenced the overall incidence of toxicity for the HA10 test. However, the absolute incidence of toxicity differed somewhat between areas. For example, the incidence of toxicity for the Great Lakes samples in the HA10 test (n=313) was 14% at mean quotients of <0.1, 68% at mean quotients of >1.0, and 78% at mean quotients of >5.0. This compares to an incidence of toxicity for the entire database (n=654) of 19% at mean quotients of <1.0 there was about a 14% higher incidence of toxicity for samples from the Great Lakes compared to the entire database for the HA10 test. The lower incidence of toxicity for samples from non-Great Lakes areas was primarily due to the lower incidence of toxicity at mean quotients >1.0 for samples from the states of New York (7%, n=15), Oregon (40%, n=20), and Washington (50%, n=34).

In the HA10 test, there was also a higher incidence of toxicity at mean quotients of <0.1 for a limited number of samples from the lower Mississippi River (89%, n=9) and from the Temblader River in California (75%, n=4) compared to the entire database. These samples contributed to the 22% incidence of toxicity that was observed for samples from non-Great Lakes areas at quotients of <0.1 (n=89). By comparison, only a 14% incidence of toxicity was evident for samples with quotients of <0.1 for Great Lakes areas (n=51). One of the reasons for the difference between non-Great Lakes samples and Great Lakes samples may be due to

contaminants associated with agricultural practices in the non-Great Lakes areas (i.e., herbicides or pesticides) that contributed to the toxicity of these samples independently of elevated metals, PAHs, or PCBs.

In the HA28 test, there was also an increase in the incidence of toxicity with an increase in the mean quotient within most of the regions, basins, and areas (Table 4). The incidence of toxicity for samples from each of the Great Lakes and within areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database. Similarly, the incidence in toxicity for samples from the Great Lakes was consistent with the incidence in toxicity for samples from non-Great Lakes areas. Therefore, no one area influenced the overall incidence of toxicity within the HA28 test. There were a relatively high number of non-toxic samples (n= 51 of 53) with low mean quotients from the upper Mississippi River. The incidence of toxicity for the Great Lakes samples (n=42) was 13% at mean quotients of <0.5, 71% at mean quotients of 0.5 to <1.0, and 100% at mean quotients of >1.0. This pattern is consistent with incidence of toxicity for the entire database (n=151) of 17% at mean quotients of <0.5, 56% at mean quotients of 0.5 to <1.0, and 97% at mean quotients of >1.0 (Table 4).

In the CS10 test, there was typically an increase in the incidence of toxicity with an increase in the mean quotient within most of the regions, basins, and areas (Table 5). The incidence of toxicity for the Great Lakes samples (n=463) was 14% at mean quotients of <0.1, 56% at mean quotients of >1.0, and 71% at mean quotients of >5.0. This pattern is consistent with incidence of toxicity for the entire database (n=611) of 20% at mean quotients of <0.1, 52% at mean quotients of >1.0, and 68% at mean quotients of >5.0. The incidence of toxicity for samples from each of the Great Lakes and within the areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database; however, there were some exceptions to this pattern. A lower incidence of toxicity was observed at mean quotients of >1.0 for samples from the St. Mary's River (41%, n=17), Fox River and Green Bay (0%, n=8), Menominee River (38%, n=8), and White Lake Montague (25%, n=8). However, at mean quotients of >5.0, there was a more consistent pattern in the incidence of toxicity in the CS10 test. An exception to this pattern at mean quotients >5.0 was observed for samples from the St. Mary's River (38%, n=8) and White Lake Montague (25%, n=4). Similarly, a lower incidence in toxicity was observed at mean quotients of >5.0 for samples from the state of Washington (50%, n=4).

In the CS10 test, there was also a higher incidence of toxicity at mean quotients of <0.1 for a limited number of samples from Sheboygan Harbor (75%, n=4), from the lower Mississippi River (89%, n=9), and from the Trinity River (67%, n=3). The samples from the Lower Mississippi River and Trinity River contributed to the 53% incidence of toxicity that was observed for samples from non-Great Lakes areas with quotients of <0.1 (n=19). By comparison, only a 14% incidence of toxicity was evident for samples with quotients of <0.1 for Great Lakes areas (n=99). Again, one of the reasons for the difference between non-Great Lakes samples and Great Lakes areas (i.e., herbicides or pesticides) that contributed to the toxicity of these samples independently of elevated metals, PAHs, or PCBs.

In summary, there was generally an increase in the incidence of toxicity with increasing mean PEC quotients within most of the regions, basins, and areas for all three tests (Tables 3 to 5). For the HA10 and HA28 tests, the incidence of toxicity for samples from each of the Great Lakes and within the areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database. However, the relationship between the incidence in toxicity and mean quotients in the CS10 test was more variable among geographic areas compared to either the HA10 or HA28 test. The results of these analyses indicate that the consensus-based PECs can be used to reliably predict toxicity of sediments on both a regional and national basis.

Future analyses planned for the database

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This paper presents results of the first analyses completed on the entire freshwater sediment database. Some of the additional analyses planned for the database (beyond the scope of this paper) are listed below:

- Approaches for designating samples as toxic. In the present study, samples were designated as toxic in the three tests based on a significant reduction in survival or a sublethal endpoint (typically growth) relative to a control or reference sediment. This designation of toxicity utilized the results of statistical analyses presented in each of the original studies. Alternatively, Long et al. (1998a) classified sediments in a marine amphipod database as either marginally toxic (significantly reduced relative to the control) or as highly toxic (significantly reduced relative to the control with a reduction greater than a minimum significant difference; MSD). The MSD was established by Long et al. (1998a) using a power analysis of data from 10-day marine amphipod tests (Thursby et al. 1997). Long et al. (1998a) and Field et al. (1999) reported reduced variability in the relationship between toxicity and sediment contamination when toxicity was evaluated using a standardized approach. Future analyses of the freshwater database will compare relationships between toxicity and contamination using marginally toxic versus highly toxic sediments. This classification may be based on a power analysis to establish an MDS for each endpoint in each of the three tests. Alternatively, the MDS for each endpoint may be established using results of round-robin testing (USEPA 2000). This latter procedure is currently being investigated for use in classifying the toxicity of freshwater samples in a revision to the report on the incidence and severity of sediment contamination in surface waters of the United States (USEPA 1997; Scott Ireland, USEPA, Washington, DC, personal communication). Classification of sediments as either marginally or highly toxic may help to reduce the variability in the observed relationship between toxicity and mean quotients. Additionally, this analysis may help to address the higher incidence of toxicity at mean quotients of <0.1 in the HA10 and CS10 tests compared to the HA28 tests.
 - **Logistic-regression modeling of the freshwater database.** Field et al. (1999) described a procedure for evaluating matching marine sediment chemistry and toxicity data using logistic regression models. These models can be used to estimate the probability of observing an effect at any contaminant concentration. The models were developed for marine amphipods using a large database (n=2524) from a variety of geographic areas.

The results of preliminary analyses using these techniques indicate that the freshwater database may have too few samples to adequately develop these regression models. However, evaluations are ongoing to determine how well the regression models developed from the marine database can be used to predict responses of organisms in the freshwater database.

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- **Optimal list of analytes for broad scale application and test of the relative efficacy of the mean versus the sum PEC quotient.** Further comparisons are needed of approaches for calculating the PEC quotient. Calculation of a mean quotient may include analytes which have limited toxicological importance. The significant toxicological contribution of a few chemicals may be averaged out by the use of the mean quotient. Analyses need to be conducted to identify an optimal list of analytes for broad scale application and to test the relative efficacy of the mean versus the sum PEC quotient. Even if the mean quotient continues to be the best at predicting toxicity, identification of an optimal list of analytes would be useful. If a consistent set of analytes is applied, the sum and the mean quotient will mathematically equivalent.
- **Influence of sediment grain size and ammonia on the incidence of toxicity.** Data on grain size and pore-water ammonia, pH, and water hardness were obtained for some studies evaluated in this paper. Future analyses of the database will evaluate the influence of either grain size or ammonia on the response of the test organisms in the HA10, HA28, or CS10 tests. ASTM (2000) and USEPA (2000) provide the following guidance for dealing with the influence of grain size or ammonia in toxicity tests with freshwater sediments.

Natural physico-chemical characteristics such as grain size or organic carbon can potentially influence the response of test organisms. ASTM (2000) and USEPA (2000) summarize results from a variety of studies and conclude *H. azteca* can tolerate a wide range in grain size and organic matter in 10- to 42-day tests with sediments. Larvae of *C. tentans* in 10-day tests were tolerant of a wide range of grain size if ash-free dry weight was used to account for the influence of inorganic material in the gut. The content of organic matter in sediments does not appear to affect survival of *C. tentans* larvae in sediments; but, may be important with respect to larval growth. Future analyses of the database will evaluate potential relationships between grain size or organic carbon of sediments on the toxicity in the HA10, HA28, or CS10 tests.

Ammonia in pore water may contribute to the toxicity of some sediments in fresh water. The toxicity of ammonia to *C. tentans* is dependent on pH whereas the toxicity of ammonia to *H. azteca* is also dependent on water hardness (ASTM 2000, USEPA 2000). Water-only LC50 values may provide suitable screening values for potential ammonia toxicity; however, higher concentrations may be necessary to actually induce ammonia toxicity in sediment exposures, particularly for *H. azteca* due to avoidance. ASTM (2000) and USEPA (2000) cite studies which describe procedures for conducting toxicity identification evaluations (TIEs) for pore-water or whole-sediment samples to determine if ammonia is contributing to the toxicity of sediment samples. Future analyses of the

database will evaluate potential relationships between pore-water ammonia, pH, and water hardness on toxicity in the HA10, HA28, or CS10 tests. These analyses will focus on the HA28 test due to limited data available for the HA10 or CS10 tests.

Guidance manual for conducting an integrated assessment of sediment

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contamination. Work is in progress to develop a guidance manual for USEPA Great Lakes National Program Office that can be used to assess sediment quality and determine the need for remediation at a site. Specifically, this guidance manual will describe procedures for combining results of sediment toxicity, bioaccumulation, benthic communities surveys, and sediment chemistry in an integrated evaluation of ecological risk. Development of this guidance manual is being coordinated with ongoing efforts to develop similar guidance by the Department of the Interior and by Environment Canada. Additional analyses of the freshwater database will be used to help develop this guidance. Specifically, these analyses will focus on determining which endpoints provide the most sensitive and cost effective measures of sediment contamination.

Summary

The primary objectives of this study were to: (1) evaluate the ability of consensus-based PECs to predict toxicity in a freshwater database for field-collected sediments in the Great Lakes basin; (2) evaluate the ability of these PECs to predict sediment toxicity on a regional geographic basis elsewhere in North America; and (3) compare approaches for evaluating the combined effects of chemical mixtures on the toxicity of field-collected sediments. When mean quotients were calculated using an approach of equally weighting up to 10 reliable PECs (PECs for metals, total PAHs, total PCBs, and sum DDE), there was an increase in the incidence of toxicity with an increase in the mean quotient in all three tests. A consistent increase in the toxicity in all three tests occurred at mean quotients of >0.5. However, the overall incidence of toxicity was greater in the HA28 test compared to the short-term tests. The reason for the higher incidence of toxicity in the HA28 test compared to the short-term tests may be due to the duration of the exposure or the sensitivity of growth in the longer HA28 test. However, comparisons of the sensitivity between these tests needs to be made with some caution. There were very few samples in the database where tests were conducted using splits of the same samples. Therefore, the differences observed in the responses of organisms may also be due to differences in the types of sediments evaluated in the individual databases for each test. Nevertheless, it appears that longer-term tests in which survival and growth are measured tend to be more sensitive than shorter-term tests, with acute to chronic ratios on the order of 6 indicated for *H. azteca*.

We were also interested in determining the predictive ability of PEC quotients for major classes of compounds. Therefore, we evaluated the incidence of toxicity based on a mean quotient for metals, a quotient for total PAHs, or a quotient for total PCBs. Different patterns of toxicity associated with the other procedures for calculating quotients were observed. For example in the HA28 test, a relatively abrupt increase in toxicity was associated with elevated PCBs alone or with elevated PAHs alone, compared to the pattern of a gradual increase toxicity observed with quotients calculated using a combination of metals, PAHs, and PCBs. These analyses indicate that the different patterns in toxicity may be the result of unique chemical signals associated with individual contaminants in samples. While mean quotients can be used to classify samples as toxic or non-toxic, individual quotients might be useful in helping to identify substances that may be causing or substantially contributing to the observed toxicity.

We chose to make comparisons across geographic areas using mean quotients calculated by equally weighting the contribution of the three major classes of compounds (metals, or PAHs, or PCBs). This approach assumes that these three diverse groups of chemicals exert some form of joint toxic action. Use of this approach also maximized the number of samples that were used to make comparisons across geographic areas. Generally, there was an increase in the incidence of toxicity increasing with mean PEC quotients within most of the regions, basins, and areas for all three toxicity tests. The incidence of toxicity for samples from each of the Great Lakes and within the areas of each Great Lake was relatively consistent with the overall pattern of toxicity in the entire database for the HA10 and HA28 tests. However, the relationship between the incidence in toxicity and mean quotients in the CS10 test was more variable among geographic areas compared to either the HA10 or HA28 test. Results of these analyses indicate that the

PECs developed using a database from across North America can be used to reliably predict toxicity of sediments on a regional basis.

One of the primary goals of sediment quality assessments is to evaluate the effects of contaminated sediments on benthic communities in the field (Ingersoll et al. 1997). Swartz et al. (1994) evaluated sediment quality conditions along a sediment contamination gradient of total DDT using information from 10-day toxicity tests with amphipods, sediment chemistry, and the abundance of benthic amphipods in the field. Survival of amphipods (*Eohaustorius estuarius*, *Rhepoxynius abronius*, and *H. azteca*) in laboratory toxicity tests was positively correlated to the abundance of amphipods in the field and negatively correlated to total DDT concentrations. The toxicity threshold for amphipods in 10-day sediment toxicity tests was about 300 μ g total DDT/g organic carbon. The threshold for reduction in abundance of amphipods in the field was about 100 μ g total DDT/g organic carbon. Therefore, correlations between toxicity, contamination, and the status of benthic macroinvertebrates in the field indicate that 10-day sediment toxicity tests can provide a reliable indicator of the presence of adverse levels of sediment contamination in the field. However, these short-term toxicity tests may be under protective of sublethal effects of contaminants on benthic communities in the field.

Similarly, Canfield et al. (1994, 1996, 1998) evaluated the composition of benthic invertebrate communities in sediments in a variety of locations including the Great Lakes, the upper Mississippi River, and the Clark Fork River in Montana. Results of these benthic invertebrate community assessments were compared to SQGs (ERMs) and 28-day sediment toxicity tests with *H. azteca*. Good concordance was evident between measures of laboratory toxicity, SQGs, and benthic invertebrate community composition in extremely contaminated samples. However, in moderately contaminated samples, less concordance was observed between the composition of the benthic community and either laboratory toxicity tests or SQGs. The laboratory toxicity tests better identified chemical contamination in sediments compared to many of the commonly used measures of benthic invertebrate community structure. As the status of benthic invertebrate communities may reflect other factors such as habitat alteration in addition to effects of contaminants, the use of longer-term toxicity tests in combination with SQGs may provide a more sensitive and protective measure of potential toxic effects of sediment contamination on benthic communities compared to use of 10-day toxicity tests.

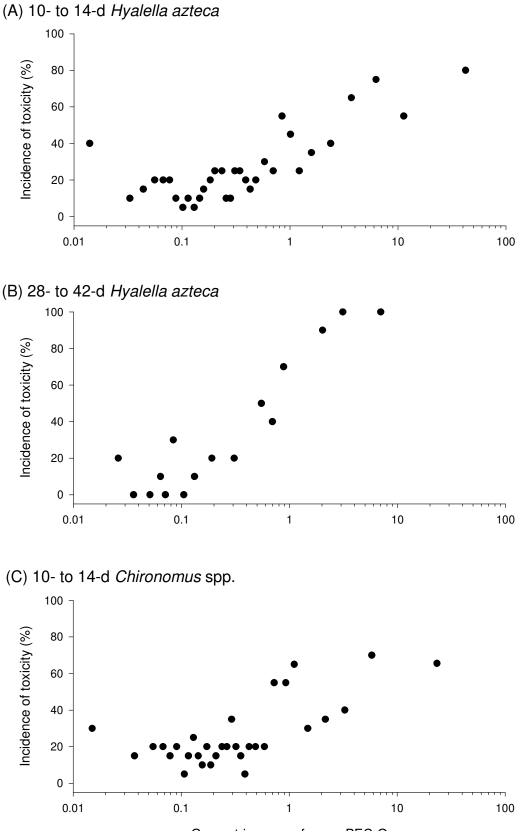
This paper presents results of the first analyses completed on the entire freshwater sediment database. Some of the additional analyses planned for the database that are beyond the scope of this paper include: (1) comparing approaches for designating samples as toxic; (2) evaluating logistic-regression models; (3) identifying a list of optimal analytes for broad scale application and testing the relative efficacy of the mean verses the sum PEC quotient; (4) evaluating the influence of grain size and ammonia on the incidence of toxicity; and (5) developing a guidance manual for conducting an integrated assessment of sediment contamination.

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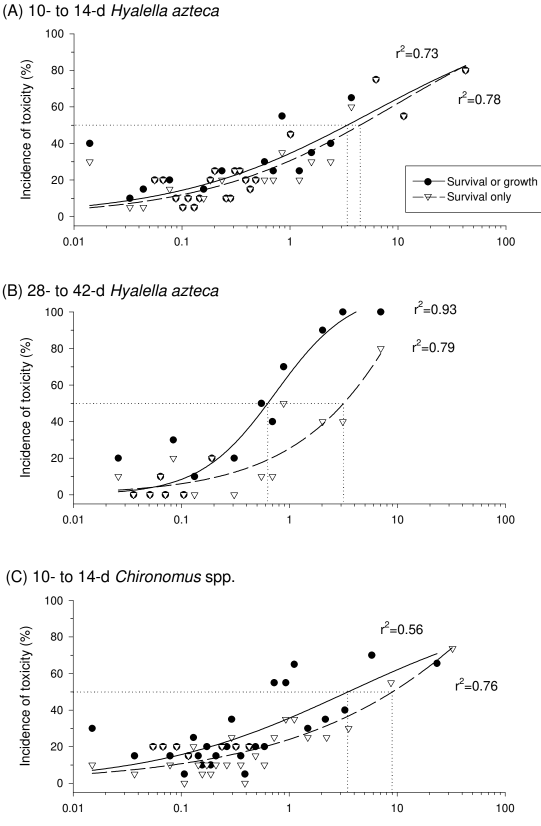
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Geometric mean of mean PEC-Q

Figure 1. Relationship between the geometric mean of the mean PEC quotients and the incidence of toxicity in the three tests, based on survival or growth. The average is based on 20 samples/data point in Figures 1A and 1C (except for the data point for the highest average of the mean quotients where n=19 to 30), and 10 samples/data point for Figure 1B (see Appendix 1 for detail).



Geometric mean of mean PEC-Q

Figure 2. Relationship between the geometric mean of the mean PEC quotients and the incidence of toxicity in the three tests, based on survival or growth, or based on survival alone. The dotted line represents a 50% incidence of toxicity. See legend for Figure 1 for additional detail.

Substance	PEL	SEL	TET	ERM	PEL-HA28	Consensus-Based PEC
Metals (in mg/kg DW)						
Arsenic	17	33	17	85	48	33.0*
Cadmium	3.53	10	3	9	3.2	4.98*
Chromium	90	110	100	145	120	111*
Copper	197	110	86	390	100	149*
Lead	91.3	250	170	110	82	128*
Mercury	0.486	2	1	1.3	NG	1.06
Nickel	36	75	61	50	33	48.6*
Zinc	315	820	540	270	540	459*
Polycyclic Aromatic Hydroca	rbons (in µg/kg D	W)				
Anthracene	NG	3700	NG	960	170	845
Fluorene	NG	1600	NG	640	150	536
Naphthalene	NG	NG	600	2100	140	561*
Phenanthrene	515	9500	800	1380	410	1170*
Benz[a]anthracene	385	14800	500	1600	280	1050*
Benzo(a)pyrene	782	14400	700	2500	320	1450*
Chrysene	862	4600	800	2800	410	1290*
Fluoranthene	2355	10200	2000	3600	320	2230
Pyrene	875	8500	1000	2200	490	1520*
Total PAHs	NG	100000	NG	35000	3400	22800*
Polychlorinated Biphenyls (ir	n µg/kg DW)					
Total PCBs	277	5300	1000	400	240	676*
Organochlorine Pesticides (in	uµg/kg DW)					
Chlordane	8.9	60	30	6	NG	17.6
Dieldrin	6.67	910	300	8	NG	61.8
Sum DDD	8.51	60	60	20	NG	28.0
Sum DDE	6.75	190	50	15	NG	31.3*
Sum DDT	NG	710	50	7	NG	62.9
Total DDTs	4450	120	NG	350	NG	572
Endrin	62.4	1300	500	45	NG	207
Heptachlor Epoxide	2.74	50	30	NG	NG	16.0
Lindane (gamma-BHC)	1.38	10	9	NG	NG	4.99

Table 1. Sediment quality guidelines that reflect probable effect concentrations (PECs; i.e., above which harmful effects are likely to be observed; MacDonald et al. 2000a). An "*" designates a reliable PEC (>20 samples and >75% correct classification as toxic).

PEL = Probable effect level; dry weight (Smith et al. 1996).

SEL = Severe effect level, dry weight (Persaud *et al.* 1993).

TET = Toxic effect threshold; dry weight (EC & MENVIQ 1992).

ERM = Effects range median; dry weight (Long and Morgan 1991).

PEL-HA28 = Probable effect level for Hyalella azteca; 28-day test; dry weight (USEPA 1996).

NG = No guideline.

PEC quotient ¹	Incidence of toxi <0.1	city (%) based or 0.1 to <0.5	1 mean PEC quo 0.5 to <1.0	tients (number of 1.0 to <5.0	f samples in parer >1.0	theses) >5.0	Total number of samples
<u>Hyalella azteca 10- to 14-0</u>	lay tests						
Mean - all	20 (102)	16 (336)	36 (80)	52 (103)	57 (152)	67 (49)	670
Mean - metals	20 (104)	19 (354)	39 (72)	63 (72)	63 (51)	62 (21)	623
Fotal PAHs	20 (178)	26 (160)	23 (47)	37 (54)	57 (103)	80 (49)	488
Fotal PCBs	26 (109)	21 (91)	46 (49)	51 (47)	60 (77)	73 (30)	326
Mean - MPP (and)	19 (79)	26 (89)	38 (34)	49 (35)	66 (64)	86 (29)	266
Mean - all (select1)	19 (36)	23 (128)	32 (38)	67 (42)	70 (64)	77 (22)	266
Mean - all (select2)	22 (46)	22 (121)	33 (40)	64 (39)	73 (59)	90 (20)	266
Mean - metals (select2)	23 (40)	24 (139)	33 (45)	81 (31)	86 (42)	100 (11)	266
Total PAHs (select2)	25 (123)	33 (76)	35 (20)	49 (33)	64 (47)	100 (14)	266
Total PCBs (select2)	20 (98)	25 (61)	47 (43)	47 (34)	59 (64)	73 (30)	266
Mean - MPP (or)	18 (147)	16 (288)	37 (73)	41 (92)	54 (162)	71 (70)	670
<u>Hyalella azteca_28- to 42-</u>	day tests						
Alean - all	8 (51)	11 (47)	57 (28)	NC^{2}	91 (34)	NC	160
/Iean - metals	8 (50)	20 (51)	62 (37)	NC	86 (22)	NC	160
Total PAHs	17 (98)	61 (46)	56 (9)	NC	86 (7)	NC	160
Total PCBs	4 (26)	6 (35)	17 (12)	NC	97 (36)	NC	109
Jean - MPP (and)	4 (45)	6 (18)	50 (18)	NC	100 (28)	NC	109
Aean - all (select1)	5 (44)	5 (17)	50 (22)	NC	100 (26)	NC	109
Aean - all (select2)	5 (40)	5 (21)	50 (22)	NC	100 (26)	NC	109
Aean - metals (select2)	5 (40)	25 (24)	60 (33)	NC	100 (12)	NC	109
Fotal PAHs (select2)	8 (57)	64 (37)	55 (9)	NC	100 (6)	NC	109
Fotal PCBs (select2)	4 (26)	6 (35)	17 (12)	NC	97 (36)	NC	109
Mean - MPP (or)	10 (63)	13 (39)	56 (27)	NC	97 (31)	NC	160
<u>Chironomus</u> spp. 10- to 1	4-day tests						
Aean - all	20 (90)	19 (333)	33 (95)	44 (75)	51 (114)	64 (39)	632
Aean - metals	22 (88)	23 (338)	25 (89)	39 (61)	44 (84)	57 (23)	599
Fotal PAHs	14 (178)	33 (133)	57 (28)	67 (33)	74 (53)	85 (20)	392
Fotal PCBs	46 (91)	22 (49)	36 (33)	53 (51)	58 (74)	70 (23)	247
Mean - MPP (and)	29 (21)	35 (78)	35 (26)	50 (34)	60 (52)	78 (18)	177
Mean - all (select1)	29 (7)	37 (98)	29 (34)	47 (17)	66 (38)	81 (21)	177
Mean - all (select2)	33 (6)	35 (98)	33 (36)	61 (23)	68 (37)	79 (14)	177
Mean - metals (select2)	8 (12)	43 (107)	22 (36)	75 (12)	82 (22)	90 (10)	177
Fotal PAHs (select2)	27 (64)	33 (73)	77 (13)	85 (20)	81 (27)	71 (7)	177
Fotal PCBs (select2)	48 (58)	23 (31)	34 (32)	35 (34)	48 (56)	68 (22)	177
Mean - MPP (or)	20 (121)	17 (313)	43 (63)	43 (88)	52 (132)	68 (44)	629

Table 2. Incidence of sediment toxicity within ranges of PEC quotients (calculated using various approaches) for freshwater tests based on survival or growth.

Mean - all: based on the PECs for each of the reliable metals, total PAHs, total PCBs, or sum DDE listed in Table 1 (n = 1 to 10 quotients/sample).

Mean - metals: based on the PECs for reliable metals listed in Table 1 (n = 1 to 7 quotients/sample).

Total PAHs: based on the PEC for total PAHs listed in Table 1 (n = 1 quotient/sample).

Total PCBs: based on the PEC for total PCBs listed in Table 1 (n = 1 quotient/sample).

Mean - MPP (and): based on reliable PECs for the average metals quotient, the total PAH quotient, and the total PCB quotient (n = 3 quotients/sample).

Mean - all (select1): based on the PECs for each of the reliable metals, total PAHs, total PCBs, or sum DDE listed in Table 1 using only samples with measured metals, total PAHs and total PCBs (n = 4 to 10 quotients/sample).

Mean - all (select2): based on the PECs for each of the reliable metals, total PAHs, or total PCBs listed in Table 1 using only samples with measured metals, total PAHs and total PCBs (n = 3 to 9 quotients/sample).

Mean - metals (select2), total PAHs (select2), or total PCBs (select2): based on the average quotient for metals, the quotient for total PAHs, or the quotient for total PCBs using only samples with measured metals, total PAHs and total PCBs.

Mean - MPP (or): based on the reliable PECs for the average metals quotient, total PAH quotient, or total PCB quotient (n = 1 to 3 quotients/sample).

²NC: Not calculated.

Table 3. Incidence of sediment toxicity within ranges of PEC quotients for the *Hyallela azteca* 10- to 14-day test in various geographic areas within the database based on survival or growth. PEC quotients were calculated as the mean of the average quotients for metals, a quotient for total PAHs, or a quotient for total PCBs ("Mean - MPP (or)" from Table 2). The numbers of samples are included within the parentheses.

Region	Basin	Area	Toxicity	Incidence of toxicity (%) based on mean PEC quotients								
			(%)	<0.1	0.1 to <0.5	0.5 to <1.0	1.0 to <5.0	>1.0	>5.0			
North America		All (with controls)	28 (670)	18 (147)	16 (288)	37 (73)	41 (92)	54 (162)	71 (70)			
		All (w/o controls)	29 (654)	19 (140)	17 (279)	37 (73)	41 (92)	54 (162)	71 (70)			
Great Lakes		All	35 (313)	14 (51)	18 (137)	47 (38)	60 (50)	68 (87)	78 (37)			
	Lake Erie	All	63 (30)	(0)	0 (4)	67 (15)	75 (8)	82 (11)	100 (3)			
		Buffalo River	100 (5)	(0)	(0)	100 (3)	100 (2)	100 (2)	(0)			
		Clinton River	67 (12)	(0)	(0)	63 (8)	75 (4)	75 (4)	(0)			
		Maumee River	46 (13)	(0)	0 (4)	50 (4)	50 (2)	80 (5)	100 (3)			
	Lake Huron	Saginaw River	10 (10)	(0)	0 (6)	0 (2)	50 (2)	50 (2)	(0)			
	Lake Michigan	All	60 (90)	50 (6)	40 (25)	33 (9)	69 (26)	76 (50)	83 (24)			
		Burns Waterway	0 (3)	(0)	0 (1)	(0)	(0)	0 (2)	0 (2)			
		Fox River	0 (6)	0 (1)	0 (3)	0 (1)	0 (1)	0 (1)	(0)			
		Indiana Harbor	71 (41)	0 (1)	25 (4)	29 (7)	77 (13)	90 (29)	100 (16)			
		Lake Michigan	0 (4)	(0)	0 (4)	(0)	(0)	(0)	(0)			
		Menominee River	80 (5)	(0)	(0)	(0)	80 (5)	80 (5)	(0			
		Sheboygan Harbor	72 (18)	75 (4)	75 (12)	(0)	0 (1)	50 (2)	100 (1			
		Waukegan Harbor	67 (3)	(0)	0 (1)	100 (1)	(0)	100 (1)	100 (1			
		Waushara County	0 (2)	(0)	(0)	(0)	0 (2)	0 (2)	(0			
		White Lake Montague	75 (8)	(0)	(0)	(0)	100 (4)	75 (8)	50 (4			
	Lake Ontario	All	19 (26)	(0)	19 (21)	100 (1)	0 (1)	0 (4)	0 (3			
		Genesee River	38 (13)	(0)	33 (12)	100 (1)	(0)	(0)	(0			
		Oswego River	0 (7)	(0)	0 (7)	(0)	(0)	(0)	(0			
		Sturgeon Lake	0 (6)	(0)	0 (2)	(0)	0 (1)	0 (4)	0 (3			
	Lake Superior	St. Louis River	18 (157)	9 (45)	12 (81)	38 (11)	38 (13)	55 (20)	86 (7			
Non-Great Lakes		All	24 (341)	22 (89)	16 (142)	26 (35)	19 (42)	39 (75)	64 (33)			
Lower Mississippi River		Tennessee	89 (9)	89 (9)	(0)	(0)	(0)	(0)	(0)			
Chesapeake Bay		All	50 (12)	50 (6)	40 (5)	100 (1)	(0)	(0)	(0			
State of California		All	38 (8)	75 (4)	0 (1)	0 (1)	0 (2)	0 (2)	(0			
		San Francisco Bay	0 (4)	(0)	0 (1)	0 (1)	0 (2)	0 (2)	(0)			
		Temblader River	75 (4)	75 (4)	(0)	(0)	(0)	(0)	(0)			
State of Florida		Miami-Dade County	22 (88)	12 (49)	27 (30)	57 (7)	50 (2)	50 (2)	(0)			
State of New York	Hudson River	New York Lakes	4 (28)	0 (5)	0 (6)	0 (2)	7 (14)	7 (15)	0 (1			
State of Oregon	Columbia River	Lower Columbia River	15 (60)	0 (1)	3 (29)	0 (10)	0 (5)	40 (20)	53 (15			
State of Texas		Trinity River	15 (40)	0 (5)	17 (35)	(0)	(0)	(0)	(0			
State of Washington		All*	27 (92)	0 (10)	14 (35)	23 (13)	28 (18)	50 (34)	75 (16			
-		Columbia River*	14 (35)	0 (10)	18 (17)	100 (2)	0 (4)	0 (6)	0 (2			
		Puget Sound	35 (57)	(0)	11 (18)	9 (11)	36 (14)	61 (28)	86 (14)			
Nova Scotia, Canada		All	100 (4)	(0)	100 (1)	100 (1)	100 (1)	100 (2)	100 (1)			

* One sample from British Columbia included in this total.

Table 4. Incidence of sediment toxicity within ranges of PEC quotients for the *Hyalella azteca* 28- to 42-day test in various geographic areas within the database based on survival or growth. PEC quotients were calculated as the mean of the average quotients for metals, a quotient for total PAHs, or a quotient for total PCBs ("Mean - MPP (or)" from Table 2). The numbers of samples are included within the parentheses.

Region	Basin	Area	Toxicity	Incidence of toxicity (%) based on mean PEC quotients						
Region	D ubin	Aita	(%)	<0.1	0.1 to <0.5	0.5 to <1.0	1.0 to <5.0	>1.0	>5.0	
North America		All (with controls)	35 (160)	10 (63)	13 (39)	56 (27)	96 (25)	97 (31)	100 (6	
		All (w/o controls)	37 (151)	10 (63)	17 (30)	56 (27)	96 (25)	97 (31)	100 (6	
Great Lakes		All	79 (42)	(0)	13 (8)	71 (7)	100 (22)	100 (27)	100 (5	
	Lake Erie	Buffalo River	80 (5)	(0)	(0)	67 (3)	100 (2)	100 (2)	(0	
	Lake Huron	Saginaw River	30 (10)	(0)	0 (6)	50 (2)	100 (2)	100 (2)	(0	
	Lake Michigan	All	96 (27)	(0)	50 (2)	100 (2)	100 (18)	100 (23)	100 (5	
		Indiana Harbor	100 (4)	(0)	(0)	(0)	100 (1)	100 (4)	100 (3	
		Waukegan Harbor	96 (23)	(0)	50 (2)	100 (2)	100 (17)	100 (19)	100 (2	
Non-Great Lakes		All	21 (109)	10 (63)	18 (22)	50 (20)	67 (3)	75 (4)	100 (1	
Upper Mississippi River		All	4 (53)	4 (45)	0 (8)	(0)	(0)	(0)	(0	
State of Alabama	Gulf of Mexico	Mobile Bay	0 (5)	(0)	0 (4)	(0)	0 (1)	0 (1)	(0	
State of Maine	Kennebec River	Brunswick	25 (4)	0 (2)	50 (2)	(0)	(0)	(0)	(0	
State of Maryland	Chesapeake Bay	Aberdeen	75 (4)	100 (1)	0 (1)	100 (2)	(0)	(0)	(0	
State of Montana	Pend Oreille River	Clark Fork River	62 (13)	50 (2)	50 (2)	57 (7)	100 (1)	100 (2)	100 (1	
State of Texas		All	20 (15)	15 (13)	50 (2)	(0)	(0)	(0)	(0	
	Gulf of Mexico	Galveston Bay	60 (5)	50 (4)	100 (1)	(0)	(0)	(0)	(0	
		Rio Grande	0 (5)	0 (5)	(0)	(0)	(0)	(0)	(0	
		Trinity River	0 (4)	0 (3)	0 (1)	(0)	(0)	(0)	(0	
		Austin	0 (1)	0 (1)	(0)	(0)	(0)	(0)	(0	
Washington D.C.	Chesapeake Bay	Chesapeake Bay	40 (15)	(0)	33 (3)	36 (11)	100 (1)	100 (1)	(0	

Table 5. Incidence of sediment toxicity within ranges of PEC quotients for the *Chironomus* spp. 10- to 14-day test in various geographic areas within the database based on survival or growth. PEC quotients were calculated as the mean of the average quotients for metals, a quotient for total PAHs, or a quotient for total PCBs ("Mean - MPP (or)" from Table 2). The numbers of samples are included within the parentheses.

Region	Basin	Area	Toxicity	Inc	idence of toxicity (%) based on mean	PEC quotients		
			(%)	<0.1	0.1 to <0.5	0.5 to <1.0	1.0 to <5.0	>1.0	>5.0
North America		All (with controls)	27 (629)	20 (121)	17 (313)	43 (63)	43 (88)	52 (132)	68 (4
		All (w/o controls)	28 (611)	20 (118)	17 (298)	43 (63)	43 (88)	52 (132)	68 (4
Freat Lakes		All	30 (463)	14 (99)	20 (211)	51 (45)	47 (70)	56 (108)	71 (
	Lake Erie	All	45 (73)	21 (19)	38 (26)	71 (17)	70 (10)	64 (11)	0
		Buffalo River	40 (5)	(0)	(0)	67 (3)	0 (2)	0 (2)	
		Clinton River	100 (12)	(0)	(0)	100 (8)	100 (4)	100 (4)	
		Elmira, Ontario, Canada	33 (6)	20 (5)	100 (1)	(0)	(0)	(0)	
		St. Clair River	39 (44)	21 (14)	47 (19)	33 (6)	75 (4)	60 (5)	0
		Wheatly	0 (6)	(0)	0 (6)	(0)	(0)	(0)	
	Lake Huron	All	46 (48)	25 (4)	47 (19)	67 (6)	45 (11)	42 (19)	38
		Saginaw River	10 (10)	(0)	0 (6)	0 (2)	50 (2)	50 (2)	
	Lako Mishigan	St. Mary's River	55 (38) 37 (89)	25 (4)	69 (13)	100 (4)	44 (9)	41 (17)	38
	Lake Michigan	All Burns Waterway Harbor	0 (3)	33 (9)	23 (30)	20 (5)	33 (27)	49 (45)	72 (
		East River	0 (3)	(0)	0 (1)	(0)	(0)	0 (2) 0 (0)	0
		Fox River & Green Bay	0 (18)	0 (1)	0 (6)	0 (3)	0 (8)	0 (8)	0
		Grand River	33 (3)	(0)	0 (1)	(0)	0 (8)	50 (2)	100
		Indiana Harbor	100 (14)	(0)	(0)	(0)	100 (6)	100 (14)	100
		Kenosha Harbor	0 (1)	(0)	(0)	0 (1)	(0)	(0)	100
		Kewanee Harbor	0 (2)	0 (2)	(0)	(0)	(0)	(0)	
		Lake Michigan	20 (5)	0 (1)	25 (4)	(0)	(0)	(0)	
		Menominee River	27 (11)	0 (1)	0 (2)	(0)	29 (7)	38 (8)	100
		Port Washington Harbor	0 (2)	(0)	0 (2)	(0)	(0)	(0)	-
		Sheboygan Harbor	56 (18)	75 (4)	50 (12)	(0)	0 (1)	50 (2)	100
		Waukegan Harbor	67 (3)	(0)	0 (1)	100 (1)	(0)	100 (1)	100
		White Lake Montague	25 (8)	(0)	(0)	(0)	25 (4)	25 (8)	25
	Lake Ontario	All	17 (89)	11 (19)	5 (56)	50 (4)	71 (7)	80 (10)	100
		Hamilton Harbor	60 (5)	(0)	(0)	100 (1)	50 (4)	50 (4)	
		Lake Ontario	3 (63)	7 (15)	2 (47)	0 (1)	(0)	(0)	
		Niagra River	63 (8)	(0)	0 (2)	50 (2)	100 (3)	100 (4)	100
		Oswego River	27 (11)	25 (4)	29 (7)	(0)	(0)	(0)	
		Sturgeon Lake	100 (2)	(0)	(0)	(0)	(0)	100 (2)	100
	Lake Superior	All	22 (164)	8 (48)	16 (80)	31 (13)	47 (15)	65 (23)	100
		St. Louis River	21 (155)	9 (47)	14 (76)	27 (11)	46 (13)	67 (21)	100
		Black River	57 (7)	0 (1)	67 (3)	100 (1)	50 (2)	50 (2)	
		Lake Superior	0 (1)	(0)	(0)	0 (1)	(0)	(0)	
		Wisconsin State	0 (1)	(0)	0 (1)	(0)	(0)	(0)	
Ion-Great Lakes		All	22 (148)	53 (19)	12 (87)	22 (18)	28 (18)	33 (24)	50
ower Mississippi River	B 10	Tennessee	89 (9)	89 (9)	(0)	(0)	(0)	(0)	
State of Montana	Pend Oreille River	Clark Fork River	15 (13)	0 (2)	0 (2)	14 (7)	100 (1)	50 (2)	0
State of New York	Hudson River	New York Lakes	9 (23)	0 (5)	0 (5)	50 (2)	9 (11)	9 (11)	
State of Texas		Trinity River	13 (72)	67 (3)	10 (68)	0 (1)	(0)	(0)	
State of Washington		All	30 (27)	(0)	18 (11)	29 (7)	40 (5)	44 (9)	50 100
Nova Scotia, Canada		All	75 (4)	(0)	100 (1)	0 (1)	100 (1)	100 (2)	100

	10- to 14-day Hyalella azteca					28- to 42-day Hyalella azteca					10- to 14-day Chironomus spp.					
Ν	Survival or	or growth Survival only			Ν	Survival o	r growth	Survival only			Survival or gro	owth		Survival only		
-	Geomean	Toxic	Geomean	Toxic	-	Geomean	Toxic	Geomean	Toxic	Ν	Geomean	Toxic	Ν	Geomean	Toxic	
	of mean Q	(%)	of mean Q	(%)		of mean Q	(%)	of mean Q	(%)		of mean Q	(%)		of mean Q	(%)	
20	0.014	40	0.014	30	10	0.026	20	0.026	10	20	0.015	30	20	0.015	10	
20	0.033	10	0.033	5	10	0.036	0	0.036	0	20	0.037	15	20	0.037	5	
20	0.044	15	0.044	5	10	0.051	0	0.051	0	20	0.055	20	20	0.055	20	
20	0.056	20	0.056	20	10	0.064	10	0.064	10	20	0.068	20	20	0.068	20	
20	0.067	20	0.067	20	10	0.071	0	0.071	0	20	0.079	15	20	0.079	10	
20	0.077	20	0.077	15	10	0.084	30	0.084	20	20	0.091	20	20	0.091	20	
20	0.088	10	0.088	10	10	0.105	0	0.105	0	20	0.107	5	20	0.107	0	
20	0.102	5	0.102	5	10	0.132	10	0.132	0	20	0.117	15	20	0.117	15	
20	0.114	10	0.114	10	10	0.191	20	0.191	20	20	0.130	25	20	0.130	20	
20	0.130	5	0.130	5	10	0.307	20	0.307	0	20	0.144	15	20	0.144	10	
20	0.146	10	0.146	10	10	0.550	50	0.550	10	20	0.157	10	20	0.157	5	
20	0.159	15	0.159	10	10	0.696	40	0.696	10	20	0.173	20	20	0.173	10	
20	0.183	20	0.183	20	10	0.881	70	0.881	50	20	0.188	10	20	0.188	5	
20	0.202	25	0.202	25	10	2.022	90	2.022	40	20	0.210	15	20	0.210	10	
20	0.235	25	0.235	20	10	3.119	100	3.119	40	20	0.239	20	20	0.239	20	
20	0.257	10	0.257	10	10	7.000	100	7.000	80	20	0.265	20	20	0.265	10	
20	0.282	10	0.282	10						20	0.294	35	20	0.294	25	
20	0.309	25	0.309	25						20	0.322	20	20	0.322	20	
20	0.344	25	0.344	25						20	0.357	15	20	0.357	10	
20	0.390	20	0.390	20						20	0.390	5	20	0.390	0	
20	0.429	15	0.429	15						20	0.426	20	20	0.426	20	
20	0.483	20	0.483	20						20	0.489	20	20	0.489	15	
20	0.582	30	0.582	20						20	0.590	20	20	0.590	10	
20	0.701	25	0.701	20						20	0.728	55	20	0.728	25	
20	0.847	55	0.847	35						20	0.931	55	20	0.931	35	
20	1.013	45	1.013	45						20	1.116	65	20	1.116	35	
20	1.224	25	1.224	20						20	1.490	30	20	1.490	25	
20	1.582	35	1.582	30						20	2.164	35	20	2.222	25	
20	2.386	40	2.386	30						20	3.281	40	20	3.555	30	
20	3.711	65	3.711	60						20	5.837	70	20	8.779	55	
20	6.276	75	6.276	75						29	23.400	66	19	32.608	74	
20	11.339	55	11.339	55												
30	42.350	80	42.350	80												
Equati	on:	y=a/[1+(x/	$(x_0)^{b}$]													
a	114.017		122.465			111.803		447.821			99.488			397.523		
b	-0.479		-0.488			-1.247		-0.639			-0.474			-0.366		
x _o	5.687		9.535			0.745		81.279			3.442			1804.374		
		nt at 50% i	ncidence of toxic	ity												
	3.4		4.5	-		0.63		3.2			3.5			9.0		

Appendix 1. Summary of the data presented in Figures 1 and 2.