Comparison of Bulk Sediment and Sediment Elutriate Toxicity Testing Methods

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Abstract  Numerous methods exist for assessing the potential toxicity of sediments in aquatic systems. In this study, the results from 10-day bulk sediment toxicity test methods using *Hyalella azteca* and *Chironomus tentans* were compared to results from 96-hour *Pimephales promelas* and *Ceriodaphnia dubia* renewed acute toxicity tests conducted using elutriate samples prepared from the same sediments. The goal of the study was to determine if the results from the elutriate tests were comparable to those obtained from the bulk sediment tests. Of the 25 samples analyzed, 16 were found to be toxic to at least one of the species tested, in either elutriate or bulk sediment tests. The *C. tentans* 10-day bulk sediment test was the most sensitive, with 12 sediment samples exhibiting toxicity to this species, while the *H. azteca* bulk sediment test and *C. dubia* 96-hour elutriate test were the least sensitive, exhibiting toxicity in only 7 of the 25 sediments tested. The *P. promelas* elutriate test found 8 of the 25 sediments to be toxic. Based on the total number of sites found to show toxicity, results from testing indicate 96-hour elutriate tests show a level of sensitivity comparable to 10-day bulk sediment tests in assessing toxicity quantitatively. However, the methods did not always find toxicity at the same sites, suggesting that the ability of elutriate tests to predict toxicity (quantitatively) is not statistically correlated with bulk sediment methods. This would indicate that a suite of toxicity test methods would provide the most complete measure of site condition; however, in circumstances where bulk sediment testing is not feasible, elutriate tests can provide a practical and credible alternative for toxicity assessment.

Introduction

Bulk sediment toxicity tests are routinely used to assess the level and extent of contamination in...
bottom sediments. While reliable, these tests can be resource intensive, requiring significant
time and materials. The purpose behind this study was to compare the results from bulk
sediment toxicity tests using *Hyalella azteca* and *Chironomus tentans* to the results obtained
from sediment elutriate tests using conventional *Ceriodaphnia dubia* and *Pimephales promelas*
ambient water toxicity testing methods. Sediment elutriate tests offer a considerable cost savings
and may have broad application where bulk sediment testing is not feasible. Such a case would
be an ambient toxicity monitoring program (ATMP) conducted by a U.S. Environmental
Protection Agency (USEPA) Regional Office or a state agency. The purpose behind most
ATMPs is to provide a measure of the baseline condition in an ecological system and to measure
any changes experienced in the system (Hall et al. 2000). These programs need to be cost
effective to be implemented. The resources required to include bulk sediment toxicity testing
preclude the use of this endpoint in most ATMPs. Use of the sediment elutriate test as a
surrogate endpoint could provide a cost-effective means to include sediment testing in an ATMP.

In this study, USEPA Region 6 (Arkansas, Louisiana, New Mexico, Oklahoma, and
Texas) sought assistance from the USEPA Office of Research and Development (ORD), through
the Regional Methods Initiative (RMI) Program, to conduct side-by-side bulk sediment and
acute sediment elutriate toxicity tests. Region 6, in cooperation with states and tribes, has been
conducting aquatic toxicity tests with ambient water samples and sediment elutriate as part of a
regional ATMP since 1990

[http://www.epa.gov/earth1r6/6wq/ecopro/watershd/monitrng/toxnet/index.htm](http://www.epa.gov/earth1r6/6wq/ecopro/watershd/monitrng/toxnet/index.htm). Using the
sediment elutriate test has allowed USEPA Region 6 to include a cost-effective sediment toxicity
endpoint as part of the ATMP. The objective of the study was to assess the use of sediment
elutriate tests as a feasible alternative to conventional bulk sediment tests. Numerous test organisms have been compared in the past to assess the toxicity associated with contaminated sediments with varying degrees of success. Ten day exposures conducted by Cairns et al. (1984) yielded 50% lethal concentrations (LC50s) of 38 and 39 µg/L, respectively, for *Hyalella azteca* and *Chironomus tentans* using copper spiked sediments. Ten day exposures performed by Suedel et al. (1993) revealed LC50s of 45 and 32 µg/L, respectively, for *H. azteca* and *C. tentans* using fluoranthane spiked water. However, species’ sensitivities have also shown differences in the past. Water spiked with zinc displayed LC50s of 73 µg/L for *H. azteca* versus 1,125 µg/L for *C. tentans* in experiments conducted by Phipps et al. (1995). Both species were tested to assess sediment toxicity in samples collected from 11 sites in the Keweenaw Waterway in Michigan by West et al. (1993) and revealed only a marginal level of agreement at 55%. These same types of discrepancies have been noted in 7-day *P. promelas* and *C. dubia* toxicity test exposures from point-source discharge effluents in National Pollutant Discharge and Elimination System (NPDES) permitting studies (Stewart et al. 1990). These limitations aside, *Hyalella azteca* and *Chironomus tentans* are the species recommended by USEPA for use in sediment toxicity testing (USEPA 2000), and *Ceriodaphnia dubia* and *Pimephales promelas* are the species recommended for use in ambient water quality monitoring (USEPA 2002), therefore, they were selected for use in this study.

**Materials and Methods**

**Sediment Samples**

The sediment toxicity samples provided by USEPA Region 6 for this study were collected by
state and tribal agencies that participate in the Region 6 Ambient Toxicity Monitoring Program. USEPA Region 6 scientists pre-screened a number of waterbody sites that had been sampled previously in the program, as well as other sites being sampled by state water quality agencies, to ensure that the samples selected for use in this study were from sites observed as being toxic or having a strong potential for toxicity.

The sediment samples were collected from freshwater lakes and streams located in Texas, Oklahoma, and New Mexico over a two year period from early 2002 (01/02) through late 2003 (12/03). When the USEPA Region 6 Laboratory in Houston, Texas received the sediment samples, they were homogenized and split into two sub-samples. One sub-sample was retained by Region 6. A second 4-L sediment sub-sample was shipped overnight to the USEPA-National Exposure Research Laboratory (NERL) in Cincinnati, Ohio for use in both bulk sediment testing and acute toxicity testing with sediment elutriate samples. These sediment samples were logged in at the Cincinnati facility and stored at 4°C until tested. Sediment samples were held for no longer than two weeks before being used to prepare elutriate samples and for no longer than eight weeks before being used in a bulk sediment toxicity test.

Sediment Elutriate and Bulk Sediment Testing Water

Moderately hard reconstituted water (MHRW), with a hardness of 100 mg/L CaCO₃, was used to prepare the sediment elutriate samples and as the control for the 96-h acute sediment elutriate tests. MHRW was also used as the overlying water in the bulk sediment tests. MHRW was prepared from a standard formula (USEPA 2002) using reagent grade chemicals and Super-Q® (Millipore Corporation, Billerica, MA) ultrapure water. The water was prepared at least three
days prior to the start of the test to allow sufficient time for stabilization.

Sediment Elutriate Preparation

Sediment elutriate samples were prepared based on procedures described in the American Society for Testing and Materials Guide E 1391 (ASTM 2000) and USEPA-U.S. Army Corps of Engineers (USEPA-USACOE 1998). The samples were mixed in a 1:4 (v/v) ratio of sediment to water and placed on a rotary shaker table for 1 h, at a speed of 100 rpm. After mixing for an hour, the samples were centrifuged at 3000 rpm for 20 min, to separate the water from the sediment. The aqueous fraction (elutriate sample) was poured off and stored in a cubitainer at 4°C for use in acute toxicity testing within 24 h of preparation. The remaining subsample was retained and stored at 4°C for use in the corresponding bulk sediment toxicity tests. This sediment elutriate mixing procedure differs from that described for use in the Region 6 ATMP. The procedure used here is the standard procedure described by USEPA-USACOE 1998 and is more widely used. The decision was made to use a standard elutriate preparation method for this comparison. Another paper will compare the USEPA-USACOE elutriate method used here with the method described in the Region 6 ATMP.

Toxicity Tests with Sediment Elutriate Samples

Ninety-six (96)-h static-renewal acute toxicity tests were conducted with the elutriate samples,
using standard USEPA methods (USEPA 1988, 2002). A test temperature of 25 ± 1°C and photoperiod of 16 h light:8 h dark were maintained during testing with both *C. dubia* and *P. promelas*. Routine chemical parameters (pH, dissolved oxygen, conductivity, and temperature) were measured in fresh test solution at test startup (0 h) and test solution renewal (48 h), and on a composite aliquot of old test solution (at 48 and 96 h), for each test treatment and control sample. No additional chemical analyses were performed with bulk sediment samples or elutriate water samples.

The *C. dubia* test procedure used four replicate test chambers (30-ml plastic cups) for each test treatment, with five animals per replicate (20 organisms per control or treatment), and a test solution volume of 25 ml. The less than 24-h old *C. dubia* neonates were obtained from in-house cultures maintained at the USEPA-Cincinnati facility. The *C. dubia* were fed 0.1 ml each of *Selenastrum capricornutum* (freshwater algae) and YCT (yeast, trout chow, cerophyll mixture) during holding, 2 h prior to test start, and 2 h prior to test solution renewal at 48 h.

The *P. promelas* acute toxicity test used four replicate test chambers (250-ml plastic cups) for each test treatment, with 10 animals per replicate (40 organisms per control or treatment), and a test solution volume of 200 ml. The *P. promelas* used in testing were supplied from an in-house culture maintained at the USEPA Cincinnati facility. The *P. promelas* were 2- to 10-days old (± 24-h age range) at the start of the test. This age range is more restrictive than the 1 to 14 day old range recommended in USEPA 2002 and was selected as a means to reduce test variability. The fish were fed 0.2 ml newly hatched brine shrimp (GSL Brine Shrimp, Ogden, UT) during holding, 2 h prior to the start of the tests and 2 h prior to test solution renewal at 48 h.
The 10-day static-renewal bulk sediment toxicity tests with *H. azteca* and *C. tentans* were conducted using standard USEPA sediment testing methods (USEPA 2002). The tests for both species were conducted at 23 ±1°C, using 100 ml of sediment and 175 ml of MHRW as the overlying water. The control sediment was a commercially available washed grade 40 white silica sand that is typically used as a landscape material. The sand was acid washed then rinsed with deionized water prior to use. It was supplemented by the addition of 1% liquid alfalfa for use in testing. This control sediment has been extensively used in testing conducted for the Environmental Monitoring and Assessment Program-Great Rivers Ecosystems project (EMAP-GRE) as well as in the assessment and remediation of contaminated sediments (Tabak et al. 2005). Each control or sediment treatment used six replicate 400-ml beakers as test chambers, with 10 animals in each replicate (60 organisms total per concentration). The photoperiod during testing was 16 h light:8 h dark and a water change of two volume additions (350 ml) was performed daily, using a modified Zumwalt renewal system (Zumwalt et al. 1994). Temperature was measured on a daily basis and routine physical/chemical parameters (pH, dissolved oxygen, conductivity, and temperature) were measured on initial setup (day 0) and final takedown (day 10) for each test concentration or control sample. The sediment testing conducted under the EPA Region 6 TOXNET program provides a screening level assessment. Therefore the majority of the toxicity results do not have accompanying chemical analysis data. Such data would be generated as part of an intensive special study to evaluate spatial and temporal characteristics of toxicity at a designated location.

The *C. tentans* and *H. azteca* used in testing were from in-house cultures maintained at the
USEPA-Cincinnati facility. Prior to testing, both species were held at 25±1°C and fed daily.

At the start of testing, the *C. tentans* were third instar larvae (10-day old) and the *H. azteca* were 7- to 10-days old. The feeding regimes for both species followed standard USEPA guidance (USEPA 2000). Each species was fed 1.0 ml YCT (yeast, trout chow, cerophyll mixture) daily throughout the duration of the test.

**Data Analysis**

The endpoint values for these tests were determined using procedures and statistical methods established by USEPA (2000, 2002). Mortality differences in the samples compared to the corresponding test controls were analyzed for both the sediment elutriate and bulk sediment tests, using the t-test function ($\alpha=0.05$) in Lotus 1-2-3 (IBM, Armonk, NY). Each sample tested included a corresponding control sample for comparison. Any sample with survival significantly less than the control ($p<0.5$) was classified as being toxic.

Statistical analyses were performed on the results of these toxicity bioassays to establish associations between the results from elutriate test methods and bulk sediment methods to determine if the use of elutriate testing is just as applicable in determining toxicity as bulk sediment methods. Comparisons between the elutriate versus bulk sediment methods were made by McNemar’s test of symmetry ($p=0.05$) in Systat 11 (Systat Software, San Jose, CA) to derive whether the probability of rating a sample as toxic is similar between the two methods being compared. Cohen’s Kappa measure of agreement (values between 0 and 1) in Systat 11 was also performed on elutriate versus bulk sediment methods to determine the strength of association between each method’s ability to detect toxic effects.
Results

A total of 25 sediment samples were tested for toxicity in this study (Tables 1 and 2). All arrived in good condition and the tests were started for each sample within the established time parameters. Of the 25 samples tested, 16 were found to be toxic to at least one of the species tested in either the elutriate tests or the bulk sediment tests (Table 2, Figs 1 and 2). Of these 15 samples, six (NM3-A, TX1-A, TX2-A, TX2-C, TX4-A, and TX7) were found to be toxic to just those species used in the bulk sediment tests (*C. tentans*, *H. azteca*, or both); three (TX1-B, TX1-D, and TX3-C) were found to be toxic to just those species used in the elutriate tests (*C. dubia*, *P. promelas*, or both); and seven (NM1-A, NM1-B, TX1-C, TX3-A, TX4-B, TX8-A, and TX8-B) showed toxicity to at least one species used in both the elutriate and bulk sediment tests. The remaining samples (NM2, NM3-B, OK1-A, OK1-B, TX2-B, TX3-B, TX5-A, TX5-B, and TX5-C) showed no toxicity to the species tested in either method.

Sediment Elutriate Tests

Test acceptability for the *C. dubia* and *P. promelas* sediment elutriate test methods is defined as ≥ 90% survival in the controls (USEPA 2002). All elutriate tests conducted in this study met or exceeded the control survival acceptability criteria. All elutriate tests met or exceeded this control survival acceptability criterion (Fig. 3). Control survival for the *C. dubia* tests ranged from 90% to 100%. Control survival for the *P. promelas* tests ranged from 95% to 100%.

The *C. dubia* elutriate tests revealed 7 of the 25 samples were toxic (Table 2, Fig. 1), and the *P. promelas* elutriate tests revealed 8 of the 25 samples were toxic (Table 2, Fig 1). Five samples were determined to be toxic by both methods (NM1-A, NM1-B, TX1-B, TX8-A, and...
Samples TX1-D and TX4-B were found to be toxic to just *C. dubia*, while samples TX1-C, TX3-A, and TX3-C were determined to be toxic to only *P. promelas*. In total, the combined elutriate tests determined 10 of the 25 samples tested were toxic to at least one of the species tested, with little difference in the overall sensitivity between the two species (Table 2, Fig. 1).

**Bulk Sediment Tests**

Test acceptability for the *C. tentans* test method is defined as ≥ 70% survival in the control, while the acceptability for the *H. azteca* test method is defined as ≥ 80% survival in the control (USEPA 2000). All bulk sediment tests conducted in this study met or exceeded the control survival acceptability criteria (Fig. 3). Control survival in the *C. tentans* tests ranged from 70% to 85%. Control survival in the *H. azteca* tests ranged from 90% to 100%.

The *H. azteca* bulk sediment tests revealed 7 of the 25 samples were toxic, while *C. tentans* bulk sediment tests yielded toxicity for 12 of the 25 samples (Table 2, Fig. 2). One sample (TX1-A) was found to be toxic to only *H. azteca*, and six samples (NM3-A, TX2-A, TX3-A, TX4-A, TX4-B, and TX7) were found to be toxic to only *C. tentans*. The remaining six samples (NM1-A, NM1-B, TX1-C, TX2-C, TX8-A, and TX8-B) were toxic to both species. In total, of the 25 samples tested using the bulk sediment method, 13 were found to be toxic to at least one of the species tested (Table 2, Fig. 2). The *C. tentans* appear to be slightly more sensitive than the *H. azteca*.

McNemar’s test of symmetry indicated no significant differences in designating a site toxic between elutriate and *H. azteca* bulk sediment tests (S=1.2857, p =0.4531) or between elutriate
and *C. tentans* bulk sediment tests (S=0.50, p =0.7266). Cohen’s Kappa test revealed no significant agreement between elutriate and *H. azteca* bulk sediment tests (K=0.386, p=0.0618) or between elutriate and *C. tentans* bulk sediment tests (K=0.3548, p=0.0820).

**Discussion**

The purpose of this study was to assess the utility of using sediment elutriate tests as a tool for monitoring sediment condition as part of a long-term ambient toxicity monitoring program. Results from this study indicate that the 10-day bulk sediment toxicity tests conducted with *C. tentans* were the most sensitive, with 12 of the 25 sediment samples exhibiting toxicity to this species (Table 2, Fig. 2). The *H. azteca* bulk sediment tests (Table 2, Fig. 2) and *C. dubia* elutriate tests (Table 2, Fig. 1) were the least sensitive, exhibiting toxicity in only 7 of the 25 sediments, while the *P. promelas* elutriate tests found 8 of the 25 sediments to be toxic (Table 2, Fig. 1). Two of the sites that were sampled and tested multiple times (OK1 and TX5) did not exhibit toxicity with either the bulk sediment or elutriate tests.

The results from the statistical comparison of elutriate testing data and bulk sediment testing data reveal interesting trends. McNemar’s test of symmetry indicated there was no significant difference between the ability of an elutriate test or bulk sediment test to predict the toxicity of a sample. Cohen’s Kappa measure of agreement suggested that both elutriate tests and bulk sediment tests alone lacked the qualitative ability to predict toxicity in a given sample. This could be due to a variety of factors, including the sensitivity of the species tested and the toxic components found in the samples. Based on the results of both data analysis methods, this data set indicates limited differences between the use of a sediment elutriate test or a bulk
sedi
tment test in predicting the toxicity in a given sediment sample. The 64% agreement between the test methods in detecting toxicity would seem to validate the performance of the elutriate test method. The lack of agreement in the remaining samples could simply be due to interspecies differences in sensitivity to various toxicant components of the sediments.

As Burton et al. (1996) note, all methods have inherent variability which must be taken into account when interpreting test results. The methods used in this study did not always find toxicity at the same sites, indicating that both elutriate and bulk sediment tests have built-in biases. The decision becomes which method provides data adequate for the scope of the project or ATMP (Hall et al. 2000, OSPAR Commission 1997). The elutriate tests can be effective in identifying acutely toxic sites; however, the assessment of elutriate toxicity alone is not sufficient to assess the overall potential hazards of contaminated sediments in some cases (Burton et al. 1996, Liß and Ahlf 1997, Ahlf and Wild-Metzko 1992, Burton 1992). Research has shown the results from elutriate tests can correlate well to bulk sediment metals contamination (Finlayson et al. 2000, Callier et al. 2009) and bulk sediment organic contamination (Karbe 1992). In a major study conducted as part of the Bremerhaven Workshop, sediments were collected from 16 sites located in the North Sea and a total of 11 different toxicity tests were conducted, with 20 toxicity endpoints being measured (Chapman et al. 1992). The results from those toxicity tests indicated that the 10-day amphipod test with bulk sediment and the 48-h oyster larval abnormal development test with sediment elutriate most clearly reflected the toxicity gradient across the samples and best corresponded with the chemical analysis and in-situ community data (Chapman et al.1992). Other researchers have shown that the results from elutriate tests correspond well to impacts noted in the in-place benthic
community (Callier et al. 2009). Based on the ability of elutriate tests with Daphnia magna to
determine toxic sites in Izmir Harbor in western Turkey, the elutriate test with D. magna has
been proposed as a low-cost, efficient method to screen for sediment toxicity (Yegane et al.
2008). Others have made this same proposal, based on the ease of conducting elutriate tests and
the associated resource savings (Marin et al. 2001).

The original purpose behind the design of the sediment elutriate test is another factor to
consider when determining which method to use. This test method was originally designed to
assess the impact of re-suspension of sediment contaminants due to dredging and the release of
the dredged material back into an aquatic environment (ASTM International 2000, USEPA
1988). The re-suspension of sediments is not limited to dredging. Flood events can cause
significant quantities of bottom sediment to be re-suspended (Mucha et al. 2004), as can boat and
ship traffic in harbors, rivers, and recreational lakes and reservoirs (Sousa et al. 2007). The
water bodies tested in this study are all large enough to support recreational use, including boat
traffic. They are also subject to high flow levels and flood events. These factors indicate the
elutriate tests would be an appropriate screening tool to use to monitor for the effects of this
type of activity, as well in these systems.

The data presented in this study indicate that sediment elutriate and bulk sediment tests
show a comparable level of sensitivity, based on the total number of sites found to be toxic with
each method. Other researchers have reached these same conclusions (see Finlayson et al. 2000,
Chapman et al. 1992). The two methods did not always find toxicity at the same sites, and both
were shown to have built-in biases. One probable cause for these biases would be interspecies
differences in tolerance to toxicants or combinations of toxicants present in the sediments. This
not only exists between the bulk and elutriate test species, but between the species used in each type of test as well. *C. dubia* are considered to be more sensitive to many types of toxicants than are *P. promelas*. The sensitivity roles change when the toxicant is ammonia or hydrogen sulfide, in which case *P. promelas* is more sensitive than *C. dubia*. The same differences can be seen with *C. tentans* and *H. azteca*. As a burrowing species *C. tentans* has an increased level of contact with the sediment and therefore with the toxic components of the sediment. *H. azteca* is more epibenthic, so it would be affected by those toxic components that are absorbed into the water column. None of these are necessarily good or bad (false positives, false negatives), since neither type of test is an absolute barometer of toxicity. Both bulk sediment and elutriate tests are imperfect and have their strengths and weaknesses which result in limitations for each method. The fact that, for a few tests, the elutriate tests showed toxicity when the bulk sediment did not could indicate greater sensitivity for certain toxicants. Whenever possible, the use of a which would indicate that a suite of toxicity test methods would provide the most complete measure of site condition. However, elutriate test methods do provide an efficient, cost-effective alternative to bulk sediment toxicity tests and can be used as a screening tool to monitor for sediment toxicity. A more intensive assessment of a site found to have consistently toxic sediment through elutriate testing may likely require the use of additional environmental measures, including bulk sediment testing, to determine the level and extent of toxicity.

For future research, two changes to the elutriate study described here could help to improve the overall utility of the test method. Work conducted with different sediment-to-water ratios used in preparing elutriates has shown potential to better reflect the toxicity of bulk sediments (Novelli et al. 2006). The addition of a chronic or sub-chronic endpoint would increase the
usefulness of the data, as well. For instance, adding a *D. magna* 4-day survival and growth test (Lazorchak et al. 2009), or using it in place of the *P. promelas* acute test would provide a sensitive sub-chronic endpoint. These improvements would increase the relevance of sediment elutriate tests as a surrogate for bulk sediment testing.

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sensitivity of three benthic invertebrates to copper-contaminated sediments from the

through the inner part of Izmir Bay with Daphnia magna Straus, 1820. EU J Fish Aquat
Table 1  Index of sampling sites used in elutriate and bulk sediment testing. Numbers in parenthesis indicate number of samples collected at each site.

<table>
<thead>
<tr>
<th>Site name (# of visits)</th>
<th>Sample IDs</th>
<th>State</th>
<th>Location</th>
</tr>
</thead>
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<td>Willow Creek (2)</td>
<td>NM1-A</td>
<td>New Mexico</td>
<td>Lat 35°45’27”</td>
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<tr>
<td></td>
<td>NM1-B</td>
<td></td>
<td>Long 105°40’17”</td>
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<td>NM2</td>
<td>New Mexico</td>
<td>Lat 33°13’35”</td>
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<td></td>
<td></td>
<td></td>
<td>Long 108°14’30”</td>
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<tr>
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<td>NM3-A</td>
<td>New Mexico</td>
<td>Lat 33°09’01”</td>
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<tr>
<td></td>
<td>NM3-B</td>
<td></td>
<td>Long 107°10’56”</td>
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<td>Black Bear Creek (2)</td>
<td>OK1-A</td>
<td>Oklahoma</td>
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<tr>
<td></td>
<td>OK1-B</td>
<td></td>
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<td>TX1-B</td>
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<td>TX1-C</td>
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<td>TX1-D</td>
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<tr>
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<td>Sample IDs</td>
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<tr>
<td>Bryan Municipal Lake (2)</td>
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<td>Lat 30°38′27″</td>
</tr>
<tr>
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<td>TX4-B</td>
<td></td>
<td>Long 96°21′37″</td>
</tr>
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<td>Lake Palestine (3)</td>
<td>TX5-A</td>
<td>Texas</td>
<td>Lat 32°12′01″</td>
</tr>
<tr>
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<td>TX5-B</td>
<td></td>
<td>Long 95°27′41″</td>
</tr>
<tr>
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<td>TX5-C</td>
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<td></td>
</tr>
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<td>TX7</td>
<td>Texas</td>
<td>Lat 35°44′32″</td>
</tr>
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<td></td>
<td>Long 101°20′30″</td>
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<tr>
<td>Alligator Bayou (2)</td>
<td>TX8-A</td>
<td>Texas</td>
<td>Lat 29°52′39″</td>
</tr>
<tr>
<td></td>
<td>TX8-B</td>
<td></td>
<td>Long 93°58′44″</td>
</tr>
</tbody>
</table>
Table 2  Summary of sediment elutriate samples and bulk sediment samples found to be toxic. An X indicates a test where the sample was determined to be toxic with that species; a blank cell indicates no toxicity effect.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Sample Collection Date</th>
<th>96-hour acute sediment elutriate tests</th>
<th>10-day bulk sediment tests</th>
</tr>
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<tbody>
<tr>
<td>TX1-A</td>
<td>2/11/02</td>
<td>Ceriodaphnia dubia X</td>
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<tr>
<td>TX2-A</td>
<td>2/20/02</td>
<td>Pimephales promelas X</td>
<td>X</td>
</tr>
<tr>
<td>NM1-A</td>
<td>3/13/02</td>
<td>Hyalella azteca X</td>
<td>X</td>
</tr>
<tr>
<td>TX1-B</td>
<td>7/25/02</td>
<td>Chironomus tentans X</td>
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</tr>
<tr>
<td>TX3-A</td>
<td>8/26/02</td>
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<td></td>
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<tr>
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<td>8/26/02</td>
<td></td>
<td></td>
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<tr>
<td>TX1-C</td>
<td>9/23/02</td>
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<td>11/4/02</td>
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