Use of artificial sediment to assess toxicity of chromium on *Chironomus xanthus*, *Danio rerio* and *Poecilia reticulata*

Utilização de sedimento artificial na avaliação da toxicidade do cromo sobre *Chironomus xanthus*, *Danio rerio* e *Poecilia reticulata*

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Abstract: Aim: Artificial sediments with simple formulations were used to assess the toxicity of Cr to *C. xanthus*, *D. rerio* and *P. reticulata*. Methods: The organisms were exposed to two sediment formulations: one without organic matter (S0) and one with 1% organic matter (SIII), both contaminated with Cr at concentrations of 90, 180, 360, 720 and 1440 mg/kg for the assays with *C. xanthus*, and 375, 750, 1500 and 3000 mg/kg for those with the two fish species. Chromium was obtained from the potassium dichromate (K₂Cr₂O₇). Results: The results of the acute toxicity tests demonstrated a reduction of up to four times in toxicity with the presence of OM, consisting of biodegradable detritus, even at the low concentration used (1%), and that *C. xanthus* was most sensitive to the metal. The reduction in the chromium toxicity was indicated by the LC₅₀ values, which were higher in SIII (1234.43 mg/kg; 2263.54 mg/kg; 2244.48 mg/kg) than in S0 (340.56 mg/kg; 1731.04 mg/kg; 1733.55 mg/kg) for *C. xanthus*, *D. rerio* and *P. reticulata*, respectively. Conclusions: Because of the repeatability of the results obtained, the artificial sediments with simple formulations were satisfactory to assess toxicity and can thus provide reliable results in ecotoxicological studies.

Keywords: Artificial sediment, Chromium, *C. Xanthus*, *D. Rerio*, *P. Reticulata*.

Resumo: Objetivo: Sedimentos artificiais com formulações simples foram utilizados para avaliar a toxicidade do Cr em *C. xanthus*, *D. rerio* e *P. reticulata*. Métodos: Os organismos foram expostos a duas formulações: sem matéria orgânica (S0) e com 1% de matéria orgânica (SIII), contaminadas com Cr nas concentrações de 90; 180; 360; 720 e 1440 mg/kg para os ensaios com *C. xanthus*, e 375; 750; 1500 e 3000 mg/kg para os peixes. Resultados: Os resultados dos ensaios agudos demonstraram uma redução de até quatro vezes na toxicidade frente à presença de OM, consistindo de detritos biodegradáveis, mesmo em concentração baixa (1%), e que *C. xanthus* foi a mais sensível ao metal. A redução na toxicidade do Cr frente à presença de OM foi verificada nos valores de CL₅₀, que estiveram mais elevados em SIII (1234.43 mg/kg; 2263.54 mg/kg; 2244.48 mg/kg) em relação a S0 (340.56 mg/kg; 1731.04 mg/kg; 1733.55 mg/kg) para *C. xanthus*, *D. rerio* e *P. reticulata*, respectivamente. Conclusões: Os sedimentos artificiais com formulações simples foram satisfatórios na avaliação da toxicidade e asseguram resultados confiáveis em estudos ecotoxicológicos.

Palavras-chave: Sedimento artificial, cromo, *C. Xanthus*, *D. Rerio*, *P. Reticulata*.

1. Introduction

Sediment tends to accumulate trace metals in aquatic ecosystems (Soares et al., 1999). This characteristic is aggravated in water bodies that receive inflows of metals, either steadily or intermittently, and can result in severe effects on the benthic biota (Doig and Liber, 2006). Mediated by physical, chemical and biological processes, metals can return to the water column and thus pose a risk
to the planktonic and nektonic communities, as well as triggering toxic effects in the other components of the food web (Adams, 1995).

Despite the importance of the sediment compartment, the assessment and quantification of the toxic potential of metals in natural sediments are still very difficult, since metals’ physical and chemical characteristics are highly variable, as are the routes for absorption and elimination of these pollutants by aquatic organisms (Edwards and Simkiss, 2005). Therefore, the number of reported experiments with artificial sediments in the laboratory has been increasing (Naylor and Rodrigues, 1995; Clément and Cadier, 1998; Kemble et al., 1999; Ribeiro et al., 1999; Verrhiest et al., 2002; Feiler et al., 2004; Roman et al., 2007). These studies permit evaluating the effects of various types of pollutants on aquatic organisms under pre-established conditions.

There are many advantages of using artificial sediment, among them: (i) the possibility of exercising rigorous control, with well-known characteristics and properties (size and proportion of particles, chemically specific surfaces), to assess and predict how natural sediments will react to contact with various types of pollutants; (ii) the absence of native microfauna and contaminants, to enable good reproducibility; (iii) the possibility of supporting the survival and growth of various organisms (plants and animals); (iv) the possibility of comparison with the results of other laboratory tests as well as field tests with natural sediments; (v) the ability to be stored without great difficulties (Suedel and Rodgers, 1994); and (vi) the ability to provide information on the behavior of substances individually or in complex mixtures in function of the substrate composition.

In a previous study we tested simple formulations of artificial sediments, aiming to reduce the costs of obtaining the fractions (by avoiding the need for sample collection in the field and storage before use in the laboratory) and to facilitate testing because of the ease of preparing the sediments. Our results demonstrated that the formulations prepared were satisfactory for use in ecotoxicological tests, since they were not found significant ecotoxicological effects on survival, growth, and morphology of gills ments and bioaccumulation of metals in benthic and nektonic species of this study. Thus, the present study aimed to test the feasibility of these formulations in assessing the ecotoxicity of a metal widely studied, chromium. In this way, the results could be demonstrated widely known to the present study, thereby, increasing the reliability in using the artificial sediment.

Chromium (Cr) is a metal widely utilized for various purposes, such as metallurgy, textiles (dyes), leather tanning, wood preservation and fertilizers (HSDB 2000, cited in Silva and Pedrozo, 2001). Its toxicity to aquatic organisms has been amply demonstrated over many decades (Stevens and Chapman, 1984; Dave et al., 1987; Jop et al., 1987; Villaescusa et al., 1997; Ceurdassier et al., 2000; Choi and Roche, 2004; Svecevicius, 2006, 2007; Vedamanikam and Shazilli, 2008). In natural systems, Cr is present in two oxidative states, Cr\textsuperscript{3+} (trivalent) and Cr\textsuperscript{6+} (hexavalent), each with different geochemical and toxicological properties. Cr\textsuperscript{6+} is highly oxidizable and unstable in reduced environments and anoxic sediments, and is thus highly soluble and toxic (De Laune et al., 1998). In contrast, Cr\textsuperscript{3+} has low solubility and is less toxic (Wang et al., 2001).

The choice of the test organisms was based on the possible exposure routes to the organisms from the addition of chromium to the sediments, namely direct exposure (affecting benthic organisms) and indirect exposure through partition to the water column (affecting nektonic organisms). Two nektonic organisms were chosen, to compare the responses of a species native to tropical ecosystems (Poecilia reticulata) and an exotic species (Danio rerio), and the benthic organisms was Chironomus xanthus.

Therefore, the aim of this study was to assess and confirms the potential use of a simple artificial sediment formulation (sand, clay and organic matter) in acute toxicity assays with chromium using test organisms from different trophic levels.

2. Material and Methods

2.1. Artificial sediments

The artificial sediments were formulated containing fractions of large-grained sand (about 90%), kaolinite clay (10%), without organic matter (S0) and 1% of organic matter (SIII). The percentages of the fractions used were determined based on the average concentration found in natural sediment samples from water bodies in the upper Jacaré-Guaçu watershed, located in São Paulo, Brazil (21º 57' 30" and 22º 22' 30" S; 47º 42' 30" and 48º 05' 30" W).

The sand was acquired from a company specialized in materials for water filtering (Mineração Jundu Ltda.) and had grain size between 0.4 and
The chromium solutions were prepared using sodium dichromate (K₂Cr₂O₇). The Cr concentrations used in the acute toxicity tests with *C. xanthus* (90 mg/kg, 180 mg/kg, 360 mg/kg, 720 mg/kg and 1440 mg/kg) and with the two fish species (375 mg/kg, 750 mg/kg, 1500 mg/kg and 3000 mg/kg) were ascertained by preliminary tests. The test solutions were added to the different artificial sediments to compose two experimental treatments: i) addition of the contaminant with organic matter (S0); and ii) addition of the contaminant to the sediment without organic matter (SIII). Each treatment was repeated three times.

The assays with *C. xanthus* were carried out according to the method described by Fonseca and Rocha (2004), by exposing six specimens of the 4th instar in samples of sediment/water in a proportion of 1:4. Each replication was in a disposable plastic cup with capacity of 250 mL, containing 60 g of sediment and 240 mL of water. There were three replications for each treatment, for a total of 18 test organisms. The tests lasted 96 hours. A control group (B), with sediment consisting only of the sand used to rear the specimens, was included in the tests. The mortality of the organisms was determined with the aid of a fine net and magnifying glass. The specimens were only fed on the first day of the test, in the same proportions used during their rearing.

For *D. rerio* and *P. reticulata*, the tests were conducted according to the standards of the ABNT (2004), whereby five organisms were exposed in sediment/water samples in the proportion of 1:4 (Burton and MacPherson, 1994). Each replication was carried out in a disposable plastic recipient with capacity of 1000 mL, containing 100 g of sediment and 400 mL of water. There were two replications for each treatment, for a total of 10 test organisms. These tests lasted 48 hours. There was also a control group (C) with only reconstituted water. During the experiments, the responses of the fish specimens only exposed with sand (A) and in the control group...
(C) did not show significant differences in the acute and chronic effects (according to Fisher’s exact test; p>0.05 and the Kruskall-Wallis test; p>0.05, respectively), so it was decided only to continue with the C group as a control in these tests. The mortality was determined by counting the dead specimens and the organisms were not fed during the testing period.

The following parameters were measured at the start and end of the experiments with all the species tested: pH (pH meter), hardness (titulometry) and conductivity (conductivity meter). During the tests, the temperature (25 ± 2 °C) and photoperiod (12h:12h; light:dark) were controlled (Fonseca and Rocha, 2004; ABNT, 2004).

2.4. Physical and chemical analyses

In all the tests, the total content of chromium in the water (APHA, 1995) and its bioavailable Cr in the sediments (as described by Silverio et al., 2006) were determined by taking samples at the start and end of the tests, using 20-mL disposable syringes to avoid resuspension of the sediment. For this purpose, three replicas of the experiments were added.

2.5. Sequential extraction analysis

The sequential extraction analysis was performed in sediments S0 and SIII to assist in the interpretation of chromium bioavailability to aquatic organisms, using some fractions of the method adapted from Silveira et al., (2006) for tropical soils because there was no breakdown of oxides as described by these authors. The fractions were: (i) exchangeable (F1), (ii) connected to carbonates (F2) (due to the high pH of the samples), (iii) bound to organic matter (F3), (iv) bound with iron oxides, Al and Mn (F4) and (v) residual (F5). To perform extraction was sequentially weighed equivalent to 1.0 g of dry material (sediment artificial) sieved to 100 mesh and added to polycarbonate centrifuge tubes. The samples were equilibrated for 2 h with 15 ml 0.1 M CaCl2 (F1). Was added 30 mL of 1 M NaOAc (pH 5) with stirring horizontal shaker for 5 h at ambient temperature for extraction of F2. To obtain F3 samples were extracted with 5 ml of 5% NaOCl pH adjusted to 8.5. The centrifuge tubes were placed in a water bath at 90 ° C for 30 min. The samples were centrifuged, filtered, and the supernatant was collected. This procedure was repeated four times. In no F4 fractions of oxides and this fraction was obtained by use of 0.2 M ammonium oxalate + 0.2 M oxalic acid + 0.1 M ascorbic acid (pH 3) (Schwertmann, 1964). The samples were mixed with 40 mL of extraction solution in a water bath at 90 ° C for 30 min. F5 was obtained after a total digestion with HF + HNO3 + ultrapure water in a microwave oven, according to EPA Method 3052 (EPA, 1996).

Between each successive extraction sample was suspended with 5.0 ml of 0.1 M NaCl, to displace the previous solution that may have remained in the tube, and to reduce the resorption of metal (Ahnstrom and Parker, 1999). The metals were determined in each fraction by atomic absorption spectrophotometry with graphite furnace (Model Spectraa 220, Varian), and the index was calculated coating used as a criterion for control of the results the maximum tolerable error equal to ± 20%, and: Recovery (%) = (S metal contents in the fractions / total content) x 100. The degree of recovery (100 ± 20%) was used as a criterion of quality control analyzes (Silveira et al., 2008). Total concentrations were determined according to EPA Method 3052 with microwave digestion with HNO3 + HF + ultrapure water (EPA, 1996).

2.6. Evaluation of the results

The mortality were expressed as LC(1)50; 96h for C. xanthus and LC(1)50; 48h for the two fish species, obtained by using the trimmed Spearman-Karber method (Hamilton et al., 1977).

3. Results

3.1. C. xanthus

In the tests with S0 the average LC50;96h was 340.56 mg/kg, with a standard deviation of 32.7 mg/kg and coefficient of variation of 9.62% (Figure 1). In the tests with SIII it was possible to calculate the LC50;96h in only one definitive test, with a value of 1234.43 mg/kg of chromium, almost four times the figure for the average LC50;96h with S0 (Figure 1). In the other definitive tests it was not possible to calculate the LC50;96h because of the low mortality at the last concentration, suggesting LC50;96h values above 1440 mg/kg of chromium for this substrate (Figure 1).

The analysis of the chromium concentrations in the water showed levels two times higher after 96 hours in the tests with both S0 and SIII. The concentrations in the water with S0 during the tests ranged from 2.24 mg/L (90 mg/kg) to 67.44 mg/L (1440 mg/kg), while for SIII the figures were between 0.25 mg/L (90 mg/kg) and 23.93 mg/L (1440 mg/kg). Therefore, the chromium concentrations in the water with sediment S0 were five to 20 times greater than in the water with sediment SIII (Figure 2).

The determination of the chromium concentration at the start (0h) and end of the tests
as the average LC\textsubscript{50};\textsubscript{48h} determined in the tests with S0. In the tests with \textit{P. reticulata} utilizing S0, the average LC\textsubscript{50};\textsubscript{48h} was 1733.55 mg/kg and the standard deviation was 710.44 mg/kg, while with SIII the average LC\textsubscript{50};\textsubscript{48h} was 2244.48 mg/kg with a standard deviation of 453.35 mg/kg, higher than for S0 (Figure 1).

The determination of the chromium concentrations in the water in the tests with the two fish species showed that they were higher after 48h except for the initial concentrations of 3000 mg/kg (S0) and 375 and 750 mg/kg (SIII). In the tests with S0, the chromium concentrations ranged from 2.06 mg/L (375 mg/kg) to 5.25 mg/L (3000 mg/kg), while in the tests with SIII these values were 2.52 mg/L (375 mg/kg) and 99.08 mg/L (3000 mg/kg). Therefore, the chromium concentrations were from 2 to 20 times higher in the water with SIII than with S0 (Figure 4).

For the sediments, the analysis of the chromium levels at the start (0h) and end (48h) of the tests showed higher metal concentrations at the start in both S0 and SIII. The concentrations in S0 during the tests varied from 2.50 mg/kg (180 mg/kg) to 233.12 mg/kg (1440 mg/kg), while in SIII these numbers ranged from 1.52 mg/kg (1440 mg/kg at 96h) to 55.57 mg/kg (1440 mg/kg at 0h). Therefore, for the sediments the chromium concentration at the start of the tests was about five times higher in S0 than in SIII (Figure 3), while after 96h the concentration was about two times higher in SIII than in S0, except at concentrations of 1440 mg/kg and 90 mg/kg, where the chromium levels were respectively about 15 and five times higher in SIII (Figure 3).

### 3.2. \textit{D. rerio} and \textit{P. reticulata}

The average LC\textsubscript{50};\textsubscript{48h} in the tests with \textit{D. rerio} with S0 was 1731.04 mg/kg, with a standard deviation of 322.02 mg/kg and a coefficient of variation of 18.60% (Figure 1). In the test with SIII, it was only possible to calculate the LC\textsubscript{50};\textsubscript{48h} in two definitive tests (2004.40 mg/kg and 2522.69 mg/kg). Both of these values were higher as the average LC\textsubscript{50};\textsubscript{48h} determined in the tests with S0. In the tests with \textit{P. reticulata} utilizing S0, the average LC\textsubscript{50};\textsubscript{48h} was 1733.55 mg/kg and the standard deviation was 710.44 mg/kg, while with SIII the average LC\textsubscript{50};\textsubscript{48h} was 2244.48 mg/kg with a standard deviation of 453.35 mg/kg, higher than for S0 (Figure 1).

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For the sediments, the analysis of the chromium levels at the start (0h) and end (48h) of the tests showed higher metal concentrations at the start in the tests with S0 (3 to 7 times) and with SIII.
Use of artificial sediment...

3.3. Sequential extraction analysis

Most of the chromium was in the residual fraction treatments S0 and SIII (up to almost 80% of the total content of the element in treating S0). In the treatment S0 distribution of Cr was in descending order: residual > oxides > MO > carbonate > exchangeable (Figure 6). In the treatment S0 only 21.3% of Cr was distributed in fractions considered potentially available, while 59.9% in the treatment SIII this element was in these fractions (Figure 6). In the treatment of the metal fraction SIII linked to organic matter was more significant than in the treatment S0, which was expected since this treatment is the addition of organic compound in its formulation and the element affinity for organic compounds.

4. Discussion

The results of the toxicity tests indicated that the mean LC$_{50}$ for all the test organisms was two to four times higher in SIII. The benthic species...
There were higher concentrations of chromium dissolved in the water in the tests with SIII both at the start and end of the testing. The chromium concentrations in the water were directly related to the concentrations added to the sediments and also to the toxicity.

The lower toxicity in the samples with the SIII in relation to those with S0, despite the higher levels of chromium dissolved in the water with the former sediment, may have occurred because of the instability of the chemical species regarding dissolved organic carbon (DOC). Cr$^{6+}$ (highly toxic and oxidizable) is easily reduced to Cr$^{3+}$ (lower toxicity) in the presence of DOC and

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The average LC$_{50}$,48h values in the tests with chromium dissolved in the water with sediments SIII and S0 for \( D.\ rerio \) (2263.54 mg/kg; 1731.04 mg/kg) and \( P.\ reticulata \) (2244.48 mg/kg; 1377.55 mg/kg) showed that mortality occurred at respective chromium concentrations of approximately 70.0 mg/L and 3.5 mg/L for both two species. Therefore, the toxicity to chromium in the water was reduced nearly 20 times with SIII in comparison with S0.

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humic substances (HS) (Skogerbøe and Wilson, 1981; Wittbrodt and Palmer, 1995). Besides this, the organic matter (OM) and byproducts of its degradation can complex metals and make them unavailable to the organisms. In light of the high DOC concentration in SIII (18.0 mg/L), the chromium in the water could be in unavailable form (complexed with OM) or reduced form (less toxic), causing lower toxicity. However, in the present study we did not evaluate the chemical species of chromium in the experiments.

Various researchers have demonstrated the influence of DOC on the toxicity of metals. Bringolf et al. (2006) evaluated the effect of dissolved OM on the acute toxicity of zinc to Pimephales promelas larvae. They found that a minimum concentration of 11.0 mg/L of DOC was necessary to reduce the LC_{50}, 96h of zinc, and this was positively correlated with the DOC concentrations. Our results here corroborate those findings.

Brooks et al. (2007) assessed the influence of photodegraded dissolved OM on the toxicity of copper to Pimephales promelas larvae and observed that the mortality was higher in the treatments with lower density of binding sites between Cu and DOC. Brooks et al. (2007) noted that the photochemical changes of the DOC in the bonds increased the concentration of free copper ions from 31 to 85%, as a result of the reduced density of the binding sites between the elements, triggering greater toxicity. Nakayasu et al. (1999) carried out a comparative study of reduction of Cr^{6+} in the presence of HS and their precursors and found different results between the compounds with high molecular weight (HS and SF) and those with low molecular weight (GA and TA). The authors found that the kinetics of reduction of Cr^{6+} to Cr^{3+} was two times higher for compounds with low molecular weight, with the reduction capacity in the following order: SH and SF<TA<GA.

The sequential extraction analysis confirmed the considerations described, since the same results demonstrated the highest bioavailability of chromium in sediments S0. According to Alomary and Belhadj (2007) the bioavailability of an element decreases with decreasing exchangeable for residual phases. The authors conducted a study in order to investigate the distribution of metals in sediment fractions of the Mediterranean Sea, and found that most metals predominate in F5 and smaller percentages are related to F1 and F2. As to the chromium percentages were found in the following order: F2 < F1 < F3 < F4 < F5, with about 5% in F1; 5% by F2; 8% by F3; 25% and 55% by F4 and F5. Although the values are related to marine sediments, results are similar to those observed in the present study.

Our results detected the distribution of chromium in order F1 < F2 < F3 < F4 < F5 in S0 and F1 < F2 < F4 < F3 < F5 in SIII. The values detected in F1 (1.9% and 2.8% for S0 in SIII) show higher bioavailability in S0. In the SIII chrome on the F3 was 29.9% while in S0 this figure was 7.8%. This fraction is represented by the portion that is not easily released to the environment, since the metal is bonded to the crystal structures of minerals from the sediment (Savvides et al., 1995). Thus, the percentages of chromium attached to this fraction, as verified in this study (78.7% and 40.1% for S0 in SIII), suggests that in the absence of MO and therefore fewer binding sites for complexation, chromium binds to mineral components of sediments in greater proportion. In fact, the fraction F3 in SIII was second in descending order presented above, other than S0. Accordingly, Förstner (1987) reported the existence of a time-dependent intensity adsorption compounds and suggested that the adsorption sites high energy (less numerous) are filled first, and become limiting in few, and that all the energy of the sediment decreases as these sites are filled less energy.

Another possible explanation for the higher chromium levels in the water in the tests with sediments containing organic matter is the characteristic of the sediment, i.e., although the sand fraction was high (~80%), the height of the sediment in the test recipients was under 3.0 cm. These conditions could have favored partition of the chromium from the sediments to the water at low OM concentrations (1%) (Eimers et al., 2002).

The chromium concentrations in the water in the tests with S0 were lower than in the tests with SIII. In this case the toxicity found at low chromium concentrations is likely mainly related to the metal’s tendency to remain in hexavalent form (highly toxic and oxidizable) in S0 (Nakayasu et al., 1999). The cation exchange capacity of kaolinite clay (30-150 mmol /kg) means there are additional binding sites, mainly of oxides of Fe and Al, perhaps mediating the partition of the chromium in S0 (Eimers et al., 2002).

The bioavailability of metals is related to various factors, such as the characteristics of the contaminant, chemical composition of the sediment, time of adsorption with the metal,
temperature, pH, redox potential and behavior and physiology of the organisms (feeding habit, capacity to metabolize compounds, among others) (Baudo et al., 1990). Therefore, relating the bioavailability of chromium only to the MO content can produce an unrealistic estimate.

In the present study we found an average $LC_{50}$ of 48h (in relation to the concentration dissolved in the test water) of 86.04 ± 48.71 mg/L of chromium for P. reticulata. For D. rerio this value was found to be 42.35 ± 15.34 mg/L (Nakagome et al., 2007). These levels are near those found for the SIII sediments (70.0 mg/L), but very different than for the S0 sediments (3.5 mg/L).

Some studies have sought to standardize the sensitivity range of potassium dichromate for these species, varying with exposure time and the life stage of the organisms (Cruz et al., 2008; Bertoletti, 2009). The results found by these authors were varied, which can be explained because sensitivity is an intrinsic factor of each organism and is directly associated with the genetic characteristics of the individual, resulting in variations in the findings between laboratories and even within a single brood.

Considering the average $LC_{50}$ of 96h for C. xanthus (greater than 1440.0 mg/kg; 340.56 mg/kg) in SIII and S0, respectively, it was possible to observe that in SIII the chromium concentration in the water was from 5 to 20 times lower. Furthermore, the concentration of bioavailable chromium in the sediments was some 5 times higher in S0, demonstrating the influence of low MO concentrations on the bioavailability of chromium and thus on its toxicity. These results show that chromium has different effects in this system because of the various forms of interaction of the organisms with the medium (fishes and benthic organisms).

Benthic organisms, such as Chironomidae larvae, are burrowers (Bird et al., 1995) and because they feed on detritus, the metal in the particles poses a high exposure risk (Martinez et al., 2003). However, the physical and chemical conditions in the digestive tract of these organisms can change the toxicity of the metal (Tessier and Campbell, 1987). Besides this, the uptake of metals by benthic organisms can result from exposure to interstitial and surface water as well (Lee et al., 2000).

Fish species, on the other hand, interact with the contaminant directly from the compounds dissolved in the water or particles ingested while feeding by contact with the sediment. Fishes of the Poeciliidae and Cyprinidae families are omnivores and the Poeciliidae can also feed on detritus (Meffe and Stelson Junior, 1989). The differences in the sensitivity of organisms can thus be a reflection of the different ways of interacting with the contaminated medium (Baudo et al., 1990).

The acute toxicity of chromium to C. xanthus, D. rerio and P. reticulata was reduced with the presence of biodegradable OM detritus, even at low concentrations (1%) and demonstrated that the artificial sediments with simple formulations tested here were adequate to assess the toxicity of chromium. Because of the ease of preparation, the repeatability of experiments and the satisfactory results in the toxicity assessment, artificial sediments with the specifications used in this study should be used more often in laboratory studies to investigate the toxicity of chemical substances and may contribute to the establishment of standardization for such an experiment.

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