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HAL Id: hal-01717786
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Submitted on 26 Feb 2018

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<td>Wiley - Manuscript type:</td>
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<td>Date Submitted by the Author:</td>
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<td>Caillat, Amélie; CEREGE UMR 7730 CNRS – Aix-Marseille Université, FR ECCOREV, Ciffroy, Philippe; Electricité de France (EDF), Division Recherche et Développement, Laboratoire National d’Hydraulique et Environnement (LNHE) Grote, Matthias; Electricité de France (EDF), Division Recherche et Développement, Laboratoire National d’Hydraulique et Environnement (LNHE) Rigaud, Sylvain; CEREGE UMR 7730 CNRS – Aix-Marseille Université, FR ECCOREV, Garnier, Jean-Marie; CEREGE UMR 7730 CNRS – Aix-Marseille Université, FR ECCOREV,</td>
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<td>bioavailability, copper, ecological risk assessment, plant uptake, sediment toxicity</td>
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**Abstract:**

The assessment of the potentially harmful effects of metals on biota depends on the speciation and bioavailability of the metals. In this study, we investigated copper accumulation and toxicity in the aquatic plant *Myriophyllum aquaticum* after exposure to artificial sediments varying in peat and/or ferric hydroxide content and spiked with Cu (5 to 200 mg.kg-1). Modeling of the kinetic DGT (diffusive gradient in thin film) measurements revealed fast and slow copper resupply from the solid phase for sediment formulated with and without peat, respectively. *M. aquaticum* proved to be sensitive to copper, as the copper accumulation and growth differed depending on the sediment composition and copper concentration. Comparing the copper accumulation in *M. aquaticum* to total dissolved, free and CDGT (concentration in solution derived from DGT measurements) copper concentrations revealed that CDGT concentrations were a better predictor of the accumulation than the others. However, the relatively weak correlation observed (r2=0.6) and the fact that plant uptake does not increase proportionally to fluxes to DGT suggest that copper uptake in plants was not diffusion limited. Thus, the free copper concentrations near the root surface were sufficient to meet the plant’s demand during the experiment. Furthermore, labile complexes that continuously resupply the Cu2+ pool may also contribute to the...
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Abstract

The assessment of the potentially harmful effects of metals on biota depends on the speciation and bioavailability of the metals. In this study, we investigated copper accumulation and toxicity in the aquatic plant *Myriophyllum aquaticum* after exposure to artificial sediments varying in peat and/or ferric hydroxide content and spiked with Cu (5 to 200 mg kg\(^{-1}\)). Modeling of the kinetic DGT (diffusive gradient in thin film) measurements revealed fast and slow copper resupply from the solid phase for sediment formulated with and without peat, respectively.

*M. aquaticum* proved to be sensitive to copper, as the copper accumulation and growth differed depending on the sediment composition and copper concentration. Comparing the copper accumulation in *M. aquaticum* to total dissolved, free and \(C_{\text{DGT}}\) (concentration in solution derived from DGT measurements) copper concentrations revealed that \(C_{\text{DGT}}\) concentrations were a better predictor of the accumulation than the others. However, the relatively weak correlation observed \((r^2=0.6)\) and the fact that plant uptake does not increase proportionally to fluxes to DGT suggest that copper uptake in plants was not diffusion limited. Thus, the free copper concentrations near the root surface were sufficient to meet the plant’s demand during the experiment. Furthermore, labile complexes that continuously resupply the Cu\(^{2+}\) pool may also contribute to the concentrations available for plant uptake. In the range of copper concentrations investigated here, saturation of uptake processes as well as toxicity are considered responsible for the poor DGT prediction of plant uptake.

Keywords: Bioavailability, *Myriophyllum aquaticum*, Diffusive gradient in thin film, Copper, Artificial sediment.
Introduction

Sediments have been recognized as a major sink of metals in aquatic systems, and the ultimate concern of such metal contamination in sediments is the impact of these metals on aquatic organisms. It is now well known that the total content of contaminants is not a good predictor for risk assessment as not all fractions are available to organisms [1]. Speciation and bioavailability are necessary conditions for assessing and predicting the food chain transfer and phytotoxicity of contaminants and their potentially harmful effects. The bioavailability of a compound can be assessed by biological methods (bioassay, bioaccumulation) and/or chemical measurements. Different procedures for biological tests on sediment can be used. Tests can be performed on the elutriate phase, the interstitial water or the whole sediment; the latter represents a more realistic type of exposure that can be simulated in the laboratory [2]. Among aquatic organisms, the widely used for bioassays on the whole sediment were nematodes (*Caenorhabditis elegans*), bacteria (*Arthrobacter globiformis*), fish embryos (*Danio rerio*), oligochaetes (*Tubifex tubifex*) and midge (*Chironomus tentans*) [3, 4]. Because macrophytes are considered indicators of water pollution at the same level as fish and macrobenthos, the integration of a primary producer (plant) that is rooted in the sediment in a test battery is desirable (European Water Framework Directive; EU-WRRL 2000). Therefore, a new sediment contact test with the *Myriophyllum aquaticum* (water milfoil) has been developed and will be normalized [5]. *M. aquaticum* is a submerging plant that extracts nutrients from sediment. The growth of the *Myriophyllum aquaticum* is affected by the origin of the sediment and the presence of toxicants [5]. *M. aquaticum* has also been used for studies of the bioaccumulation of heavy metals [6, 7] on plants collected in situ and plants grown hydroponically. Recently, *M. aquaticum* has been used in standardized sediment contact tests, and a strong correlation was found between toxic effects and metal sediment concentration [8]. Metal bioavailability can be estimated by various chemical and physico-chemical methods. In particular, Zhang and coworkers developed the so-called “diffusive gradient in thin films” (DGT) technique that is expected to mimic bioaccumulation processes and has been adapted for assessing metal bioavailability in soil and sediments [9-11]. DGT is a priori well adapted for non-equilibrium conditions where the bioavailability of metals is mainly controlled by the kinetic resupply of free metals in a solution from solid phase labile pools. Indeed, similar to a plant root, DGT induces a local decrease in the concentration of metal at the sediment/DGT interface, and metals bound to the sediment particles may be released in the interstitial water and become available [12]. To improve the interpretation of DGT experiments, the use of a conceptual model with sorption/desorption and diffusion reactions in the sediment is required. Two dynamic models have been developed; these models are known as DGT-Induced Fluxes in Soils and Sediments (1D-DIFS) [13] and DGT-PRObabilistic Fitting of parameters for Soils and Sediments (DGT-PROFS) [14]. The DGT-PROFS model has been developed to enhance the DIFS model for interpreting DGT kinetics experiments.
performed on formulated sediments [14, 15]. Previously, the link between DGT measurements and the bioaccumulation in plants showed contrasting results [9]. A strong correlation has been found between DGT-labile copper in various soils and copper accumulation in a terrestrial plant (Lepidium heterophyllum)[1]. However, DGT accurately predicted the bioavailability of Zn and Cu in spinach and ryegrass only at nontoxic concentrations [16]. Indeed, in cases of toxicity, plant absorption will be disturbed, whereas the DGT measurement will not be affected.

The aim of our study was to assess the relationships between speciation (given by different approaches, including DGT probes), bioaccumulation and toxic effects on M. aquaticum. Copper, considered a potentially extremely toxic threat to aquatic biota [17], was selected for this study because few studies have been performed comparing plant accumulation of copper and DGT measurements [9]. Therefore, we performed a set of experiments in parallel coupling bioassay and DGT approaches to compare the "real bioavailability" measured by the plant and the "potential bioavailability" estimated by the DGT tool.

Considering that the speciation of elements and therefore their bioavailability are linked to the solution and particulate phase compositions, we investigated the behavior of copper on formulated sediments of different compositions to reduce the complexity of the system. The sediment composition varied by (i) the addition of different components having more or less strong affinities toward copper (peat, iron oxides...) and (ii) the sediment copper spiking concentrations. Furthermore, the EDTA that is used in the standard solution for the bioassay (M. aquaticum) was investigated at different concentrations to assess its impact on copper bioavailability.

1. Material and Methods

1.1 Sediment and solution composition

Solid phase composition: Four different artificial sediments were prepared according to a slightly modified protocol of the standard OECD 207-218 protocol. The first sediment, named the “basic sediment” (SB), is formulated as follows: 77% acid-washed sand, 22% kaolinite clay, and 1% calcite. The other formulated sediments differed from the SB composition by the presence or absence of iron oxides or by the presence or absence of peat. The second sediment called SG corresponds to the basic sediment coated with 2 g.kg\(^{-1}\) goethite. The third sediment named SP corresponds to the basic sediment amended with 5% peat (equivalent to the so-called standard OECD 207-218). The fourth sediment named SPG and is composed of both peat and goethite.

The quartz and clay were obtained from VWR-France, and the crushed calcite was purchased from La Marchande de Couleurs, (Lyon-France). The peat is composed of blond sphagnum and was purchased from ASB Grünland Helmut Aurenz GmbH (Porschestr.4, D-71634 Ludwigsburg, Germany). The iron oxides were synthesized according to the procedure described in [18].
Solution composition: The solution used for the humidification of the sediment is the nutrient solution recommended for the sediment contact test in the standard for the M. aquaticum bioassay (Steinberg revised medium, ISO 20079). This nutrient solution is composed of the following (µM): KNO₃ (3462), Ca(NO₃)₂ (1866), KH₂PO₄ (662), K₂HPO₄ (75), MgSO₄ (67), H₃BO₃ (2), FeCl₃ (5), MnCl₂ (2), ZnSO₄ (1), Na₂MoO₄ (0.2) and EDTA (4.5). Because EDTA is known to be a complexing agent for metals, we evaluated the impact of this compound on copper availability by varying the EDTA concentration (0 or 0.45 µM). A “(E)” was added to the name of the sediment to denote the addition of EDTA.

Preparation of the sediments: The dry components of the sediment were mixed and humidified with the nutrient solution for 4 hours. The sediments were then spiked with copper by the addition of a Cu(NO₃)₂ solution to obtain four different copper-sediment concentrations: 5, 15, 50 and 200 mg.kg⁻¹ (values added to the name of the sediment; see below). After the metal addition, the sediments were agitated for 24 hours, and the pH was regularly checked and adjusted if necessary. The sediments were distributed into 15 ml beakers for the bioassay and into 110 ml beakers for the DGT experiments; the sediments were then equilibrated for 7 days at a controlled temperature (24 ± 0.5 °C). Only the combinations of some of the different components of the sediment were tested: SB5, SB15, SB50, SB50(E); SP5, SP15, SP50, SP50(E), SP200(E); SG50, SG50(E), SG200(E); SPG50, SPG50(E) and SPG200(E).

1.2 Myriophyllum aquaticum experimental protocol

Plant material and bioassay experiment

The stock culture of Myriophyllum aquaticum was obtained from the UFZ institute (HemlholtZ Zentrum Für Umweltforschung). During the culture period before the bioassay, the plants were grown in an artificial sediment formulated following the OECD 207-218 protocol and saturated with the Steinberg medium (DIN EN ISO 20079).

Prior to the start the bioassay experiment, plants whorls were selected on the basis of their fresh biomass (25 ± 6 mg) by cutting whorls from 21-day-old plants. Each whorl was then placed into a beaker containing 15 ml of sediments. For each test, 25 whorls were exposed to copper-contaminated sediments, and 10 whorls were exposed to the control sediment (without copper). All tests were performed in a growth chamber (exposure light/dark cycle, 16/8 h at 24 ± 1°C at a light intensity of 60-75 µE.m⁻².s⁻¹). The pots were irrigated every 48 h with the semi-concentrated Steinberg medium ISO 20079 (draft ISO/DIS 16191). After 14 days of exposure, the plants were harvested and washed in ultrapure water, washed with an EDTA solution (1 mM) and finally rinsed three times with ultrapure water [19]. The final fresh weight of each plant was recorded. The pH of the sediments was measured at the beginning and the end of the experiment. After pooling several beakers, the interstitial water was recovered by centrifugation of the sediments (3500 rpm, 20 min), filtered at 0.45 µm and acidified with 1% HNO₃ (suprapur).
**Effect (toxicity) assessment:**

The growth rate (GR for control and GR for test) during the exposure period was calculated from the initial and final weight (M and M, respectively) of each fresh whorl (eq. 1). This parameter has been shown to be the most suitable for the test due to the low coefficient of variation of this parameter [5]. The criterion of toxicity was expressed as the percentage of inhibition and is given by (eq. 2).

\[ GR = \frac{(\ln M_{\text{final}} - \ln M_0)}{\Delta t} \quad \text{(eq. 1)} \]

\[ I(\%) = \left( \frac{GR - GR_t}{GR} \right) \times 100 \quad \text{(eq. 2)} \]

**Copper accumulation:**

After harvesting, the plants (i.e., roots and leaves) were dried at 60 °C until a constant weight was reached. To have sufficient biomass for analysis, the plants were pooled (eight per pool) to obtain three replicates per experiment. Samples were digested in a 5:1 ratio of HNO3 and H2O2 in a microwave oven according to a similar protocol to this of Bervoets and co-workers [20] and the copper concentration in the digests were measured by ICP-OES or SAA furnace. In addition, blanks were used, and a reference material (BCR60) was used to control the efficiency of the mineralization.

**1.3 DGT kinetic experimental protocol**

DGT kinetic experiments were performed using DGT units (piston-type with a 2-cm-diameter window loaded with a Chelex-100 resin and with an open pore gel (0.76 mm) purchased from DGT Research (Lancaster, UK; [http://www.dgtresearch.com](http://www.dgtresearch.com))). The DGT units were gently pressed into plastic beakers containing 120 ml of sediment until the shoulder of the piston was in contact with the surface of the sediment housing. The devices were retrieved from the sediments at different contact times: 4, 8, 24, 96 hours and 14 days. Three replicates were performed for each time. After retrieval, the DGT units were thoroughly rinsed with ultrapure water and stored at 4 °C before the elution of the DGT resins with 5 mL of 1 M HNO3 (Merck, suprapur). For each contact, the pore waters of each beaker were extracted following the same procedure as that for bioassay experiments. pH and Eh measurements in the sediment were performed each time.
1.4 Modeling

DGT-PROFS modeling

The accumulation kinetics of metals to DGT were interpreted using the DGT-PROFS model, which is described in detail in [14]. Only the main features of this model are provided here. Briefly, the DGT-PROFS model assumes that trace metals in sediments can be distributed between three separate phases: pore water (where speciation influences the diffusion coefficient value) and weak and strong sorption sites on the particulate phase. The accumulation of metals on DGT is governed by the adsorption-desorption kinetics and the diffusion of metals within the pore water of the sediment and the gel. Only labile metals (free metals and small inorganic complexes able to dissociate in the gel) are assumed to diffuse in the DGT gel. Thus, diffusion of the different metal species through the sediment and gel layer (i.e., free, inorganic and humic species) generates ‘apparent’ diffusion coefficients in the sediment and in the diffusive gel lower than those of the free metals [21]. The diffusion coefficient can range between a minimum value, corresponding to humic complexes, and a maximum value, corresponding to free ions. For DGT-experimented sediment, the DGT-PROFS model was allowed to fit the following parameters: $\alpha_{\text{weak}}$, the proportion of particulate metal associated to weak sites before DGT deployment; $k'_{\text{des1}}$, the desorption rate from weak particulate sites; $k'_{\text{des2}}$, the desorption rate from strong particulate sites; and $D_{\text{sed}}$, the diffusion coefficient of metals in sediment. These parameters can be obtained by fitting the DGT experimental measurements using a probabilistic approach, which represents parameters by probability density functions (PDFs) (i.e., with indications of their uncertainty) instead of single values [14].
2. Results

2.1 Physico-chemical characteristics of the different sediments:

The variation of the total pore-water copper concentration ($C_{(iw)}$) over the duration of a given test ranged between 3% to 10% except in the cases of SG50, SG50(E) and SG200(E), which had variations of 13, 14 and 19%, respectively. These features allow us to consider the equilibration with the solid phase to be complete after a sediment incubation time of 7 days (Table 1).

The average values of the ($C_{(iw)}$) varied from 14.4 µg.l$^{-1}$ to 725 µg.l$^{-1}$ for the DGT experiments and from 24 to 1510 µg.l$^{-1}$ for the bioassays (Table 1). Generally, for the same composition of sediment, copper pore-water concentrations are higher in pots containing plants than in pots containing DGT. This result can mainly be explained by the higher evaporation due to the smaller and less hermetic pots used for the plant exposure. Indeed, a decrease in the supernatant water thickness was observed in the bioassay pots, which was partly compensated by using irrigation as recommended by the protocol (draft ISO/DIS 16191). According to the sediment composition, the lowest Cu$_{(iw)}$ concentrations were found in sediments coated with goethite, and higher concentrations were found in sediments formulated with peat. Sediments formulated with both goethite and peat show intermediate values.

The presence of EDTA in the sediment results in much higher Cu$_{(iw)}$ concentrations compared to tests without EDTA.

The pH values ranged from 5.76 to 7.17 and from 5.17 to 7.22 in the different sediments tested in beakers with DGT probes and M. aquaticum, respectively (Table S1 Supporting Information). The lowest pH values were observed in the sediments formulated with peat; these values are in agreement with the naturally low pH of peat and its good buffering capacity against pH changes, which makes it difficult to obtain a pH around neutrality. The pH did not vary significantly during a given experiment (≤0.1 unit pH). The redox potential shows that all the experiments were performed under oxic conditions (Table S1 Supporting Information). The DOC concentrations in the “SP” sediments are very high and show that a part of the peat was dissolved and solubilized in pore water. In the case of the sediment composed of organic matter and goethite, the DOC concentration is slightly lower than in “SP”; we can suppose that a part of the dissolved carbon was fixed on oxides. Finally, in the bioassay experiment, there is an increase in the DOC concentrations, which is certainly related to the evaporation process. The copper distribution between the solid phase and the pore water can be assessed by a partition coefficient (Kd). For the DGT experiment, the Kd values (Table 1) are on the same order of magnitude in the different sediments, with values ranging from 100 to 500 except for GS-50, for which there is a significant increase in the Kd value up to around 2900. In the presence of EDTA, Kd exhibited no marked differences.
2.2 Sediment contact test with Myriophyllum aquaticum

The growth rates of *Myriophyllum aquaticum* (Figure 1) exposed to control sediments agree with the standard for this bioassay (growth rate ≥ 0.09 day\(^{-1}\)), which demonstrates the applicability of the bioassay in sediments with different physico-chemical characteristics. For the different control sediments, lower growth rates are observed in the presence of iron oxide (SG and SPG) when compared to the basic sediment (SB). Conversely, the highest growth rate was observed for the sediment formulated with peat (SP - OECD reference sediment), which is in agreement with the results obtained by [5]. Furthermore, the growth rates for the control sediments containing EDTA were systematically higher than those in the control sediments without EDTA. This variation in growth rates leads to a strong variation in the increase of plant biomass over the 14 days of exposure. For example, with the control plants exposed to the sediments SPG and SP, the biomass increased by approximately 250% and 600%, respectively. It was observed that growth (rate) inhibition (I (%)) depends on the copper concentration for a given sediment and on the sediment composition (Figure 1). The inhibition range from 6 to 20% for the sediment containing peat (OECD reference sediment) spiked from 5 to 50 mgCu.kg\(^{-1}\) (SP5, SP15, SP50), and that from 23 to 34% for the basic sediment spiked from 5 to 50 mgCu.kg\(^{-1}\) (SB5, SB15, SB50). The presence of EDTA induced an inhibition ranging from 16 to 100%. For the different sediments formulated without EDTA and spiked at 50 mg.kg\(^{-1}\), there is a higher toxicity in the sediments formulated without peat, SB and SG (I=34 and 32%, respectively), than in the peat-containing sediments, SP and SPG (I=25 and 15%, respectively). The increase in plant biomass for the spiked sediments SG200E and SP15 was 4 and 570%, respectively, which demonstrated the wide range of toxic effects in the sediments tested.

2.3 Copper internalized concentration in *M. aquaticum*

Similar to the growth inhibition, the copper bioaccumulation in *M. aquaticum* depends on the copper concentration and the sediment composition (Table 1). For example, when comparing the basic sediment spiked with either 5 or 50 mg Cu.kg\(^{-1}\) (SB5 and SB50), the Cu accumulation in plants was observed to increase from 44 to 124 µg.g\(^{-1}\). For sediments formulated with iron oxide or peat, the bioaccumulated copper concentrations varied from 72, 32 and 29 µg.g\(^{-1}\) for SP50, SG50 and SPG50, respectively. The presence of EDTA homogenizes the copper bioaccumulation among sediments with concentrations of approximately 60 µg.g\(^{-1}\).

2.4 Copper Flux to DGT

Sediment composition has a strong influence on copper fluxes as indicated by the comparison of the measured fluxes of copper to the DGT resin (expressed as mg.cm\(^{-2}\).s\(^{-1}\)) over the duration of the test for four of the different formulated sediments tested (Figure 2). For the four experiments performed without EDTA, the highest fluxes to DGT were obtained in the basic sediment spiked with 50 mg
(Cu).kg\(^{-1}\) (SB50), and significantly lower fluxes were observed in the other sediments formulated with peat and goethite and spiked with 50 mg (Cu).kg\(^{-1}\) (SG50, SP50 and SPG50). In addition, two curves in the fluxes are observed depending on the composition of the sediment: (i) a rapid decrease of the flux during the first 96 hours of deployment followed by a continuous decrease with a lower slope during the rest of the test and (ii) a decrease of the flux during the initial hours followed by a stabilization of the flux from 24 h to the end of the experiment. The mean and the 5\(^{th}\) and 95\(^{th}\) percentile simulated fluxes using the DGT-PROFS model are also given in the Figure 2. The good agreement between the simulated and measured fluxes enhances the relevance of the modeling and fitting procedure.

The fluxes of copper to DGT for the other sediments are available in the supporting information (Figure S1). Briefly, the shape of the fluxes remains the same for each sediment composition regardless of the variation of total copper concentrations or the addition of EDTA. However, an increase in the spiking concentration as well as the addition of EDTA leads to an increase in the flux values. The SB50 (E) and SG50 (E) experiments were not simulated because too much uncertainty exists for certain data.

3. Discussion

One aim of the study was the comparison of the copper accumulation in plants and in the passive sampler DGT. However, due to the different constraints of the exposure conditions between the bioassay and DGT experiments, the physico-chemical conditions diverged for a given type of sediment. It was therefore necessary to correct the DGT data to better mimic the conditions of the bioassay. For this purpose, assuming that the key parameters \(\alpha_{\text{weak}}, k_{\text{des1}}, k_{\text{des2}},\) and \(D_{\text{sed}}\) obtained from the DGT-PROFS modeling were constant for a given test, the copper pore-water concentration for a given bioassay was used to simulate the corresponding accumulation in DGT under this slightly modified condition (designated as corrected values hereafter).

3-1 Copper lability and resupply from solid phase

Figure 3 presents the theoretical DGT fluxes relative to the solution concentration (using DIFS modeling) as a function of the Kd for fast (full line, Tc 10 s) and slow desorptions (Tc 1000 s) obtained from Degryse and co-workers [9] for DGT sampling at 24 h, in which the data of the present study were added. Three different trends can be observed according to the kinetic characteristics: slow, intermediate and fast kinetics of desorption. Fast desorption kinetics were obtained for sediments composed without peat, and slow desorption kinetics were obtained for sediments containing peat. However, as the DIFS model considers all complexes in the solution to be fully labile, the resupply from the solid phase may be underestimated in the presence of complexes in the solution that are not fully labile as copper binds to organic matter. Therefore, the desorption kinetics from sediments...
containing peat may be underestimated. Indeed, this representation gives information on the solid
phase behavior for a given time (24 h) and does not take into account its evolution over the time of the
experiment. Therefore, the description of the fluxes’ evolution over time gives supplementary
information.

Two different trends of the fluxes of copper to the DGT were observed for the different sediments
tested: constant and decreasing fluxes (Figure 2 and Figure S1 in Supporting Information). Harper and
co-workers [13] proposed a classification of fluxes to quantify the “copper lability” according to four
typical cases. In our study, we observed the “partially sustained case” for all sediments containing peat
(SP and SPG) and the “non-steady state case” for all basic sediments and for the sediments coated
with goethite (SB and SG). For sediments belonging to the partially sustained case, the desorption
capacity of the metal from the solid phase is sufficient to maintain a constant flux of copper to the
DGT device (at least over the time experiment). For sediments belonging to the non-steady state case,
the copper flux to the DGT decreases over time, indicating that the resupply from the solid phase is
slow. Considering the intensity of the fluxes, significantly high fluxes during the 24 first hours were
obtained in the basic sediments. The basic sediments are principally composed of quartz and kaolinite.
We can therefore assume that the high initial flux is due to a high initial concentration of the available
copper in solution ($C_{iw}$). The following decline of the fluxes may be related to a slow resupply due to
copper adsorption to strong bonding-energy sites (kaolinite) [22]. In the case of the sediments coated
with goethite, similar curves were observed but with significantly lower fluxes. Indeed, it is well
known that iron oxides have a high affinity for metals and that the addition of iron increases the
fraction of copper associated to strong sites. As the pH of the sediment is lower than the zero point
charge of goethite (7.9), goethite is positively charged [18]. Under these conditions, copper adsorption
on goethite results from the formation of inner-sphere complexes (chemisorptions), leading to a strong
fixation. The sediments containing peat also lead to significantly lower fluxes than in the basic
sediment. The lower values of fluxes can be related to a lower available pool of copper in the system.
Moreover, these lower values are also due to a lower velocity of organic-copper complexes in the pore
water ($D_{sed}$) and the fact that organo-metallic complexes composed of fulvic or humic acids are only
partially labile [23]. The slight decrease of the fluxes followed by a rapid stabilization over time
suggests that the peat have the ability to resupply copper easily to the pore water. Concerning the other
set of experiments in the supporting Information, the same behavior is observed for the same sediment
composition with higher fluxes in the presence of EDTA and for increasing copper spiking
concentrations related to higher available pools of dissolved copper in the sediments.

All of these qualitative features are confirmed by the values of the DGT-PROFS model parameters
that were fitted for each experiment (Table S2 in Supporting Information). The $\alpha_{weak}$ parameter, which
represents the proportion of particulate metal associated to weak sites, is much lower for the SG50
sediment (0.06) than for the SP50 or SPG50 sediments (0.37 and 0.2), which confirms that the ability
to resupply copper to the water phase is lower with iron oxide addition. The lowest value of the $D_{sed}$
3.2 Comparison of different tools to assess copper bioavailability

To relate the "actual bioavailability" measured by the plant *M. aquaticum* to the "potential bioavailability" estimated by different chemical measurements in the sediment, including DGT measurements, we first investigated whether a significant linear correlation could be observed between the copper concentration in plant and (i) the total dissolved copper concentration, (ii) the free copper concentration, and (iii) the mean labile concentration derived from DGT (*C*$_{DGT}$) in the sediment (Figure 5). The comparison between the total dissolved copper concentrations with the bioaccumulation in the *M. aquaticum* results in a poor correlation, with a $R^2$ of 0.05 indicating that the total dissolved concentration does not represent the readily bioavailable fraction for plants. The free copper concentration in the pore water was obtained for each sediment by using the speciation program WHAM VI [24] (details in Table S3 in Supporting Information). The comparison between the free copper concentrations and the copper concentration in plant also showed a poor correlation, with a $R^2$ of 0.08. Finally, the comparison between the DGT measurement and the copper concentration in plants resulted in an improved, but still poor, correlation, with a $R^2$ of 0.6 (Figure 3).

One reason for the above poor correlations might be the presence of growth inhibition in the sediments tested depending on the composition and copper content of the sediments (Figures 1 and 4). Indeed, in the literature, good correlations between the DGT-measured labile copper and the accumulation in some plants were obtained only in the cases of plants exposed to non-toxic concentrations [9, 16, 25]. Assuming that the growth rate is maximal for low copper concentrations until a threshold where toxic effects arise and that a zero growth rate is reached at high copper concentrations, a theoretical relation between the DGT labile concentrations and the growth rate can be drawn (Figure 4; mathematical relation (Eq. S1) in supporting information). This predicted relation between the DGT and the growth rate agrees with our experimental results (Figure 4) and shows that all the sediments tested are in the range of concentrations inducing a toxic effect on the plant, with high growth rate variations. Indeed, *M. aquaticum* growth began to decrease when the DGT was above 92 μM. However, there is a high variation in the DGT and growth rate measurements between the different sediments tested, which is certainly related to the difference in the copper uptake by the plants according to the toxicity of the sediment compositions.

The relationship between the plant uptake and the fluxes recorded by the DGT can help to determine the ability of DGT to mimic the plant uptake. Degryse and co-workers [9], reported such relationships for the uptake of Zn and Cd by watercress and spinach, respectively (Figure 5). Plotting our results for copper in the case of *M. aquaticum* on Figure 5 shows that the plant uptake fluxes do not increase in a
directly proportional manner with DGT fluxes, i.e., a 10-fold increase in DGT flux does not result in a
10-fold increase in plant uptake fluxes or plant concentrations. Considering that DGT mimics the
action of the plant only under diffusion-limited conditions, the absence of correlation suggests that Cu
uptake is not diffusion-limited in our case. If there is no diffusion limitation, labile complexes (that are
not taken up) will not contribute to the plant uptake, but the uptake will be governed by the free metal
ion activity and the activity of competing ions (e.g., Ca$^{2+}$, Mg$^{2+}$, and H$^+$; cf. BLM concept) [9]. In the
present study, the free Cu (Cu$^{2+}$) was low, especially in the presence of peat (Table 1). The apparently
low Cu$^{2+}$ should promote conditions for a diffusion limitation that are not observed. In fact, there is
most likely a large contribution of labile complexes leading to a continuous availability of sufficient
Cu$^{2+}$ for the plant uptake, which results in no diffusion limitation conditions.

Moreover, this study showed that in the wide range of sediments tested, the normalized bioassay was
sensitive to copper and was able to accumulate it. These data provide support for other studies that
have shown a strong sensitivity of *M. aquaticum* toward sediments contaminated with metals [8]. Until
now, the *M. aquaticum* was commonly used in toxicity tests on pesticides [26, 27]; thus, these results
show the interest of this new bioassay for the assessment of risks associated with metal contamination
and the need to go further in the knowledge of its sensitivity towards metals. The recently normalized
sediment biotest using *M. aquaticum* appears to be a sensitive tool for the assessment of copper-
contaminated sediments; however, further research is needed to assess this test’s sensitivity toward
other metals.

**Supporting Information.**

Details on physico-chemical characteristics of pore water, flux of copper to DGT versus time and
modeling using the DGT PROFS model, DGT PROFS model parameters, WHAM VI modeling of the
different sediments investigated and on the plant growth rate modeling.

**Acknowledgements:** This study was funded by EDF (Eléctricité de France) and the Agence National
de Recherche (ANR, FRANCE; ANR-07-ECOT-0794C0111). The PhD grant of Amélie Caillat was
funded by EDF with a grant attributed by ANRT (CIFRE 874/2009)
References


Table 1: Copper pore-water concentrations (C_{iw}; µg.L^{-1}) in DGT and bioassay experiment, free copper concentration (Cu^{2+}; µg.L^{-1}) and copper accumulated in M. aquaticum in the different sediments (µg.g^{-1}) for the different sediments studied (see text for nomenclature). “DGT” stands for DGT experiment, and “Bioassay”: stands for bioassay experiment. Cu^{2+} was calculated with WHAM VI (Tipping (24), see details in the Table S2 in Supporting Information). The values of experimental data represent average values: for all the DGT n=15; for the bioassay experiment, there are less replicates because the measurements was only performed at the end of the experiment; for C_{iw}, n=2 or n=1 (*) and, for bioaccumulation, n=3.

<table>
<thead>
<tr>
<th>Sediment</th>
<th>C_{iw} &quot;DGT&quot; (µg.L^{-1})</th>
<th>C_{iw} &quot;bioassay&quot; (µg.L^{-1})</th>
<th>Cu^{2+} &quot;DGT&quot; (µg.L^{-1})</th>
<th>Cu bioaccumulated (µg.g^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>SB-5</td>
<td>44 ± 2</td>
<td>58 ± 4.6</td>
<td>3,7E-02</td>
<td>44 ± 4</td>
</tr>
<tr>
<td>SB-15</td>
<td>40 ± 3</td>
<td>60 ± 4</td>
<td>2,5E-02</td>
<td>49 ± 3</td>
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<tr>
<td>SB-50</td>
<td>96.4 ± 10.1</td>
<td>157*</td>
<td>2,6E+00</td>
<td>124 ± 8</td>
</tr>
<tr>
<td>SP-5</td>
<td>14.4 ± 1</td>
<td>34*</td>
<td>4,1E-06</td>
<td>50 ± 8</td>
</tr>
<tr>
<td>SP-15</td>
<td>38 ± 0.6</td>
<td>50 ± 1</td>
<td>1,4E-04</td>
<td>71 ± 3</td>
</tr>
<tr>
<td>SP-50</td>
<td>104.1 ± 6.8</td>
<td>144.2*</td>
<td>2,2E-03</td>
<td>72 ± 6</td>
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<tr>
<td>SG-50</td>
<td>17.3 ± 2.3</td>
<td>23.9*</td>
<td>1,9E-01</td>
<td>32 ± 5</td>
</tr>
<tr>
<td>SPG-50</td>
<td>83.5 ± 2.9</td>
<td>84.9*</td>
<td>7,2E-04</td>
<td>29 ± 5</td>
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<tr>
<td>SB-50(E)</td>
<td>236 ± 18</td>
<td>275 ± 5</td>
<td>5,5E-02</td>
<td>61.3 ± 11</td>
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<td>SP-50(E)</td>
<td>295 ± 8.1</td>
<td>560 ± 9</td>
<td>2,8E-03</td>
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<td>SG-50 (E)</td>
<td>148 ± 21.4</td>
<td>190*</td>
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<tr>
<td>SPG-50 (E)</td>
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<td>402*</td>
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<td>SP-200(E)</td>
<td>620 ± 25.7</td>
<td>1510 ± 61</td>
<td>1,5E-01</td>
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<tr>
<td>SG-200 (E)</td>
<td>725 ± 141</td>
<td>299 ± 2</td>
<td>8,3E+01</td>
<td>163 ± 14</td>
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<td>SPG-200 (E)</td>
<td>478 ± 30</td>
<td>510 ± 7</td>
<td>8,5E-02</td>
<td>60 ± 12</td>
</tr>
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</table>
**Figure Captions**

**Figure 1**: Growth rate (days\(^{-1}\)) and growth inhibition (%) of *Myriophyllum aquaticum* after exposure to control and spiked sediments. “SB” corresponds to the basic sediment, “SG” to the sediment coated with goethite, “SP” to the sediment formulated with peat and “SPG” to the sediment formulated with peat and goethite. “E” represents sediments formulated with EDTA. The errors were calculated from n=10 for controls and n=25 for contaminated samples.

**Figure 2**: Copper fluxes resupply (mg.cm\(^{-2}\).s\(^{-1}\)) from particulate phases in sediments SB, SP, SG, SPG spiked at 50 mg.kg\(^{-1}\) (see text and Figure 1 for nomenclature) as a function of time. The replicates of the measured fluxes are represented by the three different open symbols. The full line represents mean flux calculated by the DGT PROFS model. The upper and lower dotted lines represent the 5\(^{th}\) and 95\(^{th}\) percentiles respectively.

**Figure 3**: Flux to DGT relative to copper pore-water concentration (F\(_{\text{DGT}/C_{\text{pw}}}\)) as a function of K\(_d\) (L.kg\(^{-1}\)) for all the sediments studied in the presence of peat (full triangle) or absence of peat (empty diamond); (see text and Figure.1 for nomenclature). The full and dotted lines represents theoretical data for slow (dotted line) and fast desorption kinetics obtained from\(^8\).

**Figure 4**: Relationship between growth rate and DGT copper concentrations (μM) for all the sediments studied in the presence of peat (full triangle) or absence of peat (empty diamond); (see text and Figure.1 for nomenclature). The full line represents the theoretical relation between the corrected DGT concentration (see text) and the growth rate (mathematical equation (Eq. S1) is presented in Supporting Information) based on several assumptions developed in the text.

**Figure 5**: Relationship between fluxes to DGT (pmol.cm\(^{-2}\).s\(^{-1}\); 14 days, corrected values; see text) and copper accumulation in plants (mmol.kg\(^{-1}\)) for all the sediments studied in the presence of peat (full triangle) or absence of peat (empty diamond); (see text and Figure.1 for nomenclature). The full line represents the theoretical limiting diffusive flux. The dotted line and the black circles represent the theoretical plant demand and the theoretical plant uptake respectively\(^8\). The circles represent zinc uptake by watercress (*Lepidium sativum*) and DGT measurements determined for a range of soils sampled near galvanized structures\(^26\). The crosses represent cadmium uptake by spinach and DGT measurements at varying NaCl doses at background cadmium concentration or soils amended by Cd-salt\(^27\).
Figure 1

Figure 2