DGT-induced copper flux predicts bioaccumulation and toxicity to bivalves in sediments with varying properties

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Abstract
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Keywords
induced, copper, flux, predicts, dgt, properties, sediments, varying, bioaccumulation, toxicity, bivalves, CMMB

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DGT-INDUCED COPPER FLUX PREDICTS BIOACCUMULATION AND TOXICITY TO BIVALVES IN SEDIMENTS WITH VARYING PROPERTIES

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ABSTRACT: Many regulatory frameworks for sediment quality assessment include consideration of contaminant bioavailability. However, the ‘snap-shots’ of metal bioavailability provided by analyses of pore waters or acid-volatile sulfide - simultaneously extractable metal (AVS-SEM) relationships do not always contribute sufficient information. The use of inappropriate or inadequate information for assessing metal bioavailability in sediments may result in incorrect assessment decisions. The technique of diffusive gradients in thin films (DGT) enables the in situ measurement of metal concentrations in waters and fluxes from sediment pore waters. We utilised the DGT technique to interpret the bioavailability of copper to the benthic bivalve *Tellina deltoidalis* in sediments of varying properties contaminated with copper-based antifouling paint particles. For a concentration series of copper-paint contaminated sandy, silty-sand and silty sediment types, DGT-probes were used to measure copper fluxes to the overlying water, at the sediment-water interface and in deeper sediments. The overlying water copper concentrations and DGT-Cu fluxes were shown to provide excellent exposure concentration-response relationships in relation to lethal effects occurring to the copper-sensitive benthic bivalve, *T. deltoidalis*. The study demonstrates the strength of the DGT technique, which we expect will become frequently used for assessing metal bioavailability in sediments.

Keywords: DGT, copper, antifouling paint, sediment, bivalve, bioaccumulation, ecotoxicology.
Introduction

The technique of diffusive gradients in thin films (DGT) was developed to enable the in situ measurement of metal concentrations in waters and fluxes from sediment pore waters. In a DGT device, dissolved metal species diffuse through a polyacrylamide gel layer and become trapped in a gel impregnated with a metal-exchange resin, which acts as a metal sink. The removal of metals from sediment porewaters causes the concentration to decline immediately adjacent to the device. This localised decline could disturb the dynamic equilibrium between the sediment and metal in solution and induced release of metals to solution, the extent of which will depend on the rate of metal resupply from the sediment solid phase to the porewater. If there is rapid resupply, metal concentrations in pore waters may be calculated from the DGT-accumulated metal concentration. However, when resupply from the sediments is limited the DGT-flux provides information on the relative rate of remobilisation of metals from sediments to the pore water. Hence the DGT directly measures the flux of metal from the sediment during the deployment time, which reflects the concentration in the porewater, its diffusional transport and the supply from the solid phase to solution. It can be interpreted simply as the average porewater concentration at the interface of the device over the deployment period. The ability of DGT measurements to provide information on the localised remobilisation of metals has been utilised to create high resolution depth profiles of metals within sediments. Such studies have demonstrated the heterogeneity of sediment environments; including the existence of microniches and to characterise zones of metal remobilisation.

Existing methods for estimating the bioavailability of metals in sediments to benthic organisms have numerous limitations. For example, the usefulness of chemical extractions that provide information on metal lability varies between metals and the degree of contamination and for the organism being studied. Further to this, the equilibrium partitioning relationships between metals and acid-volatile sulfide (AVS) become less appropriate for organisms that create oxic/sub-oxic micro-environments at burrow walls within sulfidic sediments, and pore water analyses provide non time-integrated ‘snap-shots’ of concentrations that change rapidly upon sediment disturbance.

The DGT-accumulated metal concentration has been determined to consist of free metal ions, metal ions present as simple inorganic complexes, and labile organic complexes. As these metal fractions represent the most bioavailable metals, the time-integrated measure of the bioavailable metal provided by the DGT technique is proving to be a powerful tool for assessing metal bioavailability in waters. In sediments, the bioavailability of metals to
benthic organisms is influenced by the characteristics of the metal, the properties of the sediments and the varying exposure pathways of the organisms.\textsuperscript{25-28} The DGT technique was used by Wegener et al.\textsuperscript{29} to demonstrate that sediment bioturbation and feeding by the oligochate \textit{Tubifex} modified porewater metal concentrations within surface sediments. The release of metals from sediment particles to pore water will be more rapid for sediments that contain reactive forms of metals when compared to sediments that contain more inert forms of metals, hence differences in DGT-induced metal fluxes are expected to provide a useful measure of the bioavailability of the metals in sediments. Roulier et al.\textsuperscript{30} observed total and DGT-accumulated metal concentrations both provided strong relationships with Cu and Pb bioaccumulation from sediments by \textit{Chironomus riparius}, however, only the DGT technique provided an explanation of different \textit{organismal} uptake for sediments with similar total metal concentrations.

In the present study we utilise the DGT technique to interpret the bioavailability of copper to the benthic bivalve \textit{Tellina deltoidalis} in sediments contaminated with copper-based antifouling paint. Copper-based paints are increasingly used by marine industries to reduce biofouling and the abrasion of paint from structures can result in considerable copper concentrations in sediments.\textsuperscript{31} Concentration series of copper-paint contamination were prepared for three sediments, representing sandy, silty-sand and silty environments. DGT-induced copper fluxes were measured in the overlying water, at the sediment-water interface (SWI) and in deeper sediments. The DGT-induced copper flux at the SWI was compared with copper bioaccumulation and toxicity to the bivalve.

**Material and Methods**

**General methods.** All glass and plastic-ware for analyses were usually new and were cleaned by soaking in 10\% (v/v) HNO\textsubscript{3} (BDH, Analytical Reagent grade) for a minimum of 24 h, followed by thorough rinsing with deionized water (Milli-Q, 18 M\textOmega/cm). All chemicals were analytical reagent grade or equivalent analytical purity. Water pH, salinity, temperature and dissolved oxygen measurements were made with probes calibrated according to manufacturer instructions (WTW). Methods for sediment particle size (by wet sieving through 63 \textmu m nylon sieves followed by gravimetry), total organic carbon (TOC, by high temperature TOC analyzer), porewater (PW) extraction (centrifugation at 800 g for 5 min), and dissolved ammonia analyses are described in Spadaro et al.\textsuperscript{32}. Methods for analyses of total recoverable metals (TRM, by microwave assisted aqua regia), dilute acid extractable metals (AEM, 1 M HCl) and acid-volatile sulfide (AVS) (all determined on sub-samples of the same homogenised sediment) are described previously.\textsuperscript{33} Biota tissues were freeze-dried

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before microwave-assisted (MARS 5, CEM) acid/peroxide extraction (3:1 HNO₃:H₂O₂, 80 ºC for 50 min). Dissolved metal concentrations in acid-digests of water, biota and sediment samples were determined by inductively coupled plasma - optical emission spectrometry (ICP-OES, Varian 730-ES) as described in Simpson et al. The metal concentrations of the acid-digests of DGT gels were determined by inductively coupled plasma-mass spectrophotometry (ICP-MS, Agilent 7500ce). As part of the quality assurance, analyses of filter and acid-digest blanks, replicates for 20% of samples, analyte sample-spikes and the certified reference materials (CRMs) were performed. Replicates were within 20% and recoveries for spikes and CRMs, PACS-2 for sediment (National Research Council Canada, NRCC) and DORM-3 for biota (fish protein, NRCC), were within 85-115% of expected values. The limits of reporting for the various methods were less than 10% of the lowest measured values.

**Test media and organisms.** Clean seawater was collected from Port Hacking, Sydney, Australia, membrane filtered (0.45 µm), and acclimated to a room temperature of 21±1 ºC. Where necessary, the salinity of the filtered seawater was adjusted to the test salinity of 30 PSU using Milli-Q water. Surface sediments (0-3 cm depth) were collected from three intertidal estuarine locations for use as controls and for preparing sediment with a range of particles sizes as described in Strom et al. Three sediments were prepared with 10% (sandy), 30% (silty-sand) and 60% (silty) <63 µm particles, respectively. *T. deltoidalis* with shell lengths of 5-12 mm were collected from Lane Cove River (NSW, Australia) and maintained as described previously.

**Copper-based paint spiked sediments.** Copper-based antifouling paint (13% copper(I) oxide/4% zinc oxide; Hempnet, Hempel) was dried at 45 ºC for 24 h, snap-frozen in liquid nitrogen, crushed using a Teflon beaker and mortar, and the <180 µm particles isolated using a nylon sieve. Before TRM analyses, the paint samples were heated in a porcelain crucible at 500 ºC for 4 h to destroy the organic matrix. Copper concentration series of paint-spiked sediments were prepared by adding the required amounts of paint to the three sediments. The sediments remained at pH 7.7±0.2 without pH-adjustment and were mixed and equilibrated for a period of 30 days. Copper concentrations are frequently 100-1000 mg/kg in sediments near marinas and aquaculture leases that use copper-based paints on boats, nets and other marine structures. An initial concentration series of 0, 200 and 1000 mg Cu/kg was prepared and tested. After 8 months (stored refrigerated) the 1000 mg Cu/kg sediment from each particle size composition, was diluted with control sediment of the same composition to create sandy sediments with 0, 50, 100, and 200 mg Cu/kg, and silty-sand and silty sediments...
with 0, 100, 200, and 400 mg Cu/kg. These dilution series were mixed and equilibrated for a
further 30 days to achieve dissolved-particulate copper partitioning similar to that occurring in
field-contaminated sediments. At the completion of the equilibration process, sub-samples
of the copper-paint spiked sediments taken for duplicate analyses of pH, dissolved porewater
metals, particulate metals (TRM and AEM), particle size distribution, TOC, and AVS.

Diffusive gradients in thin films (DGT). Plastic DGT assemblies (24×4×0.5 cm, with
1.8×15 cm windows) were purchased from DGT Research (http://www.dgtresearch.com/).
DGT gels were prepared following the procedures recommended by DGT Research (details
are provided in Section S1 of the Supporting Information.). The DGT probe cross-section
comprised: backing plate, 0.45 µm filter membrane, 0.25 mm Chelex gel, 0.5 mm diffusive
gel, 0.45 µm filter membranes, and a front window plate. All components had been carefully
prepared to minimise metal contamination. Deoxygenated DGT probes were gently inserted
into the sediments to a depth of 4 cm with care to ensure a good contact between the sediment
and the DGT membrane. Deployments times of 24 h (DGT equilibration) were selected
based on Garmo et al., as we used 0.5 mm diffusive gel thickness in environmentally
relevant PW-Cu concentrations in the presence of humic acids. After 24 h of deployment,
DGT probes were gently removed from the beakers and position of the sediment-water
interface (SWI) and overlying water depth were recorded. The probes were immediately
rinsed with Milli-Q water to remove sediment particles, then held in clean plastic bags at 4ºC
until disassembly. DGT devices were disassembled (within 10 days of retrieval) and resin gel
slices cut using a Teflon coated blade to obtain the desired vertical profile. For the first DGT
deployment there was a 1.5 cm slice at the SWI (from -0.75 cm to +0.75 cm), two 2-cm slices
in the overlying water above the SWI-slice and 2-cm slice in the sediment above the SWI-
slice. For the second DGT deployment there was a 1-cm slice at the SWI (-0.5 cm to 0.5 cm),
and 1 cm slices above and below the SWI-slice. The mid-point of the slice was set as the
location in data analysis and Figure 1. Slices from below the SWI were washed with Milli-Q
water to remove any sediment particles. Each slice was weighed, then eluted with 1 M HNO₃
for 16-24 h before analysis by ICP-MS. Undeployed DGT probes were analysed as handling
blanks and their copper concentrations were less than 5% of the lowest measured
concentration.

Bivalve bioaccumulation bioassays. The test sediments (400 mL) and overlying water
(500 mL) were added to 1 L Pyrex beakers (washed sequentially with phosphate-free
detergent, 1% HNO₃ and Milli-Q water) five weeks prior to the start of the tests and kept in a
temperature controlled lab (21±3°C, normal day light conditions). The beakers were
bubbled with air during this period and throughout the tests, and developed stratified redox profiles. On the day of test commencement the overlying water was removed to a level of 2 cm above the sediment surface, 10 bivalves were added to each beaker, and then new seawater was added. Water pH (pH 8.2±2), salinity (31±1‰), temperature (21±1°C,) and dissolved oxygen concentration (>80% saturation), and water levels were monitored throughout the tests. The overlying water was changed 3 times per week after sub-sampling for analyses of dissolved copper and ammonia. The DGT probes were inserted on days 8 and 15, and removed 24 h after deployment. The tests were terminated after 30 days and surviving organisms were counted, and allowed to depurate overnight in clean seawater before refrigeration. The bivalve shells were opened using a Teflon coated razor blade and the soft tissue transferred to a pre-weighed 50 mL polycarbonate vial using plastic tweezers and then frozen at -20°C before analyses.

Statistical analyses. At the end of the bioaccumulation bioassays the percent survival in tests relative to survival in the controls was recorded. All statistical analyses were carried out using the software Toxcal (Version 5.0.23, TidePool Scientific Software). Unless otherwise stated $p = 0.05$ was the level of significance. Effect concentration causing x% of the lethality to the bivalve (LC$_x$) were estimated from concentration-response data and corresponding copper concentration using linear interpolation as described previously.$^{35,37}$ Correlations (Pearson’s product-moment) were performed on the particulate metal concentrations using statistical software (NCSS).

Results and Discussion

Properties of the paint-spiked sediments. The TRM copper and zinc concentrations in the paint were 9.3±0.7 mg/kg and 3.0 ±0.2 %, respectively. These compared to concentrations of 13±1 %-Cu and 4.0±1 %-Zn, specified by the manufacturer$^{43}$ and were similar to the compositions of most copper-based antifouling formulations.$^{44}$

The properties of the paint-spiked sediments are shown in Table 1. The pH of the sediments ranged from 7.7-7.9; exhibiting no relationship with spike concentration or sediment type. The sandy, silty-sand and silty sediments had TOC concentrations of 0.7, 1.6, 3.1%, respectively, and particle size distributions within 10% of the nominal 10, 30 and 60% <63 µm particles. Less than 4% of each sediment contained 63-180 µm size fraction and the remainder was mostly 180-1100 µm sized sand. The AVS concentrations were <0.5 µmol/g for all the sediments. For total copper concentrations (TR-Cu) were within 25% of the nominal concentrations. Dilute acid-extractable copper concentrations (AE-Cu) comprised
48% to 94% (mean±SD = 68±10%) of the TR-Cu, and the copper extractability of the 200 mg/kg AE-Cu remained relatively constant over the 8-months between the 1st and 2nd series. This range of copper extractability was significantly greater than we have found for field-collected sediments contaminated with copper-based antifouling paints, where we have measured AE-Cu was 6±9% of TR-Cu for 67 samples (unpublished results). Porewater copper concentrations (PW-Cu) increased with increasing spiked-copper concentrations, and very high PW-Cu (6±0.9 mg/L) was measured for the 1000 mg Cu/kg sandy treatment. The silty-sand and sand PW-Cu concentrations were markedly greater in the 2nd series (0, 100, 200, 400) than in the 1st series (0, 200, 1000) (Table 1). This is most likely related to the Cu released from the copper-based paints over the 8 months, which would have been removed more effectively from the silty-sediment porewaters than the silty-sand and sand due to the higher concentrations of TOC and finer particles. The time-averaged dissolved copper concentrations in the overlying water (OW-Cu) also increased with increasing spiked-copper concentration. However, the 1st and 2nd silty-sand series did not display anomalously high values, as observed for PW-Cu.

**Copper profiles in sediments by diffusive gradients in thin films (DGT).** The principal characteristics of the DGT-Cu flux depth profiles on days 8 and 15 for the three sediment types (Figure 1) were: (i) peaks at the sediment-water interface (SWI), (ii) increased DGT-induced copper flux with increasing spiked-copper concentration, and (iii) lower DGT-induced copper flux in more silty sediments (sand > silty-sand > silt). The peak in the DGT-induced copper flux near to the SWI is consistent with previous studies\(^9\) and has generally been attributed to its release from organic matter as it is oxidised.\(^{45,46}\) Depth profiles of Fe and Mn in the pore water were not obtained for the present study. However subsequent experiments using sediments with similar properties to the silty-sand that had been spiked with a copper mineral phase (unpublished results) indicated that the expected porewater Fe and Mn profiles were established rapidly after sediment disturbance (from collection and spiking) and 1 month equilibration (See Figure S1, Supporting Information). The mobilisation of copper through the reductive dissolution of Fe and Mn oxyhydroxide phases near the SWI is likely to be a major process influencing the copper fluxes.\(^{19}\) The very slow rate of copper release to the pore water and overlying water in the siltier sediments was consistent with the greater concentrations of TOC and fine particles, which provide copper-binding sites, in the sediments.\(^{12,35}\)

Several anomalies existed in the DGT-Cu flux profiles for the 2nd deployment (Figure 1f). The 50 mg Cu/kg sand treatment did not exhibit a DGT-Cu flux peak at the SWI, but
exhibited unexpectedly high copper mobilisation in the deeper sediments. For the 100 and 200 mg Cu/kg sand treatments and 200 and 400 mg Cu/kg silty-sand treatments, the position of the DGT-Cu flux peaks was below the SWI. We believe that burrowing, deposit-feeding activity, and mortality of the bivalves was responsible for several of these observations. The movement of sediments by the bivalves may both increase the flux of copper at the SWI\textsuperscript{34,36} and deplete PW-Cu concentrations in the surface, thereby causing the DGT-Cu flux peak to become located at greater depth within the oxic/sub-oxic zone of the sediments. Bivalve mortalities (% initial) were less than 5% in controls, but 34% and 79% in 100 and 200 mg/kg sandy treatments, respectively, and 30% and 50% in 200 and 400 mg/kg silty-sand treatments, respectively. It is likely that significant mortality occurred within the early stages of the copper exposure\textsuperscript{47}, and the decomposition of the organisms will have increased the availability of labile organic matter and stimulated microbial activity near the dead organisms. These processes may have also increased copper mobilisation from particulate organic phases, or decreased copper mobilisation through production of sulfide and formation of copper-sulfide phases.\textsuperscript{10,12} No vertical movement in the DGT-Cu flux peak for the silty treatments was consistent with low toxicity and the low DGT-induced fluxes of PW-Cu in these sediments. As the anomalies were not present during the 1\textsuperscript{st} DGT deployments, the changes within the sediments that caused the new observations may have taken many days to occur. The reason for the highly elevated DGT-Cu flux in the 50 mg Cu/kg sand treatment is not fully understood, however, Sochaczewski et al.\textsuperscript{4} found that DGT maxima could arise from a microniche – a local source with elevated concentration in the porewater, where the intensity of the maxima was related to the distance between the source and the DGT. In this study it is likely that the base of the DGT probe in the 50 mg Cu/kg sand treatment was in close proximity to some copper-based paint particles.

**Copper bioaccumulation by *T. deltoidalis***. Due to the high level of lethality, tissue-Cu concentrations could not be determined for all 1000 mg Cu/kg treatments, the sandy and silty-sand treatments with 200 mg Cu/kg, and the silty-sand treatments with 400 mg Cu/kg (Table 1). For each sediment type, positive relationships existed between tissue-Cu concentrations and the sediment copper concentrations (See Figures S2a&b, Supporting Information). The properties of the sediments influenced the degree of copper bioaccumulation, with tissue-Cu concentrations increasing in the order sandy > silty-sand > silty when compared based on total recoverable Cu or dilute acid-extractable Cu concentrations. A number of studies have demonstrated the importance of sediment properties in influencing the partitioning of metals.
between the porewaters and sediment phases, and the influence of these two major exposure routes on the bioaccumulation of metals like copper.

For each sediment type, linear relationships existed between tissue-Cu and the time-averaged OW-Cu concentration ($r^2=0.96$ for combined data, Figure S2c). Consistent with dissolved copper in the overlying water being the predominant exposure pathway for *T. deltoidalis*, these relationships appeared to be largely independent of sediment type. While this bivalve is recognised as being a deposit feeder and accumulated copper from both dissolved and particulate exposure routes, previous studies have found that copper accumulation is mostly attributed to exposure from the overlying water. The peak DGT-Cu flux, which occurred near the SWI, also provided strong relationships ($r^2=87$ for combined data) with tissue-Cu concentrations in *T. deltoidalis*. Consistent with the observations by Roulier et al., the DGT-Cu concentration provided a better method for predicting copper bioaccumulation than the total copper measurements, i.e. it was less influenced by sediment properties.

Copper lethality to *T. deltoidalis*. The lethality of the sediments increased with copper concentrations and increased in the order sandy > silty-sand > silt (Figure 2). Dissolved ammonia concentrations were less than 5 mg/L, which is well below the level known to effect *T. deltoidalis*. Consistent with copper bioaccumulation results, copper concentration-response models could be created based on the TR-Cu or AE-Cu concentrations for the individual sediments (Figures 2a,b), and also based on OW-Cu for the three sediment collectively (Figure 2c). There also appeared to be a useful relationship between survival and the tissue-Cu concentrations in *T. deltoidalis* (Figure 2d), however, because tissue-Cu concentrations were not determined for organisms that had died, inadequate data exists for the higher exposure concentrations.

Five different measures of copper exposure were used to calculate lethal copper concentrations (LC50, LC20, and LC10) for the 30-day exposures of *T. deltoidalis*: TR-Cu and AE-Cu calculations used the separate sand, silty-sand and silt treatments, and OW-Cu, Tissue-Cu and DGT-Cu flux at the SWI used the combined data from all treatments (See Table S1, Supporting Information). As indicated in Figures 2a and b, the LC50 values (concentration causing 50% lethality) increased in the order sand < silty-sand < silt, when calculated based on TR-Cu and AE-Cu concentrations. While neither TR-Cu nor AE-Cu provided a suitable representation of the bioavailable copper concentration, the OW-Cu concentrations were useful for describing the toxicity in all the sediments. LC50 and LC10
values of 27 (25-30) and 18 (13-26) µg OW-Cu/L were calculated (Table S1). These results are consistent with the LC50 value of 33±1 µg/L determined by Strom et al. for copper-spiked sediment with a range of properties.

The use of tissue concentrations to express the toxicity of metals is usually more useful for non-essential metals like Cd, Hg, Pb, than essential metals like Cu and Zn. However, when the history of the metal exposure is controlled and known, tissue metal concentrations provide both direct evidence of exposure and metal bioavailability to the organisms and also useful expressions for toxicity for essential metals such as copper. For the 30-day laboratory exposures, owing to the rapid assimilation, the copper accumulation can be treated as metabolically bioavailable, and was used to calculate an LC10 for tissue-Cu of 500 mg/kg (dry weight) (Table S1).

A strong concentration-response relationship was also observed when the peak DGT-induced copper flux at the SWI (Figure 1) was used as a measure of the copper exposure concentration to T. deltoidalis (Figure 2e). Based on the DGT-Cu flux at the SWI, LC50 and LC10 values of 31 (24-42) and 15 (6-35) µg Cu/m²/h were calculated using all the data (Table S1). This relationship has a high degree of environmental significance in terms of the use of DGT for quantifying the risk posed by metal contaminated sediments. Many regulatory frameworks for sediment quality assessment incorporate procedures for determining a contaminants’ bioavailability. However, while metal bioavailability is frequently assessed when the total concentration exceeds the sediment quality guideline value, there are many inadequacies with the methods currently utilised. Pore water metal analyses provide non time-integrated ‘snap-shots’ that are unlikely to reflect the actual exposure for many benthic organisms. It is generally not practical to make multiple field measurements of dissolved metal concentrations, which would allow time-averaged exposures to be accurately assessed. Furthermore, there are numerous artifacts that make porewater analyses problematic in both oxic/sub-oxic and sulfidic sediments. The much used relationships between acid-volatile sulfide (AVS) and simultaneously extractable metals (SEM) can over-emphasise the importance of metal-sulfide binding, as the sediment-microenvironments surrounding many benthic organisms are oxic/sub-oxic either owing to the proximity to the SWI or due to irrigation of burrows with oxygenated waters. Consequently, porewater or AVS-SEM measurements may not always provide adequate information on metal bioavailability. While metal concentrations in organism tissues provide a direct measure of exposure, when organisms are collected from the field their exposure history is not usually known and a significant fraction of the accumulated metal is likely to have been converted to forms that are
not metabolically available.\textsuperscript{49,58} This limits the use of lethal body concentration approach for assessing metal bioavailability.\textsuperscript{12,59}

The DGT-Cu flux – bioaccumulation – toxicity relationships shown in Figure 2 demonstrate that DGT-induced metal flux measurements provide a very useful measure of metal bioavailability in sediments with varying properties. We believe the time-integrated metal fluxes derived from laboratory or field deployments of the relatively simple DGT device may provide a more reliable measure of metal bioavailability than porewater and AVS-SEM analyses in many sediments. While, in some respects the accumulation of metals by the DGT probe simulates the bioaccumulation in sedentary benthic organism, the method does not directly assess metal bioavailability arising from dietary exposure (i.e. ingestion of particles). It has yet to be determined whether the DGT-induced metal flux may also provide useful information on the lability of metals on ingested particles. Although the limitations of using the DGT technique for measuring metal fluxes from sediments requires further assessment, we expect the technique will become more frequently used for assessing the bioavailability and potential ecotoxicological effects of metals in sediments.

**Bioavailability of Cu(I)oxide-based paints in sediments.** The source of bioavailable copper in the tested sediments was copper-based antifouling paint particles and was chosen because of the increasing use by marine industries and concerns about the effects to biota within sediments contaminated with these materials.\textsuperscript{31} The release of copper from the paint particles to the pore water and overlying seawater, along with the bioaccumulation by the bivalves confirmed the presence of bioavailable copper in sediments. In the present study, the measurements of dilute acid-extractable copper provided no obvious advantage of total copper measurements for predicting the bioavailability of copper in the three sediments. This observation may be due, in part, to the use of sediments that were all artificially copper-contaminated and equilibrated for the same duration before testing, i.e. the partitioning between TR-Cu and AE-Cu forms was the similar for all sediments. For naturally contaminated sediments, which form over longer duration, dilute acid-extractable metal measurements and analyses of metal concentrations of the fine sediment fraction are frequently demonstrated to provide a better measure of metal bioavailability than analyses of total metal concentrations.\textsuperscript{12,60} Both the TR-Cu and AE-Cu measurements indicted that the bioavailability of the copper was significantly lower in the more silty sediments.

The toxicity to *T. deltoidalis* in the laboratory tests indicates copper-based paint contamination may cause significant effects to benthic organisms (Figure 2). However, as the
majority of the copper exposure to *T. deltoidalis* was from the overlying water compartment, the same paint-contaminated sediments may not cause this degree of toxicity if assessed in situ, i.e. when situated with a deep water column. The copper extractability from the paint-spiked sediments was also significantly greater than we have found for field-collected sediments contaminated with copper-based antifouling paints, and this may also contribute to an overestimation of the bioavailability compared to paint-contaminated sediments in the field.

For *T. deltoidalis* a large portion of the copper exposure was from the overlying water compartment, which was controlled by the DGT-Cu flux from the sediments in these laboratory tests. In a field location the in situ copper exposure and toxicity of the same paint-contaminated sediments would be lower due to the greater dilution of the released copper within the larger water column. For organisms that receive a greater exposure from the sediment pore waters, or may ingest fine paint particles within the sediments, the in situ exposure may be more important.

**Acknowledgements**

Ian Hamilton and David Spadaro are thanked for assisting with collection and handling of bivalves and undertaking the bioassays. Robert Jung is thanked for assisting with paint and bivalve sample analyses.

**Supporting Information**

Supporting Information includes further information on the preparation of the DGT probes, profiles of porewater Mn and Fe concentrations, and relationships between copper bioaccumulation and a table lethal copper concentrations for the bivalve is provided for each of the different copper exposures. This information is available free of charge via the internet at [http://pubs.acs.org/](http://pubs.acs.org/)

**References**


Table 1. Copper concentrations in sediments and waters and copper bioaccumulation by *T. deltoidalis* in the Cu(I)oxide-paint spiked sediment types

<table>
<thead>
<tr>
<th>Nominal</th>
<th>TR-Cu</th>
<th>AE-Cu</th>
<th>PW-Cu</th>
<th>OW-Cu</th>
<th>Tissue-Cu</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>mg/kg</td>
<td>mg/kg</td>
<td>µg/L</td>
<td>µg/L</td>
<td>µg/g</td>
</tr>
<tr>
<td></td>
<td>Mean SD</td>
<td>Mean SD</td>
<td>Mean SD</td>
<td>Mean SD</td>
<td>Mean SD</td>
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<tr>
<td>Sandy (10% &lt;63 µm) Series 1</td>
<td>0 a</td>
<td>5 2</td>
<td>3 1</td>
<td>4.4 11</td>
<td>2.6 3.1</td>
</tr>
<tr>
<td>200 a</td>
<td>200 15</td>
<td>134 5</td>
<td>5.8 5</td>
<td>45 16</td>
<td>ND ND</td>
</tr>
<tr>
<td>1000 a</td>
<td>1000 75</td>
<td>690 130</td>
<td>6000 900</td>
<td>1450 300</td>
<td>ND ND</td>
</tr>
<tr>
<td>50</td>
<td>38 7</td>
<td>29 3</td>
<td>12.7 3</td>
<td>7.5 1.8</td>
<td>275 ND</td>
</tr>
<tr>
<td>100</td>
<td>109 8</td>
<td>60 9</td>
<td>17.2 5</td>
<td>14 3.3</td>
<td>314 5 ND</td>
</tr>
<tr>
<td>200</td>
<td>176 13</td>
<td>107 5</td>
<td>115 47</td>
<td>34 5.1</td>
<td>1370 ND</td>
</tr>
<tr>
<td>Silty-sand (30% &lt;63 µm) Series 1</td>
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<td>10 4</td>
<td>7 2</td>
<td>0.2 3</td>
<td>1.1 1.5</td>
</tr>
<tr>
<td>200 a</td>
<td>200 15</td>
<td>134 15</td>
<td>2.4 2</td>
<td>19 6.5</td>
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<tr>
<td>1000 a</td>
<td>1000 75</td>
<td>670 150</td>
<td>4.6 2</td>
<td>76 28</td>
<td>ND 58 ND</td>
</tr>
<tr>
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<td>12 1</td>
<td>11 7</td>
<td>0.5 0</td>
<td>1.7 2.5</td>
<td>154 21</td>
</tr>
<tr>
<td>100</td>
<td>109 8</td>
<td>128 16</td>
<td>35.4 6</td>
<td>23 5.2</td>
<td>ND ND</td>
</tr>
<tr>
<td>200</td>
<td>159 8</td>
<td>212 73</td>
<td>62.1 21</td>
<td>26 5.4</td>
<td>ND 58 ND</td>
</tr>
<tr>
<td>Silty (60% &lt;63 µm) Series 1</td>
<td>0 a</td>
<td>30 10</td>
<td>20 4</td>
<td>0.3 1</td>
<td>0.9 0.8</td>
</tr>
<tr>
<td>200 a</td>
<td>200 15</td>
<td>134 15</td>
<td>7.2 3</td>
<td>15 6</td>
<td>499 58 40</td>
</tr>
<tr>
<td>1000 a</td>
<td>950 75</td>
<td>640 150</td>
<td>6.4 6</td>
<td>49 12</td>
<td>ND ND</td>
</tr>
<tr>
<td>0</td>
<td>27 3</td>
<td>13 7</td>
<td>0.5 0</td>
<td>0.5 0.6</td>
<td>134 58 84</td>
</tr>
<tr>
<td>100</td>
<td>143 39</td>
<td>96 11</td>
<td>5.3 6</td>
<td>1.7 1.1</td>
<td>194 58 84</td>
</tr>
<tr>
<td>200</td>
<td>234 11</td>
<td>113 20</td>
<td>1.1 1</td>
<td>2 1</td>
<td>276 58 42</td>
</tr>
<tr>
<td>400</td>
<td>426 91</td>
<td>250 77</td>
<td>3.9 3</td>
<td>7.7 2.2</td>
<td>414 58 72</td>
</tr>
</tbody>
</table>

TR-Cu = total recoverable copper (n=2). AE-Cu = dilute acid-extractable copper (n=2).
ND = not determined due to inadequate tissue mass due to poor survival.
PW-Cu (n=2, at start and finish) and OW-Cu (n=10, time average of samples approximately every 3 days) are dissolved copper in porewater and overlying water, respectively.

a First concentration series.
Figure 1. Depth profiles of the 24-h DGT copper flux on days 8 and 15 for the three copper concentration series: 0, 50, 100, 200 mg Cu/kg for the sandy, and 0, 100, 200, 400 mg Cu/kg for the silty-sand and silty series. Error bars have been omitted for clarity (Table S1 of Supplementary Information).
Figure 2. Relationships between survival of the bivalve, *Tellina deltoidalis*, and different copper exposures: (a) total recoverable copper, (b) dilute acid-extractable copper, (c) dissolved copper in overlying water, (d) bivalve tissue-Cu concentrations, and (e) peak DGT-induced Cu flux at the sediment water interface (DGT-Cu flux). Data presented for three sediment types for the different Cu-spike concentrations (mean ±SD, n=2). The lines represent log-logistic concentration response curves calculated for the three different sediment types for (a) and (b) and for the combined data for (c), (d) and (e). The LC50, LC20 and LC10 values for each relationship are provided in Table S1 of the Supporting Information.
Copper flux at sediment-water interface, \( \mu g \text{ Cu/m}^2/\text{h} \)

Benthic bivalve survival, %

- Sandy sediment
- Silty-sand
- Silty

The graph shows the relationship between copper flux and benthic bivalve survival across different sediment types.
SUPPORTING INFORMATION:

DGT-INDUCED COPPER FLUX PREDICTS BIOACCUMULATION AND TOXICITY TO BIVALVES IN SEDIMENTS WITH VARYING PROPERTIES

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S1 Methodology:

S1.1 Preparation of diffusive gradients in thin films (DGT) probes

Plastic DGT assemblies with open windows of 1.8 × 15 cm and overall dimensions of 24 × 4 × 0.5 cm were purchased from DGT Research (Lancaster, UK). All glass and plasticware for DGT probes preparation were cleaned by soaking in detergent (commercial detergent diluted in tap water) for 24 h, then in 10% (v/v) HNO₃ (70%, AR grade, Ajax Finechem Pty Ltd) for 24 h and rinsed thoroughly with MQ water. All glass and plasticware for DGT probes analysis were cleaned by soaking in 10% (v/v) HNO₃ for 24 h and rinsed thoroughly with MQ water.

DGT gels were prepared from a stock solution comprising of 15% (w/v) acrylamide (40% acrylamide solution, Electrophoresis Purity Reagent, Bio-Rad Laboratories) and 0.3% (w/v) DGT cross-linker (2% aqueous solution DGT Cross Linker, DGT Research, Lancaster, U.K.). The gel chemical polymerization for diffusive gels (0.50 mm-thick) was initiated by adding 75 μL of 10% (w/v) freshly made APS (98+% ammonium persulfate, for analysis ACS, Acros Organics) to 10 mL of stock solution and catalysed by adding 25 μL of TEMED (99% N,N,N,N-tetramethylethylenediamine, Molecular Biology tested, Sigma). The solution was stirred for 3½ minutes, then immediately cast between a pair of glass plates separated by a 0.5 mm plastic spacer. The gel solution was left to polymerize for 1 h at 45°C.

Wet Chelex resin was prepared by mixing 2 g of dry Chelex resin with 10 mL of MQ water, then allowing the resin to settle and withdrawing the overlying water with a pipette. Chelex gels (0.25 mm-thick) were prepared by adding 5 mL of stock solution to 2 g of wet chelating resin, then 25 μL of initiator (10% (w/v) freshly made APS) and 7.5 μL of catalyst (TEMED). The solution was mixed for 3 minutes, then immediately cast between a pair of glass plates separated by a 0.25 mm plastic spacer. The solution was then left to polymerize for 1 h at 45°C.

The resulting diffusive and Chelex gels were removed from the glass plates and hydrated in Milli-Q water for 24 h, replenishing the water three times to remove all unreacted chemicals. Diffusive gels were stored in
0.01 M NaNO₃ (AR grade, Chem Supply) at room temperature and Chelex gels were stored in Milli-Q water in a refrigerator at 4°C until use for probe construction. Gels were cut with a Teflon coated razor blade using a plastic rectangular strip of the desired dimensions in order to fit in the DGT device. Acid-cleaned 0.45 µm filter membranes were cut and stored in MQ water. Probes were assembled by laying a wet filter membrane on the base, overlaying a Chelex gel layer, then a diffusive gel layer and finally another wet filter membrane on the DGT backing plates and closing the devices with the front window plates. Care was taken to ensure no air bubbles were trapped within the layers. The DGT units were kept in sealed clean plastic bags containing few drops of MQ water to avoid gel drying and stored in a refrigerator until deployment.

S1.1 DGT probes deployment, retrieval and analysis

To prevent the introduction of oxygen into the sediments during the deployment, DGT probes were deoxygenated for 24 h prior to deployment by immersing them in a 0.05 M NaCl (>95.5%, Sigma) solution saturated with nitrogen gas (continually bubbling to remove the dissolved oxygen). DGT devices were immediately gently inserted into the test beakers to a depth of 4 cm with care to ensure a good contact between the sediment and the DGT membrane. After 24 h of deployment, DGT probes were gently removed from the beakers and both sediment and overlying water levels were noted. Devices were immediately rinsed with MQ water to remove all remaining sediment particles. Probes were put in clean plastic bags and kept in a cool room (4°C) until disassembly.

DGT devices were disassembled (within 10 days of retrieval) and resin gel slices cut using a Teflon coated blade to obtain the desired vertical profile: two 2-cm slices in the overlying water, a one 1-cm slice at the sediment-water interface (SWI) and one 2-cm slice in the sediment. Some of the slices from below the SWI had sediment particles adhered to them which were removed by washing with MQ water. Each slice was weighed and put into a 5-mL vial, then eluted with 1 mL of 1 M HNO₃ for 16-24 h before analysis by ICP-MS. Undeployed DGT probes were analysed as handling blanks and their copper concentrations were less than 5% of the lowest measured concentration.

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### Table S1. Lethal copper concentrations for the bivalve, *Tellina deltoidalis* (Figure 2)

<table>
<thead>
<tr>
<th></th>
<th>TR-Cu, mg/kg</th>
<th>TR-Cu, mg/kg</th>
<th>TR-Cu, mg/kg</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(Sand)</td>
<td>(Silty-sand)</td>
<td>(Silt)</td>
</tr>
<tr>
<td>LC50 (95% CL)</td>
<td>130 (110-156)</td>
<td>355 (244-518)</td>
<td>530 (---)</td>
</tr>
<tr>
<td>LC20 (95% CL)</td>
<td>102 (78-136)</td>
<td>223 (120-415)</td>
<td>490 (---)</td>
</tr>
<tr>
<td>LC10 (95% CL)</td>
<td>88 (63-132)</td>
<td>170 (67-414)</td>
<td>460 (---)</td>
</tr>
<tr>
<td></td>
<td>AE-Cu, mg/kg</td>
<td>AE-Cu, mg/kg</td>
<td>AE-Cu, mg/kg</td>
</tr>
<tr>
<td></td>
<td>(Sand)</td>
<td>(Silty-sand)</td>
<td>(Silt)</td>
</tr>
<tr>
<td>LC50 (95% CL)</td>
<td>75 (61-91)</td>
<td>178 (93-339)</td>
<td>350 (---)</td>
</tr>
<tr>
<td>LC20 (95% CL)</td>
<td>54 (39-77)</td>
<td>87 (35-224)</td>
<td>300 (---)</td>
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<tr>
<td>LC10 (95% CL)</td>
<td>44 (29-74)</td>
<td>57 (16-217)</td>
<td>280 (---)</td>
</tr>
<tr>
<td></td>
<td>OW-Cu, µg/L</td>
<td>Tissue-Cu, mg/kg</td>
<td>DGT-Cu flux, ug Cu/m²/h</td>
</tr>
<tr>
<td></td>
<td>(All treatments)</td>
<td>(All treatments)</td>
<td>(All treatments)</td>
</tr>
<tr>
<td>LC50 (95% CL)</td>
<td>27 (25-30)</td>
<td>1000 (770-1300)</td>
<td>31 (24-42)</td>
</tr>
<tr>
<td>LC20 (95% CL)</td>
<td>21 (17-27)</td>
<td>650 (380-1100)</td>
<td>19 (11-36)</td>
</tr>
<tr>
<td>LC10 (95% CL)</td>
<td>18 (13-26)</td>
<td>500 (230-1100)</td>
<td>15 (6-35)</td>
</tr>
</tbody>
</table>

**LC50 (95% CL)** = concentration causing 50% lethality (measured after 30 days).

**LC20 and LC10** represent 20% and 10% effect concentrations, respectively.

95% CL = 95% confidence limit (--- = not possible to calculate 95% CL).

TR-Cu = total recoverable copper concentration (aqua regia, mg/kg).

AE-Cu = dilute acid-extractable copper concentration (1-M HCl, mg/kg).

OW-Cu = overlying water copper concentrations (30-day time averaged concentration, µg/L)

Tissue-Cu = copper concentrations after 30 days in surviving bivalves (mg/kg, dry weight)

DGT-Cu flux = Peak DGT-induced Cu flux at the sediment-water interface (ug Cu/m²/h)

Effects thresholds for TR-Cu and AE-Cu were calculated separately for sand, silty-sand and silt treatments.

Effects thresholds for OW-Cu, Tissue-Cu and DGT-Cu flux were calculated using the combined data from treatments.
Figure S1. Porewater Mn and Fe concentrations within silty-sand that had been spiked with a copper mineral phase (unpublished results), confirming that the expected porewater Fe and Mn profiles develop following sediment disturbance and one month equilibration following spiking. The three porewater metal profiles (●, ■, ▲) are from three DGTs from separate experiments.
Figure S2. Relationships between copper bioaccumulation by bivalve, *Tellina deltoidalis*, and different copper exposures: (a) total recoverable copper (TR-Cu), (b) dilute acid-extractable copper (AE-Cu), (c) dissolved copper in overlying water (OW-Cu, time averaged) and (d) peak DGT-induced Cu flux at the sediment-water interface (DGT-Cu flux). Data presented for three sediment types for the different Cu-spike concentrations (mean ±SD, n=2).

**Correlations** (*r*^2^) between copper bioaccumulation by bivalve and copper concentrations for each of the sediment types

<table>
<thead>
<tr>
<th>Sediment type</th>
<th>(a) TR-Cu</th>
<th>(b) AE-Cu</th>
<th>(c) OW-Cu</th>
<th>(d) DGT flux</th>
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<tbody>
<tr>
<td>Sand</td>
<td>0.804</td>
<td>0.852</td>
<td>0.959</td>
<td>0.994</td>
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<td>0.995</td>
<td>0.991</td>
<td>0.886</td>
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<tr>
<td>Silt</td>
<td>0.574</td>
<td>0.650</td>
<td>0.860</td>
<td>0.718</td>
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<tr>
<td>Combined data</td>
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<td>NA</td>
<td>0.962</td>
<td>0.865</td>
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