A mesocosm study of oxygen and trace metal dynamics in sediment microniches of reactive organic material

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Deposition of particulate organic matter (POM) induces diagenetic hot spots at the sediment-water interface (SWI). Here we explore the effects of intensive POM degradation for metal mobilization at the SWI. By using a combined planar optode-DGT (diffusive gradient in thin-films) sensor we obtained simultaneous measurements of dissolved O2 and trace metal dynamics around an aggregate of reactive organic matter placed on the SWI of a sediment mesocosm. The aggregate induced a rapid, highly localized, decrease in O2 concentration, resulting in an anoxic feature at the SWI. Co-located with this feature, we observed intense Fe and Mn mobilization, removal of Co, Ni and Zn and found evidence for the concurrent release and precipitation of Pb within a small confined volume. We also identified two small microniches in the anoxic sediment below the SWI, defined by elevated trace metal mobilization. Differences between the metal release rates in these two microniches indicate that they were formed by the mineralisation of different types of organic matter buried in the sediment. Our results provide direct empirical evidence for the potential importance of POM-induced reactive microniches when considering the fluxes of metals from and within aquatic sediments, and suggest that other elements' cycles may also be affected.

Deposition of reactive organic material at the sediment water interface (SWI) as particles formed from various biological processes in the water column is a well-recognized component of aquatic carbon fluxes1–3. Following settlement on the SWI, they can bring about rapid rates of organic material mineralisation within discrete volumes of sediment4. These localized “microniches” can make a significant contribution to organic material turnover in surface sediments, and promote a high degree of variability in the cycling of terminal electron acceptors (TEA) and associated elements4–6. They have received significant interest in the literature7–9, with much work focusing initially on sulphidic microniches8,10 and more recently localized sinks of reactive nitrogen6,11. These studies have suggested that microniches can make a significant contribution to elemental cycling in aquatic environments, however direct analysis of these processes has been limited by the availability of suitable analytical methods.

Oxidation of reactive organic matter induces changes in redox conditions and the associated solubilisation of Fe and other trace metals has been studied in a variety of benthic environments12–14. Understanding of these processes has been built on increasingly sophisticated experimental and numerical modelling techniques15–18. However, the level of insight that can be gained from many experimental investigations is limited by methodology that effectively averages solute concentrations across various volumes of sediment. This means that critical information about microniches may often be overlooked1. Interpretation of transient chemical gradients occurring at mm or sub mm spatial scales due to small-scale heterogeneity in sediment processes is greatly facilitated if measurements are made in two dimensions (2D) with an appropriate temporal and spatial resolution. As small-scale

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changes in solutes usually reflect complex interactions between different elements, combined sensors for multiple analytes have the potential to enhance understanding of coupled processes. Diffusive gradients in thin-films (DGT) and planar optodes are two techniques that have been employed to provide high-resolution measurements of elemental fluxes and their distributions in sediments. DGT measurements have advanced our understanding of phosphorus, trace metal and sulphide cycling\textsuperscript{10, 19–21}, while planar optodes have provided valuable information on spatial and temporal O\textsubscript{2}, pH, pCO\textsubscript{2} and NH\textsubscript{4}\textsuperscript{+} dynamics\textsuperscript{22–26}. Recently, the two techniques have been used together in a combined DGT-planar optode sensor to measure dissolved O\textsubscript{2} and trace element mobilization simultaneously. This has provided insight into how their biogeochemical cycles are interrelated in different environments. Williams, et al\textsuperscript{27} identified the release of Fe, Pb and As associated with O\textsubscript{2} efflux by rice roots growing in an anoxic soil, while Stahl, et al\textsuperscript{28} showed trace element mobilization linked to the presence of relict organic matter and increasing O\textsubscript{2} concentrations of burrow structures in an otherwise anoxic heterogeneous sediment.

Previous work where fresh organic matter was deployed at the SWI used numerical modelling to suggest that the resulting microniches may make a significant contribution to redox cycling of nitrate and other TEAs following localized oxygen depletion\textsuperscript{6}. We now seek to advance this knowledge with experiments designed to test the effects of reactive organic matter deposition on the biogeochemical cycling of Fe and Mn in an otherwise oxic sediment. We hypothesize that localized oxygen depletion induced by the rapid microbial mineralisation of organic matter would result in the reductive dissolution of reactive Fe and Mn (oxyhydr)oxide minerals, and the concomitant release of other trace elements (e.g. Co, Ni, Cu, Zn and Pb). To test this hypothesis, we placed three aggregates of organic matter at evenly spaced intervals on the SWI of a sediment mesocosm and used the combined DGT-planar optode sensor to make high-resolution measurements of real-time O\textsubscript{2} dynamics and mobilization of trace metals simultaneously at these locations.

Results and Discussion

O\textsubscript{2} dynamics across the SWI during the deployment. Optode measurements showed a rapid localized O\textsubscript{2} depletion at the locations of all three aggregates following their deployment. However, we primarily focus our subsequent investigations on the area around the central aggregate formed from freeze-dried \textit{Spirulina} material (Fig. S1), as action of biota at the surface of the sediment visibly disturbed the two other aggregates of organic matter. This activity did not affect the central aggregate. The central \textit{Spirulina} aggregate slowly disintegrated at its deployment location during the experiment. This was due to the slow colonisation and degradation of the aggregate by micro- and meiofauna that physically fragment and rework the aggregate.

The O\textsubscript{2} concentration decreased rapidly at the centre of the aggregate (Fig. 1). This mimics the dynamic observed in previous investigations of microbial colonisation on parcels of organic matter and is thought to arise from the oxidation of organic matter, resulting in an intense demand for the most energetically favourable TEA (O\textsubscript{2})\textsuperscript{6, 11, 29}. The anoxic isoline, initially beneath the sediment-water interface (SWI), gradually rose upwards towards the centre of the aggregate (Fig. 1a), reaching a distance of 2.2 mm above the original SWI (herein we define the threshold for ‘anoxia’ as the calculated O\textsubscript{2} method detection limit by the planar optode in this experimental setup, 3.0 µM; see Supporting Information, S.iii.). The anoxic microniche area measured at the probe interface reached a maximum size of ~14 mm\textsuperscript{2} after 44 h, corresponding to approximately half of the aggregate’s cross-sectional area at its centre. Following the emergence of anoxic conditions, the mineralisation of organic matter is thought to be conducted mainly by facultative microorganisms that then employ nitrate as the TEA\textsuperscript{6, 14}. Given the low concentration of nitrate in the sediment porewater (1.13 µM, Table S1), and the rapid rate
of mineralisation at the location of the aggregate, we believe that the reduction of less energetically favourable TEAs, Mn(IV) and Fe(III), began relatively quickly following anoxia. The location at the centre of the aggregate's initial position adjacent to the probe had been anoxic for ~32 h when the high-resolution DGT (HR-DGT) was retrieved (Fig. 1b). We estimate that <5% of the organic C in the aggregate was mineralized during the deployment (Supporting Information S1.vi).

The O₂ concentrations below the SWI decreased rapidly after deployment of the combined sensor and reached a steady state within 100 min. The average background O₂ penetration depth (i.e. depth to anoxic sediment below SWI) and diffusive O₂ uptake at 43 h were determined from profiles extracted from the O₂ images (Fig. S1), these were 3.08 ± 0.28 mm (SD, n = 3) and 7.82 ± 0.24 mmol m⁻²d⁻¹ (SD, n = 3): typical for a coastal marine sediment39.

**Table 1.** Maximum (Max.) and average fluxes of metals in the bulk oxic and anoxic sediment (the former excludes \( I_2 \), the latter includes \( S_A \) and \( S_B \)) and the three microniches. The standard deviation of each average is shown in the adjacent brackets. All fluxes are reported as nmol cm⁻²s⁻¹. Superscript letters A, B and C indicate where the average flux of metal across the microniche is significantly different from the other microniches (p < 0.05), different letters indicate significant differences between microniches. Where theflux of a particular metal was not sufficiently elevated above background to qualify it as a 'microniche' for that particular metal (e.g. Ni for \( I_2 \) or \( S_B \)), the metal fluxes from the measurement points that define the corresponding Fe microniche were used to enable comparison. These are indicated in italic script, where applicable. Otherwise, the number of measurement points (n) is indicated in brackets next to the maximum measured flux for that metal in that microniche. Locations where the mass of metal/unit area is below the detection limit for that metal are designated as “not detected” (N.D.), these were treated as zero concentrations in the calculations of the averages and standard deviations.

| Metal | Oxic Sediment (n = 276) | Anoxic Sediment (n = 3496) | \( I_2 \) (n Fe = 53) | \( S_A \) (n Fe = 11) | \( S_B \) (n Fe = 18) |
|-------|------------------------|---------------------------|----------------------|----------------------|----------------------|
| Fe    | Max. Average           | 24.28 ± 0.02              | 80.36 ± 0.13         | 384.32 ± 0.65        | 37.65 ± 0.10         |
|       |                        | (n = 9) 16.31             | 13.70 ± 0.09         | 28.40 ± 0.12         | 2.04 ± 0.10          |
| Mn    | Max. Average           | 9.65 ± 0.06               | 9.76 ± 0.09          | 0.53 ± 0.04          | 1.10 ± 0.04          |
|       |                        | (n = 9) 8.13              | 0.28 ± 0.04          | 1.00 ± 0.03          | 1.30 ± 0.03          |
| Co    | Max. Average           | 1.66 ± 0.04               | 0.72 ± 0.03          | 1.79 ± 0.04          | N.D.                |
|       |                        | (n = 9) 1.56              | 0.28 ± 0.03          | 1.07 ± 0.03          | N.D.                |
| Ni    | Max. Average           | 2.31 ± 0.02               | 1.80 ± 0.02          | 0.44 ± 0.02          | 1.44 ± 0.02          |
|       |                        | (n = 9) 2.28              | 1.40 ± 0.02          | 0.28 ± 0.02          | 1.72 ± 0.02          |
| Cu    | Max. Average           | 2.28 ± 0.02               | 2.01 ± 0.03          | 1.79 ± 0.02          | N.D.                |
|       |                        | (n = 9) 2.12              | 1.15 ± 0.02          | 1.07 ± 0.02          | N.D.                |
| Zn    | Max. Average           | 15.96 ± 0.03              | 41.81 ± 0.03         | 22.42 ± 0.13         | 25.40 ± 0.13         |
|       |                        | (n = 9) 13.13             | 14.95 ± 0.03         | 28.28 ± 0.13         | 26.53 ± 0.13         |
| Pb    | Max. Average           | 0.54 ± 0.02               | 0.08 ± 0.02          | 0.94 ± 0.01          | 0.06 ± 0.01          |
|       |                        | (n = 9) 0.44              | 0.03 ± 0.02          | 0.19 ± 0.01          | 0.07 ± 0.01          |

**Interpreting DGT flux measurements in the sediment mesocosm.** The chelating resin in the HR-DGT probe binds trace metals and thus acts as a planar sink for dissolved trace metals in the sediment and the overlying water. The removal of dissolved metal results in a diffusive flux into the resin from the adjacent environment. The size of that flux is directly related to the dissolved concentration of metal at the interface of the probe and the sediment, as long as the capacity of the chelating resin to bind that metal is not exceeded. When the rate of metal uptake by the HR-DGT exceeds the rate at which the metal can be resupplied from the surrounding sediment, the concentration at the interface, and therefore the flux into the HR-DGT, will progressively decrease. However, when localized processes operating in the sediment close to the HR-DGT interface (e.g. reductive dissolution) are rapid and sustained, the mass of solute taken up by the DGT probe can be used to provide an indication of the average metal concentration in the sediment porewater adjacent to the probe during its deployment30.

The binding of divalent trace metals to the ultra-thin HR-DGT gels used in this work has been shown to be directly related to the metal solution concentration at the device interface when the bound mass is less than 71 nmol cm⁻²31. The highest total mass of metals measured at any single location in this experiment was 67 nmol cm⁻². The metal accumulated at any given location in the planar HR-DGT probe is presented here (Fig. 2) for the most part as a time-averaged flux (the mass of metal accumulated by certain area of gel, averaged over the deployment time) to reflect the total supply of metal to the HR-DGT from the sediment and is forthwith expressed as “HR-DGT flux”.

**Distribution of trace metal mobilization below the SWI.** To consistently define distinct reference zones, we used areas in the oxic upper 1 mm beneath the SWI apparently unaffected by the aggregates (‘oxic sediment’, total area: 38.6 mm²) and the anoxic sediment 6 mm below (‘anoxic sediment’, total area: 517 mm²) (Fig. S1). The average background HR-DGT fluxes of Mn, Fe, Co, Ni, Cu, Zn and Pb within oxic and anoxic reference parts of the sediment are presented in Table 1. Background HR-DGT fluxes of Cu, Ni and Zn in the oxic sediment were higher than in the anoxic sediment, while the HR-DGT fluxes of Fe were higher in the anoxic sediment. HR-DGT fluxes of Co, Mn and Pb in the oxic and anoxic sediment were broadly similar (Table 1). The average HR-DGT fluxes of all metals were generally smaller than those observed in previous studies19, 20, 32, 33.
which is to be expected given the combination of a relatively long deployment time of 44 h and the thinner material diffusive layer used here.\textsuperscript{31,34}

Depth profiles of O\textsubscript{2} availability and trace metal release from the sediment showed general patterns broadly similar to those observed in other measurements of benthic trace metal mobilization.\textsuperscript{19,35–37} (Fig. S2). There was little release of Mn, Fe and Co to the HR-DGT near the SWI, but we observed a gradual increase with increasing depth. The HR-DGT fluxes of Fe measured in these profiles are closely related to the duration of anoxia at the corresponding points in the sediment (Supporting Information S.vii; Figs S2 and S3). This supports our earlier assessment that nitrate reduction makes a relatively small contribution to the overall organic matter mineralisation in this sediment. It also suggests that reductive dissolution is driving the release of Fe over time. Conversely, HR-DGT fluxes of Cu and Zn are highest at the SWI and decrease with depth. Such near surface mobilisation has been linked to mineralisation of homogeneously distributed organic matter on the SWI.\textsuperscript{19, 20} Although the unprecedented Pb fluxes are very low, they vary systematically and show a unique double maximum below the sediment water interface, meriting further study.

Caution should be exercised when comparing these trace metal profiles to those obtained for marine sediments using traditional techniques, which provide measurements of porewaters extracted from relatively large volumes of sediment (10–100 mL).\textsuperscript{12, 13} Our measurements at 0.28 mm intervals extend in total to only 16 mm into the sediment whereas the work of Klinkhammer,\textsuperscript{31, 34} provided data at 2–3 cm intervals, while the high resolution measurements of Shaw,\textsuperscript{14} were at best 2.5 mm intervals. In our system, redox gradients, as exemplified by the Fe, Mn and O\textsubscript{2} measurements, are tighter than those observed for the less reactive sediments of previous work. They do confirm more precisely, however, the previous suggestions of mobilisation of Cu at the sediment water interface and extend the observation to Zn. There is also visual evidence of some association between Ni and Co and Fe and Mn, which was found in some previous extended-scale profiles\textsuperscript{13} and attributed to Co and Mn being released from iron and manganese oxides as they are reductively dissolved.

The two-dimensional Fe and Mn HR-DGT flux data presented in Fig. 2 shows that there are three distinct areas where their fluxes are elevated: one at the SWI (I\textsubscript{s}), the ‘surface microniche’ and two below the SWI (S\textsubscript{A} and S\textsubscript{B}, forthwith referred to collectively as ‘subsurface microniches’) (Table 1; Figs 2 and S4). Numerical modelling has shown that such well-defined features are likely to have arisen from locally increased porewater concentrations; desorption from small sediment volumes with high concentrations of exchangeable sorbed metal would be expected to show less steep gradients across the HR-DGT.\textsuperscript{38} The standard deviations of HR-DGT fluxes of each trace metal in the anoxic sediment were used to provide a systematic definition of niches of increased metal supply. Areas where the HR-DGT fluxes of metals were two standard deviations above their respective averages in the anoxic background, and consisted of five or more contiguous measurement points, were defined as being indicative of a ‘metal microniche’. We used the results from the redox-active metals, Fe and Mn, to designate three separate features with sufficient numbers of distinct measurements to give confidence that a localized process is supplying metal for uptake by the HR-DGT, rather than a single isolated measurement that may have been caused by an experimental or analytical artefact.\textsuperscript{4} This approach limited the detection of microniches to those that supply solute to areas of HR-DGT greater than 0.7 mm\textsuperscript{2}, but provided features with sufficient measurement points to enable comparisons of trace metal release rates. We did not observe any differences in trace metal release that could be associated with the brief oxygenation during the sensor mounting and immediately following the start of the deployment.

**Trace metal fluxes at the surface microniche.** I\textsubscript{s} covered an area of 7.41 mm\textsuperscript{2} on the HR-DGT and overlapped with the position of the *Spirulina* aggregate and the observed O\textsubscript{2} depletion. The average Fe HR-DGT flux within it was ~11 times higher than the average background in the oxic sediment and higher than in the two subsurface microniches (p < 0.05, Table 1). Approximately 64% of the microniche area was observed below the SWI at the position of the *Spirulina* aggregate (Fig. 2; Fig. S4). Average Pb and Mn HR-DGT fluxes within I\textsubscript{s} were also higher than their respective fluxes in the top 1 mm of the sediment (6.3 × 1.44 ×, respectively) and significantly higher than those from the subsurface microniches (Table 1). Given the small proportion of the aggregate mineralised during the deployment (<5%), we do not believe that metal in the aggregate made a significant contribution to the HR-DGT fluxes measured at I\textsubscript{s}.

Previous studies have reported high dissolved Fe concentrations below the SWI in oxic sediments subject to episodic POM deposition,\textsuperscript{35, 36} while others have demonstrated how the deposition of organic material as a homogeneous treatment onto the SWI resulted in mobilization of Mn, Co and As.\textsuperscript{37} In these studies, bioturbation, stabilisation of reduced Fe(II) by organic matter ligands and changing redox status in the sediment, associated with organic matter mineralisation were suggested as possible causes. Our results show that the localized O\textsubscript{2} depletion associated with a single aggregate of organic matter appears to be driving the co-located release of Fe, Mn and Pb above background levels across the SWI, and provide an alternative, complementary, explanation to the observations in these studies. This explanation could also account for the heterogeneity in dissolved Fe and Mn concentrations reported by Homoky,\textsuperscript{13} The total mass of Fe bound by the HR-DGT resin within the area of I\textsubscript{s} above the SWI (2.67 mm\textsuperscript{2}) was 548 pmol. We believe that this Fe originated mainly from the reductive dissolution of Fe (oxyhydr)oxide minerals in the underlying sediment, following the emergence of localized anoxia. Given the small scale of the microniche and its position adjacent to the DGT device, DGT will have captured most, but not all of the flux of Fe attributable to the microniche. If we make the highly conservative assumptions that (1) this mass of Fe is the total amount that was released from the sediment to the water overlying the SWI, and (2) the anoxic microniche only lasted for the observed 32 h that I\textsubscript{s} was anoxic, then the average rate of Fe release from the sediment during that time would have been at least 17 pmol h\textsuperscript{−1}. Homoky,\textsuperscript{37} (and references therein) report a range of Fe fluxes from sediments between 4.22–416.67 nmol m\textsuperscript{−2}h\textsuperscript{−1}. Using our estimated average Fe release rate induced by the *Spirulina* aggregate, the previously reported range of Fe fluxes could be matched by between 248–2,510 similar aggregates (0.042–5.86 mol C m\textsuperscript{−2}, Supporting Information, S1.vi) deposited on one square
metre of sediment. Over the 32 h, this corresponds to a range of particulate organic C (POC) fluxes between 1.32 and 183.06 mmol m$^{-2}$ h$^{-1}$. While previous studies suggest that these POC deposition rates are most representative of either shallow coastal conditions\(^3\), or periods of peak production under Arctic sea ice\(^4\), lower rates (such as those observed over deep sea sediments\(^3\)) may also make a significant contribution to Fe release rates elsewhere (c.f. Homoky, et al.\(^3\)). The HR-DGT is likely to have sampled only a part of the total Fe released from the sediment and it is highly likely the localized anoxic conditions would have persisted longer had we not ended the experiment, therefore our calculated Fe release rate is probably an underestimate.

Lalonde, et al.\(^4\) showed that around 21.5% of the organic C in sediments is incorporated with reactive Fe minerals. In our experiment, the reductive dissolution of Fe solid phases at the SWI may liberate further reactive organic C\(^4\), thus creating a partially self-sustaining transient diagenetic hot spot. Using the previously quoted organic C:Fe ratio of 4\(^4\), the above rates of Fe mobilization can be used to estimate a corresponding range of dissolved organic C release rates of 0.017–1.67 nmol m$^{-2}$ h$^{-1}$, which could make a significant contribution to previously reported rates of DOC diffusion from marine sediments\(^4\). Moreover, given that the organic matter existing in mineral-organic associations in sediments is often enriched in nitrogen\(^4\) and the affinity that species of phosphorus and arsenic can have for Fe (oxyhydr)oxide minerals\(^4\), these types of anoxic micro-features may also make a, hitherto unappreciated, contribution to the cycling of these elements at the SWI.

When considering these estimates, the following caveats should be considered. The organic matter content of this sediment (4.65%wt, Table S1) is more representative of a coastal sediment and higher than many deep-sea marine sediments\(^4\), 47, 48. Therefore, the local populations of microorganisms may have been adapted to rapidly mineralise the aggregate organic matter. The ambient temperature in this experiment was between 5–15°C higher than in most benthic environments, resulting in faster rates of organic matter mineralisation, but this will be offset to some extent by decreased diffusion coefficients of dissolved TEAs (O$_2$, NO$_3^-$, SO$_4^{2-}$) at the lower temperatures\(^5\), and often lower ambient dissolved O$_2$ concentrations\(^5\); these would increase the likelihood of anoxic microniche formation. Moreover, we expect that spatial and temporal variations in the quantity (and quality) of the organic matter that is deposited on the sediment surface, as well as seafloor environments (e.g. different ambient O$_2$ and/or NO$_3^-$ concentrations) would significantly affect the sizes and durations of anoxic microniches and the extent to which various TEAs are reduced therein. Further experiments, aiming to capture the complete cycles of transient anoxia, are clearly required to quantify accurately the cycling of Fe (and other elements) within microniches in different sediments. We believe processes of elemental mobilization observed and implied here are likely to make the most significant difference in relatively shallow oxic sediments where organic matter turnover is dominated by microbial mineralisation at the SWI, and in environments that experience (possibly seasonal) periods of high organic matter deposition\(^4\), 36.

A comparison of the relative fluxes of Fe and Mn from $I_x$ shows that their rates of mobilisation are closely related across the microniche (Table 2). The size of the Mn microniche at $I_x$ (1.12 mm$^2$) is somewhat smaller than that of Fe, presumably due to the lower Mn content of the sediment compared to Fe (Table S1). The correlation between Fe and Pb fluxes is also statistically significant ($p < 0.01$)(Table 2), an association that is well established\(^4\).
It is therefore likely that reductive dissolution of Fe (oxyhydr)oxide minerals is driving the Pb release across $I_{A}$. Stahl et al.\textsuperscript{28} found a peak Pb release of ~1.5 fmol cm\textsuperscript{-2} s\textsuperscript{-1} (~1.6 × higher than at $I_{A}$) during a 6 h deployment at the location of an artificial worm burrow which was closely associated with increased O$_2$ concentrations inside it. They suggested that Pb was released by local oxidation of Pb sulphide minerals. This process is unlikely to have determined Pb mobilization here, because O$_2$ was diminished at $I_{A}$.

The relationship between Fe and Pb, while statistically significant, is not consistent across $I_{A}$. When the fluxes of the two metals are compared against each other, there appear to be two antagonistic mechanisms operating: one mobilizes both Fe and Pb at closely related rates, while another limits Pb flux into the HR-DGT when Fe fluxes are high (Fig. 3). This relative limitation can be identified by a feature defined by six contiguous measurement points at the centre of the $I_{A}$ microniche: a possible 'reaction hotspot' (Figs 2 and 3). Rapid organic matter mineralisation has been linked to the formation of sulphide, S(-II), 'hotspots' in otherwise sulphide-free sediments\textsuperscript{27}. Given that Pb is thought to react with S(-II) much faster than Fe or Mn\textsuperscript{52}, the likely explanation for the reduced Pb flux in the heart of the microniche is that, at some point during the deployment, the concentrations of Pb$^{2+}$ and S(-II) increased past a level where the rate of PbS precipitation exceeded the rate of Pb$^{2+}$ release by Fe mineral dissolution.

When compared to their average fluxes in the top 1 mm of the sediment, the fluxes of Co, Ni and Zn within the area of $I_{A}$ (as defined by Fe measurements) were lower than in background oxic sediment, indicating that there was a process removing these metals from solution (Table 1; Fig. 2). The lower flux of these metals to the HR-DGT resin may be caused by different processes working independently or together. A localized decrease in H$^+$ ion activity caused by their consumption in the dissimilatory reduction of Fe minerals\textsuperscript{53} could promote immobilization of these metals; however, such an effect has not been demonstrated at this spatial resolution and under these conditions. Zn is also known to react rapidly with S(-II)\textsuperscript{52}. Given the higher concentrations of Zn in the pore water, it is possible that a high rate of ZnS mineral formation introduced a localized sink for S(-II) within the $I_{A}$ microniche: a possible 'reaction hotspot' (Figs 2 and 3). Rapid organic matter mineralisation has been linked to the formation of sulphide, S(-II), 'hotspots' in otherwise sulphide-free sediments\textsuperscript{27}. Given that Pb is thought to react with S(-II) much faster than Fe or Mn\textsuperscript{52}, the likely explanation for the reduced Pb flux in the heart of the microniche is that, at some point during the deployment, the concentrations of Pb$^{2+}$ and S(-II) increased past a level where the rate of PbS precipitation exceeded the rate of Pb$^{2+}$ release by Fe mineral dissolution.

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### Trace metal fluxes from subsurface microniches

The location of the two subsurface microniches, $S_{A}$ and $S_{B}$, are shown in Fig. 2. They were not part of the experimental design, but the presence of such features has been attributed to bioturbation\textsuperscript{6}. Within them, the fluxes of all metals were higher than the background, with the exception of Cu. Microniche $S_{A}$ can be defined by above background fluxes of Mn, Fe and Zn; microniche $S_{B}$ by fluxes of Mn, Fe, Co, Ni and Zn (Table 1).

Microniches $S_{A}$ and $S_{B}$ are comparable in terms of size (Fe microniche sizes: 1.54 and 2.52 mm$^2$, respectively) and amounts of Fe and Mn mobilized. The Fe microniches were defined by a slightly larger number of points than the corresponding microniches of the other metals, therefore these Fe points were used to analyse the correlations between all metal fluxes at these locations. The simultaneous mobilization of Mn and Co in marine sediments has

### Table 2

| Surface Microniche, $I_{A}$ | Fe | Co | Ni | Cu | Zn | Pb |
|-----------------------------|----|----|----|----|----|----|
| Mn                          | 0.866 | 0.124 | 0.100 | 0.004 | 0.058 | 0.287 |
| Fe                          | ×   | 0.322 | 0.248 | 0.008 | 0.006 | **0.490 (0.839)** |
| Co                          | −   | ×   | **0.887** | 0.017 | 0.164 | 0.297 |
| Ni                          | −   | −   | ×   | **0.002** | 0.122 | 0.173 |
| Cu                          | −   | −   | −   | ×   | **0.000** | 0.067 |
| Zn                          | −   | −   | −   | ×   | **0.102** | 

| Subsurface Microniches, $S_{A}$ and $S_{B}$ | Mn | Fe | Co | Ni | Zn | Pb |
|--------------------------------------------|----|----|----|----|----|----|
| Mn                                         | ×   | **0.958** | 0.962 | 0.512 | 0.537 | 0.140 |
| Fe                                         | **0.962** | ×   | **0.951** | 0.484 | 0.568 | 0.096 |
| Co                                         | **0.986** | ×   | **0.955** | **0.632** | 0.531 | 0.191 |
| Ni                                         | **0.961** | **0.933** | **0.981** | ×   | **0.065** | 0.712 |
| Zn                                         | **0.761** | 0.825 | 0.774 | **0.811** | ×   | 0.033 |
| Pb                                         | **0.593** | **0.588** | 0.604 | **0.672** | **0.704** | ×   |

| $S_{A}$ (n = 11; $p < 0.01$ for $R^2 > 0.735$) | $S_{B}$ (n = 18; $p < 0.01$ for $R^2 > 0.530$) |

It is therefore likely that reductive dissolution of Fe (oxyhydr)oxide minerals is driving the Pb release across $I_{A}$. Stahl et al.\textsuperscript{28} found a peak Pb release of ~1.5 fmol cm$^{-2}$ s$^{-1}$ (~1.6 × higher than at $I_{A}$) during a 6 h deployment at the location of an artificial worm burrow which was closely associated with increased O$_2$ concentrations inside it. They suggested that Pb was released by local oxidation of Pb sulphide minerals. This process is unlikely to have determined Pb mobilization here, because O$_2$ was diminished at $I_{A}$.

The relationship between Fe and Pb, while statistically significant, is not consistent across $I_{A}$. When the fluxes of the two metals are compared against each other, there appear to be two antagonistic mechanisms operating: one mobilizes both Fe and Pb at closely related rates, while another limits Pb flux into the HR-DGT when Fe fluxes are high (Fig. 3). This relative limitation can be identified by a feature defined by six contiguous measurement points at the centre of the $I_{A}$ microniche: a possible ‘reaction hotspot’ (Figs 2 and 3). Rapid organic matter mineralisation has been linked to the formation of sulphide, S(-II), ‘hotspots’ in otherwise sulphide-free sediments\textsuperscript{27}. Given that Pb is thought to react with S(-II) much faster than Fe or Mn\textsuperscript{52}, the likely explanation for the reduced Pb flux in the heart of the microniche is that, at some point during the deployment, the concentrations of Pb$^{2+}$ and S(-II) increased past a level where the rate of PbS precipitation exceeded the rate of Pb$^{2+}$ release by Fe mineral dissolution.

When compared to their average fluxes in the top 1 mm of the sediment, the fluxes of Co, Ni and Zn within the area of $I_{A}$ (as defined by Fe measurements) were lower than in background oxic sediment, indicating that there was a process removing these metals from solution (Table 1; Fig. 2). The lower flux of these metals to the HR-DGT resin may be caused by different processes working independently or together. A localized decrease in H$^+$ ion activity caused by their consumption in the dissimilatory reduction of Fe minerals\textsuperscript{53} could promote immobilization of these metals; however, such an effect has not been demonstrated at this spatial resolution and under these conditions. Zn is also known to react rapidly with S(-II)\textsuperscript{52}. Given the higher concentrations of Zn in the pore water, it is possible that a high rate of ZnS mineral formation introduced a localized sink for S(-II) within $I_{A}$. While Co and Ni do not react rapidly with S(-II), they are thought to be easily incorporated into sulphide minerals\textsuperscript{54,55}. Given that Pb is thought to react with S(-II) much faster than Fe or Mn\textsuperscript{52}, the likely explanation for the reduced Pb flux in the heart of the microniche is that, at some point during the deployment, the concentrations of Pb$^{2+}$ and S(-II) increased past a level where the rate of PbS precipitation exceeded the rate of Pb$^{2+}$ release by Fe mineral dissolution.
et al. or modified simultaneously in small volumes. The role of anoxic microniches should be duly considered in future research. Fe and Mn (oxyhydr)oxides are lost, e.g. ZnS, possibly incorporating Ni and Co) are created, and existing ones (e.g. PbS and SnS, possibly incorporating Ni and Co) are created, and existing ones (e.g. PbS) may be indirectly linked to these localized features to a greater extent than at Ix. The combination of different porosities and types of organic matter in these microniches is also possible. As noted for Ix, the possibility that labile trace metals are rendered partially or wholly unavailable for DGT uptake by organic ligands or microorganisms may also apply here, albeit to a lesser extent than at Ix.

Figure 3. Time-averaged Fe flux vs. time-averaged Pb flux at individual analysis points inside the designated area of the surface microniche, Ix. The two linear regressions, and corresponding R² values, show the correlation between Fe and Pb fluxes with and without the points inside the reaction hotspot (dashed and solid black lines, respectively).

been previously linked to redox cycling of the Mn minerals19, 20, 32. The relatively sharp features showing elevated Fe and Mn fluxes and strong relationships between their fluxes and those of Co, suggest that this is also the case in Sx and Sp (Table 2)38. However, the significant differences between average fluxes of Co, Ni, and Zn in Sx and Sp could indicate that these redox processes operated at different rates, thereby influencing the fluxes of these metals from what otherwise appear to be similar microniches. It is also possible that different amounts of these metals were present at Sx and Sp. Previous studies have found that the reactivity of the organic matter in microniches can be an important parameter in determining conditions where and when S(-II) formation and the subsequent precipitation of metals as sulphide minerals occurs6, 51. The organic matter introduced with the Spirulina aggregate was relatively fresh and highly labile. If oxidation of organic matter in the subsurface microniches is driving the release of metals here, they are likely to have been present in the sediment for some time before the start of the deployment and therefore not as fresh and or labile as at Ix. As the surrounding environment is anoxic, this less reactive organic matter can still achieve similar reducing conditions to Ix. The good correlation between Fe and Mn, and the poor correlation between Fe and Ni, Zn and Pb in Sp are indicative of this, while in Sx the correlations are consistently strong between all the metals that could be measured. This could indicate that the organic matter in Sx is more reactive than in Sp and that precipitation of insoluble sulphide minerals reduced the fluxes of some metals in Sx. However, there are other possible reasons for the differences observed between these two sub-surface microniches. Microniches that have a low porosity will have reduced rates of diffusion within them and therefore are more likely to contain more reducing conditions than their surrounding environment6. 21. The combination of different porosities and types of organic matter in these microniches is also possible. As noted for Ix, the possibility that labile trace metals are rendered partially or wholly unavailable for DGT uptake by organic ligands or microorganisms may also apply here, albeit to a lesser extent than at Ix.

Organic matter deposition at the SWI and its implications for elemental cycling in sediments. We have shown for the first time that deposition of organic matter and the resulting highly localized anoxic conditions have brought about intense mobilization of Fe and Mn at the SWI, proving part of our initial hypothesis. However, rather than observing release of some metals (e.g. Zn, Ni and Co) as the oxides are reductively dissolved, there is a net removal, presumably due to localised formation of S(-II) within the microniche, as established elsewhere51. We expect that under faster rates of mineralisation and/or higher concentrations of sulphate, the pyritization of Pb would be more extensive and a greater removal of soluble Fe may be observed. The intense release of Fe that we measured from the anoxic microniche suggests that the cycles of other elements (and their isotopes), thought to be associated with redox-active Fe minerals (e.g. C, N, P and S, as well as trace elements such as: Co, Ni, Cu, Zn and As), may also be indirectly linked to these localized features to a greater extent than previously realized; however, their magnitude should be more accurately quantified through targeted experiments across a wider range of environments. Anoxic micro-features, such as the one seen at Ix, have previously been observed both in-situ and in laboratory mesocosms4, 6, 11, 29, 59, and modelling has shown that localised decomposition arising from settling organic matter can effect increased rates of nutrient cycling. Here we demonstrate that this may also be an important factor in processes of early diagenesis, where new mineral phases (e.g. PbS and ZnS, possibly incorporating Ni and Co) are created, and existing ones (e.g. Fe and Mn (oxyhydr)oxides) are lost or modified simultaneously in small volumes. The role of anoxic microniches should be duly considered in future studies seeking to developing elemental budgets at the SWI, such as those recommended by Homoky, et al. 37.

Our results also provide circumstantial evidence that the translocation of labile organic matter from the SWI by faunal activity can move microbial mineralisation and the associated biogeochemical cycling to deeper in the sediments. This process has been implicated previously when hotspots of trace metal and S(II) release have been
Material and Methods

Sediment handling and characterisation. Due to the difficulties associated with sampling deep-sea sediments, we chose to use a coastal sediment for this study. The sediment and overlying seawater were collected from Workington Marina, (UK) (54°38′52.54″N; 3°33′57.52″W) in April 2009 and stored in two 50 L plastic containers at 15–17°C. The O₂ concentration in the overlying water was maintained at 100% air sat. At the time of sampling, sediment was inhabited by various macrofauna species (e.g. Corophium sp. and juvenile polychaetes). In March 2010, the sediment was transferred into smaller acrylic aquaria (H × L × W: 15 × 14 × 8 cm: similar to one that had been used previously in other studies of oxygen dynamics at the SWI), using a custom-made sediment sampler. The procedure maintained a virtually intact sediment structure. The setup was left for 2–3 d before the experiment was initiated, during which the overlying water inside the aquaria was circulated using pumps to ensure a steady semi-laminar flow of 2–3 cm s⁻¹ and a constant O₂ concentration at air saturation in the overlying water during the pre-incubation. The temperature was kept constant at 19°C (±1°C). Active removal of visible fauna in the time leading up to the experiment aimed to minimize macrofauna activity during the experiments. The dissolved and solid fractions of the sediment samples underwent an extensive elemental characterisation for C, N, major cations and anions and trace metals (analyses are described in Section S1. iii). These were in good agreement with previously measured values in a marine sediment (Table S1). Five background sediment micro-electrode profiles of H₂S were measured at a resolution of 200 μm, using a Clark type H₂S sensor connected to picoammeter (FA-2000, Unisense.dk). The profiles were measured at the centre of the mesocosm to avoid interfering with the rest of the experiment: no H₂S was detected.

Preparation of Spirulina aggregates and their characterisation. To form standardized labile particulate organic material, freeze dried Spirulina powder (SigmaAldrich.com) was mixed with a 0.5% agar paste (wt./wt.) made with ultra-pure water (Milli-Q™, Millipore; resistivity 18.2 MΩ·cm), rolled into spherical aggregates and dried. After drying, the aggregates had a diameter of 2–3 mm and a weight of 5–7 mg. Carbon mineralisation rates were analysed in subsets of Spirulina material and aggregates (Supporting Information, S.i), these were within previously observed values (Supporting Information, S.vi). Their total metal content was also analysed and found to be representative of natural aggregates (Supporting Information, S.i). The Fe:C ratio in the aggregates was found to be between 24–54 μmol Fe:1 mol C, which is within the range of values previously measured in particulate organic matter collected from ocean water columns (Table S1)65–68; other metal concentrations were also representative of natural aggregates 69.

Preparation of the O₂ sensitive optode foil and optode imaging. The principles for planar optode imaging have previously been described in detail70–72. The preparation of the O₂-sensitive optode used here, and the details pertaining to the imaging, are described in detail in the Supporting Information (S1.ii). Briefly, The O₂ sensitive sensing layer was coated directly onto a fibre optic faceplate measuring 75 × 50 × 3 mm (Schott.com) and mounted into the removable sidewall of an acrylic aquarium (see later). The excitation light was delivered by a 10 W blue LED (light emitting diode; LZ4–40B200, LEDENGIN.com), with a peak wavelength of 465 nm. A CCD camera (Charged Couple Device) (SensiCam, PCO.de), mounted with a prime macro lens (Sigma 50 mm F2.8 EX DG Macro) was used to capture the emitted light. The macro lens and LED were equipped with the appropriate filters for dissolved oxygen measurements. The camera and LED operation was synchronized with a custom made trigger box controlled by the software Look@Mollif. The method detection limit (MDL) of the planar optode was determined at the end of the deployment to be 3.0 μM (see Supporting Information, S1.iv); herein this concentration is used as an operationally defined threshold for ‘anoxic’.
Ultra-thin DGT gel preparation. Ultra-thin (50 µm-thick) resin gels sheets and calibration standards for the LA-ICP-MS were prepared following the method described in Lehto, et al. The details of this procedure are provided in the Supporting Information (S.iv). Hydrated resin gel sheet was trimmed to size (approx. dimensions: 50 × 70 mm) and placed between two sheets of acid washed Nuclepore membrane (pore size: 0.4 µm; thickness: 10 µm) (Whatman, Maidstone, U.K.). Herein, these three layers, specially designed for high-resolution measurements, are referred to as “High-Resolution DGT” (HR-DGT).

HR-DGT-Planar Optode (HR-DGT-PO) sensor assembly and deployment. The acrylic aquaria used in this work were designed with a replaceable sidewall (Fig. 4) allowing easy deployment of the HR-DGT-PO sensor, while ensuring minimal disturbance to the sediment structure. Prior to deployment, the sidewall containing the fibre optic faceplate was quickly wiped with 96% ethanol and rinsed with ultra-pure water. Using a pair of acid-washed plastic forceps, a 45 × 46 mm HR-DGT gel was laid flat on top of the fibre optic faceplate, facing the optode sensor. The HR-DGT was gently attached using water resistant vinyl tape (super 33+, Scotch®). Approximately 5 mm of the gel was covered by the vinyl tape on each side, leaving an exposed internal area of 40 × 42 mm.

Prior to deploying the HR-DGT-PO sensor, the water in the aquarium was drained, after which the aquarium was tilted back 15–20° to maintain the overall sediment structure and the sidewall was replaced with the sidewall containing the HR-DGT-PO sensor, ensuring a good contact between the sediment and the HR-DGT. The process of switching the sidewall took approximately 30 s, with a view to minimizing any oxygen contamination of the sediment. The aquarium was immediately filled with water, ensuring minimal disturbance of the sediment.

Immediately after deploying the sensor, two *Spirulina* aggregates and one aggregate of commercially available fish food of a similar size were gently placed in a row, at least 10 mm apart, on the water-sediment-HR-DGT interface using clean plastic forceps. The deployment was carried out for 44 h, during which the O₂ concentration in the overlying water was monitored and the ambient temperature were kept at 19 °C (±1). The overlying water inside the aquaria was circulated and carefully flushed with air. Images of the mounted combined sensor were obtained at the start and at the end of the deployment, using ambient light to confirm the positions of the SWI and the aggregates on the optode images. Subsequent optode measurement images were recorded in darkness to avoid interference of ambient light at 20 min intervals.

At the end of the deployments, the combined sensor was retrieved by removing the sidewall as described above and the sensor was immediately rinsed thoroughly with MQ water to remove sediment particles. The HR-DGT was then cut out using a PTFE-coated razorblade and placed between two acid-washed plastic sheets for subsequent storage at 5 °C. The HR-DGT resin gels were dried within one week of the end of the deployment (see below).

LA-ICP-MS analysis of the HR-DGT. Prior to analysis, the ultra-thin resin gel of the HR-DGT was separated from the Nuclepore membranes and dried using the method described by Lehto, et al. The gel was then trimmed (42 × 24 mm) targeting the SWI to fit into the ablation cell of the laser ablation unit and mounted onto a microscope slide using double-sided tape. An area of 40 × 23.22 mm was then analysed using laser ablation inductively coupled mass spectrometer (LA-ICP-MS) following the method used by Lehto, et al. The resolution of the measurements was 280 µm along the raster line, which was 100 µm wide. The interval between the centres...
of adjacent lines was 500 μm. C¹³, Rh¹⁰³ and In¹¹³ were used as internal standards. Further details regarding the laser ablation of the gels can be found in the Supporting Information (Sv).

**Data analysis.** The location of the ablation area with respect to the planar optode image was determined by measuring the exact distances from the edges of the gel. HR-DGT and O₂ optode results are primarily focussed on a specific internal area of 15.1 × 23.22 mm (Fig. S1). The original spatial resolution of each discrete measurement point (‘pixel’) is 500 × 280 μm (HR-DGT) and 37 × 37 μm (planar optode); however, the resolution of the planar optode images was decreased to match that of the HR-DGT to allow for direct comparison between the two. All calibrated images were saved in TIFF file formats and were further processed using ImageJ. The presented trace metal images are artificially increased in size from the original 41 × 84 to 225 × 348 for visualisation purposes, while avoiding interpolation between data points; all data treatment is based on the original pixel size.

Statistical analysis of the metal fluxes from the microniches were carried out using MS Excel 2016 (correlations) and IBM SPSS Statistics for Windows (Version 22.0, Armonk, NY: IBM Corp) (one-way analysis of variance, ANOVA, with Tukey’s post-hoc test to compare means); the level of significance was 0.05, unless stated otherwise.

**Data Availability Statement.** The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

**References**

1. Alldredge, A. L. & Gotschalk, C. In situ settling behavior of marine snow. Limnol. Oceanogr. 33(3), 339–351 (1988).
2. Grossart, H.-P. & Simon, M. Limnetic macroscopic organic aggregates (lake snow): Occurrence, characteristics, and microbial dynamics in Lake Constance. Limnol. Oceanogr. 38(3), 532–546 (1993).
3. Glud, R. N. et al. Copepod carcasses as microbial hot spots for pelagic denitrification. Limnol. Oceanogr. 60(6), 2026–2036 (2015).
4. Glud, R. N. et al. In situ microscale variation in distribution and consumption of O₂: A case study from a deep ocean margin sediment (Sagami Bay, Japan). Limnol. Oceanogr. 54(1), 1 (2009).
5. Motelica-Heino, M., Naylor, C., Zhang, H. & Davison, W. Simultaneous release of metals and sulphide in lacustrine sediment. Environ. Sci. Technol. 37, 4374–4381 (2003).
6. Lehto, N., Glud, R. N., Nordi, G. à, Zhang, H. & Davison, W. Anoxic microniches in marine sediments induced by aggregate settlement: Biogeochemical dynamics and implications. Biogeochemistry 119(1–3), 307–327 (2014).
7. Emery, K. O. & Rittenberg, S. C. Early diagenesis of California basin sediments in relation to origin of oil. AAPG Bull. 36(5), 735–806 (1952).
8. Jorgensen, B. B. Bacterial sulphate reduction within reduced microniches of oxidized marine sediments. Mar. Biol. 41(7–17) (1977).
9. Stockdale, A., Davison, W. & Zhang, H. Micro-scale biogeochemical heterogeneity in sediments: A review of available technology and observed evidence. Earth-Sci. Rev. 92, 81–97 (2009).
10. Widerlund, A., Nowell, G. M., Davison, W. & Pearson, D. G. High-resolution measurements of sulphur isotope variations in sediment pore-waters by laser ablation multicollector inductively coupled plasma mass spectrometry. Chem. Geol. 291, 278–282 (2012).
11. Marzocchi, U., Thamdrup, B., Stief, P., & Glud, R. N. Effects of settled diatom aggregates on benthic nitrogen cycling. Limnol. Oceanogr. Submitted (2017).
12. Klinkhammer, G., Heggie, D. & Graham, D. Metal diagenesis in oxic marine sediments. 61(2), 211–219 (1982).
13. Shaw, T. J., Gieskes, J. M. & Jahnke, R. A. Early diagenesis in differing depositional environments: The response of transition metals in pore water. 54(5), 1233–1246 (1990).
14. Burdige, D. J., Geochemistry of marine sediments. (Princeton University Press Princeton, 2006).
15. Benninger, L. K., Allen, R., Cochran, J. & Turekian, K. Effects of biological sediment mixing on the 210 Pb chronology and trace metal distribution in a Long Island Sound sediment core. Earth Planet. Sci. Lett. 43(2), 241–259 (1979).
16. van Cappellen, P. & Wang, Y. Cycling of iron and manganese in surface sediments: a general theory for the coupled transport and reaction of carbon, oxygen, nitrogen, sulfur, iron and manganese. Am. J. Sci. 296, 197–243 (1996).
17. Krumins, V., Gehrke, M., Arndt, S., Van Cappellen, P. & Regnier, P. Dissolved inorganic carbon and alkalinity fluxes from coastal marine sediments: Model estimates for different shelf environments and sensitivity to global change. Biogeosciences 10(3), 371–398 (2013).
18. Gao, Y. et al. Response of diffusive equilibrium in thin films (DET) and diffusive gradients in thin films (DGT) trace metal profiles in sediments to pythodetritus mineralisation. Environ. Chem. 9(1), 41–47 (2012).
19. Fones, G. R., Davison, W. & Hamilton-Taylor, J. The fine-scale remobilization of metals in the surface sediment of the North-East Atlantic. Cont. Shelf Res. 24(13), 1485–1504 (2004).
20. Tankéré-Muller, S. et al. Fine scale remobilisation of Fe, Mn, Co, Ni, Cu and Cd in contaminated marine sediment. Mar. Chem. 106(1–2), 192–207 (2007).
21. Stockdale, A., Davison, W. & Zhang, H. High-resolution two-dimensional quantitative assessment of phosphorus, vanadium and arsenic, and qualitative analysis of sulfide, in a freshwater sediment. Environ. Chem. 5(2), 143–149 (2008).
22. Glud, R. N., Ramsing, N. B., Gundersen, J. K. & Klimant, I. Planar optrodes: a new tool for fine scale measurements of two-dimensional O₂ distribution in benthic communities. Mar. Ecol. Prog. Ser. 140, 217–226 (1996).
23. Zhu, Q. Z., Allen, R. C. & Fan, Y. High-performance planar pH fluorosensor for two-dimensional pH measurements in marine sediment and water. Environ. Sci. Technol. 39, 8906–8911 (2005).
24. Stromberg, N. Determination of Ammonium Turnover and Flow Patterns Close to Roots Using Imaging Optodes. Environ. Sci. Technol. 42(1630–1637) (2008).
25. Zhu, Q. & Aller, R. C. A rapid response, planar fluorosensor for measuring two-dimensional pCO₂ distributions and dynamics in marine sediments. Limnol. Oceanogr. 6(7), 326–336 (2010).
26. Glud, R. N. et al. Benthic Carbon Mineralization and Nutrient Turnover in a Scottish Sea Loch: An Integrative In Situ Study. Aquat. Geochem. 22(5), 443–467 (2016).
27. Williams, P. N. et al. Localized flux maxima of arsenic, lead, and iron around root apices in flooded lowland rice. Environ. Sci. Technol. 48(15), 8498–8506 (2014).
28. Stahl, H. et al. A combined sensor for simultaneous high resolution 2-D imaging of oxygen and trace metals fluxes. Limnol. Oceanogr. Methods 10(5), 389–401 (2012).
29. Glud, R. N. Oxygen dynamics of marine sediments. 4(4), 243–289 (2008).
30. Lehto, N. J. Principles and Application in Soils and Sediments of Diffusive Gradients in Thin-Films for Environmental Measurements, edited by William Davison (Cambridge University Press, 2016), pp. 146–173.
31. Lehto, N. J., Davison, W. & Zhang, H. The use of ultra-thin diffusive gradients in thin-films (DGT) devices for the analysis of trace metal dynamics in soils and sediments: a measurement and modelling approach. Environ. Chem. 9(4), 415–423 (2012).
32. Zhang, H. et al. Localised remobilization of metals in a marine sediment. Sci. Total Environ. 296(1), 175–187 (2002).
33. Fones, G. R., Davison, W., Holby, O., Jorgensen, R. B. & Thamdrup, B. High-resolution metal gradients measured by in situ DGT/DET deployment in Black Sea sediments using an autonomous benthic lander. Limnol. Oceanogr. 46(4), 982–988 (2001).
34. Davison, W., Zhang, H. & Warnken, K. W., Chapter 16. Theory and applications of DGT measurements in soils and sediments in Comprehensive Analytical Chemistry, edited by G. Mills R & B. Vrana (Elsevier, 2007), Vol. 48, pp. 353–378 (10 May 2007).
35. Homoky, W. B. et al. Iron and manganese diagenesis in deep sea volcanogenic sediments and the origins of pore water colloids. 75(17), 5032–5048 (2011).
36. Klar, J. K. et al. Stability of dissolved and soluble Fe(II) in shelf sediment pore waters and release to anoxic water column (2017).
37. Homoky, W. B. et al. Quantifying trace element and isotope fluxes at the ocean–sediment boundary: a review. Phil. Trans. R. Soc. A 374(2081), 20140246 (2016).
38. Sochaczewski, L., Davison, W., Zhang, H. & Tych, W. Understanding small-scale features in DGT measurements in sediments. Environ. Sci. Chem. 6, 477–485 (2009).
39. Lutz, M., Dunbar, R. & Caldeira, K. Regional variability in the vertical flux of particulate organic carbon in the ocean interior. Global Biogeochem. Cy. 16(3) (2002).
40. Boetius, A. et al. Export of Algal Biomass from the Melting Arctic Sea Ice. Science 339(6126), 1430–1432 (2013).
41. Lalonde, K., Mucci, A., Ouellet, A. & Gélinas, Y. Preservation of organic matter in sediments promoted by iron. Nature 483(7388), 198–200 (2012).
42. Kleber, M. et al. Chapter One - Mineral–Organic Associations: Formation, Properties, and Relevance in Soil Environments in Advances in Agronomy, edited by L. Sparks Donald (Academic Press, 2015), Vol. 130, pp. 1–140.
43. Alkhatib, M., del Giorgio, P. A., Gelinas, Y. & Lehmann, M. F. Benthic fluxes of dissolved organic nitrogen in the lower St. Lawrence estuary and implications for selective organic matter degradation. 10(11), 7659–7662 (2013).
44. Rozen, T. F. et al. Iron-sulfur-phosphorus cycling in the sediments of a shallow coastal bay: Implications for sediment nutrient release and benthic macroalgal blooms. Limnol. Oceanogr. 47(5), 1346–1354 (2002).
45. Bennett, W. W. et al. Inorganic arsenic and iron (II) distributions in sediment porewaters investigated by a combined DGT–colourimetric DET technique. Environ. Chem. 9(1), 31–40 (2011).
46. Ding, S. et al. In situ, high-resolution evidence for iron-coupled mobilization of phosphorus in sediments. Sci. Rep. 6, 24341 (2016).
47. Rullkötter, J. Organic matter: the driving force for early diagenesis in Marine geochemistry (Springer, 2006), pp. 125–168.
48. Goh, M. A. et al. Distribution and sources of organic matter in surface marine sediments across the North American Arctic margin. 118(9), 4017–4035 (2013).
49. Tessier, A., Rapin, F. & Carignan, R. Trace metals in oxic lake sediments: possible adsorption onto iron oxyhydroxides. Geochim. Cosmochim. Acta 49(1), 183–194 (1985).
50. Widerlund, A. & Davison, W. Size and density distribution of sulphide-producing microminerals in lake sediments. Environ. Sci. Technol. 41(23), 8044–8049 (2007).
51. Stockdale, A., Davison, W. & Zhang, H. Formation of iron sulphide at faecal pellets and other microniches within suboxic surface sediment. Geochim. Cosmochim. Acta 74, 2663–2676 (2010).
52. Morse, J. & Luther, G. W. Chemical influences on trace metal-sulfide interactions in anoxic sediments. Geochim. Cosmochim. Acta 63(19/20), 3373–3378 (1999).
53. Raiswell, R. & Canfield, D. E. The iron biogeochemical cycle past and present. Geochim. Perspect. 1(1), 1–222 (2012).
54. Atkins, A. L., Shaw, S. & Peacock, C. L. Release of Ni from birnessite during transformation of birnessite to todorokite: Implications for Ni cycling in marine sediments. 189, 158–183 (2016).
55. Stockdale, A., Davison, W., Zhang, H. & Hamilton-Taylor, J. The association of cobalt with iron and manganese (oxyhydrate)oixides in marine sediment. Aquat. Geochem. 16, 575–585 (2010).
56. Davison, W. & Zhang, H. Progress in understanding the use of diffusive gradients in thin films (DGT) – back to basics. Environ. Chem. 9(1), 1–13 (2012).
57. Gadd, G. M. & Griffiths, A. J. Microorganisms and heavy metal toxicity. Microb. Ecol. 4(4), 303–317 (1977).
58. White, C., Wilkinson, S. C. & Gadd, G. M. The role of microorganisms in biosorption of toxic metals and radionuclides. Int. Biodeterior. Biodegradation 35(1), 17–40 (1995).
59. Franke, U., Polerecky, L., Precht, E. & Huetter, M. Wave Tank Study of Particulate Organic Matter Degradation in Permeable Sediments. Limnol. Oceanogr. 51(2), 1084–1096 (2006).
60. Zhu, Q., Aller, R. C. & Fan, Y. Two-dimensional pH distributions and dynamics in bioturbated marine sediments. Geochim. Cosmochim. Acta 70(19), 4933–4949 (2006).
61. Zhu, Q. Z., Aller, R. C. & Fan, Y. A new ratiometric, planar fluorosensor for measuring high resolution, two-dimensional pCO2 distributions in marine sediments. 101, 40–53 (2006).
62. Jézéquel, D. et al. Two-dimensional determination of dissolved iron and sulphur species in marine sediment porewaters by thin-film based imaging. Thau lagoon (France) 72(3), 420–431 (2007).
63. Michaud, E., Aller, R. C. & Stora, G. Sedimentary organic matter distributions, burrowing activity, and biogeochemical cycling: Natural patterns and experimental artifacts. 90(1), 21–34 (2010).
64. Kuhl, M., Steuckart, C., Eckert, G. & Jeroschewski, P. A H2S microsensor for profiling biofilms and sediments: application in an acidic lake sediment (1998).
65. Huang, H. Small-scale oxygen fluxes and remineralization in sinking aggregates. Limnol. Oceanogr. 46(7), 1624–1631 (2001).
66. Price, N. M. & Morel, F. M. Biological cycling of iron in the ocean. 3, 1–36 (1998).
67. Twining, B. S. et al. Metal quotas of plankton in the equatorial Pacific Ocean. 58(3–4), 325–341 (2011).
68. Nuester, J., Vogt, S., Newville, M., Kustka, A. & Twining, B., The Unique Biogeochemical Signature of the Marine Diazotroph Trichodesmium. 3(150) (2012).
69. Ho, T.-Y., Wan, L.-S., You, C.-F. & Lee, D.-C. The trace metal composition of size-fractionated plankton in the South China Sea: Biotic versus abiotic sources. 52(23), 1776–1788 (2007).
70. Glud, R., Ramsing, N., Gundersen, J. & Klimant, I. Planar optodes: a new tool for fine scale measurements of two-dimensional O–2 distribution in benthic communities. 140, 217–226 (1996).
71. Holst, G. et al. A modular luminescence lifetime imaging system for mapping oxygen distribution in biological samples. 51(1), 163–170 (1998).
72. Holst, G. & Grunwald, B. Luminescence lifetime imaging with transparent oxygen optodes. Sens. Actuator B-Chem. 74(1), 78–90 (2001).
73. Fischer, J. P. & Wenzhöfer, F. A novel planar optode setup for concurrent oxygen and light field imaging: Application to a benthic phototrophic community. Limnol. Oceanogr. Methods 8(6), 254–268 (2010).
74. Flaxce, K. et al. Effects of temperature and irradiance on a benthic microalgal community: A combined two-dimensional oxygen and fluorescence imaging approach. Limnol. Oceanogr. 59(5), 1599–1611 (2014).
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Author Contributions
N.L. compiled the final manuscript and prepared the figures, as well as contributing to conducting the experiment, collection and analysis of the data, interpreting the results and writing the manuscript. M.L. contributed to the conducting of the experiment, collection and analysis of the data, interpreting the results and writing the manuscript. R.G. and W.D. coordinated the research and contributed to interpreting the results and writing the manuscript. All authors contributed to the conception and design of the experiment, the review and subsequent revisions of the manuscript.

Additional Information
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