

1 **Biogeochemical Impact of Cable Bacteria on Coastal Black Sea Sediment**

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15 **ABSTRACT**

16 Cable bacteria can strongly alter sediment biogeochemistry. Here, we used laboratory
17 incubations to determine the potential impact of their activity on the cycling of Fe, phosphorus (P) and
18 sulphur (S). Microsensor depth profiles of oxygen, sulphide and pH in combination with electric
19 potential profiling and FISH analyses showed a rapid development (<5 days) of cable bacteria,
20 followed by a long period of activity (>200 days). During most of the experiment, the current density
21 correlated linearly with the oxygen demand. Sediment oxygen uptake was attributed to activity of
22 cable bacteria and the oxidation of reduced products from anaerobic degradation of organic matter,
23 such as ammonium. Pore water sulphide was low (<5 μM) throughout the experiment. Sulphate
24 reduction acted as the main source of sulphide for cable bacteria. Pore water Fe^{2+} reached levels of up
25 to 1.7 mM during the incubations, due to the dissolution of FeS (30%) and siderite, an Fe carbonate
26 mineral (70%). Following upward diffusion of Fe^{2+} , a surface enrichment of Fe oxides formed. Hence,

27 besides FeS, siderite may act as a major source of Fe for Fe oxides in coastal surface sediments where
28 cable bacteria are active. Using μ XRF, we show that the enrichments in Fe oxides induced by cable
29 bacteria are located in a thin subsurface layer of 0.3 mm. We show that similar subsurface layers
30 enriched in Fe and P are also observed at field sites where cable bacteria were recently active and
31 little bioturbation occurs. This suggests that such subsurface Fe oxide layers, which are not always
32 visible to the eye, could potentially be a marker for recent activity of cable bacteria.

33 Key words: cable bacteria, elemental cycling, solute fluxes, iron

34 **1. INTRODUCTION**

35 Depletion of oxygen (O_2) in bottom waters (i.e. water directly above the seafloor) of coastal areas
36 is increasing worldwide, as a consequence of eutrophication and climate change (Diaz and Rosenberg
37 2008; Schmidtko et al. 2017; Breitburg et al. 2018). Low O_2 can lead to the development of coastal
38 ‘dead zones’ characterised by recurrent mortality of marine life (Rabalais et al. 2002; Diaz and
39 Rosenberg 2008). Progressive eutrophication induces a characteristic response of coastal systems with
40 transient and seasonal hypoxia ($O_2 < 63 \mu\text{M}$) transitioning into permanent anoxia ($O_2 = 0 \mu\text{M}$). In this
41 later stage, free sulphide (H_2S) may escape from the sediment and accumulate in the bottom water, a
42 condition referred to as euxinia (Diaz and Rosenberg 2008; Kemp et al. 2009; Rabalais et al. 2014).
43 As H_2S is highly toxic to higher fauna, the development of euxinia may aggravate the ecological
44 consequences. However, the presence of iron (Fe) and manganese (Mn) oxides in surface sediments
45 may delay this transition towards euxinia by removing H_2S and thus preventing an efflux of H_2S to
46 the overlying water (Kristiansen et al. 2002; Kristensen et al. 2003; Diaz and Rosenberg 2008).

47 Cable bacteria are multicellular filamentous sulphur(S)-oxidising bacteria (Pfeffer et al. 2012)
48 that strongly enhance the formation of Fe and Mn oxides and efficiently remove H_2S from surface
49 sediments (Risgaard-Petersen et al. 2012; Seitaj et al. 2015; Sulu-Gambari et al. 2016a). Cable
50 bacteria belong to the *Desulfobulbaceae* family of the Deltaproteobacteria (Trojan et al. 2016;
51 Kjeldsen et al. 2019). Cable bacteria can spatially link the oxidation of H_2S in deeper sediments to the
52 reduction of O_2 near the sediment-water interface by transporting electrons over centimetre scale

53 distances (Pfeffer et al. 2012) through a conductive fibre network that is embedded in the cell
54 envelope (Meysman et al. 2019). This spatial coupling of surficial O₂ reduction with H₂S oxidation at
55 several centimetres depth creates a suboxic zone that is devoid of any O₂ and H₂S, and provides cable
56 bacteria a competitive advantage over other S-oxidising bacteria in aquatic environments (Meysman
57 2018). Cable bacteria have been documented in a range of fresh water (Risgaard-Petersen et al. 2015;
58 Müller et al. 2016) and marine environments (Malkin et al. 2014; Burdorf et al. 2017), however, they
59 appear to be particularly active in sediments overlain by seasonally hypoxic bottom waters (Seitaj et
60 al. 2015; Burdorf et al. 2018).

61 The metabolic activity of cable bacteria establishes an electrical circuit in the sediment, which
62 involves an electron current through the cable bacteria filaments (Bjerg et al. 2018), and an ionic
63 current through the pore water in the opposite direction (Naudet and Revil 2005; Revil et al. 2010;
64 Risgaard-Petersen et al. 2012). As a consequence, an electric potential (EP) is generated in the
65 sediment, which can be used as a reliable indicator for activity of cable bacteria (Risgaard-Petersen et
66 al. 2014).

67 Cable bacteria activity additionally generates a distinct biogeochemical signature, that can be
68 assessed by pH, O₂ and H₂S depth profiling (Nielsen et al. 2010). Their activity leads to the
69 development of a suboxic zone (i.e. a zone where O₂ and H₂S are both absent), and also induces a pH
70 profile that strongly changes with depth. Cathodic O₂ reduction ($O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$) in the oxic
71 zone of the sediment results in a pH maximum (~9) due to proton consumption, whereas anodic
72 sulphide oxidation ($H_2S + 4 H_2O \rightarrow SO_4^{2-} + 10H^+ + 8e^-$) causes a pH minimum (<6.5) in the anoxic
73 zone (Fig. 1A; Nielsen et al. 2010; Meysman et al. 2015).

74 The presence of cable bacteria in sediments can strongly impact the elemental cycling of Fe,
75 Mn, Ca and S (Risgaard-Petersen et al. 2012; Seitaj et al. 2015; Rao et al. 2016; Sulu-Gambari et al.
76 2016a; van de Velde et al. 2016). Pore water acidification induced by cable bacteria activity can lead
77 to dissolution of calcium (Ca) carbonates, Fe carbonates (siderite), Mn carbonates and FeS in the zone
78 where the pH is low, thus generating high concentrations of Fe²⁺ and Mn²⁺ in the pore water

79 (Risgaard-Petersen et al. 2012; Rao et al. 2016). When these dissolved species diffuse upward this can
80 lead to strong enrichments of Fe and Mn oxides upon contact with O_2 , or for dissolved Fe^{2+} , also upon
81 contact with Mn oxides (Wang and Van Cappellen 1996; Seitaj et al. 2015; Sulu-Gambari et al.
82 2016a). These metal oxides are capable of efficiently buffering the benthic release of H_2S and
83 phosphate (HPO_4^{2-}) during periods with low bottom water O_2 . This so-called ‘firewall’ for H_2S and
84 alteration of the timing of HPO_4^{2-} release linked to this buffering can play a key role in regulating
85 water quality in seasonally hypoxic coastal systems (Seitaj et al. 2015; Sulu-Gambari et al. 2016b;
86 Hermans et al. 2019a).

87 In coastal sediments, O_2 typically penetrates to a depth of only several mm’s below the
88 sediment-water interface (Rasmussen and Jørgensen 1992; Rabouille et al. 2003; Glud 2008). This
89 also holds true for sediments inhabited by active cable bacteria (Nielsen et al. 2010; Pfeffer et al.
90 2012; Larsen et al. 2015). Hence, the oxidation of upward diffusing Fe^{2+} and Mn^{2+} is expected to take
91 place below and not at the sediment-water interface. We hypothesise that, as a consequence, in the
92 initial stages of cable bacteria activity and in the absence of bioturbation, most Fe and Mn oxide
93 enrichments will be restricted to a thin subsurface layer of the sediment. However, the sample
94 resolution and timing of the collection of solid phase data in field and laboratory studies published so
95 far do not allow an assessment of this hypothesis.

96 Cable bacteria are suggested to thrive in coastal sediments characterised by high rates of H_2S
97 production due to high rates of organic matter mineralisation (Malkin et al. 2014; Burdorf et al. 2017;
98 Hermans et al. 2019a). Laboratory and model studies have shown that the dissolution of FeS accounts
99 for 12 to 94% of the H_2S consumed by cable bacteria, while the other source is H_2S production from
100 the reduction of SO_4^{2-} (Risgaard-Petersen et al. 2012; Meysman et al. 2015; Burdorf et al. 2018). At
101 present, it is not known if cable bacteria activity can establish in sediments that are relatively low in
102 FeS and dissolved H_2S .

103 In this study, we assess whether cable bacteria activity can establish and thrive in sediments
104 that are relatively poor in FeS. Although, this will be done in a controlled incubation experiment with

105 siderite-bearing sediments from a coastal site in the Black Sea, our findings are relevant for natural
106 environments populated by cable bacteria. The metabolic activity of cable bacteria is monitored using
107 microsensor profiles of pH, O₂, H₂S and EP. We also use sediment Fe and P speciation and μ XRF of
108 resin-embedded sediments to test whether we find evidence for subsurface enrichments in Fe oxides
109 and associated P. We find a rapid establishment of cable bacteria (<5 days) and the development of an
110 Fe oxide-rich subsurface layer, with the majority of the Fe ~70% supplied through dissolution of
111 siderite induced by cable bacteria activity. The depth of the Fe oxide layer was directly related to the
112 O₂ penetration depth and we propose that such subsurface enrichments in Fe, which also can contain P
113 and Mn, can be used as a marker for recent cable bacteria activity.

114 **2. METHODS AND MATERIALS**

115 **2.1. Study Area and Experimental Set-up**

116 In September 2015, 16 sediment cores (\varnothing 10 cm) were retrieved at a coastal site on the north-
117 western shelf of the Black Sea (27 m water depth; Fig. 1B; Table 1) using a multicorer (Oktopus
118 GmbH, Germany) as described in Lenstra et al. (2019). The overlying water was discarded, and the
119 upper 10 cm of the sediment was transferred into nitrogen purged aluminium bags that were sealed
120 and stored at 4 °C for several months. The anoxic storage is expected to have led to the death of all
121 macrofauna and most meiofauna (Coull and Chandler 2001; Riedel et al. 2012). Prior to incubation,
122 the sediment was passed through a 4 mm sieve to remove large debris and homogenised.
123 Subsequently, the sediment was transferred to 18 transparent polycarbonate cores (\varnothing 6 cm; 20 cm
124 length).

125 The bottom 15 cm of these cores was filled with sediment and the upper 5 cm with overlying
126 water. The cores were placed in two aquaria filled with artificial seawater (Instant Ocean Sea Salt +
127 Ultra High Quality (UHQ) water) with a salinity of 17.9, identical to the bottom water salinity at the
128 study site. The artificial sea water contained negligible concentrations of NH₄⁺, NO₃⁻, Fe, Mn and P as
129 described in Atkinson (1997) and Hovanec and Coshland (2004). The aquaria were kept in the dark at
130 a constant temperature (~20 °C), and the water was continuously aerated by two aquarium pumps.

131 Sixteen out of eighteen cores were exposed to oxygenated overlying water in the aquaria, whereas the
132 two remaining cores served as an anoxic control treatment. The control cores were tightly sealed with
133 rubber stoppers, to prevent the growth of cable bacteria by excluding O₂ (Nielsen et al. 2010).

134 Sampling for pore water and solid-phase analyses was performed at eight time points over a
135 total incubation period of 621 days. Each time point involved a three day procedure. On the first day,
136 microsensor depth profiles of EP, O₂, pH and H₂S were obtained in two randomly selected oxic cores
137 and the two anoxic control cores (O₂ profiling was not performed in the anoxic cores). On the second
138 day, solute fluxes were measured in the same oxic cores that were used for microsensor depth
139 profiling on the previous day. On the third day, the two cores were sectioned, of which only one core
140 was processed further for pore water and solid-phase analyses. Photographs were taken at four time
141 points (day 12; 33; 170 and 621) from one oxic core to follow the visual development of the surface
142 sediment during the experiment.

143 **2.2. High-resolution Microsensor Depth Profiling**

144 High-resolution depth profiles of pH, O₂ and H₂S were obtained (50- μ m depth resolution; 3
145 replicate profiles per oxic core; 2 replicate profiles per anoxic core) using commercial micro
146 electrodes (Unisense A.S., Denmark). The O₂ sensor was re-calibrated prior to each measurement,
147 using saturated bottom water (100% [O₂]) and the deeper sediment horizons (0% [O₂]) as calibration
148 points. Calibrations of the pH and H₂S electrodes were performed as described in Hermans et al.
149 (2019b). pH values are reported on the total scale. For depth profiling of EP (500- μ m resolution; 3
150 replicates per core), micro electrodes were used that were custom built at Aarhus University, as
151 described in Damgaard et al. (2014). A robust reference electrode (Ref-RM, Unisense, A.S.,
152 Denmark) was used during EP and pH measurements. To exclude turbulence-induced variations in the
153 potential of the reference electrode during EP profiling, a silicon tube filled with foam was mounted
154 on the tip of the reference electrode.

155 **2.3. Solute Flux Measurements**

156 Solute flux incubations were performed for NH_4^+ , Fe^{2+} , Mn^{2+} , Ca^{2+} , HPO_4^{2-} and H_4SiO_4 . At
157 each time point, one core was placed outside the aquarium at 20 °C, and the isolated volume of
158 overlying water in the core was continuously aerated. Potential stratification of the overlying water
159 was prevented by actively bubbling it. Parafilm was wrapped on top of the cores to prevent
160 evaporation. Water samples of 3 mL were retrieved at 7 time points over 24 hours. The same volume
161 of fresh artificial seawater was added to the cores directly after taking each sample. The samples were
162 filtered (0.45 μm), and subsamples were taken for ammonium (1 mL) and for metals (1 mL; acidified
163 with 10 μL Suprapur® HCl (35%) per mL sample), which were stored at -20°C and 4°C respectively
164 until further analysis.

165 **2.4. Pore Water and Sediment Collection**

166 At each time point, two cores were sectioned at 0.5-1 cm resolution with an UWITEC push-
167 up pole in a nitrogen-purged glovebag, but only samples for one core were used for sediment and pore
168 water collection and analyses. Bottom water samples were retrieved from the overlying water in the
169 cores. Slices for each depth interval were centrifuged at 3500 rpm for 20 minutes for pore water
170 retrieval. Samples (1 mL) for NH_4^+ were taken and stored at -20°C until analysis. Samples (1 mL) for
171 pore water S, Fe, Mn, Ca, P and Si were also collected and acidified with 10 μL Suprapur® HCl
172 (35%) per mL sample, which were stored at 4°C until analysis. Centrifuged sediment samples were
173 freeze-dried and ground to a fine powder in a nitrogen-purged glovebox under a strictly anoxic
174 environment to prevent oxidation (Kraal et al. 2009; Kraal and Slomp 2014). Only the top 5 cm of the
175 solid-phase samples were analysed in further detail. The porosity (Supporting Information 1.1; Table
176 S1) was calculated from the weight loss upon freeze-drying, using a sediment density of 2.65 g cm^{-3}
177 (Burdige 2006). Salt corrections were performed on the solid-phase data using the gravimetric water
178 content and salinity to determine the amount of salt after freeze-drying. After freeze-drying, the salt
179 from seawater stays behind in the solid-phase fraction. To determine the actual weight of the dry
180 sediment, it is necessary to subtract the weight of the salt from the total weight of freeze-dried
181 sediment.

182 **2.5. Chemical Analysis of the Water and Sediment**

183 Concentrations of NH_4^+ in the pore water and solute flux samples were determined using the
184 phenol hypochlorite method (Koroleff 1969). The total Fe, Mn, Ca, P and Si concentrations (which
185 are assumed to represent Fe^{2+} , Mn^{2+} , Ca^{2+} , HPO_4^{2-} and H_4SiO_4) in the pore water and solute flux
186 samples were determined using Inductively Coupled Plasma-Optical Emission Spectroscopy (ICP-
187 OES, Spectro Arcos). Dissolved Fe and Mn are assumed to be present in the form of Fe^{2+} and Mn^{2+} ,
188 however some Mn^{3+} (Madison et al. 2013) or colloidal and nanoparticulate Fe and Mn might also be
189 available (Boyd and Ellwood 2010; Raiswell and Canfield 2012). Concentrations of P and S are
190 assumed to represent HPO_4^{2-} and SO_4^{2-} respectively. The colourimetric detection limit for NH_4^+ was
191 $0.5 \mu\text{M}$. The practical detection limit on the ICP-OES for Fe, Mn and P was 0.73, 0.11 and $7.30 \mu\text{M}$,
192 respectively.

193 Solid-phase Fe was fractionated into [1] labile ferric Fe (hydr)oxides and ferrous Fe ($\text{FeS} +$
194 FeCO_3), [2] crystalline Fe minerals, [3] magnetite and [4] pyrite (Supporting Information 1.2; Table
195 S2), using a combination of two operational extraction methods (Poulton and Canfield 2005; Claff et
196 al. 2010) as described by Kraal et al. (2017). Concentrations of Fe in all extracts were determined
197 using the colourimetric phenanthroline method (APHA 2005). Solid-phase S was separated into [1]
198 acid volatile sulphur (AVS; representing FeS) and [2] chromium reducible sulphur (CRS; representing
199 FeS_2 ; Table S2) using the method after Burton et al. (2006; 2008) as modified by Kraal et al. (2013).
200 Sulphide released during the S extraction was trapped as ZnS in alkaline Zn acetate traps.
201 Concentrations of S were determined by iodometric titration (APHA 2005). Solid-phase siderite
202 (FeCO_3) was determined by subtracting AVS from the labile ferrous concentrations retrieved from the
203 first step of the Fe extraction. Solid-phase P was fractionated into [1] exchangeable P, [2] citrate-
204 dithionite-bicarbonate (CDB)-P, [3] authigenic P, [4] detrital P and [5] organic P (Table S2) after
205 Ruttenberg (1992) as modified by Slomp et al. (1996). The sum of exchangeable P and CDB-P
206 represents metal bound P, as described in Hermans et al. (2019b). Concentrations of P in all extracts,
207 except CDB, were measured with the molybdenum blue colourimetric method (Murphy and Riley
208 1958). The P, Mn (assuming to represent Mn oxides; Hermans et al. 2019b) and Si (assuming to

209 represent metal oxide bound Si; Kostka and Luther III 1994; Rao et al. 2016) in CDB extracts was
210 determined using ICP-OES.

211 **2.6. Elemental Mapping of Fe, Mn, P and Ca**

212 On day 47, an undisturbed core (first 5 cm of surface sediment) was sampled for epoxy resin
213 embedding for high-resolution elemental mapping (Jilbert et al. 2008; Jilbert and Slomp 2013).
214 Sediment was carefully pushed upwards from the experimental core into a shorter (7 cm length; 1 cm
215 diameter) mini core. This mini sub-core was then transferred to an acetone bath in a argon-filled
216 glovebox and subsequently embedded with Spurr's epoxy resin as described in Jilbert et al. (2008).
217 After curing, the epoxy-embedded core was split vertically using a rock saw. The surface was
218 smoothed by applying a 0.3 μm alumina powder layer. Elemental maps of Fe, Mn, P and Ca (30 μm
219 resolution) were retrieved using a Desktop EDAX Orbis μXRF analyser (Rh tube set at 30 kV,
220 500 μA , 300 ms dwell-time, equipped with a poly-capillary lens). Similar μXRF maps for Fe, Mn and
221 P in epoxy embedded surface sediment were obtained for two field sites: (1) the Gulf of Finland, for
222 sediments collected in June 2016 as described by Hermans et al. (Submitted), and Lake Grevelingen,
223 for sediments collected in January and May 2012 as described in Sulu-Gambari et al. (2016a; 2018).

224 **2.7. Fluorescence *In-situ* Hybridisation**

225 Fluorescence *in-situ* hybridisation (FISH; Pernthaler et al. 2001) was used to microscopically
226 quantify the abundance of cable bacteria filaments, as described in Seitaj et al. (2015). FISH analysis
227 was performed on one intact sediment core retrieved at our sampling site, and the sediment cores from
228 our incubation experiment used for pore water collection at three time points (days 5, 26 and 207).
229 These cores were sectioned at 0.5 cm depth resolution for the first 2.5 cm. Each sediment slice was
230 homogenised and fixed with 0.5 mL ethanol ($\geq 99.8\%$ purity), and stored in a 2 mL Eppendorf tube at
231 $-20\text{ }^{\circ}\text{C}$. For FISH analysis, a volume of 100 μL was retrieved from the Eppendorf tubes and mixed
232 with a 1:1 solution of PBS/ethanol (500 μL). Then 10 μL of this mixture was filtered through a
233 polycarbonate membrane (type GTTP; pore size 0.2 μm , Millipore, USA). Cable bacteria were
234 classified with a *Desulfobulbaceae*-specific oligonucleotide probe (DSB706; 5-ACC CGT ATT CCT
235 CCC GAT-3') after counter staining with DAPI (1 $\mu\text{g}/\text{mL}$) under an epifluorescence microscope

236 (Zeiss Axioplan, Germany) at 100x magnification. The abundance of cable bacteria was quantified by
237 determining the length and diameter of all observed filaments in a field ($105 \times 141 \mu\text{m}$) on the filter at
238 100x magnification (200 fields per sample). Cable bacterial abundances are expressed as filament
239 length per volumetric unit (m cm^{-3}) or depth integrated per unit area of sediment surface (m cm^{-2}),
240 consistent with previous studies (Schauer et al. 2014; Malkin et al. 2017).

241 **2.8. Scanning Electron Microscopy**

242 Cable bacteria filaments were taken from surface sediments from the oxic zone (upper 2 mm)
243 after 40 days using a microscope and were transferred to a 15 mL centrifuge tube. The tube was filled
244 to a volume of ~10 mL using ultra clean water, and was subsequently centrifuged at 2100 rpm for 2
245 min, after which the water was discarded. This washing step was repeated three times. The washed
246 samples were then transferred to a sample stub, where the sediment was air-dried over-night prior to
247 gold coating. The filaments were subsequently subjected to scanning electron microscopy (SEM)
248 imaging on a Phenom ProX Desktop SEM (Phenom-World B.V., the Netherlands) to obtain high-
249 resolution images, as described in Geerlings et al. (2019). SEM images were generated under 0.1-0.3
250 mbar vacuum, and a high accelerating voltage (10 or 15 kV).

251 **2.9. Data Analysis and Calculations**

252 The diffusive uptake of O_2 was calculated from the high-resolution O_2 depth profiles using the
253 PROFILE software package (Berg et al. 1998). Total H_2S ($\Sigma\text{H}_2\text{S} = \text{H}_2\text{S} + \text{HS}^- + \text{S}^{2-}$) was calculated as
254 a function of the recorded H_2S and pH values, accounting for temperature and salinity (Millero et al.
255 1988; Jeroschewski et al. 1996).

256 The EP depth profiles were normalised by subtracting the background EP signal in the
257 overlying water from the EP depth profiles, to calculate the EP value relative to that in the overlying
258 water (Damgaard et al. 2014). The electric field in the sediment was calculated from the linear slope
259 of the EP depth profiles (average of triplicates) in the surface sediments (Risgaard-Petersen et al.
260 2014). The magnitude of the current density was subsequently calculated from the gradient in the EP,
261 the so-called electric field, using Ohm's law:

$$J = \sigma_{sed} \cdot E \quad (1)$$

262 where J represents the magnitude of the current density (mA m^{-2}), σ_{pw} is the conductivity of
 263 the sediment matrix (S m^{-1}) and E (mV m^{-1}) represents the electric field. The conductivity of the pore
 264 water was corrected for tortuosity and calculated as a function of the temperature and salinity using
 265 the equations provided by Fofonoff and Millard Jr (1983).

266 The solute fluxes were calculated as described in Glud (2008) and Rao et al. (2016):

$$J = \frac{\Delta C_{ow}}{\Delta t} \cdot \frac{V_{ow}}{A} \quad (2)$$

267 where J represents the diffusive flux ($\text{mmol m}^{-2} \text{d}^{-1}$), ΔC_{ow} represents the concentration change in the
 268 overlying water (mmol m^{-3}), Δt is the incubation time (d), V_{ow} is the volume in the overlying water
 269 (m^3) and A the surface area of sediment in the core (m^2). In our experimental setup, only those fluxes
 270 were measurable for NH_4^+ , Fe^{2+} , Mn^{2+} and HPO_4^{2-} , that were >0.08 , >0.06 , >0.01 and >0.55 mmol m^{-2}
 271 d^{-1} , respectively. However, for these four solutes, fluxes were always too low to be detected. Hence,
 272 only Ca^{2+} and H_4SiO_4 fluxes are presented.

273 Diffusive downward fluxes of SO_4^{2-} and diffusive upward fluxes of NH_4^+ , Fe^{2+} , Mn^{2+} and
 274 Ca^{2+} were calculated from linearized pore water gradients using Fick's first law (Berner 1980):

$$J = -\phi D_s \cdot \frac{dC}{dz} \quad (3)$$

275

276 The molecular diffusion coefficient was calculated as a function of pressure, salinity and
 277 temperature using the R package *marelac* (Soetaert et al. 2010) and corrected for the ambient
 278 tortuosity using the relations listed in Boudreau (1997).

279 3. RESULTS

280 3.1. Abundance of Cable Bacteria

281 Examination of the top 2.5 cm of the surface sediments using FISH showed the presence of
 282 filamentous cable bacteria (Fig. 1C; Fig. S1). The *in-situ* cable bacterial abundance in the sediment at

283 our field site was low (14 m cm^{-2}). However, after 5 days of incubation in the laboratory, the
284 abundance increased strongly (724 m cm^{-2}). At day 26 the abundance of cable bacteria was even
285 higher (1035 m cm^{-2}). After 207 days, the cable bacterial abundance in the surface sediment was low
286 again (131 m cm^{-2}). SEM imaging confirmed that filaments were indeed cable bacteria (Fig. 1D), as
287 the external surface of the filament was characterised by a parallel pattern of ridges and grooves along
288 its latitudinal axis, which is a typical feature of cable bacteria (Cornelissen et al. 2018; Geerlings et al.
289 2019).

290 **3.2. High-resolution Depth Profiles of pH, O_2 , $\Sigma\text{H}_2\text{S}$ and EP**

291 High-resolution depth profiles of pH showed the development of a distinct peak near the
292 sediment-water interface at day 5, and acidification of the pore water in the deeper sediment (Fig.
293 2A). The width of this pore water acidification zone increased with time and reached its maximum at
294 day 26, followed by a decrease in the acidification. The distinct pH peak near the sediment-water
295 interface gradually disappeared after 33 days. The depth of O_2 penetration in the sediment remained
296 constant within the first 40 days of incubation ($\sim 1.1 \text{ mm}$) and subsequently moved downwards with
297 time to 9.6 mm (Fig. 2A; Fig. 3; Fig. S2). The dissolved $\Sigma\text{H}_2\text{S}$ concentrations remained low ($<5 \mu\text{M}$)
298 throughout the experiment (Fig. 2A). The $\Sigma\text{H}_2\text{S}$ appearance depth was initially equivalent to the O_2
299 penetration depth, and shifted downwards within 5 days, creating a suboxic zone where O_2 and $\Sigma\text{H}_2\text{S}$
300 remained below detection (Fig. 2A; Fig. 3). The width of the suboxic zone remained relatively
301 constant with time ($\sim 25 \text{ mm}$; Fig. 3), and only slight decreased after 207 days.

302 The EP depth profiles indicate a rapid establishment of an electric current after 5 days (0.4
303 mV ; Fig. 2B). The time-series of depth profiles show that the EP increased and also accumulated over
304 a thicker depth horizon. At day 26 the EP reached its maximum value (1.2 mV), followed by a
305 decrease with time. Long-distance electron transport was not active in the anoxic control core (Fig.
306 S3).

307 **3.3. Diffusive Uptake of O₂ and Current Density**

308 The diffusive O₂ uptake of the sediment was highest after 5 days and gradually decreased
309 with time from ~30 to ~3.6 mmol m⁻² d⁻¹ (Fig. 4A). The current density rapidly increased from day 0
310 to day 18, from 6 to 128 e⁻ mmol m⁻² d⁻¹, and then gradually decreased with time (Fig. 4B). The
311 duplicate measurements show the same trend for the diffusive O₂ uptake and the current density,
312 which indicates that the results are reproducible.

313 **3.4. Pore Water Profiles**

314 Concentrations of NH₄⁺ were low near the sediment-water interface and increased with
315 sediment depth reaching maximum levels of up to 1.7 mM at depth in the sediment (Fig. 5). The time-
316 series suggest a gradual decrease in production of dissolved NH₄⁺ in the sediment leading to
317 decreasing concentrations with time. The pore water depth profiles of dissolved SO₄²⁻ show a decline
318 with sediment depth at all time points. However, SO₄²⁻ concentrations remained relatively constant
319 within the top 2 cm of surface sediment between day 12 and 33. Dissolved Fe²⁺, Mn²⁺ and Ca²⁺ all
320 show the development of distinct peaks in the pore water with time, and after 40 days those peaks
321 disappear again. Pore water concentrations of HPO₄²⁻ generally increased with sediment depth for all
322 time points, and concentrations within the top 2 cm were below the detection limit indicating removal.
323 Dissolved H₄SiO₄ increased with sediment depth reaching concentration of up to 1 mM.

324 **3.5. Diffusive Fluxes**

325 Calculated diffusive fluxes of NH₄⁺ into the oxic zone decreased during the incubation
326 experiment from 4.7 to 1.8 mmol m⁻² d⁻¹ (Fig. 6A; Fig. S4; Table S3). Rates of SO₄²⁻ reduction
327 estimated from the linear gradient of the decrease in pore water SO₄²⁻ in the surface sediment with
328 depth generally also showed a decrease with time (Fig. 6B; Fig. S5; Table S3). The upward diffusive
329 flux of Fe²⁺ greatly increased from day 5 to day 12 and then gradually decreased with time (Fig. 6C;
330 Fig. S6; Table S3). The upward diffusive flux of Mn showed an increase in the initial stage of the
331 experiment and reached its maximum at day 18, followed by a decrease with time (Fig. 6D; Fig. S7;
332 Table S3). The upward diffusive flux of Ca²⁺ showed no clear trend with time, however after 207 days
333 the flux became extremely low (Fig. 6E; Fig. S8). The upward diffusive flux of H₄SiO₄ also showed

334 an increase in the initial stage of the experiment, and reached its maximum at day 12, followed by a
335 decrease with time (Fig. 6F; Fig. S9).

336 **3.6. Solid-phase Profiles**

337 The surface sediment in the oxic cores became more enriched in Fe oxides with time, with
338 concentrations increasing from 53 to 485 $\mu\text{mol g}^{-1}$ (Fig. 7). The deeper sediment in the oxic cores and
339 the entire anoxic control core had low or no Fe oxides. At day 5, FeS was strongly depleted within the
340 top 1 cm of the surface sediment and was gradually lost further with time. At day 621, most of the
341 FeS within the top 2.5 cm of the surface sediment had been dissolved. The anoxic core did not show
342 such a depletion of FeS in the surface sediment and even showed a slight increase in FeS. Solid-phase
343 siderite remained rather constant with depth from day 5 to 33, but afterwards was gradually lost from
344 the surface sediment. At day 621 a large proportion of the siderite was dissolved within the top 2 cm.
345 Solid-phase siderite concentrations remained constant with depth in the anoxic control core. Solid-
346 phase depth profiles of Mn oxides, metal bound P and metal oxide bound Si all showed a gradual
347 increase in the surface sediment with time.

348 **3.7. High-resolution Elemental Mapping**

349 High-resolution desktop μXRF mapping of Fe, Mn, P and Ca of our core after 47 days of
350 incubations revealed a subsurface sediment layer highly enriched in Fe and P (Fig. 8A). Subsurface
351 enrichments in Fe, P and Mn in relatively thin layers were also observed in sediments populated by
352 cable bacteria in the Gulf of Finland and Lake Grevelingen (Fig. 8B and C). In the latter system, the
353 layers enriched in Fe, P, Mn broadened upon recolonization by macrofauna (Fig. 8D).

354 **3. DISCUSSION**

355 **4.1. Metabolic Activity of Cable Bacteria**

356 Cable bacteria in our incubation experiment demonstrated a rapid growth, since their
357 abundance greatly increased after 5 days, and reached its peak at day 26 (Fig. 1C). Such high
358 abundances are similar to those observed in previous experiments, in which FeS-rich marine
359 sediments from Aarhus Bay and Lake Grevelingen were incubated (Schauer et al. 2014; Burdorf et al.

360 2018). The activity of cable bacteria exerted a strong impact on the pore water depth profiles of pH,
361 O₂, and $\Sigma\text{H}_2\text{S}$, as evident from the development of a pH maximum near the sediment-water interface,
362 the strong pore water acidification in the deeper sediment and the development of a suboxic zone (Fig.
363 2A). These pore water depth profiles resemble the distinct biogeochemical fingerprint typical for
364 active cable bacteria, as observed in previous laboratory incubation experiments (Risgaard-Petersen et
365 al. 2012; Malkin et al. 2014; Schauer et al. 2014; Vasquez-Cardenas et al. 2015; Rao et al. 2016;
366 Burdorf et al. 2018). The widening of the suboxic zone with time (Fig. 3) is a consequence of the
367 downward expansion of the cable bacteria filament network (Schauer et al. 2014; Vasquez-Cardenas
368 et al. 2015).

369 The EP depth profiles demonstrated that long-distance electron transport by cable bacteria
370 was already active 5 days after the start of the experiment, as indicated by the increase of EP at depth
371 to 0.4 mV). With time, the EP signal increased to higher values and also accumulated over a thicker
372 depth horizon (Fig. 2B), indicating that cable bacteria activity both increased and extended to deeper
373 sediment depth, which is also a consequence of the downward expansion of cable bacteria filaments.
374 The EP reached a maximum after 18 days (1.3 mV; Fig. 2B) in concert with the highest current
375 density of $\sim 130 \text{ mmol e}^- \text{ m}^{-2} \text{ d}^{-1}$ (Fig. 4B). This maximum EP value and current density are similar in
376 magnitude to those found in sediment incubations with seawater with a similar salinity (Damgaard et
377 al. 2014). From day 18 onwards the EP and current density flux gradually decreased with time to 13
378 $\text{mmol e}^- \text{ m}^{-2} \text{ d}^{-1}$ after 207 days (Fig. 4B), which implies a decrease in the metabolic activity of cable
379 bacteria. The suboxic zone persisted long after the current density had decreased (Fig. 3).

380 To summarise, the metabolic activity of cable bacteria was likely highest between day 18 and
381 day 26 based on the cable bacterial abundances, the extent of acidification of the pore water and the
382 current density (Fig. 1C; Fig. 2A and Fig. 4B).

383 **4.2. Organic Matter Degradation**

384 Ammonium fluxes are assumed to reflect rates of anaerobic degradation of organic matter (Fig
385 6A), and the observed decline during the experiment coincides with the decrease in activity of cable

386 bacteria based on the EP profiles and current density (Fig. 2B; Fig. 4B). This suggests that the
387 availability of easily degradable organic matter plays a role in sustaining the metabolic activity of
388 cable bacteria, most likely by controlling the rate of SO_4^{2-} reduction (Nielsen and Risgaard-Petersen
389 2015).

390 Rates of SO_4^{2-} reduction estimated from the linear gradient of the decrease in pore water SO_4^{2-} in
391 the surface sediment with depth indeed also showed a decline during the experiment. We note,
392 however, that a direct measurement of SO_4^{2-} reduction rates (Fossing and Jørgensen 1989; Kallmeyer
393 et al. 2004) would provide a better indicator, because SO_4^{2-} estimated from pore water profiles are in
394 general lower than rates estimated from tracer experiments (Hermans et al. 2019a; Sandfeld et al.
395 2020). Another cause for a slight underestimation of our SO_4^{2-} reduction rates, is due to the effect of
396 the electric field imposed by cable bacteria, which is not taken into account in Fick's law. Solutes can
397 also move with respect to the fluid by electrostatic forces (Bockris and Reddy 1998). Given the
398 relatively low strength of the electric field in the cores ($<0.073\text{V m}^{-1}$ at day 18; as estimated from Fig.
399 2B), including the contribution of ionic drift to the sulphate flux would lead to SO_4^{2-} reduction rates
400 that are at most 10-20% higher.

401 The metabolic activity of cable bacteria can lead to the production of SO_4^{2-} in the suboxic zone
402 via anodic sulphide oxidation (Risgaard-Petersen et al. 2012; Rao et al. 2016). We suspect that this
403 also explains the lack of change in pore water SO_4^{2-} with depth in the upper 2 cm of the sediment in
404 our experiment between 12 and 40 days (Fig. 5). Despite relatively high SO_4^{2-} reduction rates ranging
405 from 5.4 to 17.6 $\text{mmol m}^{-2} \text{d}^{-1}$ (Fig. 6B; Table S3), pore water concentrations of $\Sigma\text{H}_2\text{S}$ remained very
406 low throughout the experiment (Fig. 2A). This is likely due to the direct consumption of $\Sigma\text{H}_2\text{S}$
407 through the activity of cable bacteria, preventing $\Sigma\text{H}_2\text{S}$ from accumulating in the pore water, or
408 alternatively, precipitation of FeS by dissolved Fe^{2+} released from the dissolution of siderite.

409 Laboratory experiments have shown that S-oxidation by cable bacteria can play a dominant role
410 in the O_2 uptake of coastal sediments (Nielsen et al. 2010; Schauer et al. 2014; Nielsen and Risgaard-
411 Petersen 2015), and model analysis predicts up to 93% of the total O_2 uptake (Meysman et al. 2015).

412 When we plot diffusive uptake of O₂ against the current density (i.e. upward flux of electrons towards
413 the oxic zone), a linear relationship – with some scatter - emerges for days 12 to 621 (Fig. 9).
414 However, the data points for day 0 and 5 during the initial stages of our experiment do not follow this
415 linear relationship. We explain these findings as follows: At day 0, the cable bacteria were not active
416 yet and other processes, such as aerobic respiration and oxidation of NH₄⁺ and other solutes (Table 2)
417 and solids (FeS) dominated the consumption of O₂. At day 5 and 12, the activity of cable bacteria and
418 the oxidation of reduced products from anaerobic degradation of organic matter both contributed to
419 consumption of O₂. From day 12 onwards, both the O₂ consumption and electron flux follow a
420 downward decrease with time (Fig. 9). If cable bacteria would account for all of the O₂ consumption,
421 a ratio between the diffusive uptake of O₂ and the current density of 1:4 is expected (Fig. 1A; Nielsen
422 et al. 2010). We find that from day 12 onwards, most data points plot rather close to the line for this
423 1:4 relationship (Fig. 9), suggesting that cathodic O₂ reduction by cable bacteria is responsible for
424 nearly all O₂ consumption in the sediment (in line with the model results of Meysman et al. 2015).
425 This however poses a problem for the nitrogen budget, because our data indicate complete removal of
426 the NH₄⁺ that diffuses upward into the oxic zone (Fig. 6A), and based on the solute fluxes, no escape
427 to the overlying water (see section 2.4). This implies substantial O₂ consumption due to nitrification
428 (Table 2). These findings can be explained, however, if we assume that at least part of the NO₃⁻ that is
429 being formed near the sediment-water interface is also used for the metabolic activity of cable
430 bacteria. It has been shown that cable bacteria can couple the oxidation of \sum H₂S to NO₃⁻ in the
431 absence of O₂ (Marzocchi et al. 2014). Our data suggest that this process may also occur in sediments
432 where O₂ is present in concert with NO₃⁻ near the sediment-water interface. However, we cannot
433 exclude release of NO₃⁻ to the water column or denitrification by other bacteria in the sediment.
434 Another explanation is that cable bacteria might consume O₂ directly above the sediment-water
435 interface, as recently has been proposed by Burdorf et al. (2018). Lastly, the current density might be
436 slightly overestimated, since it ignores other sources that can create an electric potential, such as the
437 diffusion potential (Revil et al. 2012; Nielsen and Risgaard-Petersen 2015).

438 **4.3. Impact of Cable Bacteria on Fe, Mn and S Cycling**

439 The activity of cable bacteria had a strong impact on the biogeochemistry of the surface sediment
440 in our experiment (Fig. 7). Cable bacteria activity induced an intense acidification of the pore water in
441 the suboxic zone (Fig. 2A), which led to the dissolution of Fe and Mn minerals in deeper sediment
442 layers, as can be inferred from the sharp maxima in dissolved Fe^{2+} and Mn^{2+} in the pore water
443 reaching concentrations of up to ~ 1700 and ~ 80 μM , respectively (Fig. 5). The twenty-fold higher
444 dissolved Fe^{2+} concentrations with respect to pore water Mn^{2+} can be attributed to the relatively higher
445 availability of FeS and siderite compared to the availability of Mn carbonates in the sediment that was
446 used for incubation (Lenstra et al. 2020). The peaks in dissolved Fe^{2+} and Mn^{2+} in the pore water
447 broadened over time spanning a depth of $>5\text{cm}$ (Fig. 5; Fig. S6; Fig. S7).

448 The upward diffusive flux of dissolved Fe^{2+} and Mn^{2+} was highest after 12 days, reaching values
449 of up to 3.16 and 0.16 $\text{mmol m}^{-2} \text{d}^{-1}$ respectively. Fluxes subsequently gradually decreased with time
450 (Fig. 6C and D). The continuous upward diffusion of dissolved Fe^{2+} and Mn^{2+} led to enrichments of
451 poorly crystalline Fe and Mn oxides in the surface sediment (Fig. 7). Despite high upward fluxes of
452 dissolved Fe^{2+} and Mn^{2+} towards the sediment-water interface, our solute flux incubations indicate
453 there was little escape of Fe^{2+} and Mn^{2+} to the overlying water (see section 2.4). This implies that all
454 Fe^{2+} and Mn^{2+} that diffused upward was precipitated as Fe and Mn oxides upon contact with O_2 or
455 NO_3^- (Buresh and Moraghan 1976; Kuz'minskii et al. 1994; Straub et al. 1996). Little or no escape of
456 dissolved Fe^{2+} from the sediment into the overlying water, was suggested previously for a field site
457 with active cable bacteria based on diffusive flux calculations (Lake Grevelingen; Sulu-Gambari et al.
458 2016a) and was determined in flux incubations of cores during a laboratory experiment with cable
459 bacteria (Rao et al. 2016).

460 At the start of the experiment, the sedimentary FeS content was (~ 25 $\mu\text{mol g}^{-1}$), which is not
461 unusual for coastal sediments on the north-western Black Sea margin (Wijsman et al. 2001), but is
462 low compared to sediments in eutrophic coastal systems (e.g. Morgan et al. 2012; Kraal et al. 2013;
463 Hermans et al. 2019a). The solid-phase depth profiles reveal a gradual removal of the FeS in the
464 surface sediment in our experiment over time (Fig. 7). At the end of our experiment (621 days), there

465 was no longer any FeS within the top 1.5 cm of the sediment. While approximately 90 mmol m⁻² of
466 FeS was removed from the surface sediment within the first 5 days, a total of ~240 mmol m⁻² was
467 removed after 621 days (Fig. 10; Table 3). Likely, part of the FeS that was removed from the surface
468 sediment within the first 5 days was removed through oxidation upon contact with O₂, rather than the
469 metabolic activity of cable bacteria itself. The pore water acidification associated with cable bacteria
470 activity led to a strong loss of siderite within the top 2 cm of the sediment, with a total removal of
471 ~560 mmol m⁻² during the experiment (Fig. 7; Fig. 10; Table 3; Table S5). The depletion of
472 sedimentary FeS and siderite was directly proportional to the formation of Fe oxides near the
473 sediment-water interface (Fig. 10), and accounted for 30% and 70% of the Fe oxides, respectively
474 (Table 3).

475 With these data we cannot accurately determine the role of FeS versus SO₄²⁻ reduction in
476 supplying the ΣH₂S sustaining the activity of cable bacteria throughout the experiment. This is
477 primarily related to the variability between cores, and for this type of calculation, the low temporal
478 resolution of sampling. However, we can make an estimation of the relative role of SO₄²⁻ reduction
479 and FeS dissolution in ΣH₂S production, based on the pore water profiles of SO₄²⁻ and dissolved Fe²⁺,
480 and the solid-phase mass balance of FeS and siderite (Fig. 6B and C; Table 4). This estimation shows
481 that SO₄²⁻ was mainly responsible for ΣH₂S production, accounting for 85-99% (Table 4), and thus
482 that the dissolution of FeS only played a minor role in providing ΣH₂S.

483 **4.4. Impact of Cable Bacteria on Ca, P and Si Cycling**

484 Cable bacteria activity is known to lead to dissolution of Ca carbonates, because of the strong
485 acidification of the pore water (Risgaard-Petersen et al. 2012; Rao et al. 2016). We indeed find similar
486 maxima in pore water Ca²⁺ during the experiment (Fig. 5) and a high upward flux of Ca²⁺ (up to ~18
487 mmol m⁻² d⁻¹; Fig. 6E; Fig. S8) of which a substantial fraction (up to ~55%) escapes to the overlying
488 water (Fig. S10; Table S4), which is consistent with a previous incubation experiment Rao et al.
489 (2016).

490 Pore water depth profiles of HPO_4^{2-} reveal a production at depth and removal of all upward
491 diffusing HPO_4^{2-} within the first 1-3 cm of the surface sediment (Fig. 5). A major proportion of this
492 HPO_4^{2-} is bound to Fe oxides (Fig. 7). Given that a large proportion of the Fe oxides in our sediment
493 cores derive from the dissolution of siderite, this suggests that the buffer mechanism that delays the
494 benthic release of HPO_4^{2-} through retention of P associated with newly formed Fe oxides (Sulu-
495 Gambari et al. 2016b), might also be active in systems that are relatively poor in sedimentary FeS.

496 The shape of the pore water HPO_4^{2-} profiles suggests that some of the HPO_4^{2-} is removed below
497 the zone where Fe and Mn oxides are present (Fig. 5; Fig. 7). A possible explanation could be the
498 formation of vivianite, an Fe(II) phosphate mineral. Vivianite formation in sediments typically occurs
499 when pore water levels of Fe^{2+} and HPO_4^{2-} are high and concentrations of $\Sigma\text{H}_2\text{S}$ are low (Nriagu
500 1972), as observed in our study. In our experiment, free $\Sigma\text{H}_2\text{S}$ does not accumulate in the pore water,
501 which we attribute to removal through the activity of cable bacteria and FeS formation at depth (Fig.
502 2A; Fig. 7). Hence, cable bacteria may create a geochemical niche that allows the formation of
503 vivianite in the suboxic zone. Further work with sediments with higher P concentrations would be
504 needed to assess this with direct measurement techniques, such as X-ray spectroscopy (Egger et al.
505 2015; Kraal et al. 2017; Sulu-Gambari et al. 2018). Other sediment P pools, i.e. organic, authigenic
506 and detrital P remained constant over time, indicating that the P contents determined for discrete
507 sediment slices using sequential extractions were not affected by pore water acidification as a result of
508 cable bacteria activity (Table S6).

509 Pore water H_4SiO_4 profiles show a typical increase with depth as observed upon dissolution of
510 biogenic silica in marine sediments (Aller 2014). Fluxes of H_4SiO_4 towards the sediment-water
511 interface range up to $\sim 2.8 \text{ mmol m}^{-2} \text{ d}^{-1}$ and gradually decreased with time (Fig. 6F; Fig. S9). The
512 results of the solute flux incubations indicate that most of this H_4SiO_4 escaped to the overlying water
513 (ranging from 28 to 92%; Table S4; Fig. S10). The decline in the benthic release flux of H_4SiO_4
514 contrasts with results of a previous incubation experiment by Rao et al. (2016) with similar pore water
515 concentrations of H_4SiO_4 reaching values up to $\sim 1 \text{ mM}$. In their study, the flux remained constant
516 over time, possibly because of differences in the amount of biogenic Si in the sediment. The solid-

517 phase metal oxide bound Si pool in the surface sediment increased directly proportional to the
518 formation of Fe oxides throughout the experiment (Fig. 7). Silica is known to absorb to Fe oxides
519 (Sigg and Stumm 1981; Davis et al. 2002). Hence, the results suggest that the Fe oxides formed
520 through the activity of cable bacteria captured some of the upward diffusing H_4SiO_4 .

521 **4.5. Sediment Marker for Cable Bacteria Activity**

522 Visual observations of core photographs reveal the gradual development of an orange layer (oxic
523 zone) up to 9 mm thick, overlying a grey layer (suboxic zone) and a black layer (sulphidic zone)
524 during the experiment (Fig. S11). This colour zonation is typical for sediments that have been
525 geochemically affected by cable bacteria activity, as seen both in laboratory experiments (Nielsen and
526 Risgaard-Petersen 2015) and at coastal field sites (Sulu-Gambari et al. 2016a). High-resolution
527 elemental maps of our sediments reveal the development of a ~0.3 mm thin subsurface layer highly
528 enriched in Fe oxides and associated P, 47 days after the start of the incubation (Fig. 8A). While the
529 Fe oxide layer is clearly enriched in P, we also observed a second layer enriched in P very close to the
530 sediment-water interface (Fig. 8A). This layer is located above the Fe oxide layer, and in this layer P
531 is strongly correlated with Ca. Below, we describe the formation of this layer in more detail and
532 explain why such subsurface enrichments, detected with μ XRF, may act as an additional sediment
533 marker for present or recent cable bacteria activity, also in cases where visual observations are not
534 conclusive.

535 During the experiment, O_2 penetration varied within a narrow range and was initially fixed
536 between 1 and 2 mm depth (Fig. 3A), with the layer highly enriched in Fe forming mostly at a depth
537 of 2 mm (Fig. 8A). Such a range in O_2 penetration is in accordance with observations in coastal
538 sediments (e.g. Seitaj et al. 2015). The formation of the Fe-enriched layer can be explained by rapid
539 oxidation of upward diffusing Fe^{2+} upon contact with O_2 (and possibly NO_3^- ; Fig. 6C). Directly, above
540 the Fe oxide layer a broader ~0.8 mm thick Mn oxide layer was observed (Fig. 8A). This contrast in
541 zonation between Fe and Mn is likely due to the slower oxidation kinetics of Mn^{2+} compared to Fe^{2+}
542 (Burdige 1993; Luther 2010; Learman et al. 2011).

543 While the Fe oxide layer is clearly enriched in P, we also observed a second layer enriched in P
544 close to the sediment-water interface (Fig. 8A). In this layer, P is strongly correlated with Ca. This
545 layer likely consists of carbonate fluorapatite (CFA), a Ca-P mineral, which is typically formed in
546 marine sediments (Van Cappellen and Berner 1988; Ruttenberg and Berner 1993). Possibly, the high
547 pore water pH near the sediment-water interface (resulting from cathodic O₂ reduction by cable
548 bacteria; Fig. 2A), promotes apatite formation (Bellier et al. 2006), and the elevated Ca²⁺
549 concentrations (Fig. 5) created a biogeochemical niche for the formation of CFA.

550 Such focusing of Fe, Mn, P and associated elements within a thin subsurface layer, as a
551 consequence of cable bacteria activity, also occurs in the field. This was demonstrated by Hermans et
552 al. (Submitted) in a study of a coastal site in the Gulf of Finland where cable bacteria were recently
553 active. Here, μ XRF mapping of resin embedded sediments revealed strong focusing of Fe oxides,
554 Mn(II) phosphates and Fe bound P within a 3 mm thick layer near the sediment-water interface (Fig.
555 8B). A re-assessment of the μ XRF data of Sulu-Gambari et al. (2016a; 2018) of surface sediments
556 with active cable bacteria from seasonally hypoxic marine Lake Grevelingen in January also revealed
557 similar subsurface enrichments in Fe, Mn and P (Fig. 8C). Importantly, no visual signals for cable
558 bacteria based on the colour pattern of the sediment were observed at the time.

559 Macrofaunal activity likely counteracts or prevents strong focusing of Fe oxides and associated P
560 within such a thin subsurface layer at field sites. Bioturbation, i.e. mixing of the sediment, typically
561 leads to oxidation from the sediment surface downwards (Norkko et al. 2012). Bioirrigation can
562 efficiently pump O₂ into the pore water and thereby enhance the oxidation of dissolved Fe²⁺
563 (Kristensen et al. 2012; Norkko et al. 2012), but is not expected to lead to such a sharp oxidation front
564 (Norkko et al. 2012; Hermans et al. 2019a). This is also evident from high-resolution elemental maps
565 of the surface sediment from Lake Grevelingen in May, which shows the disappearance of the thin
566 layer highly enriched in Fe and P formed by cable bacteria in January as a consequence of
567 macrofaunal activity in May (Fig. 8D; Seitaj et al. 2015; Sulu-Gambari et al. 2016b).

568 We conclude that the focusing of Fe, Mn and associated P within a thin layer below the sediment-
569 water interface is likely a consistent feature in sediments populated by active cable bacteria and may
570 act as an additional sediment marker for present or recent cable bacteria activity, both in laboratory
571 experiments and at field sites, also in cases where visual observations are not conclusive. Focussing of
572 Fe and Mn oxides in the surface sediment is not exclusively tied to the activity of cable bacteria, and
573 can also occur in the absence of cable bacteria. However, the upward fluxes of Fe^{2+} and Mn^{2+} in
574 sediments populated by cable bacteria, are higher due to active dissolution of Fe and Mn minerals at
575 depth (e.g. Risgaard-Petersen et al. 2012; Rao et al. 2016). Hence, within the same time frame
576 following an environmental perturbation (such as a transition to oxic bottom waters after a period of
577 anoxia or mixing of the sediment), more Fe^{2+} and Mn^{2+} can oxidise upon contact with O_2 near the
578 sediment-water interface and stronger enrichments of Fe and Mn minerals will be observed. Hence,
579 focusing of Fe and Mn oxides in subsurface sediments is likely more prominent and stronger in
580 sediments populated by active cable bacteria compared to sediments where no cable bacteria are
581 active under such conditions. Macrofaunal activity within natural environments likely counteracts or
582 prevents strong focusing of Fe oxides and associated P within such a thin subsurface layer. When
583 using standard techniques for sediment sampling (i.e. core slicing and chemical analysis of these
584 slices), these layers may be missed due to the relatively coarse depth resolution. Hence, μXRF
585 mapping of epoxy embedded sediment is recommended.

586 **4.6. Cable Bacteria Activity at the Field Site**

587 We can only speculate about the possible *in-situ* relevance of cable bacteria at the coastal site
588 in the western Black Sea where the sediment for our incubation was collected. At this site, which is in
589 a region that is subject to seasonal hypoxia (Capet et al. 2013), both bivalves (up to ~ 7200 ind. m^{-2})
590 and polychaetes (up to ~ 1700 ind. m^{-2}) were observed at the time of sampling (Lenstra et al. 2019).
591 Macrofauna can inhibit the activity of cable bacteria through bioturbation by physically cutting and
592 damaging the filaments, rendering them unable to transport electrons (Malkin et al. 2014). Recent
593 work has shown, however, that in some cases, cable bacterial communities can also thrive in
594 sediments with macrofauna (Burdorf et al. 2017; Malkin et al. 2017; Aller et al. 2019). In a study of

595 bivalve reefs, cable bacteria were found to efficiently remove highly toxic $\Sigma\text{H}_2\text{S}$, which is beneficial
596 for bivalves (Malkin et al. 2017). Cable bacteria can also be abundant in bioturbated deposits, when
597 associated with stable subdomains of the bioturbated zone, such as worm tubes (Aller et al. 2019). In
598 such settings, a more complex precipitation pattern, e.g. along tube linings is observed (Aller et al.
599 2019), than described here for laboratory experiments with defaunated sediments and field sediments
600 with an impoverished macrofaunal population (Fig. 8A). Further field studies are required to assess
601 the role of cable bacteria at our field site, preferably including an assessment of the burrow structures.

602 **Conclusions**

603 The results of our laboratory incubation (with a total duration of 621 days) show that cable
604 bacteria can potentially strongly impact the Fe, Mn, P and S dynamics in coastal sediments. The
605 strong acidity of the pore water associated with the activity of cable bacteria, which was monitored
606 using microsensor profiling of the EP during the experiment, led to dissolution of FeS and siderite and
607 formation of Fe and Mn oxides and Ca-P in mineral form near the sediment surface. Our experimental
608 results provide conclusive evidence for siderite dissolution driven by cable bacteria activity as a
609 source of Fe that can form an Fe oxide-enriched surface layer. Both FeS and SO_4^{2-} reduction provided
610 the $\Sigma\text{H}_2\text{S}$ required by cable bacteria to sustain their activity. Pore water $\Sigma\text{H}_2\text{S}$ was always low (<5
611 μM). Using μXRF mapping of epoxy embedded sediment, we show that the activity of cable bacteria
612 led to the development of a thin subsurface sediment layer (0.3 mm) that was highly enriched in Fe
613 and P. The position of this layer in the sediment was directly proportional to the O_2 penetration depth
614 during the experiment. We show that a similar layer highly enriched in Fe and P was also formed in
615 sediments of field locations populated by cable bacteria (i.e. marine Lake Grevelingen and the
616 brackish Gulf of Finland). We suggest that such layers, which are not necessarily visible by eye, may
617 be used as a marker of cable bacteria activity in sediments with low macrofaunal activity.

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630 **Code availability.** All data, if not directly available from the tables and supplementary information,
631 will be made available in the PANGAEA database. In the meantime data are available upon request to
632 the main author.

633 **Author contributions.** MH and CPS designed the experiment. MH carried out the experiment and
634 analysis. All authors interpreted the data. MH and CPS wrote the paper with comments provided by
635 NRP and FJRM.

636 **Competing interests.** The authors declare that they have no conflict of interest.

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886 **TABLES AND FIGURES**

887 **Table 1.** Key site characteristics: latitude, longitude, water depth, bottom water O₂ concentration, *in-situ* O₂ uptake, *in-situ*
 888 O₂ penetration depth in the sediment, porosity and salinity. These data were retrieved from Lenstra et al. (2019). Our study
 889 site is station 9 in Lenstra et al. (2019).

Black Sea (Station 9)		Unit
Latitude	44°34.9'	N
Longitude	29°11.4'	E
Water depth	27	m
Bottom water O ₂	92	μM
O ₂ uptake	25.8 ± 1.77	mmol m ⁻² d ⁻¹
O ₂ penetration depth	2.25	mm
Porosity	0.86	-
Salinity	17.881	-
Avg. organic carbon content (0-0.5 cm)	1.8%	

890
 891 **Table 2.** Mass balance of O₂ consumption. The diffusive uptake of O₂ as calculated from the O₂ depth profiles (column 1)
 892 was compared to the potential O₂ demand from the oxidation of NH₄⁺, Fe²⁺ and Mn²⁺ (column 2-4). The O₂ consumption of
 893 the oxidation of NH₄⁺, Fe²⁺ and Mn²⁺ was determined based on the stoichiometry of NH₄⁺, Fe²⁺ and Mn²⁺ oxidation with O₂
 894 as described in Reed et al. (2011). The oxidation of dissolved Fe²⁺ and Mn²⁺ only played a minor role in the total O₂
 895 consumption during the experiment, contributing only 0.9 to 3.8% and 0.1 to 0.4%, respectively.

	O₂ [mmol m⁻² d⁻¹]	Potential O₂ Demand			e⁻ [mmol m⁻² d⁻¹]
		NH₄⁺ [mmol m⁻² d⁻¹]	Fe²⁺ [mmol m⁻² d⁻¹]	Mn²⁺ [mmol m⁻² d⁻¹]	
Day 5	-23.35	9.42	0.21	0.05	82.68
Day 12	-23.24	8.46	0.89	0.09	111.94
Day 18	-21.10	8.04	0.70	0.08	127.97
Day 26	-23.00	7.58	0.63	0.07	97.55
Day 33	-22.80	5.06	0.62	0.05	84.16
Day 40	-19.60	4.88	0.60	0.06	76.31
Day 207	-6.90	3.52	0.08	0.01	13.10
Day 621	-3.25	N/A	0.03	0.01	9.47

896

897 **Table 3.** Mass balance of Fe. Time-series of the depth integrated (0-5 cm) increase in Fe oxides and the depth integrated (0-5
898 cm) depletion of FeS and FeCO₃ (siderite) in mmol m⁻². All values are reported in mmol Fe m⁻². Negative values represent a
899 decrease, whereas positive values indicate an increase in the mineral pools.

	$\Delta\text{Fe oxides}$ [mmol m ⁻²]	ΔFeS [mmol m ⁻²]	ΔFeCO_3 [mmol m ⁻²]
Day 5	120	-90	-42
Day 12	170	-90	-126
Day 18	189	-105	-92
Day 26	276	-174	-99
Day 33	315	-176	-109
Day 40	412	-223	-200
Day 207	523	-236	-341
Day 621	874	-242	-566

900

901 **Table 4.** Sources of $\Sigma\text{H}_2\text{S}$ calculated from the reduction of SO₄²⁻ and the dissolution of FeS. The numbers are presented
902 either as mmol m⁻² d⁻¹ or as the relative percentage of the $\Sigma\text{H}_2\text{S}$ production. The amount of S from the dissolution of FeS
903 was estimated from the upward diffusive flux of Fe²⁺ (Fig. 6C) and the relative fraction of FeS (FeS/FeS+siderite) based on
904 the mass balance calculations (Table 3).

	S from SO ₄ ²⁻ reduction [mmol m ⁻² d ⁻¹]	S from FeS dissolution [mmol m ⁻² d ⁻¹]	S from SO ₄ ²⁻ reduction [%]	S from FeS dissolution [%]
Day 5	10.49	0.56	95%	5%
Day 12	17.60	1.48	92%	8%
Day 18	8.87	1.50	86%	14%
Day 26	11.15	1.61	87%	13%
Day 33	8.54	1.52	85%	15%
Day 40	7.57	1.25	86%	14%
Day 207	10.38	0.13	99%	1%
Day 621	5.36	0.03	99%	1%

905

906 **Fig. 1. (A)** Geochemical pore water fingerprint typical for cable bacteria activity. This fingerprint is defined by a distinct pH
907 profile (light grey line) and a sub-oxic zone that is devoid of O₂ (red line) and H₂S (blue line). The cable bacteria filaments
908 are depicted in yellow. On the background, the sediment core photograph, taken 278 days after the start of the experiment,
909 shows a distinct colour zonation where (1) the oxic zone displays an orange colour (2) the suboxic zone has a grey colour
910 and (3) the sulphidic zone has a black colour. The scale bar denotes a distance of 6 cm, with 0.5 cm intervals. **(B)**
911 Bathymetric map of the Black Sea. The purple star indicates the location of our study site (44°34.93'N, 29°11.38'E), which
912 was sampled with R/V *Pelagia* in September 2015. Further details are provided in Lenstra et al. (2019). **(C)** Volumetric
913 density of cable bacteria [m cm⁻³] in the top 2.5 cm of the sediment, for *in-situ* as well as for three time points during the
914 incubation experiment **(D)** SEM image of a cable bacteria filament that was extracted from the surface sediment after 40
915 days.

916

917 **Fig. 2. (A)** Time-series of the pore water pH (black), O₂ (red) and ΣH₂S (blue) signatures of the incubated sediment. **(B)**
918 Development of the EP depth profile in the incubated sediment over time. The dashed-line at 0 mm depth represents the
919 sediment-water interface. The blue boxes indicate the overlying water, whereas the underlying light grey boxes represent the
920 sediment. The EP depth profiles represent an average of 3 replicate measurements. The error bars indicate the minimum and
921 maximum EP values that were observed. The orange depth profiles represent duplicate measurement performed on a
922 different core.

923

924 **Fig. 3.** Time-series of the development of the oxic zone (orange), suboxic zone (light grey) and the anoxic/sulphidic zone
925 (dark grey) in the sediment. These zones were calculated from 3 replicate microelectrode depth profiles retrieved from two
926 different cores.

927

928 **Fig. 4.** Time-series of the **(A)** diffusive O₂ uptake in mmol O₂ m⁻² d⁻¹ and **(B)** current density as a consequence of long-
929 distance electron transport (e⁻) in mmol e⁻ m⁻² d⁻¹ in the sediment incubation.

930

931 **Fig. 5.** Time-series of pore water depth profiles of NH₄⁺ (orange), SO₄²⁻ (purple), Fe²⁺ (red), Mn²⁺ (green), Ca²⁺ (grey),
932 HPO₄²⁻ (blue) and H₄SiO₄ (yellow). The control core was sampled at day 621.

933

934 **Fig. 6.** Time-series of diffusive fluxes calculated from the linear gradient of the pore water profiles of **(A)** NH₄⁺, **(B)** SO₄²⁻,
935 **(C)** Fe²⁺, **(D)** Mn²⁺, **(E)** Ca²⁺ and **(F)** H₄SiO₄ in mmol m⁻² d⁻¹ towards the oxic zone of the sediment, based on the linear pore
936 water gradients (Section 1.6; Fig. S4-S9). Here, a positive value indicates an upward flux, whereas a negative value
937 represents a downward flux. N/A = not available. The control core was sampled at day 621.

938

939 **Fig. 7.** Time-series of solid-phase depth profiles of Fe oxides (red), FeS (black), siderite (grey), Mn oxides (green), metal
940 bound P (blue) and metal oxide bound Si (yellow).

941

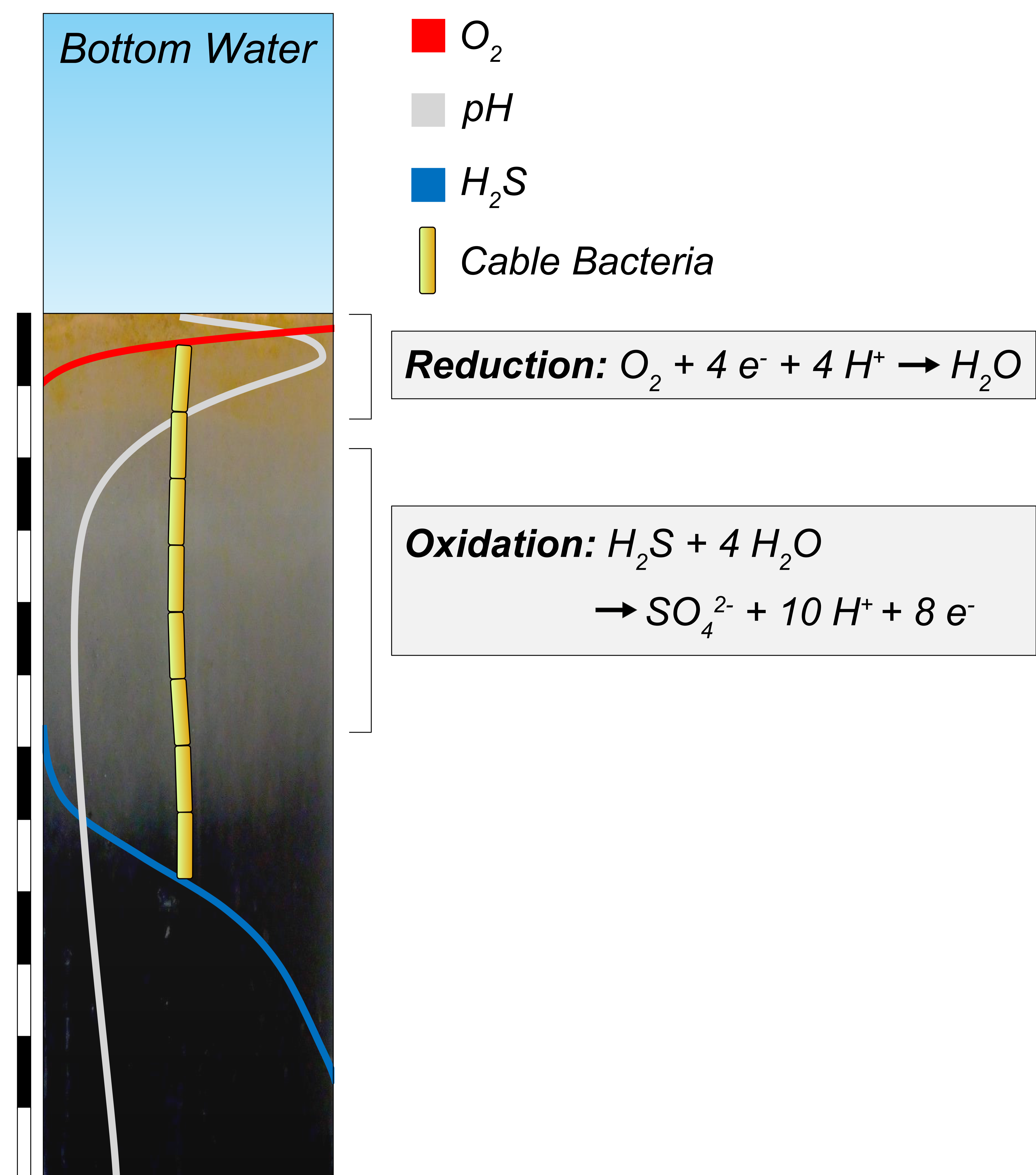
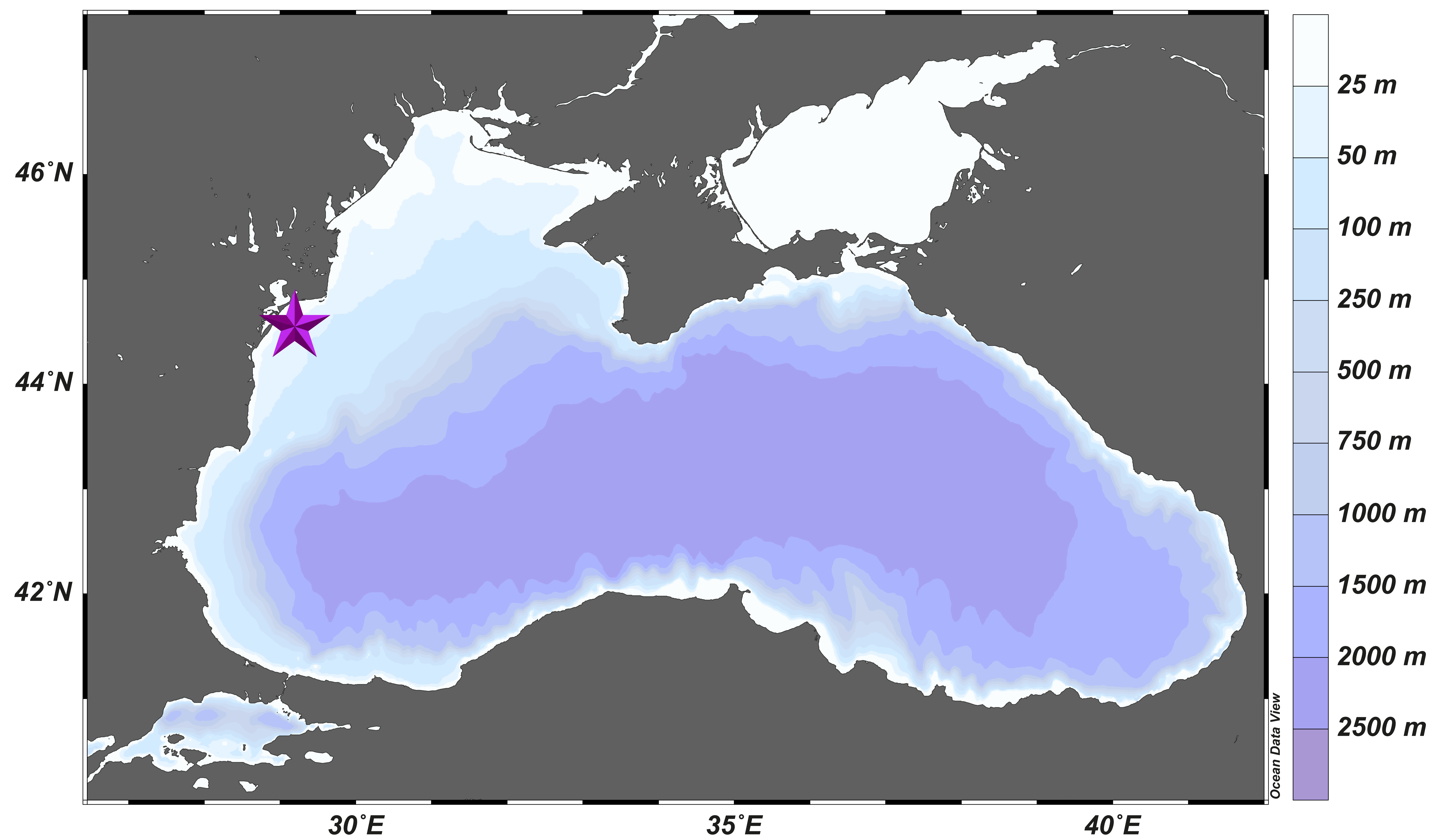
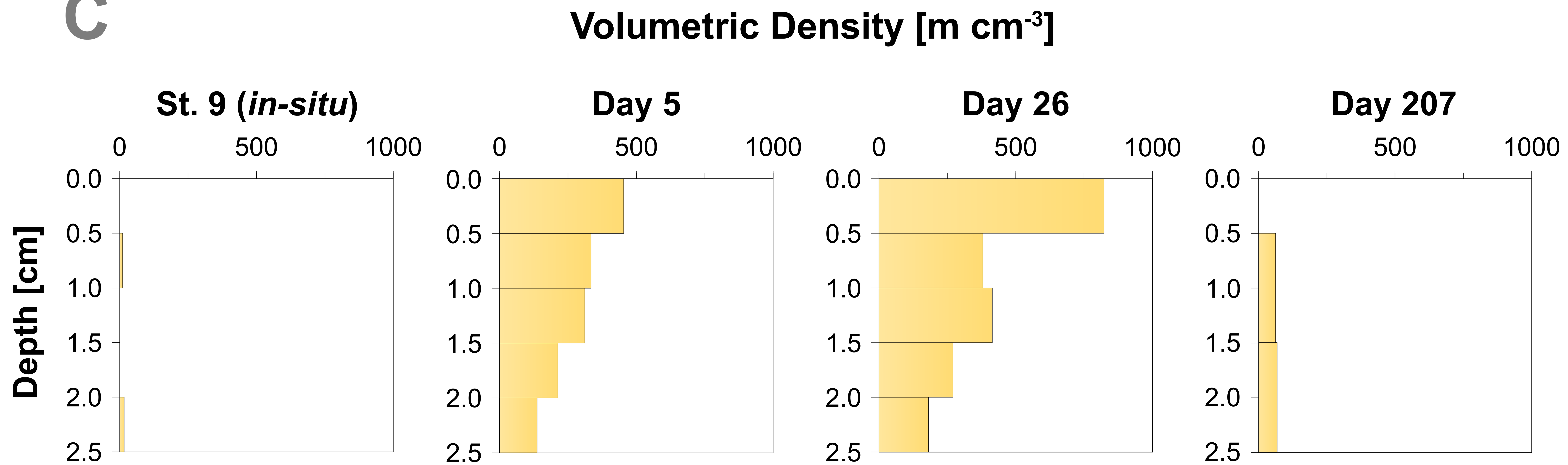
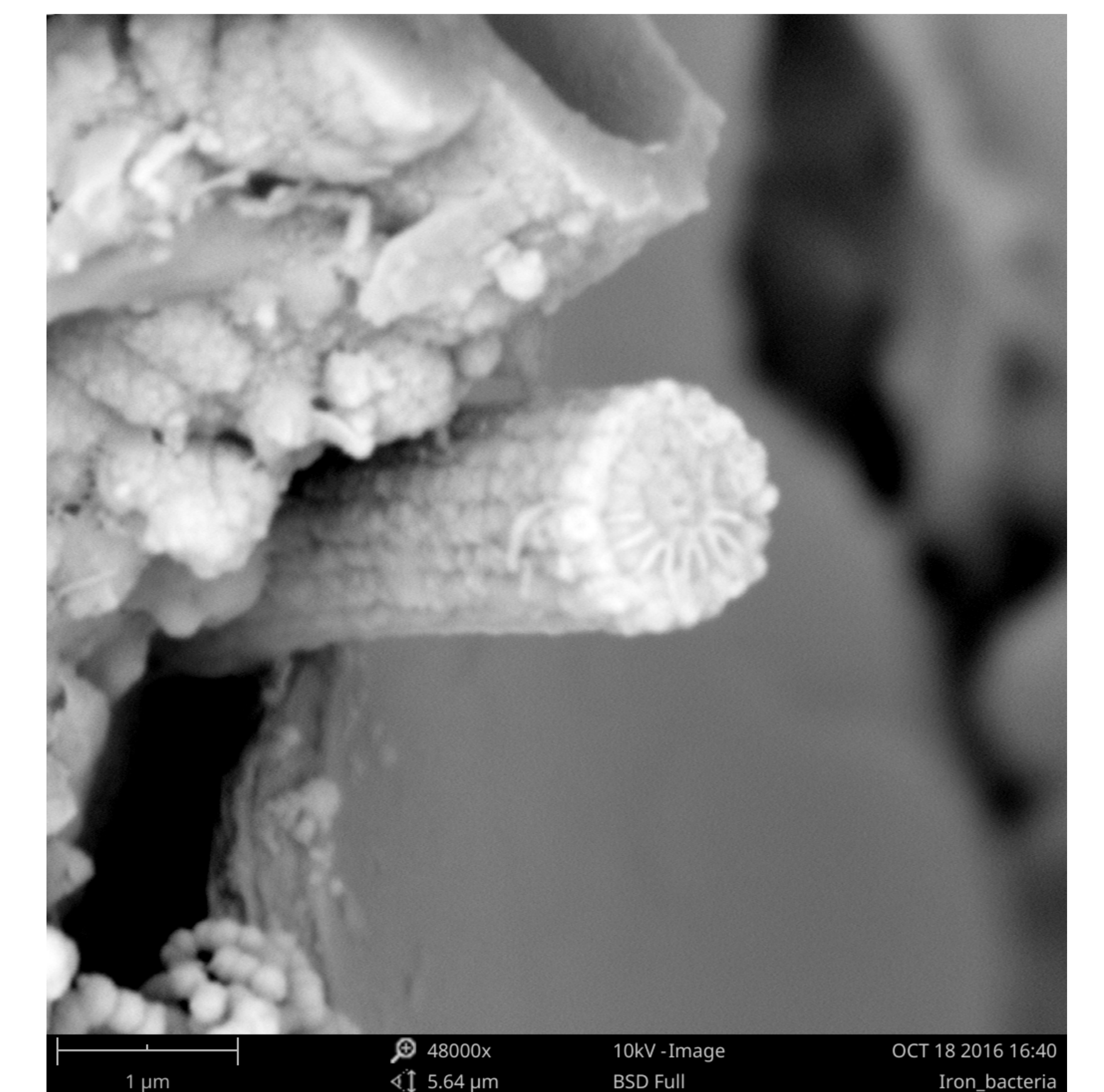
942 **Fig. 8.** High-resolution elemental maps of Fe (red), Mn (green), P (blue) and Ca (white) of surface sediments. These maps
943 are shown in true vertical orientation and the colours accentuate the relative count intensities adjusted for brightness and
944 contrast to highlight the features in the sediment. The tick marks represent 1 mm intervals. μ XRF maps of the surface
945 sediment (A) from the incubation experiment, (B) from the Gulf of Finland at site GOF5 in June (Hermans et al. Submitted),
946 (C) from Lake Grevelingen in January (when cable bacteria become active) and (D) from Lake Grevelingen in May
947 (showing the effects of bioturbation as described in Seitaj et al. (2015)).

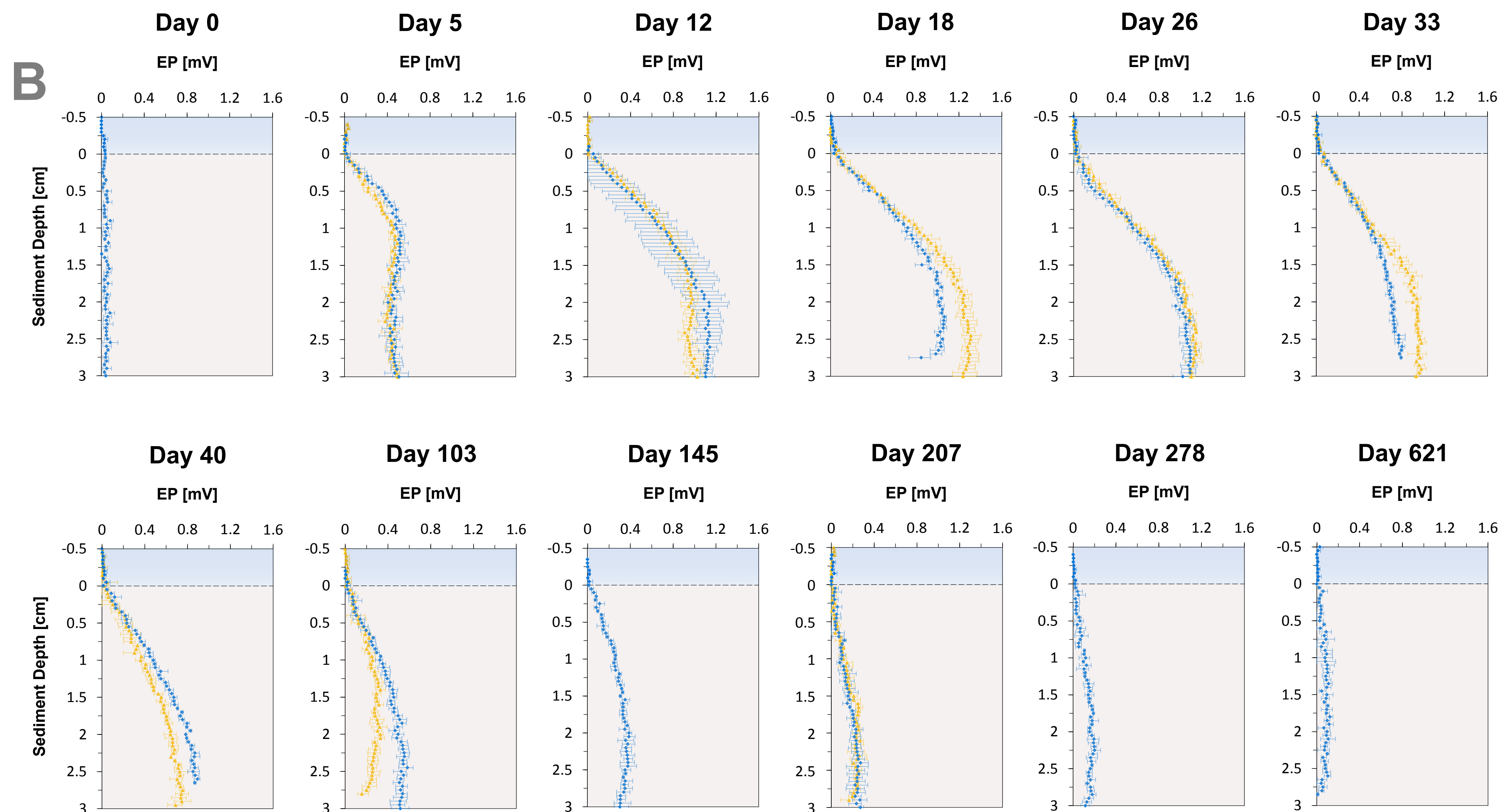
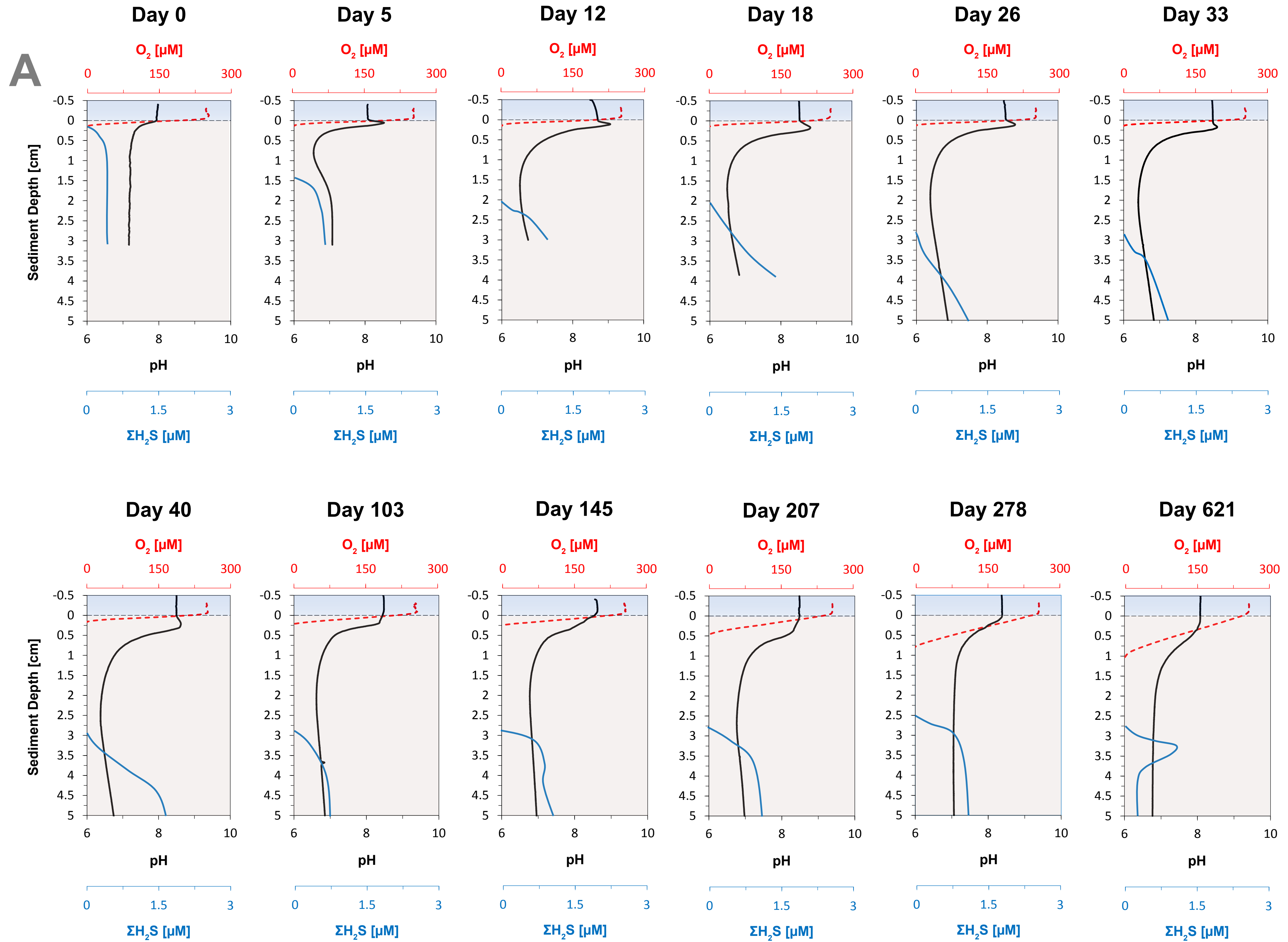
948

949 **Fig. 9.** The relationship between the diffusive uptake of O_2 ($mmol O_2 m^{-2} d^{-1}$) and the current density of long-distance
950 electron transport ($mmol e^{-} m^{-2} d^{-1}$). Red triangles are data for days 0 and 5. Green diamonds are data for all other time
951 points. The blue line represents the expected correlation between the cathodic O_2 consumption rate and the current density
952 assuming a 1:4 ratio (Nielsen et al. 2010). Here, a positive value indicates an upward flux, whereas a negative value
953 represents a downward flux.

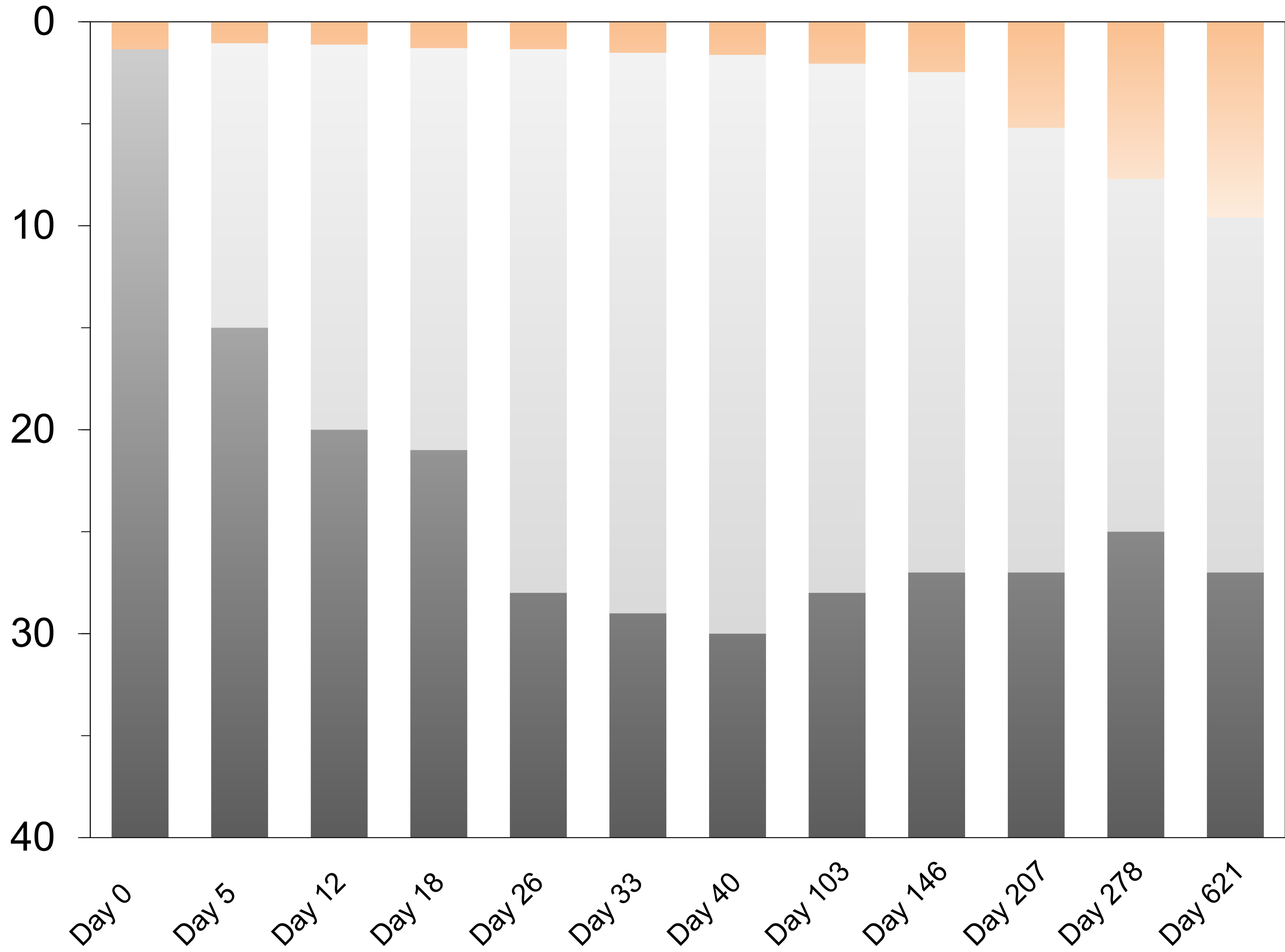
954

955 **Fig. 10.** Time-series of the depth integrated (0-5 cm) increase in Fe oxides (red) and the depletion of FeS (black) and siderite
956 (grey) in $mmol m^{-2}$. Negative values represent a decrease, whereas positive values indicate an increase in the mineral pools.

A**B****C****D**

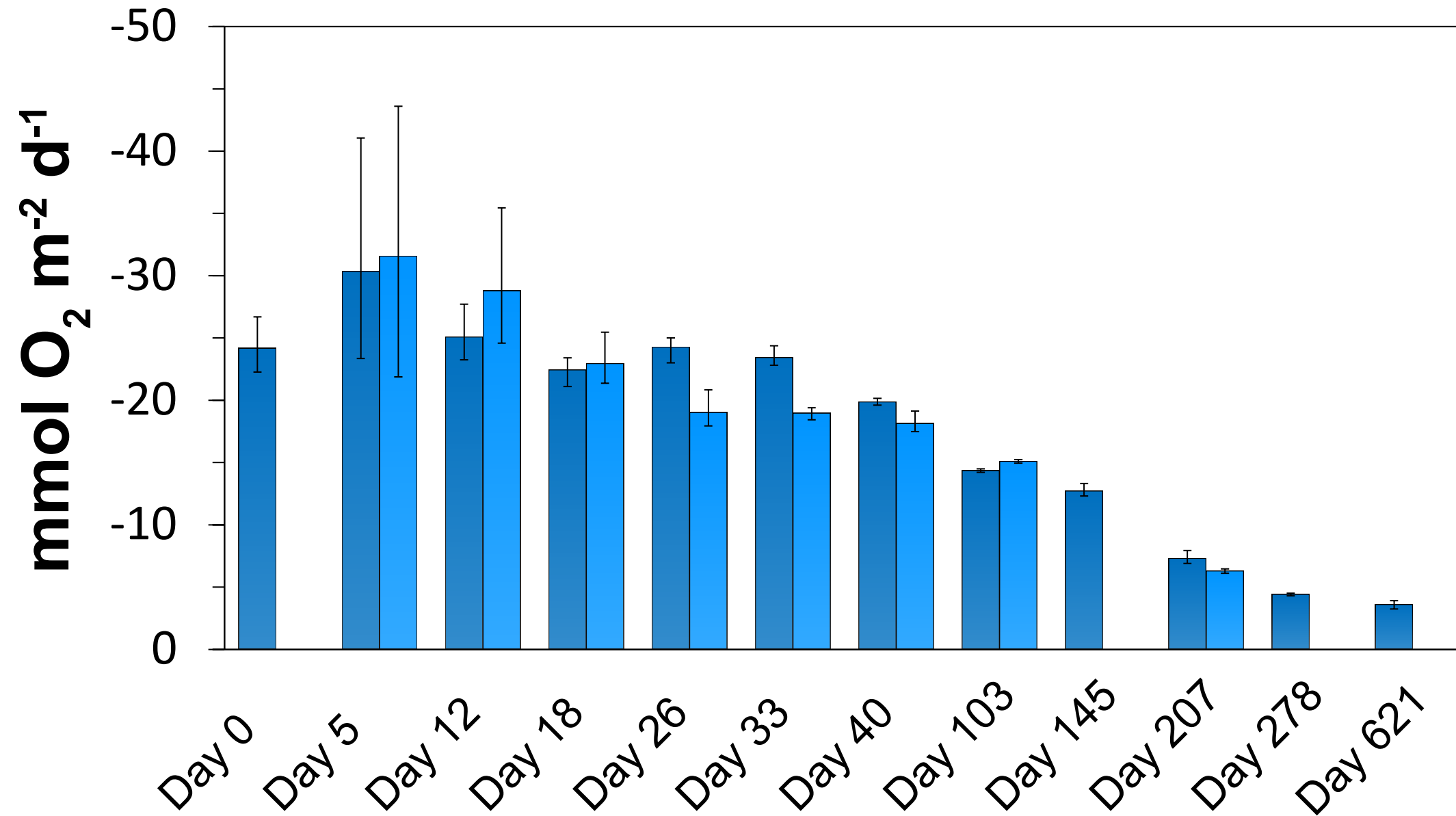


Depth Zonation [mm]

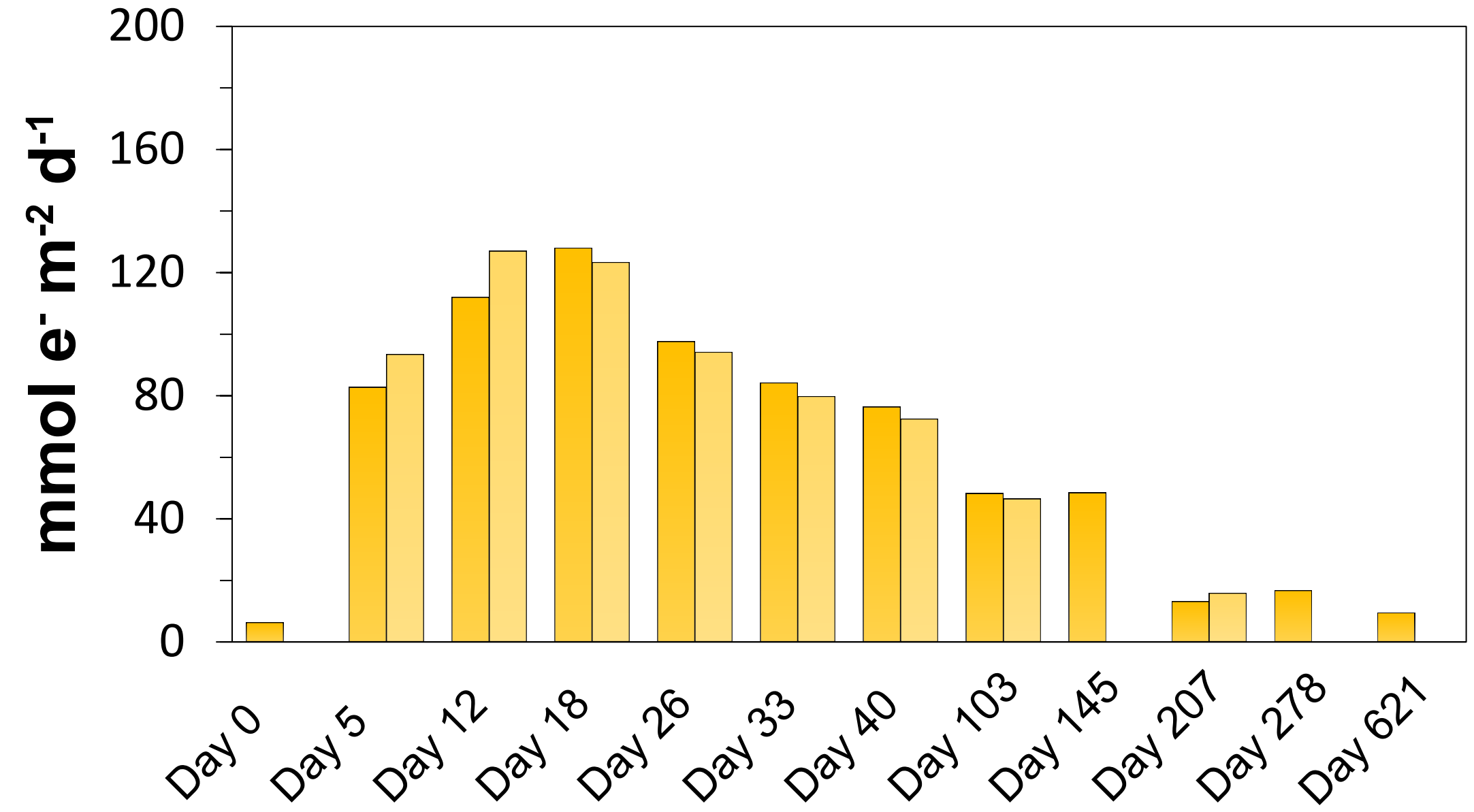


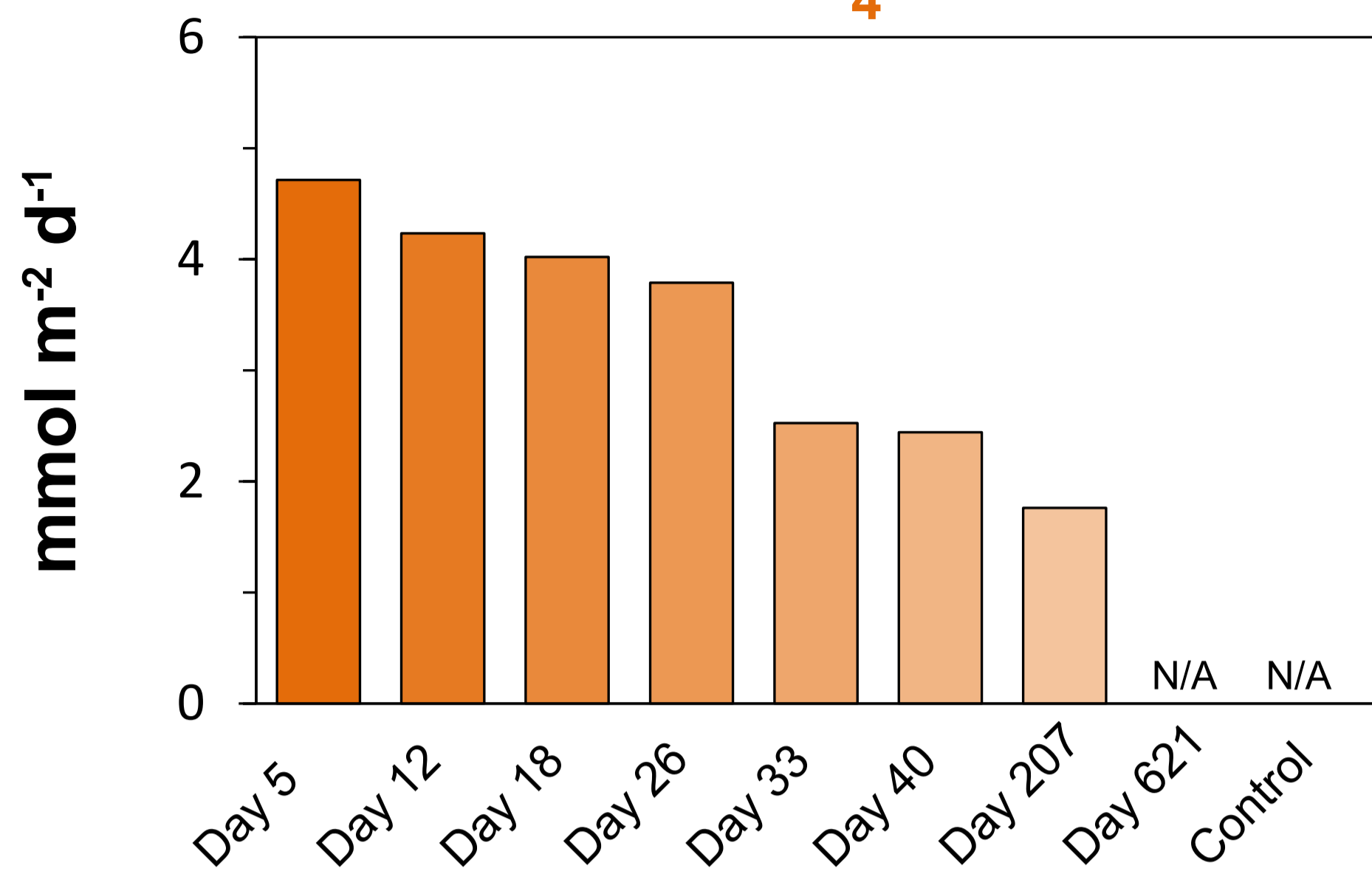
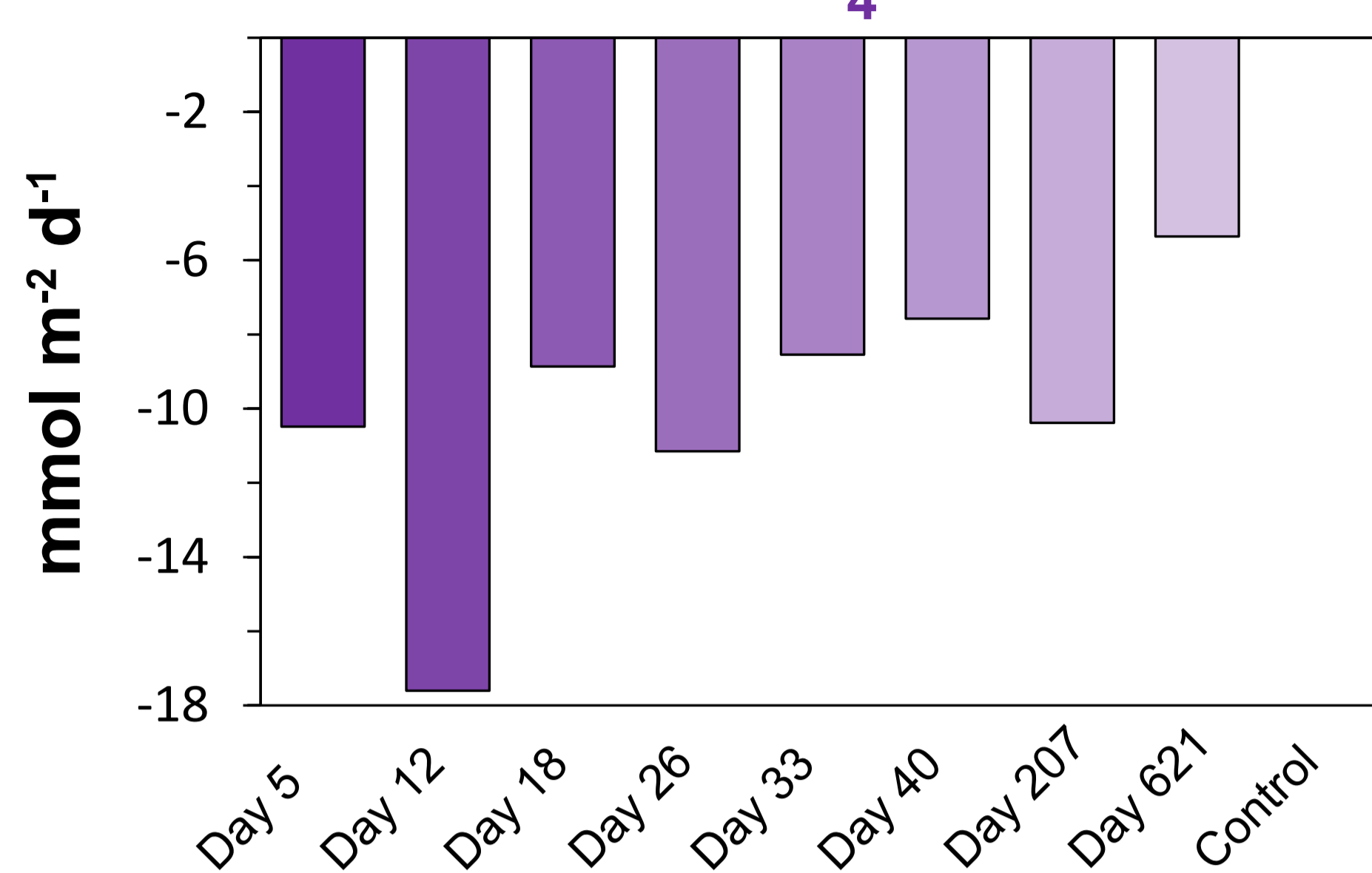
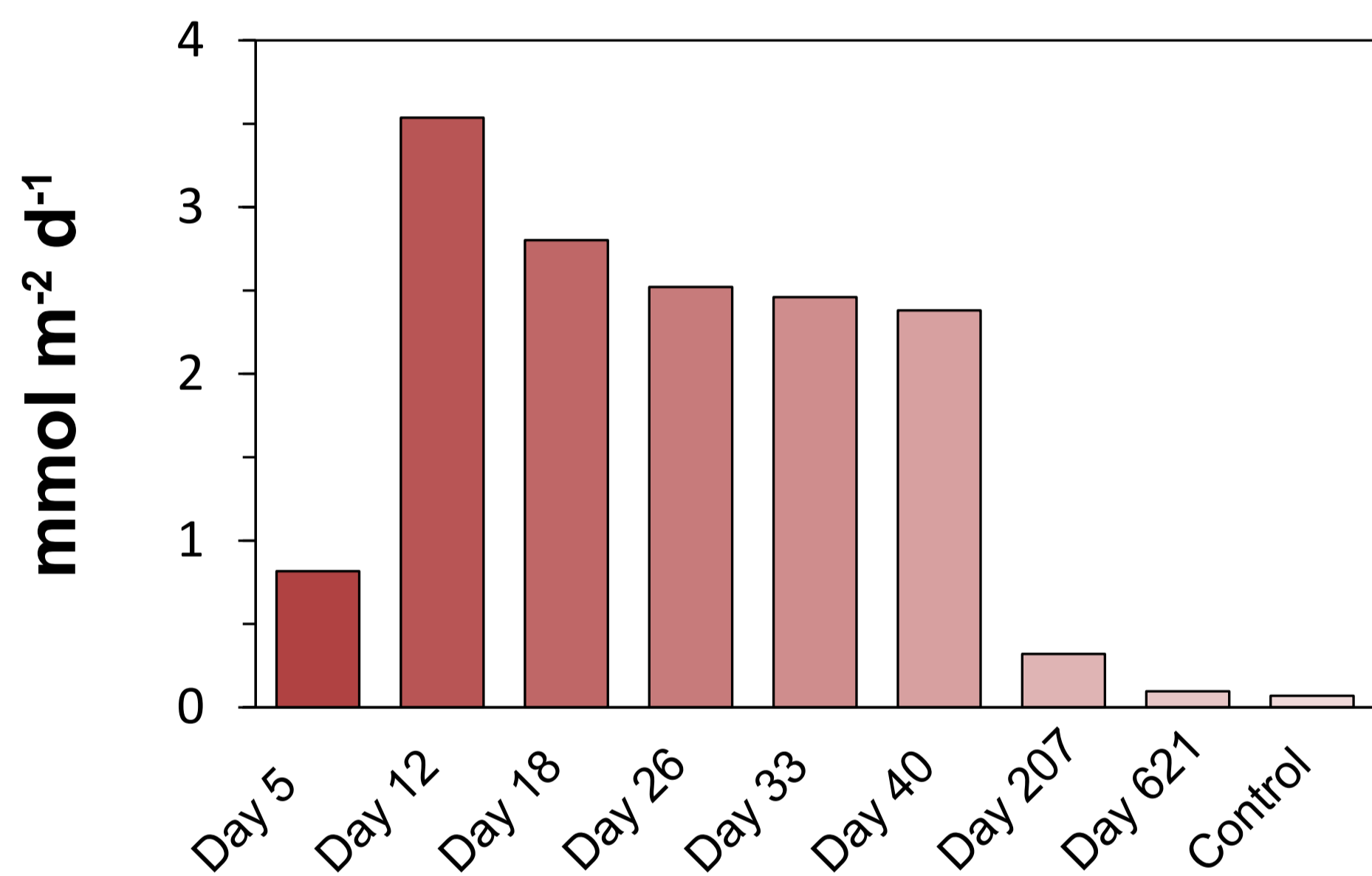
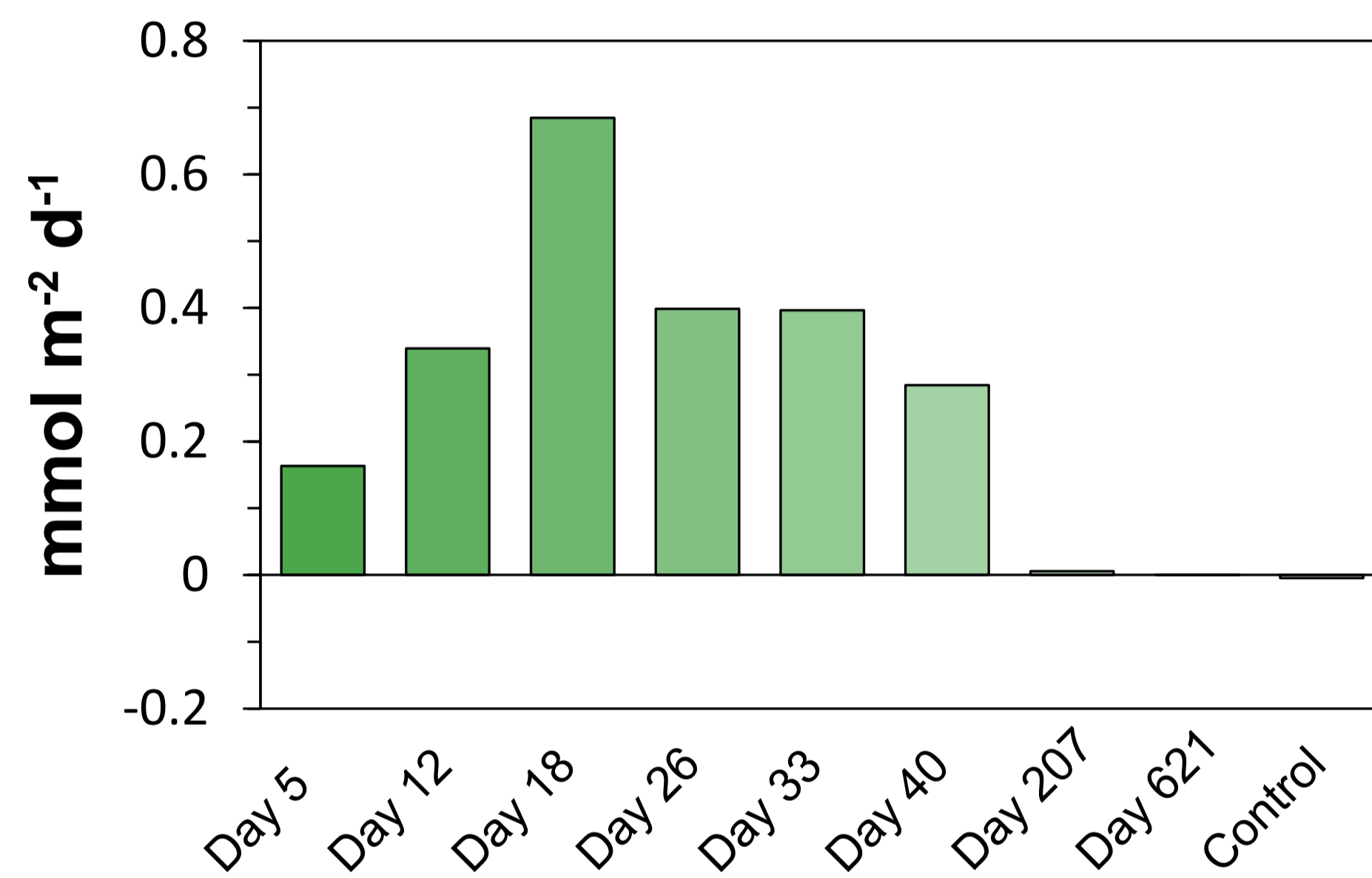
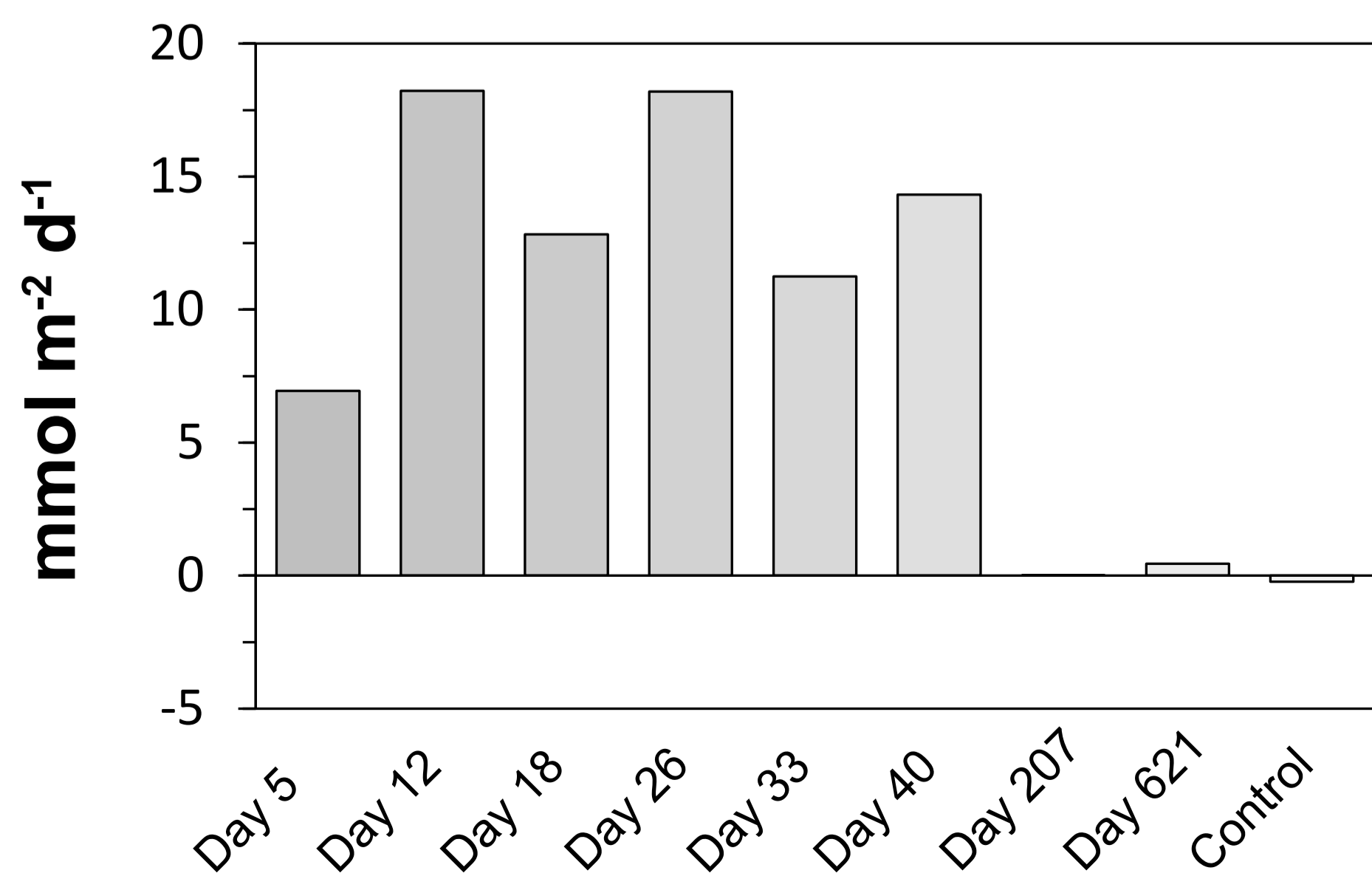
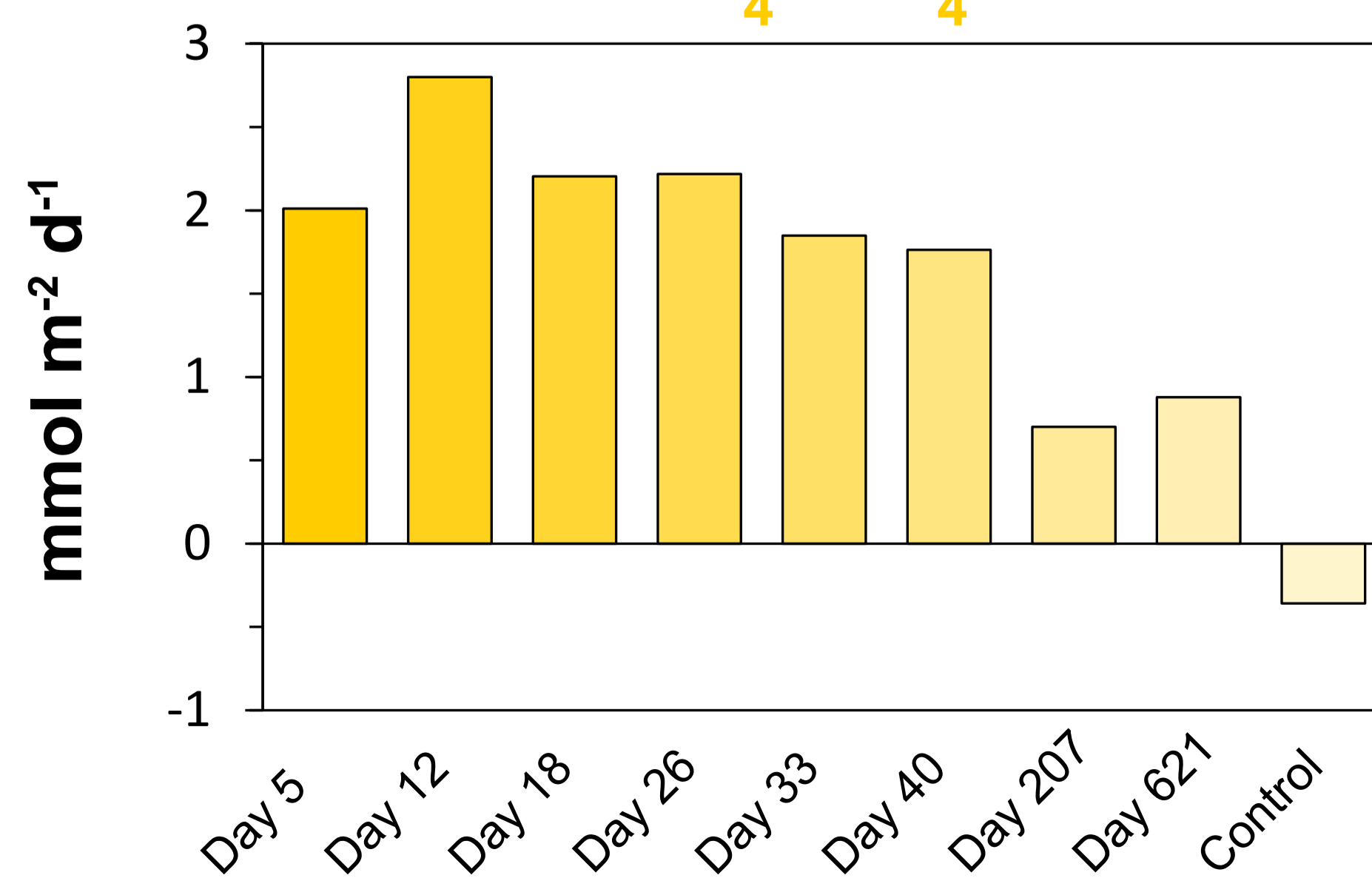
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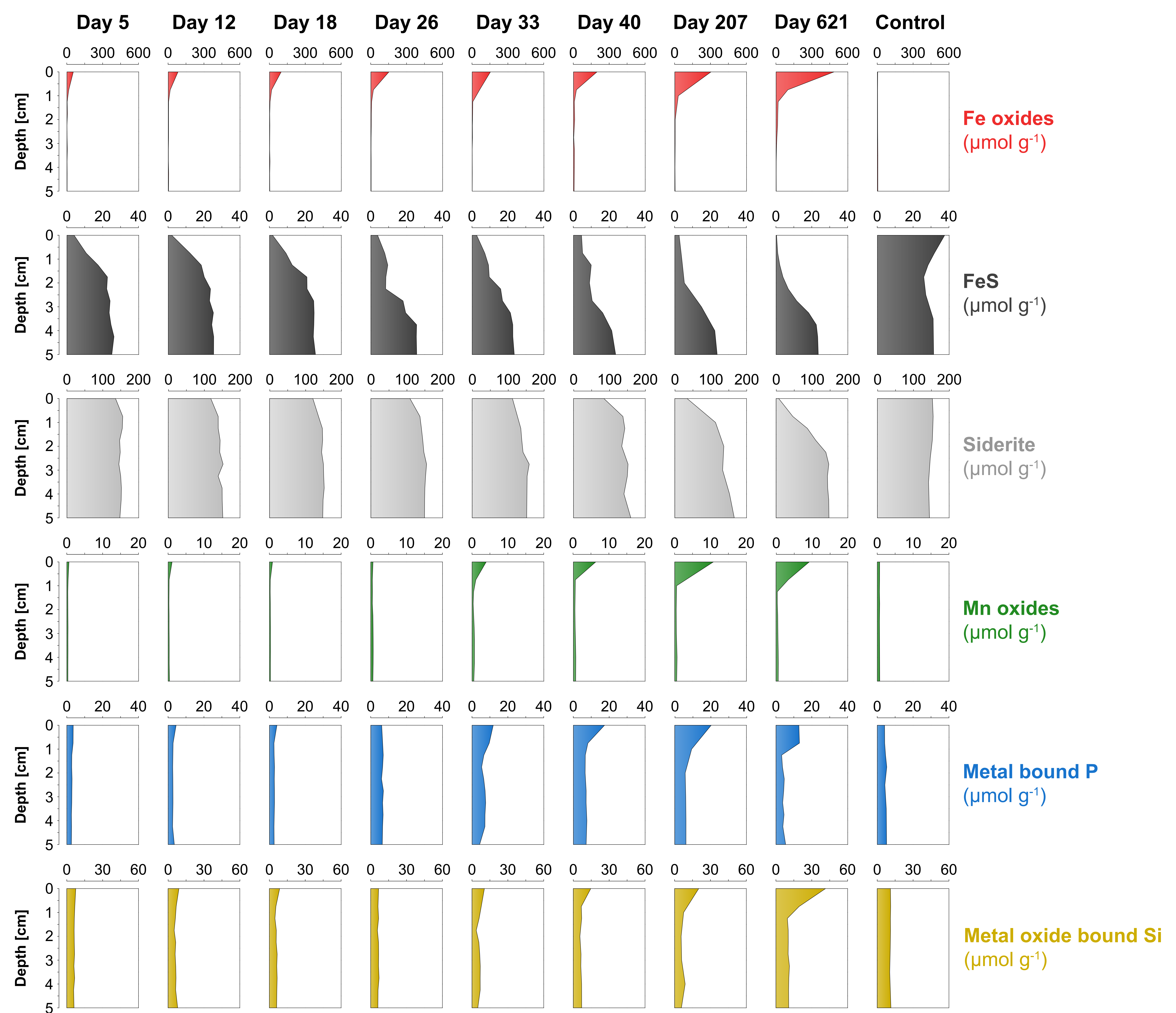
Diffusive O₂ Uptake

**B**

Current Density

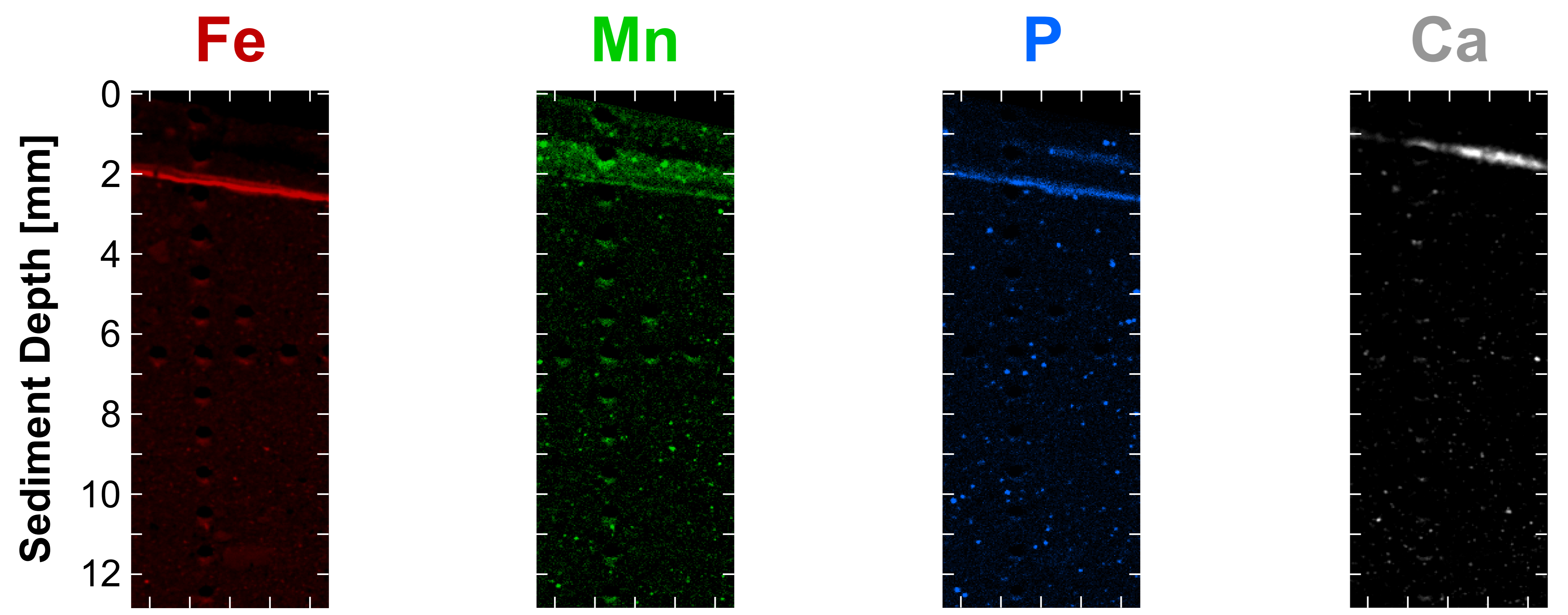


A**B****C****D****E****F**

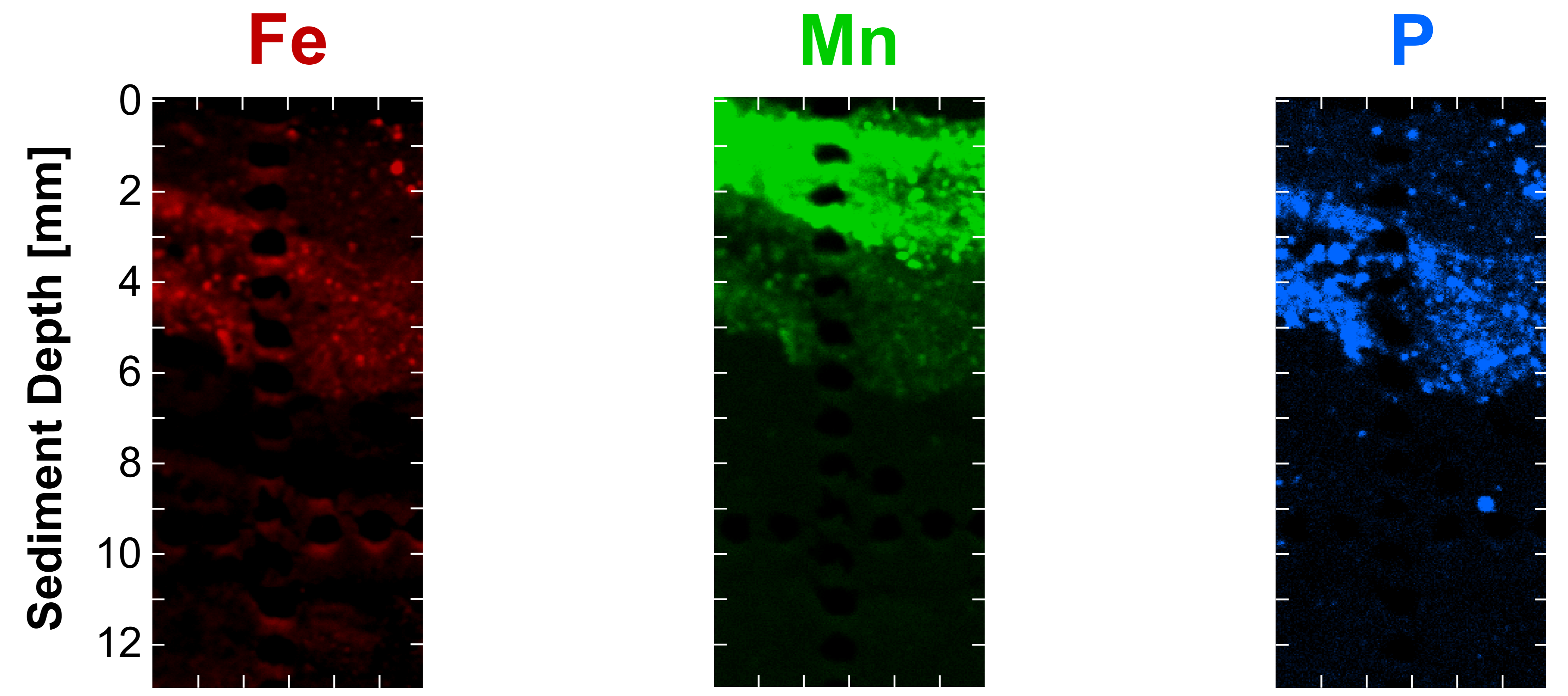


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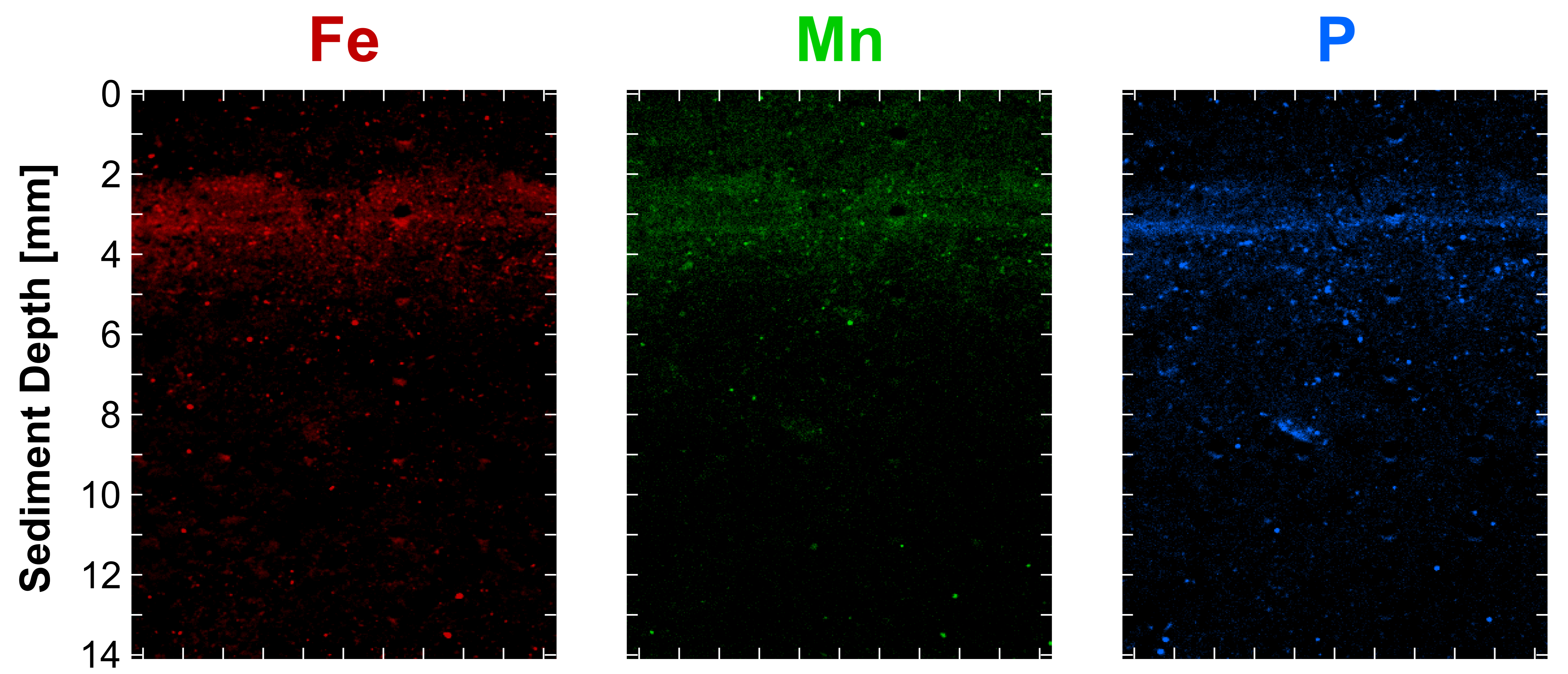
Black Sea Experiment
Cable bacteria

**B**

Gulf of Finland (GOF5)
Cable bacteria

**C**

Lake Grevelingen in January
Cable bacteria

**D**

Lake Grevelingen in May
Macrofauna

