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| 5        | Manganese and iron reduction dominate organic carbon oxidation in surface   |
| 6        | sediments of the deep Ulleung Basin, East Sea   |
| 7        |   |
| 8        |   |
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31 Abstract. Rates and pathways of benthic organic carbon (Corg) oxidation were investigated in surface sediments of the Ulleung Basin (UB) characterized by high organic carbon contents 32 (> 2.5 %, dry wt.) and very high contents of Mn oxides (> 200  $\mu$ mol cm<sup>-3</sup>) and Fe oxides (up 33 to 100 µmol cm<sup>-3</sup>). The combination of geochemical analyses and independently executed 34 35 metabolic rate measurements revealed that Mn and Fe reduction were the dominant Corg oxidation pathways in the center of the UB, comprising 45 % and 20 % of total Corg oxidation, 36 respectively. By contrast, sulfate reduction was the dominant Corg oxidation pathway 37 accounting for 50 % of total Corg mineralization in sediments of the continental slope. The 38 relative significance of each Corg oxidation pathway matched the depth distribution of the 39 respective electron acceptors. The relative importance of Mn reduction for  $C_{\mbox{\scriptsize org}}$  oxidation 40 displays saturation kinetics with respect to Mn oxide content with a low half-saturation value 41 of 8.6  $\mu$ mol cm<sup>-3</sup>, which further implies that Mn reduction can be a dominant C<sub>org</sub> oxidation 42 process even in sediments with lower MnO<sub>2</sub> content as known from several other locations. 43 This is the first report of a high contribution of manganese reduction to Corg oxidation in 44 offshore sediments on the Asian margin. The high manganese oxide content in the surface 45 sediment in the central UB was maintained by an extreme degree of recycling, with each Mn 46 atom on average being reoxidized ~3800 times before permanent burial. This is the highest 47 48 degree of recycling so far reported for Mn-rich sediments, and it appears linked to the high benthic mineralization rates resulting from the high organic carbon content that indicate the 49 50 UB as a biogeochemical hotspot for turnover of organic matter and nutrient regeneration.

51 52

53 **Keywords.** Benthic mineralization, Manganese reduction, Iron reduction, Sulfate reduction,

- 54 Ulleung Basin, East Sea
- 55

### 56 **1 Introduction**

57

Although they cover only 15 % ( $47 \times 10^6 \text{ km}^2$ ) of the ocean surface area, sediments of 58 continental margins (200-2000 m depth) are characterized by enhanced organic matter flux 59 generated either by vertical transport from the highly productive overlying water column or 60 by lateral transport from adjacent shelves, and thus play an important role in deposition and 61 mineralization of organic matter (Romankevich, 1984; Jahnke et al., 1990; Walsh, 1991; 62 Jahnke and Jahnke, 2000). Organic particles that reach the seafloor are quickly mineralized 63 64 by hydrolysis, fermentation, and a variety of respiratory processes using different electron acceptors such as oxygen, nitrate, Mn oxides, Fe oxides, and sulfate (Froelich et al., 1979; 65 Jørgensen, 2006). The partitioning of organic carbon (Corg) oxidation among the different 66 electron accepting pathways has profound influence on the distribution and the release and/or 67 retention of Mn, Fe, S and nutrients (nitrogen and phosphate) (Canfield et al., 2005; Hansen 68 et al., 2006; Jørgensen, 2006; Slomp et al., 2013). Therefore, it is particularly important to 69 elucidate the contribution of each Corg oxidation pathway in order to better understand the 70 role of sediments in biogeochemical element cycles. 71

72 The relative significance of each carbon oxidation pathway is largely controlled by the 73 combination of organic matter supply and availability of electron acceptors. In general, aerobic metabolism dominates the organic matter mineralization in deep-sea sediments that 74 75 are characterized by low organic matter content (Jahnke et al., 1982; Glud, 2008), especially 76 in organic carbon-starved deep-sea sediments with low sedimentation rates (Mewes et al., 77 2014, 2016; D'Hondt et al. 2015; Mogollón et al., 2016). In contrast, owing to high sulfate concentrations in marine sediment, sulfate reduction might account for up to 50 % of total 78 79 carbon oxidation in continental margins with high organic matter flux (Jørgensen, 1982; Jørgensen and Kasten, 2006; Bowles et al., 2014). However, in sediments where manganese 80 81 and iron oxides are abundant or rapidly recycled, microbial reduction of manganese and iron can be the dominant electron accepting processes over sulfate reduction (Sørensen and 82 Jørgensen, 1987; Aller, 1990; Canfield et al., 1993b). The significance of dissimilatory iron 83 reduction for Corg oxidation is well established in the sediments of various continental 84 margins and coastal wetlands (Thamdrup, 2000; Thamdrup and Canfield, 1996; Jensen et al., 85 2003, Kostka et al., 2002a, 2002b; Vandieken et al., 2006; Hyun et al., 2007, 2009b). 86 However, only a few locations such as the Panama Basin (Aller, 1990), the coastal 87 Norwegian trough in Skagerrak and an adjacent fjord (Canfield et al., 1993a, 1993b; 88

Vandieken et al., 2014), the Black Sea shelf (Thamdrup et al., 2000) and the continental shelf
of the northern Barents Sea (Vandieken et al., 2006; Nickel et al., 2008) are known where
microbial manganese reduction significantly contributes to carbon mineralization.

The East Sea (often referred to as Japan Sea), located in the far eastern part of the Eurasian 92 93 continental margin, consists of three major basins deeper than 2000 m, the Japan Basin, the Yamato Basin and the Ulleung Basin (Fig. 1). Compared to the other two basins, the surface 94 95 waters of the Ulleung Basin (UB) are characterized by higher phytoplankton biomass and primary production (Yamada et al., 2005; Yoo and Park, 2009), which is associated with 96 coastal upwelling (Hyun et al., 2009a). The enhanced biological production in the euphotic 97 zone of the UB is responsible for the high organic carbon content (> 2.5 % wt) in the 98 sediment, and the highest rates of Corg oxidation compared to other deep-sea sediments with 99 similar depth range (Lee et al., 2008; Hyun et al., 2010). An intriguing geochemical property 100 of the UB surface sediment is the high content of Mn oxides (> 200  $\mu$ mol cm<sup>-3</sup>) and Fe oxides 101 (up to 100 µmol cm<sup>-3</sup>) (Cha et al., 2007; Hyun et al., 2010). In accordance with these 102 geochemical findings, the suppression of sulfate reduction (Hyun et al., 2010) and 103 accumulation of Mn<sup>2+</sup> in anoxic incubation of surface sediment (Vandieken et al., 2012) 104 strongly implied that the Corg oxidation in the surface sediment of the UB is dominated by 105 106 microbial manganese and iron reduction, but actual rates and partitioning of each electron accepting pathway in Corg oxidation remain to be determined in this deep marginal sediment 107 108 underlying highly productive surface waters.

The primary objective of this paper was to characterize the sediment biogeochemistry with regard to the rate of  $C_{org}$  oxidation and partitioning of major terminal electron accepting pathways at two contrasting sites at the continental slope and rise in the UB. Here, for the first time in sediments of the Asian marginal seas, we document that Mn and Fe reduction are the dominant  $C_{org}$  oxidation pathways accounting for respectively 45 % and 20 % of total  $C_{org}$ oxidation in the center of the UB, and suggest that Mn and Fe reduction may be of greater importance in deep-sea sediments than previously recognized.

- 116
- 117 **2 Materials and methods**
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- 119 **2.1 Study site**
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<sup>121</sup> The East Sea is a marginal sea surrounded by the east Asian continent and Japanese Islands

(Fig. 1, Kang et al., 2010; Liu et al., 2010). The UB located in the southwestern part of the
East Sea is a bowl-shaped deep basin (2000–3000 m depth) (Fig. 1) delimited by continental
slopes of Korean Peninsula and the southwestern Japanese Archipelago on the west and south,
respectively, and by the Korea Plateau and the Oki Bank on the north and east, respectively
(Chough et al., 2000).

127 Shipboard experiments were conducted in June, 2009 at two sites on the continental slope 128 (Station M1, hereafter M1) and in the center (Station D3, hereafter D3) of the UB (Fig. 1, 129 Table 1). Surface sediments consist of fine-grained clay with a mean grain size less than 130 0.004 mm in diameter (Cha et al., 2007). Two stations were characterized by two contrasting 131 sediment colors. The Mn oxide-enriched surface sediment at the basin site (D3) was reddish-132 brown, whereas at the slope site (M1) it exhibited the typical gray-brown color of muddy 133 continental margin sediments (Fig. 1). Further environmental properties are listed in Table 1.

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### 135 2.2 Sampling and handling

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Sediment samples were collected with a box corer. Onboard, duplicate or mostly triplicate 137 sub-samples for geochemical analyses were collected using acrylic cores (6–9 cm in diameter 138 139 and 30-40 cm in length). The sub-cores for geochemical analyses were immediately sealed with butyl rubber stoppers and transferred to a N<sub>2</sub>-filled glove bag for sectioning and loading 140 141 into polypropylene centrifuge tubes that were then tightly capped and centrifuged for 15 min at 5000  $\times$  g. After reintroduction into the N<sub>2</sub>-filled glove bag, pore-waters were sampled and 142 143 filtered through 0.2-µm cellulose ester syringe filters (ADVANTEC, Toyo Rashi Kaisha, Ltd). One to two mL of pore water to determine  $NH_4^+$  was fixed with saturated HgCl<sub>2</sub>, and frozen. 144 For determination of  $Fe^{2+}$ , Mn,  $SO_4^{2-}$  and  $Ca^{2+}$ , 2 mL of the pore water were acidified with 12 145 M HCl and stored at 4 °C. Pore-water for sulfide analysis was preserved with Zn acetate 146 147 (20 %). Sediments for solid-phase analysis were frozen at -25 °C for future analyses.

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### 149 **2.3 Anoxic bag incubations**

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Anaerobic carbon mineralization rates and dissimilatory Mn and Fe reduction rates were determined in batch incubations based on the procedures of Canfield et al. (1993b) and Thamdrup and Canfield (1996). Sediment cores were transferred to a  $N_2$ -filled glove bag and sliced in 2-cm intervals to a depth of 10 cm. Sediment from parallel sections was pooled, 155 mixed and loaded into gas-tight plastic bags (Hansen et al., 2000). The bags were sealed without gas space, and incubated in the dark at near in situ temperature (ca. 1–2 °C) in larger 156 N<sub>2</sub> filled bags to ensure anoxic conditions. Over a period of 18 days of incubation, sub-157 samples to determine the accumulation of total dissolved inorganic carbon (DIC) and Mn in 158 159 pore water were withdrawn on days 0, 1, 3, 5, 9 and 18. Two 50-mL centrifuge tubes per bag were filled completely with sediment in a N<sub>2</sub>-filled glove bag, and pore-water was extracted 160 as described above. For DIC analysis, we collected 1.8 mL aliquots into glass vials without 161 head space, fixed with 18 µL of HgCl<sub>2</sub> (125 mM), and stored at 4 °C until analysis within 4 162 163 weeks. Samples for Mn analysis were acidified with 12 M HCl and stored at 4 °C. Sediment remaining after the collection of pore water was frozen at -25 °C for later analysis of oxalate 164 extractable solid Fe(II). 165

166

## 167 **2.4 Pore-water analyses**

168

Total dissolved inorganic carbon (DIC) and NH<sub>4</sub><sup>+</sup> were measured by flow injection analysis 169 170 with conductivity detection (Hall and Aller. 1992). Nitrate was measured spectrophotometrically (Parsons et al., 1984). Dissolved  $Fe^{2+}$  was determined by colorimetric 171 method with Ferrozine (Stookey, 1970). Dissolved  $Mn^{2+}$  and  $Ca^{2+}$  were analyzed in acidified 172 pore water by inductively coupled plasma-atomic emission spectrometry (ICP-AES, Optima 173 174 3300DV, Perkin-Elmer Co.) and flame atomic absorption spectrometer (SpectrAA 220/FS, Varian), respectively (Thamdrup and Canfield, 1996). Dissolved sulfide was determined by 175 the methylene blue method (Cline, 1969). Sulfate concentrations were measured using ion 176 chromatography (Metrohm 761). The detection limit of H<sub>2</sub>S,  $Ca^{2+}$ ,  $Mn^{2+}$  and  $Fe^{2+}$  was 3  $\mu$ M, 177 1.8  $\mu$ M, 3  $\mu$ M and 1  $\mu$ M, respectively. Reproducibility of DIC and NH<sub>4</sub><sup>+</sup> was better than 178 10 %. Precision of  $NO_3^-$  was 1–2 %. 179

180

### 181 **2.5 Solid-phase analyses**

182

Total oxalate-extractable Fe [Fe(II) + Fe(III)] was extracted from air-dried sediment in a 0.2 M oxic oxalate solution (pH 3) for 4 h (Thamdrup and Canfield, 1996), and Fe(II) was extracted from frozen sediment in anoxic oxalate (Phillips and Lovley, 1987). The total oxalate-extractable Fe and Fe(II), hereafter total  $Fe_{(oxal)}$  and  $Fe(II)_{(oxal)}$ , were determined as described for the pore-water analysis of  $Fe^{2+}$ . Oxalate-extractable Fe(III), hereafter

188 Fe(III)<sub>(oxal)</sub>, was defined as the difference between total Fe<sub>(oxal)</sub> and Fe(II)<sub>(oxal)</sub>. This fraction represents poorly crystalline Fe(III) oxides. Particulate Mn, hereafter Mn<sub>(DCA)</sub> was extracted 189 with dithionite-citrate-acetic acid (DCA; pH 4.8) for 4 h from air-dried sediment and was 190 determined by inductively coupled plasma-atomic emission spectrometry (ICP-AES, Optima 191 192 3300DV, Perkin-Elmer Co). The DCA extraction aims at dissolving free Mn oxides and authigenic Mn(II) phases. The reproducibility of the measurements was better than 10 % and 193 the detection limits was 3 µM for Mn and 1 µM for Fe. For the determination of total reduced 194 sulfur (TRS) that includes acid volatile sulfide (AVS =  $FeS + H_2S$  and small amounts of other 195 metal sulfides, see Rickard and Morse, 2005; Luther, 2005) and chromium-reducible sulfur 196  $(CRS = S^{0} + FeS_{2})$ , sediment samples were fixed with Zn acetate, and sulfide was determined 197 according to the method of Cline (1969) after a two-step distillation with cold 12 M HCl and 198 boiling 0.5 M Cr<sup>2+</sup> solution (Fossing and Jørgensen, 1989). The contents of particulate 199 organic carbon (POC) and nitrogen (PON) in the surface sediment were analyzed using a 200 CHN analyzer (CE Instrument, EA 1110) after removing CaCO<sub>3</sub> using 12 M HCl. 201

202

### 203 2.6 Oxygen micro-profiles

204

Oxygen profiles were measured at 50  $\mu$ m resolution using Clark-type microelectrodes (Unisense, OX-50) while stirring the overlying water. Microelectrodes were calibrated between 100 % air-saturated *in situ* bottom water and N<sub>2</sub> purged anoxic bottom water. Three profiles were measured at each site. The diffusive boundary layer (DBL) and sediment-water interface (SWI) were determined according to Jørgensen and Revsbech (1985). To estimate the volume-specific oxygen consumption rate, we used the PROFILE software (Berg et al., 1998).

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### 213 2.7 Rate measurements

214

The diffusive oxygen uptake (DOU) was calculated from the calibrated oxygen microprofiles.

217 
$$DOU = -D_0 \times (\Delta C / \Delta z)$$
 (1)

218

where  $D_0 (1.07 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \text{ at M1} \text{ and } 1.03 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \text{ at D3})$  is the temperature-corrected molecular diffusion coefficient estimated from Schulz (2006), and C is the oxygen concentration at depth z within the diffusive boundary layer (DBL) (Jørgensen and Revsbech,
1985).

The volume-specific  $O_2$  consumption rates exhibited a bimodal depth distribution (see results section 3.2) with activity peaks near the sediment-water interface and the oxic/anoxic interface, respectively. Thus,  $O_2$  consumption rates by aerobic organotrophic respiration was defined as the  $O_2$  consumption rate near the sediment-water interface, whereas the oxygen consumption at the oxic-anoxic interface was assigned to re-oxidation of reduced inorganic compounds (Rasmussen and Jørgensen, 1992; Canfield et al., 2005).

Total anaerobic  $C_{org}$  mineralization rates were determined by linear regression of the accumulation of total DIC with time during the anoxic bag incubations (Fig. 3) after correcting for CaCO<sub>3</sub> precipitation (Thamdrup et al., 2000). Briefly, CaCO<sub>3</sub> precipitation was calculated from decreasing dissolved Ca<sup>2+</sup> concentration during the anoxic bag incubation: 233

234 
$$\Delta \text{CaCO}_3 = \Delta[\text{Ca}^{2+}]_{\text{sol}} \times (1+K_{\text{Ca}})$$

235

where,  $K_{Ca}$  is the adsorption constant for  $Ca^{2+}$  ( $K_{Ca} = 1.6$ ) (Li and Gregory, 1974). Then the DIC production rate corrected for CaCO<sub>3</sub> precipitation was calculated as:

(2)

(3)

239 DIC production = DIC accumulation + 
$$CaCO_3$$
 precipitation

240

238

Fe(III) reduction rates were determined by linear regression of the increase in solid-phase Fe(II)<sub>(oxal)</sub> content with time during anoxic bag incubations. The dissimilatory microbial Fe(III) reduction rate was derived by subtracting abiotic Fe reduction coupled to the oxidation of sulfide produced by sulfate reduction (Gribsholt et al., 2003):

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assuming that abiotic Fe reduction coupled to H<sub>2</sub>S oxidation occurred at a stoichiometry of 2
Fe(III) per 3 H<sub>2</sub>S (Pyzik and Sommer, 1981; Melton et al., 2014):

250

$$251 2FeOOH + 3H_2S_{(produced by SR)} = 2FeS + S^o + 4H_2O (5)$$

252

Finally, to estimate the C<sub>org</sub> oxidation by microbial Fe reduction, the 4:1 stoichiometry of

iron reduction coupled to  $C_{\text{org}}$  oxidation was used from the stoichiometric equation (Canfield 254 255 et al., 1993a):

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- 257

$$CH_2O + 4FeOOH + 8H^+ = CO_2 + 4Fe^{2+} + 7H_2O$$
(6)

258

Mn reduction rates were determined from linear regression of the production of dissolved 259 Mn<sup>2+</sup> with time during the anoxic bag incubations. Similar to previous studies (e.g., Canfield 260 et al., 1993a, 1993b; Thamdrup and Dalsgaard, 2000), we assumed that accumulating 261 dissolved Mn was Mn<sup>2+</sup>. This ignores a potential contribution from Mn<sup>3+</sup>, which in some 262 cases can constitute a substantial fraction of the dissolved Mn pool at the upper boundary of 263 the zone with soluble Mn accumulation in marine sediments (Madison et al., 2013). Further 264 studies of the dynamics of soluble Mn<sup>3+</sup> are required to evaluate its potential importance in 265 anoxic incubations. Such studies pending, we find justification for our assumption in the 266 good agreement observed in the previous studies between Mn reduction rates calculated 267 based on the assumption that soluble Mn is  $Mn^{2+}$  (Eq. 7) and independent estimates of rates 268 of carbon mineralization through dissimilatory Mn reduction based on DIC or NH4<sup>+</sup> 269 accumulation. Due to strong adsorption of Mn<sup>2+</sup> to Mn oxide surfaces (Canfield et al., 1993b), 270 the Mn reduction rates were estimated after compensating for the adsorption effect of  $\mathrm{Mn}^{2+}$  to 271 Mn-oxides according to Thamdrup and Dalsgaard (2000): 272

273

Mn reduction rate = Mn<sup>2+</sup> accumulation rate  $\times (1 + K_{Mn}^{*2+} \times (1 - \Phi) \times \Phi^{-1} \times \delta_s)$ (7)274

275

where,  $\Phi = \text{porosity}$ 276

- $\delta_s$  = density of sediment 277
- 278

 $K_{Mn}^{*}^{2+} = 4.8 + 0.14 \times [Mn(IV)] \text{ (ml g}^{-1}\text{)}$ 

[Mn(IV)] = the content of Mn(IV) (
$$\mu$$
mol g<sup>-1</sup>)

280

We here assume that extracted Mn(DCA) represents Mn(IV) as observed in surface 281 sediments of another Mn-rich site (Canfield et al., 1993b; Thamdrup and Dalsgaard, 2000). 282 Small levels of Mn<sub>(DCA)</sub> remaining at depth further suggest that little Mn(II) accumulates in 283 the solid phase (see Results). Corg oxidation by dissimilatory Mn(IV) reduction was 284 calculated from the stoichiometric equation (Canfield et al., 1993a): 285

286

287

$$CH_2O + 2MnO_2 + 4H^+ = CO_2 + 2Mn^{2+} + 3H_2O$$
(8)

288

Sulfate reduction rates were determined using the radiotracer method of Jørgensen (1978). 289 Sediment cores (35 cm long with 2.9 cm i.d.) were collected in triplicate, injected 290 horizontally at 1-cm vertical interval with 5  $\mu$ L radiolabeled sulfate (<sup>35</sup>S-SO<sub>4</sub><sup>2-</sup>, 15 kBg  $\mu$ l<sup>-1</sup>, 291 Amersham) diluted in sterilized NaCl solution (3.0 %), and incubated for 12 h at in situ 292 293 temperature. At the end of the incubation, the sediment was sliced into sections, fixed in Zn acetate (20 %), and frozen at -25 °C until processed in the laboratory. The reduced <sup>35</sup>S was 294 recovered using distillation with a boiling acidic Cr<sup>2+</sup> solution according to Fossing and 295 Jørgensen (1989). Background radioactivity of <sup>35</sup>S was  $32.4 \pm 3.7$  cpm cm<sup>-3</sup> (n = 10) at site 296 D3 and 87.5  $\pm$  38.7 cpm cm<sup>-3</sup> (n = 10) at site M1. Detection limits of the SRR, estimated from 297 the double standard deviation of the blank value (i.e., 7.4 and 77.4 cpm) according to Fossing 298 et al. (2000), ranged from 0.79 to 2.62 nmol cm<sup>-3</sup> d<sup>-1</sup>. To elucidate the contribution of sulfate 299 300 reduction in anaerobic carbon oxidation, the SRRs (Fig. 5B, 5G) were converted to carbon oxidation using a stoichiometric equation (Thamdrup and Canfield, 1996): 301

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$$2CH_2O + SO_4^{2-} + 2H^+ = 2CO_2 + H_2S + 2H_2O$$
(9)

304

305 **3 Results** 

306

### 307 **3.1 Pore-water and solid-phase constituents**

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The depth distributions of  $NH_4^+$ ,  $NO_3^-$ ,  $Mn^{2+}$  and  $Fe^{2+}$  in the pore-water as well as solid phase 309 Mn, Fe and S for the two stations are shown in Fig. 2.  $NH_4^+$  concentrations at M1 increased 310 steadily with depth (Fig. 2A) whereas at D3 it decreased down to 3 cm depth before it 311 increased below (Fig. 2F). Highest concentrations of nitrate were measured at 0 to 1 cm 312 sediment depth at the two stations and concentrations decreased below a background level (< 313  $2 \mu$ M) below 1 cm at both M1 and D3 (Fig. 2A, 2F). Dissolved Mn<sup>2+</sup> concentrations differed 314 widely between the sites showing a maximum of 56 µM between 0 and 3 cm depth and not 315 exceeding 10 µM below at M1 (Fig. 2B), whereas at D3 concentrations increased to a 316 maximum of 286  $\mu$ M at 10–12 cm depth (Fig. 2G). Conversely, dissolved Fe<sup>2+</sup> concentrations 317

at M1 increased from 11  $\mu$ M at 0–0.5 cm to 32  $\mu$ M at 6–7 cm depth, and stayed constant below (Fig. 2C), whereas at D3, concentrations were uniformly low showing a slight increase to 12  $\mu$ M at 15 cm (Fig. 2H).

Extractable Mn (Mn<sub>(DCA)</sub>) contents were low (< 3  $\mu$ mol cm<sup>-3</sup>) in the upper 20 cm at the 321 slope site (M1) (Fig. 2B), but up to 200  $\mu$ mol cm<sup>-3</sup> in the upper 4 cm depth of the sediment at 322 the center of the basin (D3), with a sharp decrease to near depletion ( $\sim 1 \mu mol cm^{-3}$ ) below 10 323 cm (Fig. 2G). At the slope site (M1), contents of Fe(III)(oxal) decreased slightly with 324 increasing depth from 28 µmol cm<sup>-3</sup> near the surface to 13 µmol cm<sup>-3</sup> at 20 cm depth, 325 mirroring an increase in Fe(II)(oxal) (Fig. 2D). At the center of the basin (D3), Fe(III)(oxal) 326 increased slightly from 67 µmol cm<sup>-3</sup> at 0–0.5 cm to 90 µmol cm<sup>-3</sup> at 4–6 cm depth, and 327 decreased steeply below to 4.8  $\mu$ mol cm<sup>-3</sup> at 12–14 cm depth (Fig. 2I). Of total Fe<sub>(oxal)</sub>, 328  $Fe(III)_{(oxal)}$  comprised > 98 % at 0–2 cm and > 97 % at 0–8 cm depth at M1 and D3, 329 respectively. The fraction of Fe(III)<sub>(oxal)</sub> in Fe<sub>(oxal)</sub> then decreased to 40 % at 10–12 cm depth 330 at both sites. Acid volatile sulfur (AVS) exhibited a slight increase with depth at M1 from 0.8 331  $\mu$ mol cm<sup>-3</sup> at the surface to 7.2  $\mu$ mol cm<sup>-3</sup> at 20 cm depth (Fig. 2E), but was not detected at 332 D3 (Fig. 2J). Chromium reducible sulfur (CRS) contents at M1 increased rapidly with depth 333 from 1.9  $\mu$ mol cm<sup>-3</sup> at 0–0.5 cm to 21.8  $\mu$ mol cm<sup>-3</sup> at 20 cm depth (Fig. 2E), whereas the 334 CRS contents remained  $< 0.1 \mu mol \text{ cm}^{-3}$  at D3 (Fig. 2J). 335

336

### **337 3.2 O**<sub>2</sub> microprofiles and diffusive oxygen uptake rate

338

Oxygen penetrated less than 4 mm into the sediments (Fig. 3), and rates of diffusive oxygen 339 uptake (DOU) were 7.1 and 6.0 mmol  $O_2$  m<sup>-2</sup> d<sup>-1</sup> at M1 and D3, respectively (Table 2). 340 Oxygen consumption by aerobic respiration estimated from the O<sub>2</sub> micro-profiles (area I and 341 II in Fig. 3) was higher at the M1 in the slope site (4.0 mmol  $O_2 \text{ m}^{-2} \text{ d}^{-1}$ ) than at the D3 in the 342 center of the basin (2.5 mmol  $O_2$  m<sup>-2</sup> d<sup>-1</sup>).  $O_2$  consumption by re-oxidation of reduced 343 inorganic compounds indicated by increased activity at the oxic/anoxic interface (area III in 344 Fig. 3) accounted for 43 % and 57 % of the DOU at M1 and D3, respectively. From the 345 profiles of geochemical constituents (Fig. 2), O<sub>2</sub> consumption was mainly attributed to the re-346 oxidation of sulfide and  $Fe^{2+}$  at M1 and of  $Mn^{2+}$  at D3. 347

348

### 349 **3.3 Anoxic bag incubations**

Changes in concentrations of DIC, Ca<sup>2+</sup> and dissolved Mn<sup>2+</sup> and solid Fe(II)<sub>(oxal)</sub> contents 351 over time during anoxic bag incubations from sediment of 0-2, 2-4, 4-6 and 6-8 cm depth 352 353 intervals are presented in Fig. 4. The DIC concentrations increased linearly over time during incubations of sediment in all bags from M1 and D3, except the bag from 6-8 cm at D3. The 354 355 DIC accumulation rates were generally higher at the slope site (M1) than at the basin site (D3) (Table 4). The concentrations of  $Ca^{2+}$  decreased with time at all depth intervals of M1, 356 whereas a decrease of  $Ca^{2+}$  was observed only for the 2–4 cm depth interval at D3. The 357 decrease of  $Ca^{2+}$  indicates  $CaCO_3$  precipitation, which consequently underestimates DIC 358 accumulation, especially at M1. 359

Coinciding with high solid  $Mn_{(DCA)}$  contents (Fig. 2G), prominent  $Mn^{2+}$  accumulation appeared at 0–6 cm depth of D3, whereas no increase of  $Mn^{2+}$  was observed at M1 except a slight accumulation at 0–2 cm interval (Fig. 4). Solid Fe(II)<sub>(oxal)</sub> contents increased linearly with time at 0–4 cm depth of M1, whereas highest Fe(II)<sub>(oxal)</sub> accumulation was observed at 4–6 cm depth at D3. An increase of Fe(II)<sub>(oxal)</sub> was not discernible in the Mn-oxide-rich surface sediment (0–2 cm) of D3.

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### 367 **3.4 Sulfate reduction rates (SRR)**

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### 5.4 Sunate reduction rates (SKK)

At the slope site (M1), SRR increased from 18 nmol cm<sup>-3</sup> d<sup>-1</sup> at the surface to 97–103 nmol cm<sup>-3</sup> d<sup>-1</sup> at 1.5–2 cm depth, and decreased below to 12.5 nmol cm<sup>-3</sup> d<sup>-1</sup> at 20 cm depth (Fig. 5B). In contrast, SRR at the manganese oxide-rich basin site (D3) ranged from 1.7 to 8.7 nmol cm<sup>-3</sup> d<sup>-1</sup>, and did not vary with depth (Fig. 5G). Depth integrated SRR down to 10 cm depth was 10 times higher at M1 (4.3 mmol m<sup>-2</sup> d<sup>-1</sup>) than at D3 (0.4 mmol m<sup>-2</sup> d<sup>-1</sup>) (Table 3).

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# 375 **3.5 DIC production rates**

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Vertical profiles of the DIC production rate that were derived from the linear regression of the DIC production measured in anoxic bag incubation (Fig. 4) after correcting for CaCO<sub>3</sub> precipitation, are presented in Fig. 5C and 5H for M1 and D3, respectively. At M1, the DIC production rates decreased with depth from 280 nmol cm<sup>-3</sup> d<sup>-1</sup> (0–2 cm depth) to 69 nmol cm<sup>-3</sup>  $d^{-1}$  (8–10 cm depth) (Fig. 5C), whereas the DIC production rates at D3 were relatively similar across the upper 6 cm ranging from 86 to 136 nmol cm<sup>-3</sup> d<sup>-1</sup>, and decreased to 8–15 nmol cm<sup>-3</sup> d<sup>-1</sup> at 6–10 cm (Fig. 5H). The integrated DIC production rate within 10 cm depth of the sediment was twice as high at M1 (14.0 mmol  $m^{-2} d^{-1}$ ) as at the D3 (7.2 mmol  $m^{-2} d^{-1}$ ) (Table 4).

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### 387 **3.6 Rates of Mn and Fe reduction**

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The accumulation of  $Mn^{2+}$  presented evidence that manganese reduction was occurring in the 389 surface sediment (0-6 cm) of D3 (Fig. 4). The manganese reduction rate (MnRR) derived 390 from  $Mn^{2+}$  accumulation with correction for adsorption ranged from 7.5 nmol cm<sup>-3</sup> d<sup>-1</sup> (0–2 391 cm depth) to 198 nmol cm<sup>-3</sup> d<sup>-1</sup> (2–4 cm depth) at D3 (Fig. 5I). In contrast, MnRR at M1 was 392 indiscernible except for low activity (2.2 nmol cm<sup>-3</sup> d<sup>-1</sup>) at 0–2 cm depth (Fig. 5D). Depth 393 integrated MnRR at D3 (8.21 mmol m<sup>-2</sup> d<sup>-1</sup>) was 200 times higher than the MnRR at M1 394  $(0.04 \text{ mmol m}^{-2} \text{ d}^{-1})$  (Table 3). The iron reduction rate (FeRR), derived from Fe(II)<sub>(oxal)</sub> 395 accumulation, at M1 was highest in the 0-2 cm interval (237 nmol cm<sup>-3</sup> d<sup>-1</sup>), and then 396 decreased with depth to 38 nmol cm<sup>-3</sup> d<sup>-1</sup> at 8–10 cm depth (Fig. 5E). In contrast, Fe 397 reduction was not detected in the surface sediment at D3, but increased to its maximum rate 398 of 240 nmol cm<sup>-3</sup> d<sup>-1</sup> at 4–6 cm depth. The FeRR then decreased with depth to 12 nmol cm<sup>-3</sup> 399  $d^{-1}$  at 8–10 cm (Fig. 5J) where a few data points were adopted to derive the line of best-fit 400 regression. Depth integrated total FeRR was slightly higher at M1 (11.4 mmol m<sup>-2</sup> d<sup>-1</sup>) than at 401 D3 (7.53 mmol  $m^{-2} d^{-1}$ ) (Table 3). The ratio of microbial Fe reduction, Fe Red<sub>(microbial)</sub>, to 402 abiotic Fe reduction coupled to sulfide oxidation, Fe Red<sub>(abiotic)</sub>, ranged from 1.14 (8-10 cm at 403 M1) to 52.3 (2–4 cm at D3), which indicated that the Fe reduction at Mn- and Fe oxides rich 404 405 basin site was mostly a microbiologically mediated process (Table 3).

- 406
- 407 4 Discussion
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# 409 4.1 Partitioning of C<sub>org</sub> oxidation in accordance with the distribution of terminal 410 electron acceptors

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One of the most prominent features revealed from the vertical distributions of geochemical constituents at the basin site (D3) was that electron acceptors such as  $O_2$ , nitrate, Mn- and Fe oxides were systematically distributed with discrete zonation according to the order of decreasing energy yield for  $C_{org}$  oxidation (Fig. 5F). Such biogeochemical zones are not sharply separated in most aquatic sediments due to, e.g., sediment heterogeneity and mixing

resulting from bioirrigation, bioturbation, and bottom turbidity currents. The profiles of 417 dissolved and solid phase geochemical constituents in the sediment provide indications as to 418 specific diagenetic reactions prevailing (Froelich et al., 1979). However, reoxidation of 419 reduced inorganic compounds often mask the primary reactions involved in carbon oxidation 420 421 (Sørensen and Jørgensen, 1987; Hines et al., 1991). Together with the discrete geochemical zonation of the electron acceptors, the independently executed metabolic rate measurements 422 (Fig. 5) allowed us to evaluate the relative contribution of each terminal electron-accepting 423 424 pathway with sediment depth.

425 Previous experimental studies that have quantified pathways of anaerobic carbon oxidation in subtidal marine sediments have generally determined the contributions of Mn 426 and Fe reduction indirectly from the difference between rates of DIC production and sulfate 427 reduction converted to carbon equivalents (e.g., Canfield et al., 1993b; Thamdrup and 428 Canfield, 1996; Vandieken et al., 2006). The inferred rates of Mn and Fe reduction were 429 further supported by the depth distribution of metal oxides and patterns of  $Mn^{2+}$  and  $Fe^{2+}$ 430 accumulation in the pore water, but could not be verified because the accumulation of 431 particulate Mn(II) and Fe(II) – which represents the overwhelming fraction of the reduced 432 pools - was not quantified. Here, we combined the indirect approach with independent 433 434 determination of Mn and Fe reduction rates. Thus, we obtained two separate estimates of anaerobic carbon oxidation rates; based on DIC production and on the sum of sulfate, Fe, and 435 436 Mn reduction converted to carbon equivalents, respectively (Table 4). At M1, within the 0–10 cm depth interval, the average ratio between total anaerobic Corg oxidation rate (10.7 mmol C 437  $m^{-2} d^{-1}$ ) and the  $C_{org}$  oxidation from DIC production (14.0 mmol C  $m^{-2} d^{-1}$ ) was 0.77 (Table 4). 438 Similarly, at D3, the average ratio between total anaerobic  $C_{org}$  oxidation (6.79 mmol m<sup>-2</sup> d<sup>-1</sup>) 439 and anaerobic DIC production (7.22 mmol m<sup>-2</sup> d<sup>-1</sup>) was 0.94. There was a good agreement 440 between the two estimates with a ratio of total anaerobic  $C_{org}$  oxidation by Mn + Fe + sulfate: 441 442 DIC production for individual depth intervals of 0.8–1.2 (Table 4) with the exception at the 0-2 cm depth of slope site (M1) where the ratio was slightly lower, 0.66, possibly due to a 443 contribution from the Corg oxidation by nitrate reduction. The similarity of the two estimates 444 across all incubations spanning a range of redox conditions provides confidence in our 445 approach for calculating dissimilatory Mn and Fe reduction rates. Specifically, the good 446 agreement indicates that the underlying assumptions concerning Mn adsorption and reactions 447 of Fe(III) and sulfide are valid as first-order approximations. The general agreement further 448 supports the validity of previous determinations of dissimilatory Mn and Fe reduction rates 449

450 based on the difference between DIC production and  $SO_4^{2-}$  reduction (Canfield et al., 1993a, 451 1993b; Thamdrup et al., 2000; Vandieken et al., 2006; Vandieken et al., 2014).

To elucidate the contribution of sulfate reduction in anaerobic carbon oxidation, the SRRs 452 (Fig. 5B, 5G) were converted to carbon oxidation (Thamdrup and Canfield, 1996), and then 453 compared to the DIC production rates from anoxic bag incubation (Fig. 5C, 5H). At the slope 454 site (M1), the fraction of anaerobic Corg oxidation coupled to sulfate reduction increased with 455 depth from 48 % at 0–2 cm, to 80 % at 8–10 cm (Table 4). Thus, the excess Corg oxidation in 456 the upper layers should be coupled to other electron accepting processes. Indeed, the Corg 457 oxidation by Fe reduction (0.96 mmol  $m^{-2} d^{-1}$ ) accounted for most of the remaining anaerobic 458 Corg oxidation (12-18 % of DIC production) at 0-8 cm depth, consistent with the distribution 459 of Fe(III) decreasing from > 25  $\mu$ mol cm<sup>-3</sup> near the surface (Fig. 6, Table 4). Mn reduction 460 was of minor importance at M1 because of the low content of Mn oxide ( $< 3 \mu mol cm^{-3}$ ). 461 Carbon oxidation coupled to aerobic respiration was estimated to 3.1 mmol  $m^{-2} d^{-1}$ 462 corresponding to 18 % of the total aerobic + anaerobic oxidation, while the contributions of 463 Fe and sulfate reduction to this total were 12 % and 50 %, respectively (Table 4). As 464 mentioned above, nitrate reduction/denitrification may contribute part of the unexplained 19 % 465 of carbon oxidation, but most of this imbalance likely reflects the combined uncertainties in 466 467 the estimates of the individual pathways. Additionally, our partitioning of carbon oxidation pathways could be biased towards the anaerobic electron acceptors due to the use of the 468 469 diffusive oxygen uptake (DOU) rather than total oxygen uptake (TOU), which will exceed DOU if bioirrigation is active (Glud, 2008). Bioirrigation was not determined at our sites, but 470 471 the pore water profiles show no indication of strong irrigation (Fig. 2). An average DOU/TOU ratio of ~0.6 has been reported for sediments at 1.5-2.5 km depth (Glud, 2008). 472 473 Using this ratio, and assuming that TOU is partitioned similarly as DOU between aerobic carbon oxidation and reoxidation, aerobic carbon oxidation would account for 25 %, while Fe 474 475 and sulfate reduction would account for 11 % and 46 % of of carbon oxidation, respectively. This, the potential bias from using DOU is not expected to affect the ranking of electron 476 acceptors by quantitative importance (SO<sub>4</sub><sup>2-</sup> >  $O_2$  > Fe(III)), and, as discussed further below, 477 the partitioning of Corg oxidation at M1 falls within the range previously reported for 478 continental margin sediments. 479

In contrast to M1,  $C_{org}$  oxidation by sulfate reduction at the basin site (D3) accounted for only a small fraction (< 11 %) of anaerobic  $C_{org}$  oxidation at 0–6 cm interval and it only dominated carbon oxidation at 8–10 cm (Fig. 5H, Table 4). Oxygen and NO<sub>3</sub><sup>-</sup> were depleted

within 3.6 mm and 1 cm depth of the sediment surface, respectively (Fig. 5F), while Mn and 483 Fe(III) oxides were abundant at 0-4 cm and 0-6 cm, respectively. Consistent with the 484 abundance of electron acceptors, high rates of Mn and Fe reduction (Fig. 5I and 5J) implied 485 Mn and Fe reduction as the most significant Corg oxidation pathways to 6 cm depth. At 0-2 486 cm depth,  $C_{org}$  oxidation by aerobic respiration and Mn reduction accounted for 53 % and 43 % 487 of total  $C_{\text{org}}$  oxidation, respectively (Fig. 6). At 2–4 cm, Mn reduction accounted for 73 % of 488 total Corg oxidation and 92 % of anaerobic Corg oxidation (Table 4, Fig. 6). Its importance 489 decreased to 22 % at 4-6 cm due to lower Mn contents, while microbial Fe(III) reduction 490 491 here contributed 51 %, and the partitioning of sulfate reduction increased to 11 % (Fig. 6). Consequently, the relative distribution of each Corg oxidation pathway with depth at D3 (Fig. 6) 492 matched well with the depth distribution of the respective electron acceptors (Fig. 5F). 493 Overall, within the 10 cm depth sediment interval, Mn and Fe reduction were the dominant 494 Corg oxidation pathways comprising 45 % and 20 % of total carbon oxidation, respectively, at 495 the Mn and Fe oxide-rich site in the center of the UB (Table 4). Correction for a potential 496 underestimation of TOU, as discussed for M1, would reduce the contributions of Mn and Fe 497 reduction slightly to 41 % and 18 %, respectively. 498

499 Despite the high Fe oxide content at 0-4 cm at D3 (Fig. 5F), no solid Fe(II)(oxal) 500 accumulation was observed at this depth range (Fig. 4). This indicates that Fe(III) reduction may not occur under these Mn-oxide rich conditions. Indeed, using acombination of 16S 501 502 rRNA-stable isotope probing and geochemical analysis in three manganese oxides-rich sediments including the UB, Vandieken et al. (2012) identified bacteria related to Colwellia, 503 504 Oceanospillaceae and Arcobacter as acetate-oxidizing bacteria that potentially reduce manganese, whereas no known iron reducers were detected in the Mn-rich sediment. 505 506 Similarly, Thamdrup et al. (2000), in Mn oxide- rich Black Sea sediment, found that the 507 abundance of viable Fe-reducing bacteria in most probable number counts was low in 508 comparison to Mn reducers and the addition of ferrihydrite did not stimulate Fe reduction, which implied that Fe reduction should be outcompeted by the Mn reduction process. 509

As manganese reduction is thermodynamically more favorable than iron and sulfate reduction, the  $Mn^{2+}$  liberation (Fig. 4) is likely resulted from dissimilatory Mn reduction. Nonetheless, Mn reduction estimated from the increase of  $Mn^{2+}$  at 0–4 cm interval at D3 (Fig. 4) could be due to oxidation of Fe<sup>2+</sup> or sulfide. Fe<sup>2+</sup> may readily react with Mn oxides (Myers and Nealson, 1988; Lovley and Phillips, 1988) by the reaction  $2Fe^{2+} + MnO_2 + 4H_2O = Mn^{2+}$  $+ 2Fe(OH)_3 + 2H^+$ . However, in the Mn oxide-rich sediment of the Skagerrak, Canfield et al. 516 (1993b) revealed that the addition of Ferrozine, a strong complexation agent for  $Fe^{2+}$ , had no 517 inhibitory effect on the  $Mn^{2+}$  liberation, indicating that the chemical reaction of  $MnO_2$  with 518  $Fe^{2+}$  generated by Fe reduction was not responsible for the accumulation of  $Mn^{2+}$ .

Despite the anoxic conditions and nitrate depletion during the bag incubation, Mn 519 reduction rates at 0-2 cm depth (Fig. 5I) based on Mn<sup>2+</sup> accumulation were substantially 520 lower than the rates inferred from DIC accumulation (Fig. 5H). A similar discrepancy was 521 522 previously observed for the uppermost part of the Mn reduction zone (Thamdrup et al., 2000), and is likely explained by particularly strong sorption of  $Mn^{2+}$  to fresh Mn oxide surfaces. 523 which is not included in the adsorption coefficient used here. Low Mn<sup>2+</sup> together with the 524 rapid decrease of nitrate at 0-2 cm depth at D3 (Fig. 2F, 2G) also suggested that dissolved 525 reduced manganese might act as a reducing agent for nitrate as it was suggested by Aller et al. 526 (1998) in the Panama Basin and Mogollón et al. (2016) in the deep-sea sediment of the 527 Clarion-Clierton fracture zone in the northeast equatirial Pacific. 528

Previous estimation of denitrification in 0-2 cm depth of the UB ranged from 0.01 to 0.17 529 mmol N m<sup>-2</sup> d<sup>-1</sup> (Lee, 2009), which is equivalent to a  $C_{org}$  oxidation of 0.013–0.213 mmol C 530  $m^{-2} d^{-1}$  using the stoichiometric equation of  $4H^+ + 5CH_2O + 4NO_3^- = 5CO_2 + 2N_2 + 7H_2O$ . 531 Based on the average, the contribution of carbon oxidation by denitrification (0.11 mmol C 532  $m^{\text{-2}}~d^{\text{-1}})$  should be minor at the basin site (≤ 3 % of total  $C_{\text{org}}$  oxidation at 0–2 cm; ~1 % of 533 integrated  $C_{\text{org}}$  oxidation). This is consistent with the general consensus that  $C_{\text{org}}$  oxidation by 534 535 denitrification is of little importance in most marine sediments (Sørensen et al., 1979; Canfield et al., 1993a; Trimmer and Engström, 2011). Denitrification may be even further 536 537 suppressed in Mn-rich sediments due to competitive inhibition from Mn reduction (Trimmer et al., 2013). 538

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# 540 **4.2** C<sub>org</sub> oxidation dominated by manganese reduction in the UB

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Microbial Fe reduction has been quantified directly in sediments of various coastal oceans (Gribsholt et al., 2003; Kostka et al., 2002a, 2002b; Hyun et al., 2007, 2009b) and indirectly in deeper continental margins (Thamdrup and Canfield, 1996; Jensen et al., 2003; Kostka et al., 1999). Earlier estimation from 16 different continental margin sediments indicated that Fe(III) reduction contributed 22 % on average to anaerobic carbon oxidation (Thamdrup, 2000). Thus, the contributions from Fe(III) reduction of 12 % and 20 % of anaerobic  $C_{org}$ oxidation onthe slope (M1) and in the basin (D3) of the UB (Table 4) fall in the range of the 549 previous indirect estimates.

Unlike Fe reduction, direct estimation of manganese reduction rates is not easy, mainly 550 because of the restriction of the process to a thin surface layer (Sundby and Silverberg, 1985), 551 the rapid reduction of manganese oxides with  $H_2S$  and  $Fe^{2+}$  (Postma, 1985; Burdige and 552 Nealson, 1986; Kostka et al., 1995; Lovley and Phillips, 1988), and the adsorption of Mn<sup>2+</sup> to 553 Mn oxide surface (Canfield et al., 1993b). For that reason, only two studies, from the 554 Skagerrak and Black Sea, are available for direct comparison on the partitioning of Mn 555 reduction. The process has also been indicated to be of importance in the Panama Basin based 556 557 on diagenetic modeling (Aller, 1990) and at some Artic shelf sites where it was however not quantified separately from Fe reduction (Vandieken et al., 2006; Nickel et al., 2008). Mn 558 reduction was responsible for over 90 % of total Corg oxidationat 600 m depth in the 559 Skagerrak (Canfield et al., 1993b), and accounted for 13-45 % of anaerobic Corg oxidation in 560 the Black Sea shelf sites at 60-130 m of water depth (Thamdrup et al., 2000). To our 561 knowledge, this report of Corg oxidation dominated by Mn reduction comprising 45 % of total 562 Corg oxidation and 57 % of anaerobic Corg respiration in the center of the UB (Table 4) 563 represents the first from deep-offshore basin of the eastern Asian marginal seas. 564

The difference in partitioning of Mn reduction in Corg oxidation between the UB, Black 565 566 Sea and Skagerrak reflects the close relationship between Mn oxide content in the sediment and Mn reduction (Thamdrup et al., 2000). From the vertical distribution of electron 567 568 acceptors (Fig. 5F) and contribution of each Corg oxidation pathway with depth (Fig. 6), it is apparent that the availability of Mn(IV) largely controls the relative contribution to C 569 570 oxidation. In the Skagerrak, the Mn oxides are abundant in high content down to 10 cm depth (Canfield et al., 1993b), whereas Mn oxides in the Black Sea and the Ulleung Basin were 571 572 enriched only down to 2 cm and 4 cm, respectively (Thamdrup et al., 2000; Fig. 2G). Using the available data set for the three marine sediments, we further plotted the relative 573 574 contribution of manganese reduction to anaerobic carbon oxidation as a function of Mnoxides content to expand data from Thamdrup et al., 2000 (Fig. 7). The plot indicates 575 saturation kinetics with a close correlation between Mn oxide content and the importance of 576 Mn reduction at low contents. Curve-fitting yields a content of MnO<sub>2</sub> at 50 % of contribution 577 of manganese reduction to total  $C_{org}$  oxidation (K<sub>s</sub>) of 8.6 µmol cm<sup>-3</sup> similar to the approx. 10 578 µmol cm<sup>-3</sup> suggested before (Thamdrup et al., 2000). This indicates that Mn reduction can be 579 a dominant Corg oxidation process even at low contents of Mn oxides compared to those 580 found at UB. Manganese enrichments of this magnitude have been reported for several 581

locations on the continental margins and in deep basins (Murray et al., 1984; Gingele and Kasten, 1994; Gobeil et al., 1997; Haese et al., 2000; Mouret et al., 2009; Magen et al., 2011; Macdonald and Gobeil, 2012; Mewes et al., 2014) in addition to the relatively few places where dissimilatory Mn reduction was already indicated to be of importance, as discussed above. Thus, the process may be of more widespread significance, particularly in deep basin settings such as UB that allow geochemical focusing of manganese.

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# 589 **4.3 Source of high Mn oxide content**

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The strong enrichment of Mn in the UB surface sediment is primarily of diagenetic origin as 591 indicated by just slightly higher Mn contents at depth in the sediment at D3 (mean 1.1 µmol 592 cm<sup>-3</sup> at 10–20 cm depth) compared to M1 (0.45 µmol cm<sup>-3</sup>) (Fig. 2) combined with higher 593 sediment accumulation rates at the slope  $(0.15-0.3 \text{ cm y}^{-1})$  than in the basin  $(0.07 \text{ cm y}^{-1})$ ; Cha 594 et al., 2005). Thus, the burial flux of Mn, and thereby the net input assuming steady state 595 deposition, is similar or higher at M1 compared to D3. Furthermore, Mn is likely subject to 596 geochemical focusing in the basin as Mn depositing at shallower depths is reductively 597 mobilized and incompletely oxidized in the thin oxic surface layer, resulting in release to the 598 599 water column and net down-slope transport, as inferred in other ventilated basins (Sundby and Silverberg, 1985; Canfield at al., 1993b; Schaller and Wehrli, 1997). A diagenetic source 600 of Mn enrichment was also concluded in previous studies (Yin et al., 1989; Cha et al., 2007; 601 Choi et al., 2009). The Mn remaining and being buried at M1 likely represents unreactive 602 detrital forms to a larger extent than at D3 (Cha et al., 2007). Adopting the sediment 603 accumulation rate of 0.07 cm y<sup>-1</sup> in the UB determined at a station 50 km from D3 (Cha et al., 604 2005), the average  $Mn_{(DCA)}$  content of 1.1 µmol cm<sup>-3</sup> at 10–20 cm depth (Fig. 2G) 605 corresponds to a flux for permanent burial of 0.002 mmol  $m^{-2} d^{-1}$  or just 0.03 % of the Mn 606 reduction rate (Table 3), i.e., an Mn atom is recycled 3800 times before it finally gets buried 607 - first by stripping from the particles that settle to the seafloor and subsequently, over and 608 over, by reductive dissolution of the Mn oxides that from by reoxidation in the oxic surface 609 layer (or, potentially, in the nitrate zone; Aller et al., 1998; Mogollón et al., 2016). This is a 610 much more extensive recycling than found in the Mn sediment of Skagerrak (130–260 times; 611 Canfield et al., 1993b). The difference results mainly from a much higher burial flux of Mn 612 (as authigenic Mn[II]) in the Skagerrak (~40 µmol cm<sup>-3</sup>; Canfield et al., 1993b). The reason 613 that little, if any, authigenic Mn(II) is buried in the UB is not clear. 614

615 As noted in previous studies (Aller, 1990; Canfield et al., 1993b), high contributions of Mn and Fe reduction to carbon oxidation in off-shore sediments requires physical mixing, 616 which typically occurs through bioturbation. This is also the case for the UB, where the burial 617 flux from the oxic surface layer into the Mn reduction zone corresponded to 0.4 mmol  $m^{-2} d^{-1}$ 618 or 5 % of the Mn reduction rate (213  $\mu$ mol cm<sup>-3</sup> × 0.07 cm y<sup>-1</sup>). Bioturbation has previously 619 been inferred, but not quantified, from <sup>210</sup>Pb profiles in the UB (Cha, 2002), and thin 620 polychaete worms were observed during our sampling. Assuming bioturbation to be a 621 diffusive process, we estimate, in a similar manner as in the previous studies and based on the 622 average gradient in Mn<sub>(DCA)</sub> from 0.5-1 to 7-8 cm, that the Mn reduction rate would be 623 supported at a biodiffusion coefficient of 9.5  $\text{cm}^2 \text{ y}^{-1}$ . This value is 3.6 times lower than the 624 coefficient estimated for the Skagerrak (Canfield et al., 1993b) and consistent with estimates 625 for other sediments with similar deposition rates (Boudreau, 1994). Thus, it is realistic that 626 bioturbation drives Mn cycling in the UB. 627

Meantime, the estimated biodiffusion coefficient of (Db) of 9.5 cm<sup>2</sup> yr<sup>-1</sup> at Site D3 628 corresponds to ~2 % of the molecular diffusion coefficient of oxygen (388 cm<sup>2</sup> yr<sup>-1</sup>). Judging 629 from the absence of major fauna in the UB sediments, the mixing is brought about by small 630 organisms with each individual affecting only a small area relative to the size of our cores, 631 632 and the Db averaging many of these small but frequent events. Therefore, we see no contradiction between the presence of bioturbation and the relatively distinct redox zonation 633 634 at D3 (Fig. 5F). Similarly, Hyacinthe et al. (2001) found that well defined profiles can be observed in both sediments with low and high bioactivity in the Bay of Biscay. 635

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# 637 **4.4 The UB as a biogeochemical hotspot**

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The SRRs measured in this study  $(0.43-4.29 \text{ mmol m}^{-2} \text{ d}^{-1})$  are higher than those measured in 639 productive systems such as the Benguela upwelling system in the Southeast Atlantic (0.14– 640 1.39 mmol m<sup>-2</sup> d<sup>-1</sup>; Ferdelman et al., 1999), and even comparable to those reported at the 641 Chilean (2.7–4.8 mmol  $m^{-2} d^{-1}$ ; Thamdrup and Canfield, 1996) and Peruvian (5.2 mmol  $m^{-2} d^{-1}$ 642 <sup>1</sup>; Fossing, 1990) upwelling system at a similar depth range of 1000–2500 m. The total 643 anaerobic DIC production rates at the slope (14.0 mmol  $m^{-2} d^{-1}$ ) and basin site (7.2 mmol  $m^{-2}$ 644  $d^{-1}$ ) were also comparable to those measured at the same depth range of a Chilean upwelling 645 site  $(9.2-11.6 \text{ mmol m}^{-2} \text{ d}^{-1})$  (Thamdrup and Canfield, 1996). Since rates of benthic carbon 646 oxidation are largely controlled by the supply of organic carbon (Canfield et al., 2005), a high 647

648 organic carbon flux reflected in the high organic carbon content (> 2.5 %, dry wt.) in the sediment of the UB (Table 1) is likely to explain the high metabolic activities. A similar high 649 650 organic carbon content as in the UB is rarely found in deep-sea sediment underlying oxic bottom water at depths exceeding 2000 m, except for a Chilean upwelling site (Lee et al., 651 652 2008). This high organic carbon content in the UB is mainly associated with the combination of enhanced biological production resulting from the formation of coastal upwelling (Hyun et 653 al., 2009a), occurrence of an intrathermocline eddy resulting in the extraordinary subsurface 654 chlorophyll-a maximum (Kim et al., 2012), high organic C accumulation rates exceeding 2 g 655  $C m^{-2} yr^{-1}$  (Lee et al., 2008), and high export production (Kim et al., 2009). In addition to the 656 large vertical sinking flux, the lateral transport of the organic matter along the highly 657 productive southeastern slope of the UB also contributes to the high organic carbon content 658 (Lee et al., 2015). Consequently, high benthic mineralization resulting from the high organic 659 content in the sediment implied that the UB is a biogeochemical hotspot where significant 660 turnover of organic matter and nutrient regeneration occur. 661

Recent oceanographic observations revealed that the gradual deoxygenation and warming 662 of the bottom water of the East Sea over the last 30 years have resulted in an  $\sim 10$  % 663 decrease in dissolved oxygen and  $\sim 0.04$  °C increase in potential temperature (Kim et al., 664 2001; Gamo et al., 2011). Benthic metabolism and respiratory  $C_{\text{org}}$  oxidation coupled to 665 various TEAP in the sediments are largely controlled by the combination of O<sub>2</sub> content, 666 temperature and biological production overlying water column (Canfield et al., 2005). It is 667 thus important to monitor any changes in the rates and partitioning of Corg oxidation to better 668 understand and predict the variations of biogeochemical cycles of carbon, nutrients and 669 670 metals potentially associated with long-term climatic changes in the UB, the biogeochemical hotspot of the East Sea. 671

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### 673 5. Conclusions

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Surface sediments of the Ulleung Basin (UB) in the East Sea are characterized by a high organic carbon content (> 2.5 %, dry wt.), high contents of Fe oxides (up to 100  $\mu$ mol cm<sup>-3</sup>), and very high contents of Mn oxides (> 200  $\mu$ mol cm<sup>-3</sup>). We show that microbial Mn and Fe reduction are the dominant C<sub>org</sub> oxidation pathways, comprising 45 % and 20 % of total C<sub>org</sub> oxidation, respectively. The high Mn content results from highly efficient recycling through reoxidation with very low permanent burial of authigenic Mn(II) phases. The basin

topography may ensure that any Mn<sup>2+</sup> escaping to the overlying water returns to the sediment 681 after reprecipitation. The relative importance of Mn reduction to Corg oxidation displays 682 saturation kinetics with respect to Mn oxide content with a low half-saturation value (8.6 683  $\mu$ mol cm<sup>-3</sup>), which further implies that Mn reduction can be a dominant C<sub>org</sub> oxidation 684 685 process in sediments with lower MnO<sub>2</sub> content, and thereby that the process might be more important in continental margin and deep basin sediments than previously thought. Vertical 686 distributions of the major terminal electron acceptors such as O<sub>2</sub>, nitrate, Mn- and Fe oxides 687 were systematically zonated with discrete sequential depletion according to the order of 688 decreasing energy yield for Corg oxidation, which are not sharply separated in most aquatic 689 sediments due to, e.g., sediment heterogeneity and mixing resulting from bioirrigation, 690 bioturbation, and bottom turbidity currents. High benthic mineralization resulting from the 691 high organic carbon content in the sediment implied that the UB is a biogeochemical hotspot 692 693 where significant turnover of organic matter and nutrient regeneration occur.

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### 695 Author contribution

J-H Hyun as first author and leader of the Korean research group designed the original experiments and conducted most writing; S-H Kim, J-S Mok, and H Cho participated in onboard research activities and analytical processes; V Vandieken participated in onboard research and was actively involved in the discussion of the manuscript; D Lee, as project manager of the EAST-1 program, paid the ship-time and has participated in discussion of the results; B Thamdrup, as leader of the Danish research group, collaborated with J-H Hyun in designing the experiments and writing and discussing the manuscript.

704

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| Environmental nonemeter             | <b>M</b> 1          | D3                    |  |  |
|-------------------------------------|---------------------|-----------------------|--|--|
| Environmental parameter             | (Continental slope) | (Center of the basin) |  |  |
| Latitude                            | 36° 10' N           | 37 <sup>°</sup> 00' N |  |  |
| Longitude                           | 130° 10' E          | 131°00' E             |  |  |
| Water depth (m)                     | 1,453               | 2,154                 |  |  |
| Sediment temperature (°C)           | 1.3                 | 0.6                   |  |  |
| Pore-water salinity (psu)           | 34.2                | 34.8                  |  |  |
| Water content (%)                   | 85 (± 3.1)          | 77 (± 1.8)            |  |  |
| Porosity                            | 0.95 (± 0.03)       | 0.86 (± 0.01)         |  |  |
| Density (g cm <sup>-3</sup> )       | 1.10 (± 0.02)       | 1.12 (± 0.02)         |  |  |
| Total organic carbon (%, dry wt.)   | 3.96 (± 0.27)       | 2.66 (± 0.09)         |  |  |
| Total organic nitrogen (%, dry wt.) | 0.38 (± 0.01)       | 0.35 (± 0.01)         |  |  |

978 Table 1. Environmental settings and sediment characteristics

979 Numbers in parenthesis indicate  $\pm$  1SD of triplicate samples.

# 

# Table 2. Oxygen penetration depth (OPD), diffusive oxygen utilization (DOU) rate and O<sub>2</sub> consumption rate by aerobic respiration and re-oxidation of reduced inorganic compounds (RIC) in the pore water.

| Station | OPD<br>(mm)  | DOU   | $O_2$ consumption (mmol $O_2 \text{ m}^{-2} \text{ d}^{-1}$ ) by |                     |  |  |
|---------|--------------|---|--|---------------------|--|--|
| Station |              | $(\text{mmol O}_2 \text{ m}^{-2} \text{ d}^{-1})$ | Aerobic respiration  | Re-oxidation of RIC |  |  |
| M1      | 3.2 (± 0.20) | 7.12 (± 1.36)                                     | 4.04 (± 2.03)  | 3.07 (± 0.68)       |  |  |
| D3      | 3.6 (± 0.03) | 5.95 (± 0.16)                                     | 2.53 (± 0.72)  | 3.42 (± 0.58)       |  |  |
|         |              |   |  |                     |  |  |

Values represent averages  $\pm 1$ SD (n = 3)

|     | Depth<br>Interval<br>(cm) | SQ. <sup>2-</sup> | Mn                  | <sup>(a)</sup> Total —<br>Fe(III) Red | Fe redu                          | Fe Redaman /                       |                             |
|-----|---------------------------|-------------------|---------------------|---------------------------------------|----------------------------------|------------------------------------|-----------------------------|
| St. |                           | Red               | Red                 |                                       | <sup>(a)</sup> Abiotic<br>Fe Red | <sup>(a)</sup> Microbial<br>Fe Red | Fe Red <sub>(Abiotic)</sub> |
| M1  | 0–2                       | 1.35              | 0.04                | 4.75                                  | 0.90                             | 3.86                               | 4.28                        |
|     | 2–4                       | 1.04              | -                   | 3.02                                  | 0.70                             | 2.33                               | 3.33                        |
|     | 4–6                       | 0.84              | -                   | 1.58                                  | 0.56                             | 1.21                               | 2.16                        |
|     | 6–8                       | 0.54              | -                   | 1.25                                  | 0.36                             | 0.89                               | 2.47                        |
|     | 8–10                      | 0.53              | -                   | 0.77                                  | 0.36                             | 0.41                               | 1.14                        |
|     | Sum (0-10)                | 4.30              | 0.04                | 11.4                                  | 2.88                             | 8.70                               |                             |
| D3  | 0–2                       | 0.06              | <sup>(b)</sup> 3.19 | -                                     | -                                | -                                  | n.a.                        |
|     | 2–4                       | 0.11              | 3.96                | 1.63                                  | 0.07                             | 1.56                               | 22.3                        |
|     | 4–6                       | 0.13              | 1.05                | 4.80                                  | 0.09                             | 4.71                               | 52.3                        |
|     | 6–8                       | 0.06              | 0.01                | 0.86                                  | 0.04                             | 0.83                               | 20.8                        |
|     | 8–10                      | 0.07              | 0.00                | 0.24                                  | 0.05                             | 0.19                               | 3.80                        |
|     | Sum (0-10)                | 0.43              | 8.21                | 7.53                                  | 0.25                             | 7.29                               |                             |

990 Table 3. Depth integrated rates (mmol  $m^{-2} d^{-1}$ ) of Mn reduction, Fe reduction, and sulfate reduction and the partitioning of abiotic and 991 microbial Fe(III) reduction in total Fe(III) reduction with depth.

<sup>(a)</sup>Stoichiometric equations were used to evaluate the relative significance of abiotic and microbial Fe reduction:

993 Abiotic reduction of Fe(III) by sulfide oxidation,  $3H_2S + 2FeOOH = 2FeS + S^\circ + 4H_2O$ ; Microbial Fe(III) reduction = Total Fe(III) reduction – abiotic Fe(III) reduction.

994  $^{(b)}$  back-calculated from the C oxidation by Mn reduction in the 0–2 cm interval in Table 4 using the stoichiometric equation,  $2MnO_2 + CH_2O + H_2O = 2Mn^{2+} + HCO_3^- + 3OH^-$ .

995 '- 'indicates that the process does not occur or is regarded as negligible at the depth interval based on the OPD for aerobic respiration and geochemical profiles or anoxic

bag incubations for Mn(IV) and Fe(III) reduction

997 'n.a.' indicates that data are not available.

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|     | Depth<br>Interval<br>(cm)         | $C_{org}$ oxidation measured by                |  | <sup>(c)</sup> Total C <sub>org</sub> | Anaerobic C <sub>org</sub> oxidation<br>by dissimilatory |                            |   | Total anaerobic   | Total Anaerobic          |
|-----|-----------------------------------|--|--|---------------------------------------|--|----------------------------|---|---|--------------------------|
| St. |                                   | <sup>(a)</sup> DOU<br>(Aerobic<br>respiration) | <sup>(b)</sup> DIC prod.<br>(Anaerobic<br>respiration) | oxidation<br>(DOU + DIC)              | <sup>(d)</sup> Mn<br>Red                                 | <sup>(d,e)</sup> Fe<br>Red | <sup>(d)</sup> SO <sub>4</sub> <sup>2-</sup><br>Red | $(Mn \text{ Red} + \text{Fe Red} + \text{SO}_4^{2-} \text{ Red})$ | Anoxic DIC<br>production |
| M1  | 0–2                               | 3.11   | 5.59   | 8.70                                  | 0.02   | 0.96                       | 2.69  | 3.67  | 0.66                     |
|     | 2–4                               | -  | 3.31   | 3.31                                  | -  | 0.58                       | 2.09  | 2.67  | 0.81                     |
|     | 4–6                               | -  | 2.26   | 2.26                                  | -  | 0.26                       | 1.67  | 1.93  | 0.85                     |
|     | 6–8                               | -  | 1.50   | 1.50                                  | -  | 0.22                       | 1.08  | 1.30  | 0.87                     |
|     | 8–10                              | -  | 1.37   | 1.37                                  | -  | 0.10                       | 1.06  | 1.17  | 0.85                     |
|     | Sum (0–10)                        | 3.11   | 14.0   | 17.1                                  | 0.02   | 2.13                       | 8.59  | 10.7  | 0.77                     |
|     | (% Total C <sub>org</sub> ox)     | (18.1)   | (81.9)   | (100)                                 | (0.13)   | (12.4)                     | (50.1)  | (62.7)  |                          |
|     | (% Anaerobic Corg ox)             |  |  |                                       | (0.16)   | (15.2)                     | (61.2)  |   |                          |
| D3  | 0–2                               | 1.94   | 1.72   | 3.66                                  | <sup>(f)</sup> 1.59                                      | -                          | 0.13  | 1.72  | 1.00                     |
|     | 2–4                               | -  | 2.72   | 2.72                                  | 1.98   | 0.39                       | 0.22  | 2.58  | 0.95                     |
|     | 4–6                               | -  | 2.32   | 2.32                                  | 0.52   | 1.18                       | 0.26  | 1.96  | 0.84                     |
|     | 6–8                               | -  | 0.30   | 0.30                                  | 0.01   | 0.21                       | 0.12  | 0.33  | 1.10                     |
|     | 8-10                              | -  | 0.16   | 0.16                                  | -  | 0.05                       | 0.15  | 0.19  | 1.21                     |
|     | Sum (0–10)                        | 1.94   | 7.22   | 9.2                                   | 4.10   | 1.82                       | 0.86  | 6.79  | 0.94                     |
|     | (% Total C <sub>org</sub> ox)     | (20.6)   | (78.8)   | (100)                                 | (44.8)   | (19.9)                     | (9.41)  | (77.8)  |                          |
|     | (% Anaerobic C <sub>org</sub> ox) |  |  |                                       | (56.8)   | (25.2)                     | (11.9)  |   |                          |

Table 4. Organic carbon ( $C_{org}$ ) oxidation (mmol C m<sup>-2</sup> d<sup>-1</sup>) by each  $C_{org}$  oxidation pathway, and its partitioning in total  $C_{org}$  oxidation (% Total  $C_{ox}$ ) and anaerobic  $C_{org}$  oxidation (% Anaerobic  $C_{org}$  ox) at each depth interval within 10 cm of the sediment. Mn Red, Mn reduction; Fe Red, Fe reduction; and SO<sub>4</sub><sup>2-</sup> Red, sulfate reduction

1001  $^{(a)}$  Aerobic C<sub>org</sub> oxidation rate (= O<sub>2</sub> consumption by aerobic respiration × (106C/138O<sub>2</sub>) calculated using the Redfield ratio; O<sub>2</sub> consumption by aerobic respiration rate (= DOU - re-

1002 oxidation rates) is calculated from Table 2 that is derived from the  $O_2$  micro-profiles in Fig. 2.

<sup>(b)</sup> independently measured from the DIC accumulation rate in anoxic bag incubation experiment in Fig. 4. 1003

1004 <sup>(c)</sup> Total  $C_{org}$  oxidation = aerobic  $C_{org}$  oxidation + anaerobic  $C_{org}$  oxidation

 $\frac{(d)}{C_{\text{org}}} \text{ oxidation} = \text{defore } \frac{C_{\text{org}}}{C_{\text{org}}} \text{ defore } \frac{C_{\text{org}}}{C_{\text{org}}} \text{ defore } \frac{C_{\text{org}}}{C_{\text{oxidation}}} = \text{defore } \frac{C_{\text{oxidation}}}{C_{\text{oxidation}}} = \text{defore } \frac{C_{\text{oxidation}}}{C_{\text{oxidation}}} \text{ defore } \frac{C_{\text{oxidation}}}{C_{\text{oxidation}}} = \text{defore } \frac{C_{\text{oxidat$ 1005

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(e) Dissimilatory Fe(III) reduction = (Total Fe(III) reduction in Fig.5) – (Abiotic Fe(III) reduction coupled to  $H_2S$  oxidation;  $3H_2S + 2FeOOH = 2FeS + S^\circ + 4H_2O$ ) 1007

<sup>(f</sup>back-calculated from: DIC production rate - (C oxidation by  $SO_4^{2-}$  Red and Fe Red). See text for further discussion 1008

1009 '-' indicates that the process does not occur or is regarded as negligible based on the OPD for aerobic respiration and geochemical profiles or anoxic bag incubations for Mn and Fe Red.

1010

# Figure legends

Fig. 1. Sampling stations in the East Sea and pictures showing contrasting colors between surface sediments of the continental slope (M1) and center of the basin (D3)

Fig. 2. Concentrations of dissolved  $NH_4^+$ ,  $NO_3^-$ ,  $Mn^{2+}$  and  $Fe^{2+}$  in pore water and contents of solid phase  $Mn_{(DCA)}$ ,  $Fe(II)_{(oxal)}$ ,  $Fe(III)_{(oxal)}$ , acid volatile sulfur (AVS) and chromium reducible sulfur (CRS) in the sediment at M1 and D3.

Fig. 3. Vertical profiles of  $O_2$ . The slashed area indicates the diffusive boundary layer in the sedimentwater interface. The shaded area indicates that  $O_2$  consumption by aerobic respiration (I and II) and re-oxidation of reduced inorganic compounds (III), respectively.

Fig. 4. Changes of concentrations of DIC,  $Ca^{2+}$  and  $Mn^{2+}$  in pore water and contents of solid phase Fe(II)<sub>(oxal)</sub> during anoxic bag incubations of sediments from 0-2, 2-4, 4-6, and 6-8 cm depth at M1 and D3. Data obtained at 8-10 cm depth interval is not shown.

Fig. 5. Vertical distribution of terminal electron acceptors ( $O_2$ ,  $NO_3^-$ , Mn and Fe) and rates of sulfate reduction measured from whole core analyses, and rates of anaerobic carbon oxidation (DIC production rates), Mn reduction and Fe reduction measured from anoxic bag incubations in Fig. 4.  $C_{org}$  by sulfate reduction in panel C and H was calculated from the stoichiometry of 2:1 of  $C_{org}$  oxidized to sulfate reduced.

Fig. 6. Depth variations of partitioning of each carbon oxidation pathway in total carbon oxidation at M1 and D3

Fig. 7. The relative contribution of Mn reduction to anaerobic carbon oxidation as a function of the content of Mn(DCA) at 3 different sites. BS, Black Sea (Thamdrup et al. 2000); UB, Ulleung Basin (This study); Sk, Skagerrak (Canfield et al. 1993b).

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1 Hyun et al – Figure 2



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14 Hyun et al – Figure 4





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# 1718 Hyun et al. – Figure 5



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