



1 Composition and Vertical Flux of Particulate Organic Matter to the Oxygen Minimum Zone
2 of the Central Baltic Sea: Impact of a sporadic North Sea inflow

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12 Export efficiency.

13

14 **Abstract**

15

16 Sinking particles are the main form to transport photosynthetically fixed carbon from the euphotic
17 zone to the ocean interior. Oxygen (O₂) depletion may improve the efficiency of the biological
18 carbon pump. However, how the lack of O₂ mechanistically enhances particulate organic matter
19 (POM) fluxes is not well understood. In the Baltic Sea, the Gotland Basin (GB) and the Landsort
20 Deep (LD) exhibit permanent bottom-water hypoxia, this is on occasions alleviated by Major
21 Baltic Inflow (MBI), such as the one that occurred in 2014/2015 which oxygenated the bottom
22 waters of the GB (but not of the LD). Here, we investigate the distribution and fluxes of POM in
23 the GB and the LD in June 2015 and how they were affected by the 2015 MBI.

24 Fluxes and composition of sinking particles were different in the GB and the LD. In the GB, POC
25 flux was 18% lower at 40 m than at 180 m. Particulate nitrogen (PN) and Coomassie stainable
26 particles (CSP) fluxes decreased with depth, and particulate organic phosphorous (POP), biogenic
27 silicate (BSi), Chl *a*, and transparent exopolymeric particles (TEP) clearly peaked within the core
28 of the oxygen minimum zone (OMZ), which coincided with a high flux of manganese oxide
29 (MnOx)-like particles. Contrastingly, in the LD, POC, PN, and CSP fluxes decreased 28, 42 and
30 56% respectively from 40 to 180m. POP, BSi and TEP fluxes, however, did not decrease with
31 depth and only a slightly higher flux was measured at 110 m. MnOx-like particle flux was two
32 orders of magnitude higher in the GB relative to the LD.

33 MnOx-like particles formed after the inflow of oxygenated water into the deep GB may form
34 aggregates with POM. Our results suggest, that when the deep waters of GB were oxygenated
35 (2014/2015 North Sea inflow), not only transparent exopolymeric particles, as indicated
36 previously, but also POC, POP, BSi, and Chl *a* may bind to MnOx-like particles. POM associated
37 with MnOx-like particles may accumulate in the redoxcline, where they formed larger particles
38 that eventually sank to the seafloor. We propose that this mechanism would alter the vertical
39 distribution and the flux of POM; and it may contribute to the higher transfer efficiency of POC in
40 the GB. This is consistent with the fact that the OM reaching the seafloor was fresher and less
41 degraded in the GB than in the LD.



42 1. Introduction

43 Understanding the downward flux of organic matter (OM) from the euphotic zone is critical to
44 understand biogeochemical cycles in the ocean. Sinking particles are the primary vehicles for
45 transporting photosynthetically fixed carbon from the surface to the deep ocean (Boyd and Trull,
46 2007; Turner, 2015). It has been suggested that the transfer of particulate organic carbon (POC)
47 from the euphotic zone to the ocean interior is enhanced in oxygen minimum zones (OMZs)
48 (Cavan et al., 2017; Devol and Hartnett, 2001; Engel et al., 2017; Keil et al., 2016; van Mooy et
49 al., 2002). Possible mechanisms explaining the higher POC transfer include: i) the reduction of
50 aggregate fragmentation due to the lower zooplankton abundance within the OMZ (Cavan et al.,
51 2017; Keil et al., 2016); ii) a higher refractory nature of sinking particles (Keil et al., 2016; van
52 Mooy et al., 2002); iii) a decrease in heterotrophic microbial activity due to oxygen limitation
53 (Devol and Hartnett, 2001); and iv) the preferential degradation of nitrogen-rich organic
54 compounds (Kalvelage et al. 2013; Van Mooy et al. 2002, Engel et al. 2017). However,
55 mechanisms of how low O₂ concentration would affect the composition and fate of sinking OM,
56 and the efficiency of the biologic carbon pump in oxygen deficient basins have hardly been
57 investigated.

58 The semi-enclosed, brackish Baltic Sea is a unique environment with strong natural gradients of
59 salinity and temperature (Kullenberg and Jacobsen, 1981), primary productivity, nutrients
60 (Andersen et al., 2017), and O₂ concentrations (Carstensen et al., 2014a). New production,
61 defined as the fraction of the autotrophic production supported by allochthonous sources of
62 nitrogen (Dugdale and Goering, 1967) is considered equivalent to the particulate OM export
63 (Eppley and Peterson, 1979; Legendre and Gosselin, 1989) on appropriate timescales. In the
64 Baltic Sea, new production varies seasonally (Thomas and Schneider, 1999); spring and summer
65 are periods of elevated new production supported by the diatom-dominated spring bloom and by
66 diazotrophic cyanobacteria, respectively (Wasmund and Uhlig, 2003). Based on sediment trap
67 data, collected at 140 m in the Gotland Basin, Struck et al. (2004) reported that the highest fluxes
68 of POC occur in fall, followed by summer and spring. Using $\delta^{15}\text{N}$ they showed that during the



69 summer, N₂ fixation by diazotrophic species was the primary source (~41%) of the exported
70 nitrogen, and that the majority of the particulate OM sedimenting in the central Baltic Sea is of
71 pelagic origin.

72 OM export from the euphotic zone to the seafloor has a dual significance in the deep basins of the
73 Baltic Sea. On the one hand, it contributes to the long-term burial of POC, and consequently to
74 the removal and long term storage of CO₂ from surface waters (Emeis et al., 2000; Leipe et al.,
75 2011); on the other hand, it connects the pelagic and the benthic systems contributing to the
76 oxygen consumption and hence deoxygenation at depth. Environmental and anthropogenic
77 changes may alter the magnitude and composition of OM transferred from the surface to the
78 seafloor in the Baltic Sea (Tamelander et al. 2017). On the long term, a decrease in OM
79 downward flux may limit the oxygen depletion. However, to fully suppress hypoxia enhanced
80 ventilation would be necessary the bottom waters of the Baltic Sea.

81 The Gotland Basin (GB) and the Landsort Deep (LD) are the deepest basins of the Baltic Sea.
82 They exhibit permanent bottom-water hypoxia (Conley et al. 2002), caused by a combination of
83 limited water exchange with the North Sea through the Kattegat Strait, strong vertical
84 stratification, and high production /rem mineralization of OM due to eutrophication (Carstensen et
85 al., 2014b; Conley et al., 2009). The Baltic Sea is naturally prone to hypoxia due to physical
86 factors such as permanent salinity stratification and restricted water exchange with the ocean.
87 From the 1950s to 1970s, the hypoxic zones (<60 μmol O₂ kg⁻¹) in the Baltic Sea had expanded
88 fourfold (Carstensen et al. 2014). North Sea inflows are the primary mechanism renewing deep
89 water in the central Baltic Sea. A Major Baltic Inflow (MBI) occurred in 2014/2015 (Mohrholz et
90 al. 2015); this event ventilated bottom waters for five months between February and July 2015
91 (Holtermann et al., 2017). Saltier, denser, O₂-rich North Sea waters entered the western Baltic Sea
92 in December 2014 and reached the Gotland Basin on February 2015. This caused the intrusion of
93 O₂ to deep hypoxic waters, a substantial temperature variability, and high turbidities that may be
94 associated with redox reactions products (Schmale et al., 2016). At the time of sampling, this MBI
95 also affected the neighboring Faroe Deep; but not the LD, located further northwest. At the LD,



96 water properties did not change due to the MBI, the sulfidic layer was maintained (hydrogen
97 sulfide, H₂S concentrations of 20.7- 21.2 μM), and salinity varied between 10.6 and 10.9
98 (Holtermann et al., 2017).

99 In the GB and the LD, a permanent transition zone of about 15 to 20 m thickness separates the
100 surface oxygenated and the anoxic waters. This zone is known as “pelagic redoxcline” and it is
101 only disrupted by sporadic intrusions of saline, well-oxygenated waters from the North Sea
102 (Günter et al., 2008). In the GB, the 2014/2015 MBI oxygenated the deep water column, removed
103 the sulfidic waters in the deeper layers below the redoxcline, and created a secondary near-bottom
104 redoxcline (Schmale et al., 2016). A steep redox gradient characterizes the pelagic redoxcline;
105 here electron acceptors and their reduced counterparts are vertically segregated, and
106 biogeochemical transformations mediated by microbial processes are actively occurring (Bonaglia
107 et al., 2016; Brettar and Rheinheimer, 1991; Neretin et al., 2003). For instance, iron (Fe) and
108 manganese (Mn) undergo rapidly reversible transformations at the redox interface. Under anoxic
109 conditions, these metals are present in dissolved reduced forms Mn(II) and Fe(II); under oxic
110 conditions they form particulate oxides, when react with O₂ or nitrate. Manganese oxides (MnOx)
111 production may be microbially mediated (Neretin et al., 2003; Richardson et al., 1988), or
112 authigenic (Glockzin et al., 2014). The reduction of Mn(IV) with sulfide occurs within a scale of
113 seconds to minutes (Neretin et al., 2003), and is inhibited by nitrate (Dollhopf et al., 2000). The
114 sporadic oxygenation of the deep water of the GB combined with the release of Mn from the
115 sediments into the water column (Lenz et al., 2015) generate appropriate conditions for particulate
116 MnOx formation. MnOx particles have previously been observed in pelagic redoxclines in the
117 Baltic Sea (Glockzin et al., 2014; Neretin et al., 2003). They are amorphous or star-shaped
118 particles that can occur as single particles or form aggregates enriched in OM (Neretin et al.,
119 2003), specifically in transparent exopolymer particles (TEP) (Glockzin et al., 2014). TEP are
120 highly sticky, polysaccharide-rich particles that can enhance aggregation and the formation of
121 marine snow (Engel, 2000; Logan et al., 1995). Thus, MnOx-OM aggregates may significantly
122 contribute to the downward flux of POC. However, TEP are less dense than seawater (Azetsu-



123 Scott and Passow, 2004); therefore they could also reduce the density of marine aggregates and
124 decrease their sinking velocity if the ratio of dense particles to TEP is too small (Azetsu-Scott and
125 Passow, 2004; Engel and Schartau, 1999; Mari et al., 2017). Mixed aggregates containing MnOx
126 and TEP have reported before for the GB and LD (Dellwig et al. 2010; Glockzin et al. 2014).
127 Their sizes ranged between 0.8 and 41 μm equivalent spherical diameter, and their sinking
128 velocity (0.76 m d^{-1}) was lower than what was predicted by the Stokes' law (Glockzin et al.,
129 2014) possibly due to their star-shaped morphology and the high OM content attached to them.
130 Additionally, MnOx aggregates may affect the cycling of particle-reactive elements like
131 phosphorous and trace metals via scavenging processes (Dellwig et al., 2010). To date, there are
132 no measurements of the density of MnOx-OM aggregates, their potential ballast effect of sinking
133 OM, or their effect on the flux of particle-reactive elements in the Baltic Sea.

134 In this study, first, we characterize the amount and composition of particles sinking out of the
135 euphotic zone in two deep basins of the Baltic Sea: the GB and the LD. Second, we compare the
136 sinking fluxes of POM at two stations with different O_2 concentrations below pycnocline (70 m):
137 the GB affected by the MBI that changed the increased the O_2 concentration in the deep waters
138 (between 140 and 220 m) and the LD that was not affected by the MBI and exhibited low O_2
139 concentration and sulfidic conditions in the deep water (from 74 to 430 m). We hypothesize that
140 the different O_2 conditions in the water column of the GB compared with the LD, affected the
141 formation of MnOx rich-aggregates and subsequently OM distribution causing differences in
142 degradation and export of OM between those two stations.

143 **2. Methods**

144 *2.1. Sampling location and water column properties*

145 Samples were collected during the BalticOM cruise in the Baltic Sea onboard the *RV Alkor* from
146 June 3th to June 19th, 2015. We collected sinking particles using surface-tethered sediment traps
147 (Engel et al., 2017; Knauer et al., 1979) in the GB and the LB (Fig.1). Additionally, water column
148 samples were collected using a Niskin-bottle rosette at the locations of the trap deployments.
149 Temperature, salinity and O_2 concentration were determined at each station using a conductivity



150 temperature depth (CTD, Sea-Bird) instrument with an Oxyguard (PreSens) oxygen sensor,
151 calibrated with discrete samples measured using the Winkler method (Strickland and Parsons,
152 1968; Wilhelm, 1888).

153 *2.2. Sediment trap design and deployment*

154 We deployed two surface-tethered sediment traps for two days in the GB, and one day in the LD
155 (Fig.1). Each trap collected particles at four depths between 40 and 180 m (Table 1) to estimate
156 POM fluxes to and within the OMZ. The sediment trap consisted in five arrays of 12 acrylic
157 particle interceptor tubes (PITs) mounted in a PVC cross frame; each tube was equipped with an
158 acrylic baffle at the top to minimize the collection of swimmers (Engel et al., 2017; Knauer et al.,
159 1979). Two particle collector arrays were located at 40 m to estimate the replicability of the
160 system. The PITs were 7 cm in diameter and 53 cm in height with an aspect ratio of 7.5 and a
161 collection area of 0.0038 m². The cross frame and PITs were attached to a line that had a bottom
162 weight and a set of surface and subsurface floats. The procedures for PIT preparation and sample
163 recovery followed Engel et al. (2017). Shortly before deployment, each PIT was filled with 1.5 L
164 of seawater previously filtered through a 0.2 µm pore size cartridge. A preservative solution of
165 saline brine (50 g L⁻¹) was added slowly to each PIT and underneath the 1.5 L of filtered seawater,
166 carefully keeping the density gradient. The PITs were kept capped until deployment and again
167 immediately after recovery to avoid contamination. After recovery, the density gradient was
168 visually verified. Then, the supernatant seawater was siphoned off the PIT, the remaining bottom
169 waters (approx. 0.6 L) containing the particles were pooled together and filled-up to 10L with
170 filtered seawater. After that, the samples were screened with a 500 µm mesh to remove
171 swimmers. Subsequently, samples were split into aliquots that were processed for the different
172 biogeochemical analysis as described in Engel et al. (2017).

173 *2.3. Biogeochemical analysis*

174 Nutrients were measured in unfiltered seawater samples of the deployment stations. Ammonium
175 (detection limit 0.05 µM) was measured directly on board after Solórzano (1969). Phosphate,
176 nitrate, and nitrite (detection limit 0.04 µM) were stored frozen until their analysis; samples were



177 measured photometrically with continuous flow analysis on an auto-analyzer (QuAAtro; Seal
178 Analytical) after Grasshoff et al. (1999).

179 Particulate organic carbon (POC), nitrogen (PN), organic phosphorous (POP), and chlorophyll a
180 (Chl *a*) were determined as described in Engel et al. (2017). Aliquots of 100 to 200 ml of the
181 trapped material, and 500 ml for the sampled seawater were filtered in duplicated for each
182 parameter at low vacuum (<200 mbar), onto pre-combusted GF/F filters (8h at 500°C). After
183 filtration, the filters were stored frozen (-20°C) until analysis. Prior analysis, filters for POC-PN
184 determination were exposed to acid fumes (37% hydrochloric acid) to remove carbonates, and
185 subsequently dried for 12h at 60 °C. POC and PN concentrations were determined using an
186 elemental analyzer (Euro EA, Hechatech) after Sharp (1974).

187 POP was analyzed after Hansen and Koroleff (1999). POP was oxidized to orthophosphate by
188 heating the filters in 40 mL of deionized water (18.2MΩ) with Oxisolv (MERCK 112936) for 30
189 min in a pressure cooker. Orthophosphate was determined spectrophotometrically at 882 nm in a
190 Shimadzu UV-VIS Spectrophotometer UV1201.

191 Chl *a* was analyzed after extraction with 10 mL of 90% acetone, the fluorescence of the samples
192 was measured using a Turner fluorimeter (Turner, 10-AU) according to Strickland et al. (1972).
193 The fluorometer was calibrated with a standard solution of Chl *a* (Sigma-Aldrich C-5753).

194 Phytoplankton composition and abundance in the stations where we deployed sediment traps was
195 characterized microscopically and using a flow cytometer. Phytoplankton, > 5 µm, was counted
196 and identified in 50 ml of fixed samples (Lugol's solution, 1% final concentration) using a Zeiss
197 Axiovert inverted microscope (200x magnification). The size of the counted phytoplankton
198 species ranged from 10 to 200 µm. Phytoplankton, <20 µm, cell abundance was quantified using a
199 flow cytometer (FACSCalibur, Becton, Dickson, Oxford, UK). 2 ml samples were fixed with
200 formaldehyde (1% final concentration) and stored frozen (-80 °C) until analysis (two weeks later).

201 Cell counts were determined with CellQuest software (Becton Dickenson); pico- and
202 nanoplankton populations of naturally containing chlorophyll or phycoerythrin (*i.e.*,
203 *Synechococcus*) were identified and enumerated.



204 Biogenic silica (BSi) was determined by filtering duplicate aliquots of 50 to 100 mL onto 0.4 μ m
205 cellulose acetate filters. Samples were stored at -20°C until analysis. For the measurements, filters
206 were digested in NaOH at 85°C for 135 min; the pH was adjusted to 8 with HCl. Silicate was
207 measured spectrophotometrically according to Hansen and Koroleff (2007).

208 Transparent exopolymeric particles (TEP) and coomassie stainable particles (CSP) from trap and
209 water column were analyzed by microscopy according to Alldredge et al. (1993) and Long and
210 Azam (1996) respectively. Duplicate aliquots of 5 to 20 ml were filtered onto 0.4 μ m Nuclepore
211 membrane filters (Whatmann) and stained with 1 ml of Alcian Blue solution for TEP and
212 Coomassie brilliant blue solution for CSP. Filters were transferred onto Cytoclear® slides and
213 frozen (-20°C) until microscopy analysis. Thirty images for each filter were captured under 200x
214 magnification using a light microscope (Zeiss Axio Scope A.1) connected to a color camera
215 (AxioCam MRc). Particle number and area was measured semi-automatically using WCIF ImageJ
216 software. Image analysis of TEP and CSP were conducted after Engel (2009). Additionally, TEP
217 and CSP in water samples from the stations where we deployed sediment traps were analyzed
218 spectrophotometrically according to Passow and Alldredge (1995) and Cisternas-Novoa et al.
219 (2014) respectively.

220 MnOx-like particle abundance was determined microscopically using the same images that for
221 TEP and CSP determination and a modification of the method described above. Thirty images per
222 filter (200x) were analyzed semi-automatically using Image J. The RGB image was split in three
223 channels: red, blue and green, and the blue, instead of the red channel, was used to quantify the
224 amount of MnOx-like particles in the water column and sediment traps, in this manner the MnOx-
225 like particles were clearly visible with a negligible disruption from TEP or CSP stained blue.

226 Total amino acids (TAA) were analyzed directly in the unfiltered seawater and trapped material.
227 Samples were stored at -20°C until analysis. Duplicate samples were hydrolyzed at 100 °C in 6N
228 HCl (Suprapur® Hydrochloric acid 30%) and 11 mM ascorbic acid for 20 h. Amino acids were
229 separated and measured by high-performance liquid chromatography (HPLC), after derivatization
230 with ortho-phthaldialdehyde using a fluorescence detector (Excitation/Emission 330/445 nm)
231 (Dittmar et al., 2009; Lindroth and Mopper, 1979).



232 Total combined carbohydrates (TCHO) were determined by ion chromatography according to
233 Engel and Händel (2011). TCHO were analyzed directly in the unfiltered seawater and sediment
234 trap material. Samples were stored at -20°C until analysis. Prior to analysis, the samples were
235 desalted by membrane dialysis using dialysis tubes with 1 kDa molecular weight cut-off
236 (Spectra Por). The desalination was conducted for 4.5 h at 1°C . Then, a 2 mL subsample was
237 sealed with 1.6 mL 1M HCl in pre-combusted glass ampoules and hydrolyzed. Samples were
238 hydrolyzed for 20 h at 100°C . After hydrolysis, the subsamples were neutralized by acid
239 evaporation under N_2 atmosphere at 50°C , resuspended with ultrapure Milli-Q water and analyzed
240 by ion chromatography.

241 *2.4 Statistics*

242 A Mann-Whitney U-test was used to test for significant differences between two parameters. The
243 results of statistical analyses were assumed to be significant at p -values < 0.05 . Statistical
244 analyses were performed using Matlab software (MatlabR2014a).

245

246 **3. Results**

247 *3.1. Biogeochemistry of the water column*

248 The water column of both stations was stratified during the study. Deeper in the water column, a
249 pycnocline (halocline) coincided with the oxycline and was located between 63 and 80 m in the
250 GB and between 55 and 75 m in the LD (Fig. 2). Additionally, a seasonal thermocline was located
251 between 20 and 30 m in the GB and between 12 and 38 m in the LD. The GB had a hypoxic layer
252 ($< 60 \mu\text{mol O}_2 \text{ L}^{-1}$) between 74 and 140 m and the core of the OMZ ($< 10 \mu\text{mol O}_2 \text{ L}^{-1}$) was located
253 between 96 and 125 m. The increasing oxygen concentrations at depth ($> 140\text{m}$) can be related to
254 the MBI event in 2014/2015 that ventilated the otherwise anoxic deep layer of the GB
255 (Holtermann et al., 2017) and caused a rise in O_2 concentration from less than $40 \mu\text{mol O}_2 \text{ L}^{-1}$ at
256 140 m to $79 \mu\text{mol O}_2 \text{ L}^{-1}$ at 220 m (Fig. 2a). The MBI, however, did not reach the LD (Fig. 2b),
257 where oxygen was below the detection limit ($< 3 \mu\text{mol O}_2 \text{ L}^{-1}$) from 74 m to the bottom (430 m).
258 The vertical profile of nutrients was different at both stations (Fig. 2). In the GB, nitrate increased
259 from below the detection limit in the surface waters to $0.17 \mu\text{M}$ at 40 m. Nitrate concentrations



260 were variable within the OMZ (6 μM at 80 and 140 m, and 0.12 μM at 110 m). At 220 m nitrate
261 concentration decreased to 4.8 μM (Fig. 2a). Nitrite was below the detection limit in most of the
262 water column except for 60 m (0.09 μM) and 110 m (0.11 μM). Ammonium increased from 0.14
263 μM in the surface to 1.15 μM at 60 m; concentrations were variable in the OMZ with less than
264 0.15 μM at 80 and 140 m, and maximum concentration of 3.28 ± 0.01 μM at 110m. Vertical
265 profiles of phosphate and silicate at the GB were similar; the concentrations steadily increased
266 from the surface (0.29 μM and 10.36 μM respectively) to the OMZ (2.67 μM and 39.07 μM
267 respectively), and gradually decreased below the OMZ (Fig. 2a). Hydrogen sulfide was not
268 detectable in the GB.

269 In the LD, nitrate and nitrite concentrations were below the detection limit between the surface
270 and 250 m (<0.06 μM) (Fig. 2b). Ammonium concentrations varied between 0.06 and 0.59 μM in
271 the upper 70 m and increased to 5.97 and 8.03 μM in the OMZ. The lowest concentration
272 (0.07 μM) was measured in the surface and maximum concentration of 8.03 μM at 110 m.
273 Phosphate and silicate concentrations varied between 1.58 ± 0.04 (at 40 m) and 2.18 ± 0.80 μM
274 phosphate and between 21.75 ± 4.78 (at 40 m) and 35.67 ± 14.59 μM silicate in the upper 110 m
275 of the water column; lowest concentrations were measured at 180 m (0.22 μM and 7.4 μM
276 respectively). Highest concentrations of nitrate and nitrite (6.01 and 0.22 μM) were observed at
277 400 and 350 m, respectively. Hydrogen sulfide was measurable below 180 m, with the highest
278 concentration (3.97 μM) measured at 250 m and the lowest (0.04 μM) between 300 and 350 m
279 (Fig. 2b).

280 3.2. Particulate organic matter concentration in the water column

281 Chl *a* concentration in the upper 10 m was slightly higher in the GB (1.5-1.7 $\mu\text{g L}^{-1}$, Fig. 3b) than
282 in LD (1.4-1.2 $\mu\text{g L}^{-1}$ and 0.1-0.3 μM , Fig. 3e). This agrees with estimates of integrated total
283 primary production, which were 380 $\text{mg C m}^{-2} \text{d}^{-1}$ in the GB and 334 $\text{mg C m}^{-2} \text{d}^{-1}$ in the LD
284 (Piontek et al., unpublished). Pico- (<2 μm) and nanophytoplankton (2-20 μm) abundances, as
285 determined by flow cytometry, were higher in the upper 60 m, although detectable in the entire



286 water column. Pico- and nanophytoplankton abundances were 10% higher in GB than in LD
287 (Table 2). Phycoerythrin fluorescence, mainly from picophytoplankton (92% in GB and 96% in
288 LD), was 30% higher in GB than in LD.

289 The abundance of larger phytoplankton (>5 μm) was determined by microscopy. Microscopic
290 counts of cells showed about 63% higher phytoplankton abundance in the LD than in the GB
291 (Table 3). Filamentous, unicellular cyanobacteria dominated the large phytoplankton community
292 at both stations with up to 90% corresponding to *Aphanizomenon* sp. Cyanobacteria were 60%
293 less abundant in the GB than in the LD (Table 3). They represented 56% of the total
294 phytoplankton counts in the GB and up to 74% in the LD. Dinoflagellates (dominated by
295 *Dinophysis* sp.) were significant in both stations (19%), whereas chlorophytes (dominated by
296 filaments of *Planctonema* sp. containing cylindrical cells) were more abundant in the GB than in
297 LD (25% and 4% of the total respectively). Diatoms represented less than 1% of the
298 phytoplankton in both stations, and they were slightly more abundant at 40 m in the LD (Table 3).

299 BSi was higher in the upper 10 m (0.4-0.5 μM) and decreased with depth in the GB (Fig. 3b),
300 whereas in the LD, BSi showed a peak at 40 m and then decreased with depth (Fig. 3f).

301 Vertical profiles of POC, PN, and POP concentration were similar in the water column of the two
302 stations (Fig. 3a, d). In the GB, the concentrations were higher in the surface (POC: 40.38 ± 0.80 ,
303 PN: 3.89 ± 0.01 , and POP: $0.26 \pm 0.04 \mu\text{M}$) and decreased gradually with depth until 110 m where
304 relatively high concentrations (POC 18 ± 0.63 , PN: 2 ± 0.08 , and POP: $0.2 \mu\text{M}$) were observed.

305 The lowest concentrations were found at 180 m (POC: 11.97 ± 1.03 , PN: 1.05 ± 0.02 , and POP
306 $<0.03 \mu\text{M}$) (Fig. 3a). In the LD, POM decreased with depth from the surface (POC: 35 ± 0.99 ,
307 PN: 4 ± 0.09 , and POP: $0.2 \mu\text{M}$) to 40 m, remained relatively constant between 40 and 80 m and
308 decreased again between 110 and 250 m (Fig. 3d).

309 We observed high concentrations of TEP and CSP in the upper 10 m in both stations. The highest
310 TEP concentration was measured at 1 and 10 m at both stations, and it was slightly higher (19%)
311 in the GB than in the LD (Fig. 3c, f). TEP and CSP vertical profiles were different from each
312 other in the GB (Fig. 3c) and covaried in the LD (Fig. 3f). Like observed for POC, PN, and POP,



313 TEP concentrations showed a peak at 110 m ($50.29 \pm 6.17 \mu\text{g XG eq. L}^{-1}$) in the GB. The highest
314 concentration of CSP at this station was observed in the shallowest (1 m) sample, CSP
315 concentration decreased quickly at 10 m, and then it increased at 140 and 230 m (the deepest
316 sample ~20 m above the seafloor) (Fig. 3c). In the LD, the highest concentrations of TEP and
317 CSP were measured in surface (1 and 10 m) and at 110 m (Fig. 3f). TEP and CSP decreased with
318 depth in the first 80 m (from 53.26 ± 7.10 to $18.39 \pm 4.57 \mu\text{g XG eq. L}^{-1}$ and from 53.26 ± 7.10 to
319 $31.57 \pm 18.78 \mu\text{g BSA eq. L}^{-1}$). Both types of gel-like particles showed an increase in
320 concentration at 110 m ($49.25 \pm 4.08 \mu\text{g XG eq. L}^{-1}$ and $66.89 \pm 22.33 \mu\text{g BSA eq. L}^{-1}$
321 respectively). Below 110 m, TEP concentrations stayed relatively constant, while CSP
322 concentrations decreased at 180 m and kept relatively constant below that depth.

323 *3.3. MnOx-like particles vertical distribution in the water column*

324 Dark, star-shaped, MnOx-like particles (Glockzin et al., 2014; Neretin et al., 2003) were observed
325 below the fully oxygenated mixed layer in the GB and, in less abundance, in the LD (Fig. 4). In
326 GB, single MnOx-like particle and large aggregates were observed from 80 m to 220 m (the
327 deepest sample, approximately 28 m above the seafloor). Relatively high concentration of MnOx-
328 like particles (2×10^6 particles L^{-1}), were measured in the upper (80 m) and lower (140 m) oxycline
329 where the O_2 concentration was less than $40 \mu\text{M}$, and at 220 m (4×10^6 particles L^{-1}) (Fig. 4a). The
330 lowest abundance of MnOx-like particles (7×10^5 particles L^{-1}) was observed at 110 m, in the core
331 of the OMZ where the O_2 concentration was less than $10 \mu\text{M}$. The ESD varied between 0.6 and
332 $30.5 \mu\text{m}$ and the median was $3.0 \mu\text{m}$. The largest aggregates were observed in the upper oxycline
333 (80 m). In the LD, MnOx-like particles were less abundant, smaller and had a narrow distribution
334 in the water column than in the GB. MnOx-like particles were not detected in the fully oxic (0–40
335 m) or fully anoxic (180 to 430 m) water column. At 60 m, right above the oxycline, MnOx-like
336 particles began to appear, however, in relatively low abundance. The maximum abundance, 9×10^5
337 particles L^{-1} , was observed in the oxycline at 70 m (Fig. 4b). The ESD varied ranged between 0.6
338 and $13.4 \mu\text{m}$, the largest aggregates were observed at 70 m.



339 *3.4. Fluxes of Particulate Organic Matter*

340 Fluxes of particulate organic matter varied little with depth in the GB (Fig. 5a-c). POC flux
341 slightly increased by 18% from the shallowest (40 m) to the deepest (180 m) depth. Fluxes of PN
342 and CSP were higher at 40 and 60 m and decreased by 19 and 70% from 60 to 180 m,
343 respectively (Fig. 5a and 5c). On the other hand, fluxes of POP, BSi, Chl *a* (Fig. 5b) and TEP
344 (Fig. 6a) peaked at 110 m. Those fluxes increased by 68, 61, 44 and 68% respectively from 40 m
345 to 110 m; then they decreased by 22, 65, 27 and 19% from 110 m to 180 m. This increment of
346 fluxes at 110 m coincided with the presence of abundant MnOx-like particles associated with TEP
347 (Fig. 6a). In addition, TEP size distribution, determined by image analysis, indicated an increase
348 in large TEP at 110 m (data not shown). In contrast, in the LD, POC, PN (Fig. 5d) and CSP (Fig.
349 6d) fluxes decreased with depth. Fluxes were 28, 42 and 56% less at 180 than at 40 m. However,
350 the POP, BSi (Fig. 5e) and TEP (Fig. 6c) showed highest fluxes at 110 m.

351 MnOx-like particles were drastically less abundant in sediment trap samples from the LD than in
352 the GB and when present, only as single particles not as aggregates with TEP or CSP (Fig. 6c, d).
353 The flux of MnOx-like particles at 110 and 180 m was two orders of magnitude larger in the GB
354 than in the LD (Table 4). At both stations, and similar to the water column, MnOx-like particles
355 were not observed in sediment trap samples collected at 40 and 60 m. In the GB, MnOx-like
356 particles were present in the sediment traps at 110 m and 180 m. MnOx-like were as single
357 particles and forming aggregates with each other and other particles such as: TEP (Figure 6a, f),
358 phytoplankton cells, or detrital material. The ESD of MnOx-like particles and aggregates ranged
359 from 0.6 to 167 μm (median 2.8 μm) at 110 m and from 0.6 to 153 μm (median 3.3 μm) at 180 m.
360 In the LD, only a few, single MnOx-like particles were observed at 110 and 180 m and their size
361 ranged from 0.6 to 16.5 μm (media 1.8) at 110 m (Table 4).

362 TAA flux ranged from 371 ± 12 to $501 \pm 33 \mu\text{mol m}^{-2}\text{d}^{-1}$ in the GB and from 502 ± 84 to 785 ± 54
363 $\mu\text{mol m}^{-2}\text{d}^{-1}$ in the LD (Fig. 7a). In the GB, the flux decreased with depth whereas, in the LD, the
364 TAA flux at 40 m was lower than at 60 m and decreased with depth from 60 to 180 m (Fig. 7b).

365 The TCHO flux varied between 303 ± 8 and $428 \pm 14 \mu\text{mol m}^{-2}\text{d}^{-1}$ in the GB (Fig. 7a) and between



366 503 ± 19 and $584 \pm 8 \mu\text{mol m}^{-2}\text{d}^{-1}$ in the LD (Fig. 7b). Vertical profile of TCHO flux was similar in
367 both stations. TCHO flux increased from 40 to 110 m, where the highest TCHO flux was
368 measured, and then TCHO flux decreased at 180 m. The TCHO flux at 180 m was 22% higher
369 than at 40 m in the GB, and the same that at 40 m in the LD.

370 3.5. Chemical composition of sinking and suspended OM

371 Elemental ratios for sinking and suspended OM in the GB and the LD are shown in Table 5. The
372 POC:PN ratio of the sinking OM increased with depth at both stations. In suspended OM, this
373 ratio was more variable in the GB and decreased with depth in LD. The POC:PN molar ratio of
374 suspended and sinking OM may be compared to the classical Redfield ratio for living plankton
375 which is 106: 16: 1 for C:N:P (Redfield et al., 1963). Sinking OM was slightly above Redfield's at
376 both stations. The POC:PN ratios of the sinking OM in both GB and LD were not significantly
377 different. Contrastingly, in the suspended OM, POC:PN ratios were higher in the GB compared to
378 the LD ($p < 0.001$; Mann–Whitney U-test). In the LD the POC:PN of sinking OM was significantly
379 lower than in suspended OM ($p < 0.001$).

380 The POC:POP molar ratio of sinking OM was lower ($p < 0.05$) in the GB than in the LD; and it
381 was higher ($p < 0.01$) in sinking than in suspended OM in the LD (Table 5). The POC:BSi molar
382 ratio was lower in sinking than in suspended OM in both stations (GB: $p < 0.05$; LD: $p < 0.01$). In
383 suspended OM, the POC:BSi ratio was above Redfield ratio, whereas in sinking OM it was below
384 Redfield value (Table 5). The PN:POP molar ratio was lower in sinking OM than in suspended
385 OM in both stations ($p < 0.001$). In sinking OM this value was always below the Redfield ratio,
386 while in suspended OM, it was always above the Redfield ratio.

387 At both stations, the fraction of sinking POC composed of AA was larger than in suspended OM.
388 Similarly, the C contained in CHO made up a larger percentage in sinking OM than in suspended
389 OM (Table 5).

390 The amino acid-based degradation index (DI, Dauwe et al., 1999) in sinking OM varied from 0.1
391 to 1.14 and was higher than in suspended OM (-1.25 to -0.42). The DI was higher in the GB than
392 in the LD in sinking and suspended OM. In the sinking OM of the GB, the DI decreased with
393 depth but in the LD was more positive at 110m than at 60 m (Table 5).



394 **4. Discussion**

395 In this study, we described the results of: 1) the characterization of the surface biogeochemical
396 conditions and the amount and composition of the particles produced in the euphotic zone of two
397 deep basins in the central Baltic Sea, *i.e.*, the GB and the LD, during early summer 2015, and 2)
398 the flux of sinking particles out of the euphotic zone as well as their variation at depth in the two
399 basins. We assess the potential influence of increased O₂ concentration caused by the 2014/2015
400 MBI in the GB on the chemical composition and degradation stage of the sinking and suspended
401 OM relative to the anoxic LD.

402 *4.1 Characterization of biogeochemical conditions in GB and LD*

403 Temperature, O₂, and inorganic nutrient concentrations were similar in surface at both stations.
404 Moreover, though there were slight differences in biogeochemical conditions, such as primary
405 production, phytoplankton composition and chemical composition of POM, in the surface water
406 column, those were not significant. The concentration of Chl *a* (Fig. 3), the abundance of
407 picophytoplankton, nanophytoplankton (Table 2) and primary production (PP, Piontek et al.
408 unpublished data) were slightly higher (20, 10 and 10 % respectively) in the GB than in the LD.
409 At both stations, phycoerythrin-containing cyanobacteria were a small fraction of the pico- and
410 nano-phytoplankton. Pico-phytoplankton cell abundance (cell mL⁻¹) dominated the small
411 phytoplankton (Table 2), suggesting a significant contribution to PP and Chl *a* concentration.
412 These findings coincide with what was described previously for early summer, in the Baltic Sea
413 that indicate that this period corresponded to a low productivity transition phase characterized by
414 low Chl *a* concentration ($\leq 2 \mu\text{g L}^{-1}$) sustained mostly by nano- and picophytoplankton
415 communities (Leppänen et al., 1995) which co-existed with cyanobacteria and other
416 phytoplankton species (Kreus et al. 2015). Microscopic analysis of larger phytoplankton (>5 μm),
417 on the other hand, showed that filamentous cyanobacteria *Aphanizomenon* sp. (up to 200 μm
418 large) was the dominant type on this size fraction in the upper 40 m (Table 3). *Aphanizomenon* sp.
419 and *Nodularia spumigena*, are known to form summer blooms in the Baltic Sea, where they
420 accumulate at the sea surface of the thermally stratified water column (Bianchi et al., 2000;



421 Nausch et al., 2009; Wasmund, 1997). Cell abundance of total phytoplankton (>5 µm) were not
422 significantly different ($p=0.74$) in the GB and the LD.

423 POC, PN, POP, BSi, TEP and CSP concentrations in the surface waters were also similar at both
424 stations (Fig. 3). The concentration of TEP was higher than of CSP, both types of gel-like
425 particles were most abundant in the euphotic zone indicating a phytoplankton origin. In the
426 surface water column, TEP concentrations (48 and 62 µg X.G. Eq. L⁻¹ in the GB and the LD,
427 respectively) were 69 and 76% lower than the value previously reported for summer in the central
428 Baltic Sea in June (200 µg X.G. Eq. L⁻¹) (Engel et al., 2002). Likewise, our dissolved inorganic
429 nitrogen concentrations were below the detection limit in the surface; however phosphate
430 concentrations were higher (0.2-0.65 µM) than the ones on the Engel et al. (2002) study. Mari and
431 Burd (1998) reported that TEP concentration peaked during the spring bloom and in summer in
432 the Kattegat. TEP production may be enhance by enviromental conditions such as nutrient
433 limitation (Mari et al., 2005; Passow, 2002), which are characteristic of late summer in the Baltic
434 Sea (Mari and Burd 1998). Our samples were collected right after the peak of the spring bloom
435 (Le Moigne et al., 2017), thus, likely TEP concentrations had not reached the usually higher
436 summer value yet since phosphate remained present in the water column (potentially not limiting
437 the PP).Anoter possible explanation for the rather low concentrations of TEP could be that TEP
438 may be removed from the surface by aggregation and subsequent sedimentation during the spring
439 bloom due to the high abundance of cells and detrital particles during this time (Engel et al.,
440 2002).

441 Although the composition and amount of OM in the surface waters at the two trap stations were
442 similar, below the euphotic zone (40 m) the vertical profile of nutrients and POM concentrations
443 were clearly different; likely due to the 2014/2015 MBI (Holtermann et al., 2017) that reached the
444 deep waters of the GB. The MBI changed the vertical distribution and increase the concentration of
445 O₂ in the GB compared with the LD. In the GB the oxygen-deficient zone (O₂ <40 µmol L⁻¹) was
446 constrained between 74 and 140 m and the core of the OMZ (O₂ <10 µmol L⁻¹) between 96 and



447 125 m; below 140 m O₂ concentrations increased <40 μmol L⁻¹. In contrast, the LD maintained
448 permanent suboxic (<5 μmol L⁻¹) waters below 74 m and hydrogen sulfide was detectable at 180
449 and 250 m (Fig. 2). In the GB nitrate concentration increased possibly as a consequence of the
450 oxidation of reduced nitrogen compounds (e.g., ammonium, ammonia and organic nitrogen
451 compound like urea) (Le Moigne et al., 2017) that accumulated during the stagnation (anoxic)
452 period previous to the MBI (Hannig et al., 2007). MBIs can have a major impact on nutrient
453 recycling. For instance, phosphorous could bind to iron hydroxides and MnOx and settle down
454 during oxic conditions, building up a phosphate pool in the sediments that later on when the O₂
455 decreases close to the sediments, it may become a source of phosphate (Gustafsson and
456 Stigebrandt, 2007). In addition to changes in O₂ concentration, the MBI altered the redox
457 conditions in the GB creating a secondary redoxcline at 140 m, where the O₂ and the MnOx-like
458 particles concentration increased (Fig. 4a). One consequence of those changes is the vertical
459 extension of the layer in which MnOx aggregates could form. A previous study showed that
460 MnOx might precipitate from the water column of the GB following a MBI event (Lenz et al.,
461 2015). Scavenging of phosphate into Mn or Fe oxides had been shown in previous studies
462 (Neretin et al., 2003). Moreover, there is a downward flux of phosphate associated to particule
463 iron and MnOx in the oxic water column to the anoxic basin where particles dissolved and
464 phosphate is release (Gustafsson and Stigebrandt, 2007). This process may be responsible for the
465 decrease of phosphate concentration below 110 m in our study (Fig. 2a). In contrast, in the LD,
466 the water column remained suboxic down the sea floor (430 m), below the oxycline an increase of
467 ammonium was observed (Fig.2) which could be an indicator for anaerobic respiration of OM,
468 e.g., denitrification (Bonaglia et al., 2016; Hietanen et al., 2012). Low phosphate and silicate
469 concentrations within the mixed layer due to phytoplankton consumption gradually increased
470 below the pycnocline and decreased between 110 and 180 m.

471 In summary, although the GB and the LD had similar surface conditions in terms of
472 phytoplankton production and POM stocks, during this study, we found differences the vertical
473 concentration of POM (Fig. 3) in the GB, ventilated by the MBI, relative to the LD, a station that



474 remains suboxic. Our results suggest that differences in the vertical profile of O_2 may modify the
475 redox conditions of the water column, enhancing the formation of MnOx-like particles (Fig. 4)
476 that may aggregate with POM in the GB and changed its vertical distribution.

477 *4.2 Potential influence of O_2 concentration and redox conditions on sinking fluxes of POM in the*
478 *GB and the LD*

479 During this study, we also investigated the effect of different O_2 concentrations and redox
480 conditions on the fluxes of particles. Our measurement of carbon flux below the euphotic zone
481 (40 m) were $11.7 \pm 0.82 \text{ mmol C m}^{-2} \text{ d}^{-1}$ in the GB and $19.8 \pm 1.22 \text{ mmol C m}^{-2} \text{ d}^{-1}$ in the LD.
482 Extrapolating those measurements to annual flux we obtain $4.37 \pm 0.31 \text{ mol C m}^{-2} \text{ a}^{-1}$ in the GB and
483 $7.44 \pm 0.46 \text{ mol C m}^{-2} \text{ a}^{-1}$ in the LD. Our results from the LD are compatible with the long-term
484 annual estimations from models that varied between 3.8 to $4.2 \text{ mol C m}^{-2} \text{ d}^{-1}$ (Kreus and Schartau,
485 2015; Sandberg et al., 2000; Stigebrandt, 1991) for the Baltic Sea; however, the estimations based
486 on our results from the GB are higher than the C fluxes predicted by those models.

487 The vertical flux of POM was different the two studied stations; likely due to differences in O_2
488 concentrations that may affect POM remineralization and transport; in the GB, the POC flux
489 between 40 and 180 m showed a small increase while PN slightly decreased from the bellow the
490 oxycline (60 m) to 180 m. In the LD, the POC flux decreased greatly between 40 and 60 m, and
491 remained relatively constant between 60 and 180 m; PN flux, however, decreased with depth. In
492 the GB, and to a lower degree in the LD, we observed a distinctive peak of POP, BSi, Chl *a* and
493 TEP fluxes at 110 m. This high flux of POM coincided with the appearance of dark, star-shaped
494 particles (Fig. 6a, f) which may correspond to MnOx particles enriched in OM that have been
495 described in the GB and the LD before (Neretin et al., 2003; Pohl et al., 2004). We observed a
496 higher concentration of MnOx-like aggregates associated with TEP at 110 m in the GB. The 110
497 m sediment trap was located between the upper (80 m) and lower (140 m) oxycline where the
498 MnOx-like particles are likely formed. This corresponds to the depth range where lowest O_2
499 concentration was measured but hydrogen sulfide (H_2S) was absent in the water column, which



500 allows the presence of those aggregates also at 180 m. On the contrary, in the LD, we measured
501 H₂S at 180 m, this could explain why although those aggregates were present in this station below
502 the oxycline (*i.e.*, 70 m) at 110 m, they dissolved in sulfidic waters, thus were not as abundant,
503 and did not form aggregates with TEP (Fig.6c).

504 The presence of MnOx-containing aggregates enriched in OM (see TEP fluxes, Fig 6c) may have
505 implications for the vertical flux of C and N in a stratified system with a pelagic redoxcline like
506 the Baltic Sea. Under steady state, the upward diffusion and oxidation rate of the dissolved Mn
507 are balanced by the sinking and dissolution rate of MnOx. During the Mn-oxidation, the POM
508 could aggregate with the MnOx including particulate elements, and trace metals. Then, in the
509 sulfidic waters, slow-sinking MnOx enriched in OM will be dissolved liberating the OM and
510 altering the vertical distribution and the flux of all associated particle elements (Glockzin et al.,
511 2014). The precipitation of MnOx could be enhanced by the oxygenation of the otherwise anoxic
512 deep of the Baltic Sea caused by the 2014/105 MBI (Dellwig et al., 2018), those particles could
513 bind with phosphorous and trace metals trapping them in the redoxcline (Dellwig et al., 2010).
514 For example, in the Cariaco Basin, total particulate phosphorous reached their maximum flux in
515 sediment traps close to the redoxcline (Benitez-Nelson et al., 2004; Benitez-Nelson et al., 2007).
516 MnOx formation and scavenging of trace metal may be a relevant mechanism for transfer trace
517 metals from the oxygenated to the anoxic deep waters (Dellwig et al., 2010). Moreover, even in
518 the anoxic zone, the abundant aggregate associated bacteria (Grossart et al., 2006) could partially
519 or completely degrade the organic compounds in those particles using NO₃⁻ or Mn²⁺ as an electron
520 acceptor. This may be the reason why we observed a clear peak in the flux of POP, BSi, Chl *a*
521 (Fig. 3a, b), TEP (Fig. 6a) and TCHO (Fig. 7a) at 110 m followed by a small decrease at 180 m in
522 the GB. In the LD a smaller increment in the flux of POP, BSi (Fig. 3d), TEP (Fig. 6c) and TCHO
523 (Fig. 7b) was also observed. The vertical fluxes of those compounds coincided with the
524 abundance of MnOx particles; we assume that the MnOx aggregated not only with TEP as
525 described before (Glockzin et al. 2014) and observed in this study (Fig. 6a) but also with POP,
526 BSi, Chl *a*, and TCHO. On the other hand, nitrogen-rich compounds like PN (Fig. 3a), TAA (Fig.



527 7a), and CSP (Fig. 6a) gradually decreased with depth in the GB, suggesting that those
528 compounds were less scavenged by MnOx organic-rich aggregates.

529 Primary production (PP) in the GB was 10% higher than in LD during our study (Piontek et al.
530 unpublished data). However, the POC flux below the euphotic zone (at 40 m) was 42% higher in
531 LD than in GB and comparable at both stations at 180 m. The fraction of PP exported as POC is
532 termed export production (*e-ratio*) (Buesseler et al., 1992), and it is calculated as the POC flux
533 below the euphotic zone divided by the primary production. The *e-ratio* was calculated here
534 using the ¹⁴C based PP (Piontek et al. unpublished data) and carbon flux at 40 m (shallowest
535 sediment trap depth, considered at the base of the euphotic zone). The *e-ratio* was 0.41 in the GB
536 and 0.77 in the LD; *i.e.*, in GB 41% of the primary production was exported as POC below the
537 euphotic zone (40 m) versus 77% in the LD). This suggests that a higher proportion of the
538 primary production was remineralized in the euphotic zone of the GB compared with the LD. On
539 the other hand, the transfer efficiency of POC to the deeper water column (*i.e.* the ratio of POC
540 flux at 180 m over POC flux at 40 m) was higher in the GB (115%) than in the LD (69%). The
541 transfer efficiency of POM is largely controlled by the remineralization rate and the sinking
542 velocity of particles (De La Rocha and Passow, 2007; McDonnell et al., 2015; Trull et al., 2008).
543 The higher POC transfer efficiency in the GB than in the LD can be attributable to differences in
544 the sinking velocities of the particles in those two stations. The presence of MnOx-OM rich
545 aggregates in the GB may fast sinking organic particles that spend less time in the water column
546 limiting the opportunity of particle-attached microbes to remineralize them. Assuming that
547 MnOx had a density between 1.5 and 2.0 g cm⁻³ (Glockzin et al., 2014). The largest particles
548 measured in GB (167 μm, Table 4) will have a sinking velocity based in Stokes' law between 508
549 and 1014 m d⁻¹. If we considered a mix aggregate that is 50% TEP, density 0.9 g cm⁻³ (Azetsu-
550 Scott and Passow, 2004) and 50% MnOx (density 1.5 g cm⁻³), its density would be 1.2 g cm⁻³,
551 and its theoretical sinking velocity will be 204 m d⁻¹. This indicates that theoretically, the largest
552 mix aggregates composed of MnOx and TEP observed in the GB could reach 180 m (the location
553 of our deepest sediment trap) in less than one day. However, the average measured sinking



554 velocity of MnOx in the laboratory for particles between 2 and 20 μm was 0.76 m d^{-1} , this is
555 significantly lower than the theoretical value (Glockzin et al., 2014). Glockzin et al. (2014)
556 suggested that the star shape and the content of OM were responsible for the lower than predicted
557 sinking velocity. There is not information about the amount of OM relatively to MnOx particles in
558 those mix aggregates, or how the MnOx to OM ratio may affect the density and sinking velocity
559 of larger aggregates like the ones we observed. Due to the shape and size of MnOx-OM
560 aggregates observed in our study (Fig. 6e), we could assume those are the same type of aggregates
561 described before by Glockzin et al. (2014). Although, we did not measure the sinking velocity of
562 those aggregates, we did observe a higher abundance of them associated with TEP at 110 and 180
563 m in the GB than in the LD. The formation of these organic matter rich MnOx aggregates could
564 represent an additional mechanism (see introduction) to explain why the efficiency of the OM
565 export is different under anoxic than under oxic conditions in the Baltic Sea. The oxygenation of
566 anoxic deep water in the GB caused by the 2014/2015 MBI, may have led to an enhanced
567 precipitation of manganese, iron and phosphorous particles (Dellwig et al., 2010; Dellwig et al.,
568 2018). For example, the formation of P-rich, metal oxides precipitates occur in the anoxic waters
569 of the Black Sea (Shaffer, 1986) and Cariaco Basin (Benitez-Nelson et al., 2004; Benitez-Nelson
570 et al., 2007) where higher concentration of particulate inorganic and organic phosphorous have
571 been observed in sediment traps close to the redoxcline.

572 *4.3 Differences on composition and lability of sinking and suspended organic matter in the GB* 573 *and the LD*

574 In the sections above, we discussed how similar biogeochemical conditions and the size of the
575 surface POM pool in both the GB and the LD were. We then looked at how the sinking flux of
576 OM was affected by the different O_2 concentrations in the water column. We now focus on the
577 influence of O_2 in the chemical composition of sinking and suspended POM. Suspended or slow
578 sinking POM, that spend more time in the water column, should theoretically, show a larger
579 degree of degradation (Goutx et al., 2007). Relative to the Redfield molar ratio: 106 POC:16
580 PN:POP, OM showed an enrichment in carbon, especially in sinking particles from the LD and



581 suspended OM from the GB (Table 5). Our measured values of POC:PN (~10) and POC:POP
582 (between 89 and 506) in suspended OM coincide with the simulated ratio reported by Kreus et al.
583 (2015) immediately after the culmination of the spring bloom, those relatively high ratios are
584 consequence of the nitrogen depletion and are characteristic during the summer in the Baltic Sea.
585 The same study had suggested that POC:POP higher than Redfield ratio might lead to an
586 enhancement of particle export (Kreus et al., 2015), however, no direct observations had
587 confirmed this hypothesis. Our measurements showed that the relative higher POC:POP ratios in
588 sinking OM from LD, compared with the GB, do not lead to a higher transfer efficiency at this
589 station. Compared to the suspended OM in the LD, the POP content was lower in the GB,
590 possible related to scavenging of POP into MnOx aggregates (see section 3.4).

591 The AA based degradation index, DI (Dauwe et al. 1999) covers a wide range of alteration stages;
592 the more negative the DI, the more degraded the samples, positive DI indicates fresh organic
593 matter. In our study, the sediment trap material had a DI between 0.10 and 1.14, while suspended
594 OM has a DI between -0.26 and -1.25 (Table 4). These values coincide with what reported earlier
595 by Dauwe et al. (1999), and indicate that: first, the sinking particles collected in the sediment
596 traps were less altered (they have a more positive DI) than the suspended OM collected in the
597 CTD. Second, sinking particles from the GB were fresher than the ones from the LD, and the
598 degradation stage increased with depth in both stations. The higher contribution of AA and CHO
599 to the POC pool in sinking than in suspended OM and the AA- DI indicates that suspended OM
600 was more degraded than sinking OM. The highest degree of degradation in suspended OM and
601 sinking OM from the LD may be the result of a long time that light suspended OM or slow
602 sinking particles spend exposed to degradation in fully oxygenated surface waters than dense, fast
603 sinking particles collected in sediment traps.

604 The higher abundance of aggregates, formed by a combination of MnOx-like particles and OM,
605 observed at 110 and 180 m in the GB could act as bacteria hot spots that combined with a higher
606 O₂ concentration in the GB may increase the microbial degradation on sinking particles collected
607 in the GB. However, the AA-DI, indicated that sinking OM was less altered and therefore more



608 labile than the sinking OM in the LD. This implied that in addition to the higher transfer
609 efficiency of POC in the GB (see discussion above); the OM reaching the seafloor was fresher
610 and less degraded. This support the idea that mix aggregates composed by MnOx and OM may be
611 larger and faster sinking than the previously described by Glockzin et al. (2014). This explanation
612 is mostly speculative, and based on the observation of large mixed aggregates in the 110 and 180
613 m traps (Fig. 6, Table 4). However, as mention in the previous section, further work on directly
614 determines sinking velocity is required to prove this hypothesis.

615 **Conclusion**

616 Fluxes and composition of sinking particles were different in two deep basins in the Baltic Sea:
617 the GB and the LD during early summer 2015. The two stations had similar surface characteristics
618 and POM stock; however, at depth, the vertical profile of the O₂ concentration was different. The
619 2014/2015 MBI supplied oxygen-rich waters to the GB modifying the O₂ vertical profile and the
620 redox conditions in the otherwise permanent suboxic deep waters. This event did not affect the
621 LD allowing the comparing POM fluxes and composition under two different O₂ concentrations
622 with similar surface water conditions. Export efficiency (*e-ratio*) derived from *in-situ* PP
623 measurements and POC flux derivate from sediment traps indicated higher export efficiency in
624 LD than in GB. However, the transfer efficiency (POC flux at 180 m over POC flux at 40 m)
625 suggested that under anoxic conditions found in the LD, a smaller portion of the POC exported
626 below the euphotic zone was transferred to 180 m than under re-oxygenated conditions present in
627 the GB. Our results suggest that a new possible mechanism to explain the differences in the OM
628 fluxes under different O₂ concentration could be the formation and prevalence of aggregates
629 composed of MnOx and organic matter in the GB. Those aggregates were significantly larger and
630 more abundant in the GB compared to the LD where sulfidic waters constrained their presence.
631 We propose that after a MBI in the GB, the aggregates containing MnOx-like particles and
632 organic matter could reached the sediments relatively fast and unaltered, scavenging not only
633 phosphorous, as described previously (Dellwig et al., 2010), but also other organic compounds.
634 The remineralization of this organic matter reaching the sediments may contribute to the quick re-



635 establishment of anoxic conditions in the sediment-water interface in the GB. The relevance of
636 this process need to be further investigate in order to be included in O₂ budget and long-term
637 predictions of the MBI impact in the O₂ and OM cycles.

638 **Author Contributions**

639 C.C.N. performed deployments, analyzed samples and wrote the manuscript. F.A.C.L.M.,
640 performed deployments and contributed to the writing of the manuscript. A.E designed and
641 conducted the scientific program at sea and discussed and commented on the manuscript.

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Figure Captions

Figure 1. Monthly averaged Chl *a* distribution derived from VIIRS for June 2015 in the Baltic Sea. Black circle and “x” indicate the position of the trap deployment and the seawater collection respectively in Gotland Deep (GB) and Landsort Deep (LD). The lower panel shows the trajectory of the trap deployed at GB and LD.

Figure 2. Water column profiles at the location of the sediment trap deployments in (A) the GB, and (B) the LD. Left panel: oxygen (blue), temperature (red), and salinity (black). Middle panel: nitrate (NO₃), nitrite (NO₂), and ammonium (NH₄). Right panel: phosphate (PO₄), and silicate (Si(OH)₄). Grey lines indicate the depths at which we deployed sediment traps.

Figure 3. Particulate organic matter profiles in the water column at the location of the sediment traps deployments in the GB (A, B and C) and the LD (D, E and F). (A and D) particulate organic carbon (POC), particulate nitrogen (PN), and particulate organic phosphorous (POP). (B and E) chlorophyll *a* (Chl *a*) and biogenic silicate (BSi). (C and F) transparent exopolymeric particles (TEP) and Coomassie stainable particles (CSP). Grey lines as figure 2.

Figure 4. MnOx-like containing particles and O₂ concentration profiles in the water column at the location of the sediment traps deployments. (A) the GB and (B) the LD. Grey lines as in figure 3.

Figure 5. Particulate organic matter fluxes in the GB (A and B) and the LD (C and D). (A and C) POC, PN and O₂. (B and D) POP, Chl *a*, and BSi.

Figure 6. TEP and CSP fluxes in the GB (A and B) and the LD (C and D). In addition to the vertical distribution of the flux, each profile is complemented with images captured under the microscope (200x) at each depth. Star-shaped MnOx-like particles are clearly visible in the GB associated to TEP (A), but not with CSP (B). MnOx-like particles were significantly less abundant in the LD (C and D). (F) A larger magnification (400x) image of MnOx-like particles at 110 m showing more detail on the shape of those particles and aggregates formed with TEP.



Figure 7. Total hydrolyzable amino acids (TAA) and total carbohydrates (TCHO) fluxes in (A) the GB, and (B) the LD.



Table 1. Sediment traps deployment and recovery locations, dates, collection times and depths.

Station	Lat	Lon	Date	Station depth	Deployment time (d)	Trap depths (m)
Gotland Basin (GB)	57.21 °N	20.03 °E	08/06/2015	248 m	2	40A, 40B, 60, 110, and 180m
	57.27 °N	20.25 °E	10/06/2015			
Landsort Deep (LD)	58.69 °N	18.55 °E	15/06/2015	460 m	1	40A, 40B, 55, 110, and 180m
	58.68 °N	18.68 °E	16/06/2015			



Table 2. Abundance of chlorophyll and phycoerythrin containing pico- and nanoplankton measured by flow-cytometry in the GB and the LD.

	Depth (m)	Phytoplankton (mL ⁻¹)			Cyanobacteria-like cells (mL ⁻¹)		
		picoplankton	nanoplankton	Total	picoplankton	nanoplankton	Total
GB	1	87963	2097	90060	5225	731	5956
	10	94369	2628	96997	8795	920	9716
	40	4999	68	5067	2174	69	2243
	60	4125	35	4160	1990	42	2032
	80	599	7	606	238	15	253
	110	594	7	601	326	29	356
	140	1144	14	1158	356	2	358
	180	908	9	917	366	20	385
	220	2270	19	2289	1063	34	1097
LD	1	92359	2283	94642	834	177	1011
	10	86426	1708	88134	2990	232	3223
	40	2022	92	2114	2243	69	2312
	60	1524	62	1586	1294	24	1318
	70	908	43	951	613	17	630
	110	1735	82	1817	1181	17	1198
	180	1339	75	1415	946	34	980
	250	1593	82	1676	949	36	985
	300	1521	48	1569	1047	17	1064
	350	1608	57	1665	908	12	920
	400	1548	73	1621	1047	22	1069
430	1562	68	1631	875	19	894	



Table 3. Phytoplankton abundances analyzed microscopically in the GB and the LD, volume analyzed was 50 ml per sample.

		GB (cells mL ⁻¹)				LD (cells mL ⁻¹)			
		1 m	10 m	40 m	Total	1 m	10 m	40 m	Total
Cyanophyceae *	Total	14148	13536	0	27684	37368	32526	96	69990
Chryptophyta	Total	140	112	28	280	1400	882	56	2338
Bacillariophyceae	Total	96	94	44	234	462	112	102	676
	<i>Chaetoceros</i> sp.	58	42	24	124	434	106	26	566
	<i>Skeletonema</i> sp.	26	8	12	46	12	0	8	20
	<i>Thalassiosira</i> sp.	12	44	8	64	16	6	68	90
Dinophyceae	Total	3772	4424	1192	9388	9032	7662	1404	18098
	<i>Dinophysis</i> sp.	678	742	2	1422	450	214	4	668
	other	3094	3682	1190	7966	8582	7448	1400	17430
Chlorophyta	Total	5320	6860	28	12208	2072	1022	238	3332
	<i>Planctonema</i> sp.	5320	6860	28	12208	2072	1022	238	3332

* >90% were filamentous unicellular cyanobacteria *Aphanizomenon* sp.



Table 4. MnOx-like particles fluxes and size determined by image analysis in GB and LD.

Station	Depth (m)	MnOx-like particles ($\text{cm}^2 \text{m}^{-2} \text{d}^{-1}$)	Median size ESD (μm)	Size range ESD (μm)
GB	110	5666.1± 993.5	2.8	0.6-166.7
	180	7789.1± 954.7	3.3	0.6-152.7
LD	110	50.3±1.8	1.8	0.6-16.5
	180	2.6±0.3	1.4	1.2-9.3



Table 5. Amino acids (AA), carbohydrates (CHO) and elemental molar ratios of sinking and suspended OM in the GB and in the LD.

	Depth (m)	AA-C:POC %	CHO-C:POC %	POC:PN	POC:POP	POC:Bsi	PN:POP
GB	40	19.19	18.26	9.80	244.05	3.86	0.39
sinking OM	40	17.58	17.21	9.43	222.42	4.07	0.43
	60	15.78	17.56	9.52	231.56	2.78	0.29
	110	13.87	22.24	11.31	90.12	1.73	0.15
	180	11.13	18.47	12.68	122.87	2.97	0.23
LD	40	13.52	9.43	12.17	771.70	3.58	0.29
sinking OM	40	14.27	8.40	11.09	413.14	4.12	0.37
	55	19.10	10.97	12.43	331.81	3.03	0.24
	110	13.37	11.97	15.44	229.70	2.67	0.17
	180	14.32	12.85	15.29	341.33	4.19	0.27
GB	1	8.22	16.94	10.39	154.56	91.45	14.88
suspended OM	10	10.81	8.84	10.48	150.51	87.15	14.36
	40	4.91	2.80	9.19	88.78	133.75	9.66
	60	5.43	2.66	9.78	127.36	125.24	13.02
	80	4.67		10.43	144.92		13.89
	110	9.01	6.63	8.45	245.26		29.01
	140	5.34		10.60	283.42		26.73
	180	5.73	4.29	11.37	506.21		44.54
	220	8.57	3.35	12.06	270.78		22.45
LD	1	6.96		8.66	205.29	514.94	23.71
suspended OM	10	12.97	9.12	8.43	196.44	100.91	23.31
	40	0.00	8.88	8.09	335.66	24.48	41.51
	60	6.09	10.26	7.83	300.75	16.89	38.43
	70	7.92	10.72	7.71	291.81	247.80	37.86
	110	12.22	5.41	7.93	224.56		28.32
	180	10.12	11.32	7.02	205.33		29.23
	250	11.97	8.81	6.52	249.36		38.22
	300	10.88		6.71	136.67		20.37
	350	10.67	10.12	6.76	145.80		21.56
	400	9.99		6.18	229.53		37.16
	430	9.35	9.45	7.82	148.61		19.01

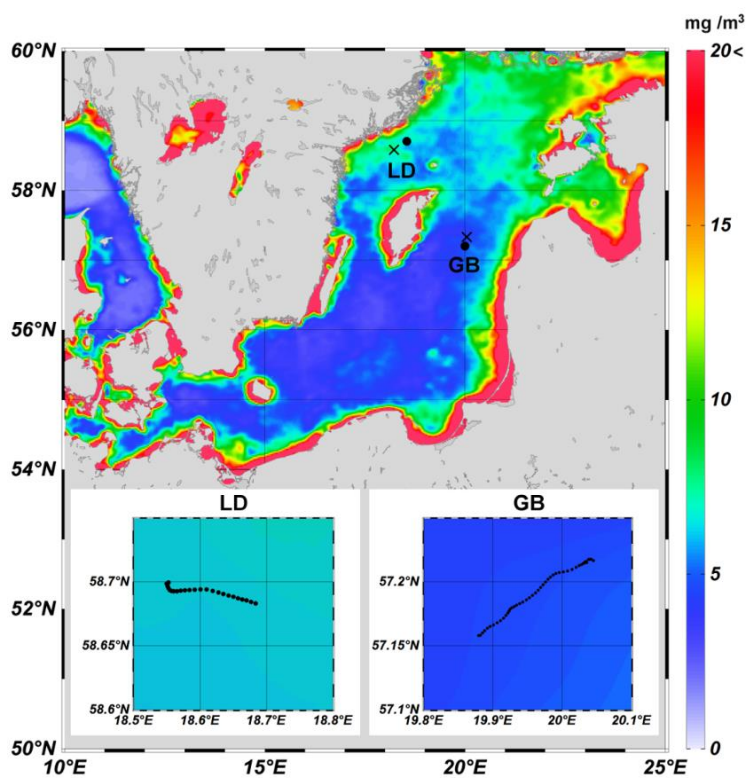


Fig. 1

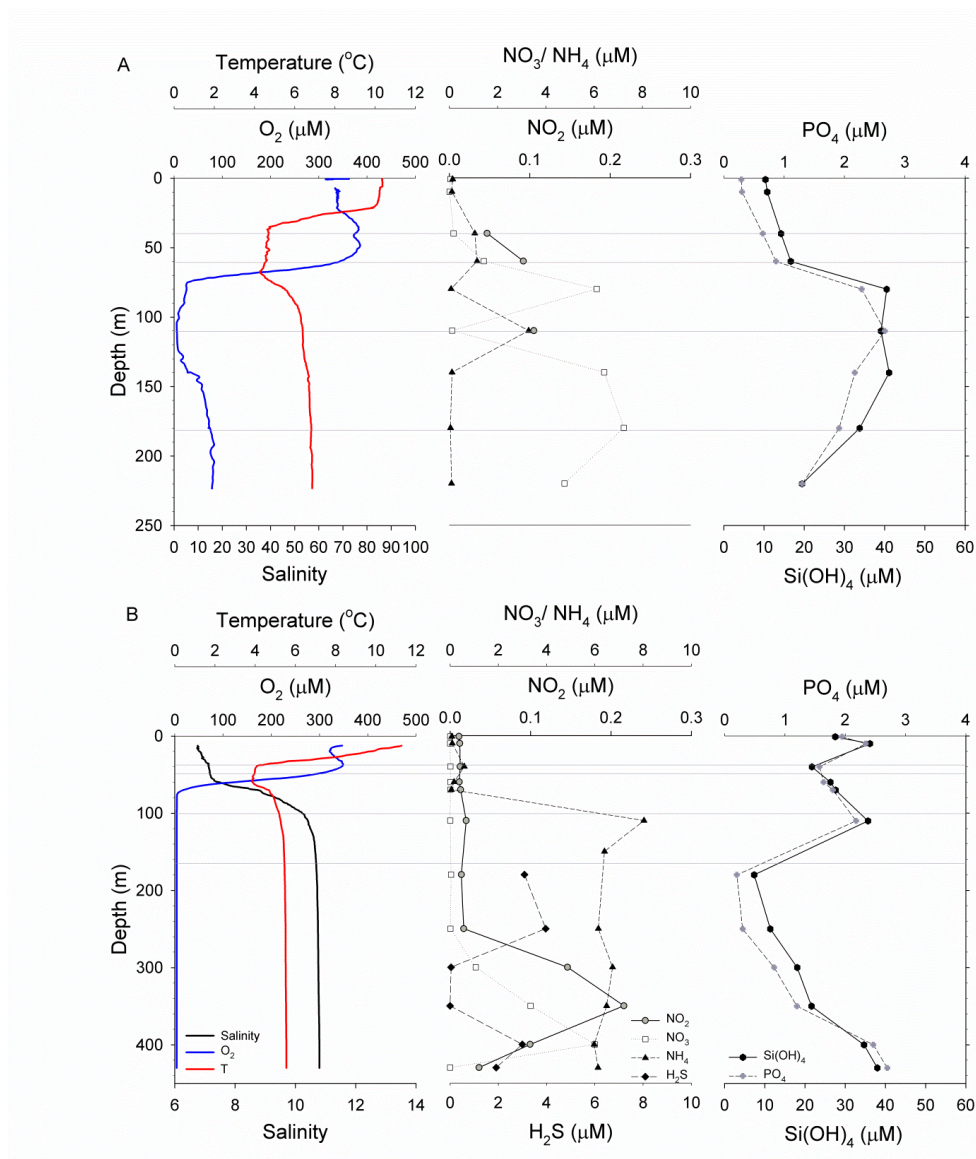


Fig. 2

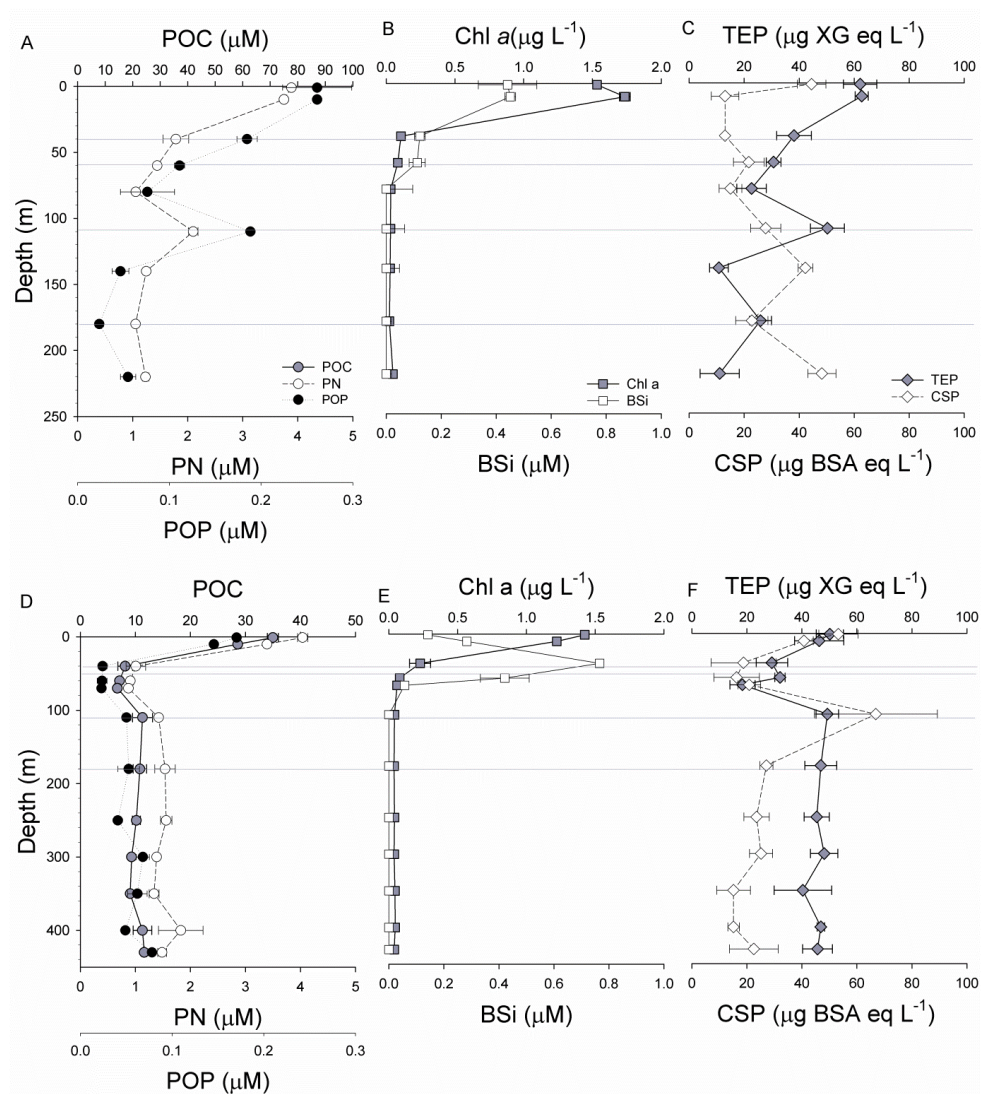


Fig. 3

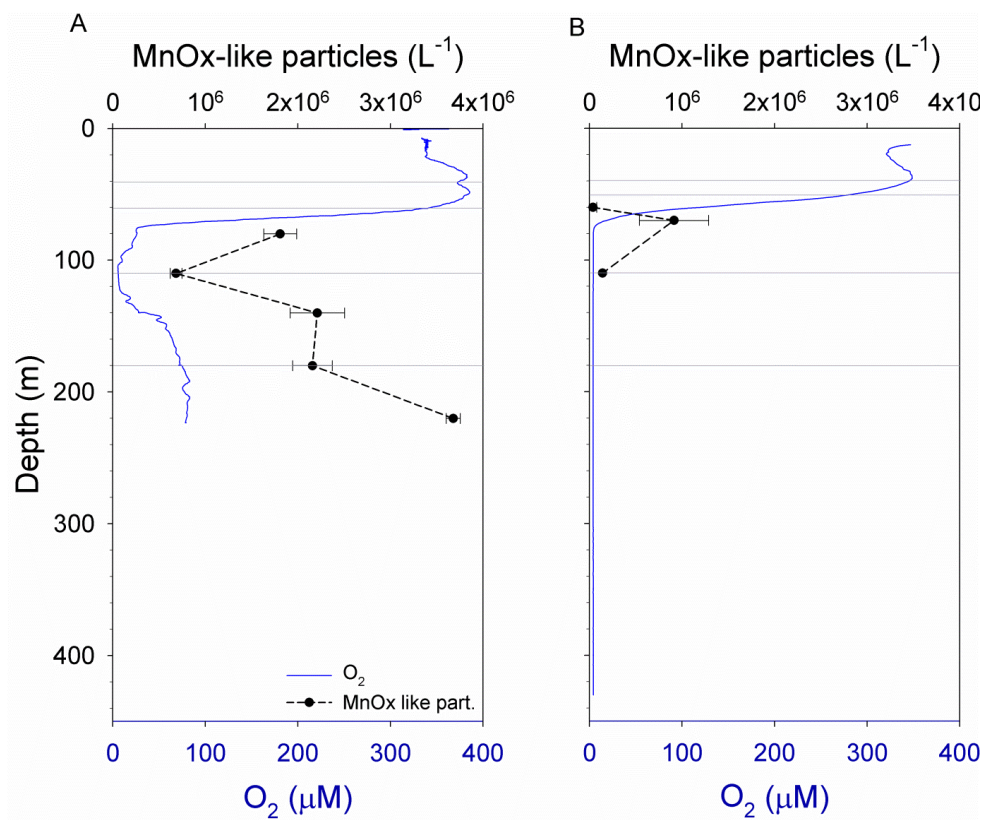


Fig. 4

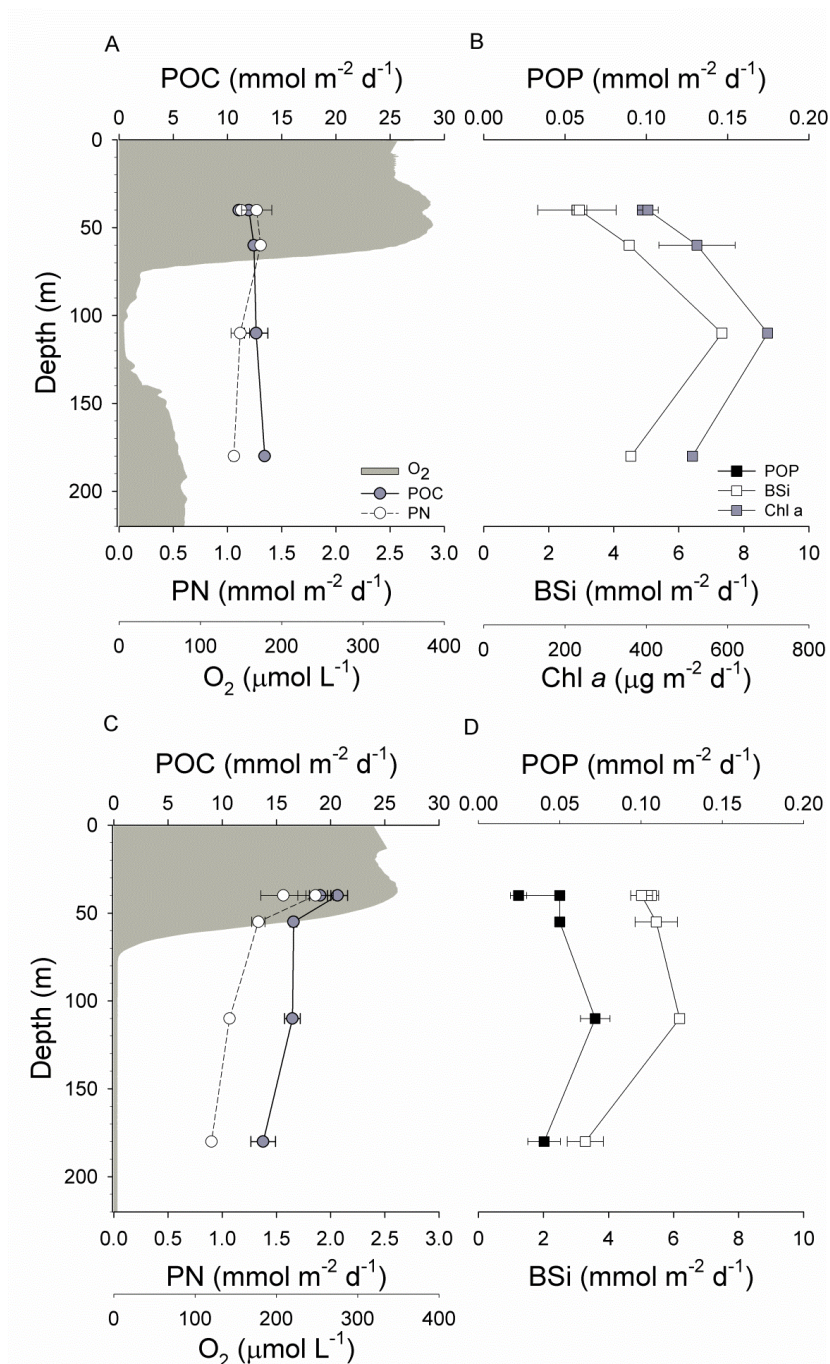


Fig. 5

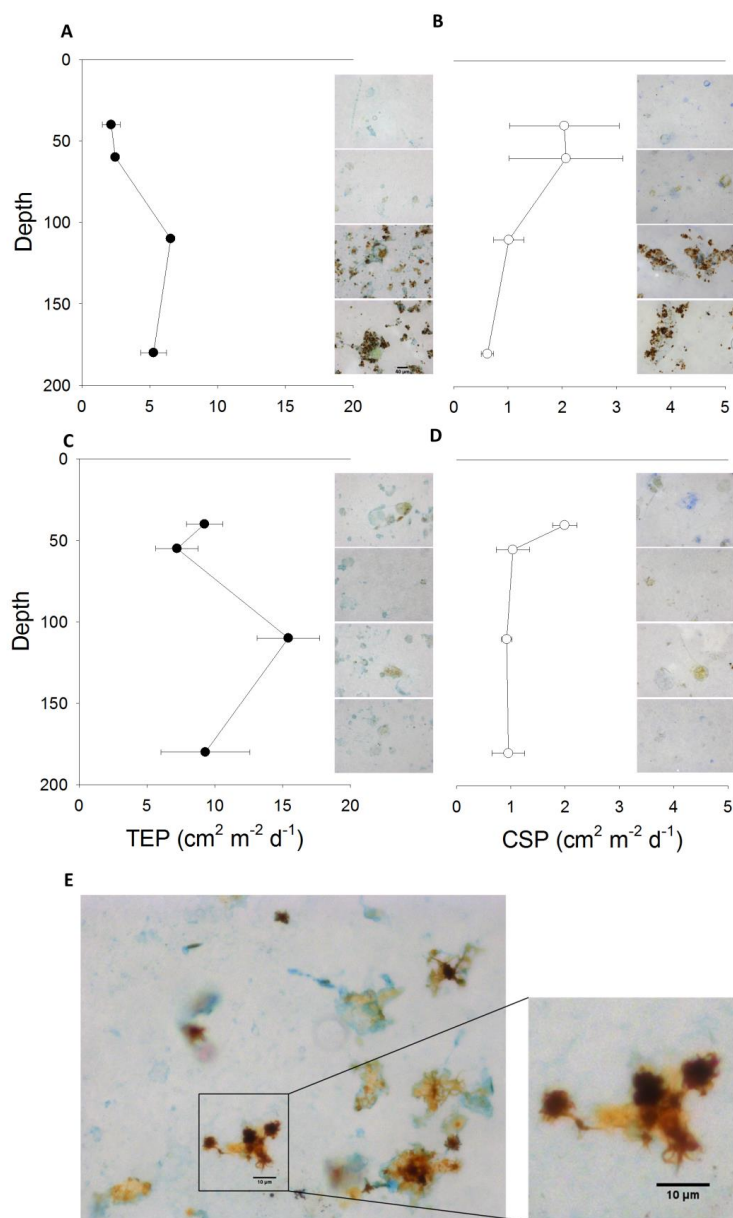


Fig. 6

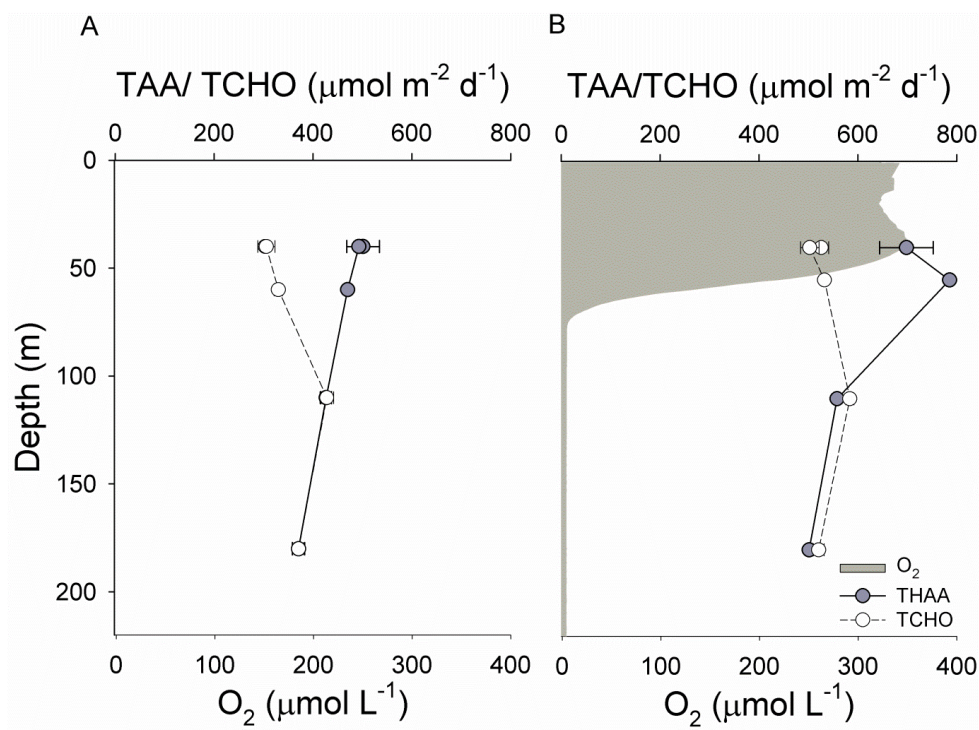


Fig. 7