

<https://helda.helsinki.fi>

Ecosystem functioning along gradients of increasing hypoxia and changing soft-sediment community types

Norkko, Joanna

2019-11

Norkko, J, Pilditch, C A, Gammal, J, Rosenberg, R, Enemar, A, Magnusson, M, Granberg, M E, Lindgren, J F, Agrenius, S & Norkko, A 2019, ' Ecosystem functioning along gradients of increasing hypoxia and changing soft-sediment community types ', Journal of Sea Research, vol. 153, 101781. <https://doi.org/10.1016/j.seares.2019.101781>

<http://hdl.handle.net/10138/308707>

<https://doi.org/10.1016/j.seares.2019.101781>

cc_by

acceptedVersion

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.

This is the AUTHORS' ACCEPTED MANUSCRIPT, for the published article, please see:
<https://doi.org/10.1016/j.seares.2019.101781>

Full citation:

Norkko J, Pilditch CA, Gammal J, Rosenberg R, Enemar A, Magnusson M, Granberg ME, Lindgren JF, Agrenius S, Norkko A (2019) Ecosystem functioning along gradients of increasing hypoxia and changing soft-sediment community types. *Journal of Sea Research* 153: 101781

***Ecosystem functioning along gradients of increasing hypoxia and
changing soft-sediment community types***

Joanna Norkko^{a,*}, Conrad A. Pilditch^b, Johanna Gammal^a, Rutger Rosenberg^c, Arvid Enemar^c, Marina Magnusson^d, Maria E. Granberg^e, J. Fredrik Lindgren^f, Stefan Agrenius^g, Alf Norkko^{a,h}

^a Tvärminne Zoological Station, University of Helsinki, 10900 Hanko, Finland

^b School of Science, University of Waikato, Private Bag 3105, Hamilton, New Zealand.

^c Department of Biological and Environmental Sciences - Kristineberg, University of Gothenburg, Kristineberg 566, 45178 Fiskebäckskil, Sweden

^d Marine Monitoring AB, Strandvägen 9, 453 30 Lysekil, Sweden

^e IVL Swedish Environmental Research Institute, Kristineberg Marine Research and Innovation Center, Kristineberg 566, 45178 Fiskebäckskil, Sweden

^f Department of Mechanics and Maritime Sciences, Chalmers University of Technology, 41296 Gothenburg, Sweden

^g Department of Marine Sciences - Kristineberg, University of Gothenburg, Kristineberg 566, 45178 Fiskebäckskil, Sweden

^h Baltic Sea Centre, Stockholm University, 106 91 Stockholm, Sweden

* Corresponding author, joanna.norkko@helsinki.fi

Declarations of interest: None.

Author contributions: JN, AN, JG, RR designed the study.

JN, CP, JG, RR, AE, MM, MG, AN conducted the field sampling.

JG, RR, AE, SA analysed the macrofauna samples.

FL analysed the meiofauna samples.

RR, MM analysed the sediment profile image data.

CP, JG, AN, FL conducted the statistical analyses.

JN drafted the manuscript and all authors contributed.

All authors have approved the final article.

Keywords. Hypoxia; nutrient cycling; structural community changes; ecosystem functioning; macrofauna; meiofauna

Highlights

1. Hypoxia decimates macrofauna, but fauna can still contribute to nutrient cycling
2. Meiofauna is less sensitive to hypoxia compared with macrofauna
3. The link between community structure and ecosystem function is mediated by context

1 **Abstract.** Marine ecosystems world-wide are threatened by oxygen deficiency, with potential
2 serious consequences for ecosystem functioning and the goods and services they provide. While the
3 effects of hypoxia on benthic species diversity are well documented, the effects on ecosystem
4 function have only rarely been assessed in real-world settings. To better understand the links
5 between structural changes in macro- and meiofaunal communities, hypoxic stress and benthic
6 ecosystem function (benthic nutrient fluxes, community metabolism), we sampled a total of 11 sites
7 in Havstensfjord and Askeröfjord (Swedish west coast) in late summer, coinciding with the largest
8 extent and severity of seasonal hypoxia in the area. The sites spanned oxic to anoxic bottom water,
9 and a corresponding gradient in faunal diversity. Intact sediment cores were incubated to measure
10 fluxes of oxygen and nutrients (NO_3^- , NO_2^- , NH_4^+ , PO_4^{3-} , SiO_4) across the sediment-water interface.
11 Sediment profile imaging (SPI) footage was obtained from all sites to assess structural elements and
12 the bioturbation depth, and additional samples were collected to characterise sediment properties
13 and macro- and meiofaunal community composition. Bottom-water O_2 concentration was the main
14 driver of macrofauna communities, with highest abundance and biomass, as well as variability, at
15 the sites with intermediate O_2 concentration. Meiofauna on the other hand was less sensitive to
16 bottom-water O_2 concentration. Oxygen was the main driver of nutrient fluxes too, but macrofauna
17 as well meiofauna were also significant predictors; DistLM analyses indicated that O_2
18 concentration, macrofaunal abundance or biomass, and meiofaunal abundance collectively
19 explained 63%, 30% and 28% of the variation in sediment O_2 consumption, NH_4^+ -flux and PO_4^{3-} -
20 flux, respectively. The study provides a step towards a more realistic understanding of the link
21 between benthic fauna and ecosystem functioning, and the influence of disturbance on this
22 relationship, which is important for management decisions aimed at protecting the dwindling
23 biodiversity in the coastal zones around the world.

24

25

26 **INTRODUCTION**

27 Marine ecosystems worldwide are threatened by oxygen deficiency, with potential serious
28 consequences for ecosystem functioning and the goods and services these ecosystems provide (Diaz
29 and Rosenberg 2008, Rabalais et al. 2014). While eutrophication and organic enrichment are the
30 main anthropogenic causes of hypoxia, the warming climate will further exacerbate the
31 deoxygenation of the oceans (Breitburg et al. 2018). This highlights the urgency of better
32 understanding how ecosystem functioning might change with increasing hypoxia, and what factors
33 and mechanisms are driving these changes.

34
35 The deleterious effects of increasing hypoxia on soft-sediment macrofaunal communities are well
36 documented, with a general decrease of large, deeper-dwelling animals and an increase of smaller,
37 fast-growing species, until anoxia decimates all macrofauna (Pearson and Rosenberg 1978, Diaz
38 and Rosenberg 1995, Gray et al. 2002, Levin et al. 2009). Through their bioturbation and
39 bioirrigation activities, macrofauna enhance oxygen penetration into the sediments influencing all
40 oxygen-dependent processes in the sediment, including organic matter mineralization through
41 stimulation of microbial activity, and nutrient cycling (Levinton 1995, Aller and Aller 1998,
42 Meysman et al. 2006, Glud 2008). These activities are reduced under hypoxic conditions due to
43 changes in the behaviour and diversity of macrofauna. Bottom-water O₂ concentrations also
44 influence biogeochemical processes at the sediment-water interface affecting nutrient
45 concentrations and speciation in the water column. In particular, the release of phosphate and
46 ammonium is enhanced under hypoxic conditions (Mortimer 1941, Ingall et al. 1993, Cowan and
47 Boynton 1996, Slomp et al. 2002, McCarthy et al. 2008, Reed et al. 2011, Jäntti and Hietanen
48 2012). Nevertheless, the step from documenting structural changes in faunal composition due to
49 hypoxia to understanding the impact on ecosystem functions (e.g., nutrient cycling) is long and we
50 have only recently begun to assess the interacting direct (e.g. chemical release of nutrients from the

51 sediment) and indirect (e.g. via effects on macrofauna) effects of hypoxia on ecosystem function in
52 natural settings (Norkko et al. 2015, Gammal et al. 2017).

53

54 The effects of hypoxia on benthic ecosystem functioning are likely to be highly context dependent,
55 posing further challenges to building a general understanding of the effects and predicting future
56 changes. Impacts will depend on the temporal and spatial scales of hypoxia (a function of mixing
57 and water exchange), the type of habitat (e.g. muddy, sandy) and faunal diversity. Places with high
58 species diversity are generally expected to tolerate stress better than low-diversity systems
59 (insurance hypothesis, e.g., Yachi and Loreau 1999), but it is important to consider biodiversity as a
60 much wider concept than just the number of species, including aspects such as species identity and
61 dominance patterns. For example, dominance by one or a few species with particular functional
62 traits may be more important than species diversity *per se* for some aspect of ecosystem functioning
63 (Chapin III et al. 1997). A dominant species with a good hypoxia tolerance may thus maintain a
64 vital process, e.g. bioturbation, when conditions deteriorate (Norkko et al. 2015, Rakocinski and
65 Menke 2016)). Thus, species identity and the prevalence of functionally important traits will be
66 important for assessing hypoxia-induced changes in ecosystem functioning.

67

68 It is difficult to directly link disturbance-induced community changes to quantifiable shifts in
69 functioning without empirical measurements. Nevertheless, some assumptions can be made. Under
70 anoxic conditions, when the macrofauna has been lost, there is no effect of the fauna on nutrient
71 cycling and therefore chemical reactions, modulated by microbes, dominate solute fluxes and
72 ecosystem function. As conditions deteriorate from normoxia, the ensuing hypoxia results in
73 different community types, representing different successional stages (Pearson and Rosenberg
74 1978, Diaz and Rosenberg 1995, Nilsson and Rosenberg 1997, Rosenberg et al. 2002). Sustained
75 hypoxia decimates big individuals, e.g., large deep-burrowing bivalves, which are particularly

76 important for nutrient cycling (Norkko et al. 2013). Thus, the macrofaunal influence on nutrient
77 cycling is likely to be reduced in a decimated community, with lower species diversity, abundance
78 and biomass. In addition, already at sub-lethal levels of hypoxic stress, behavioural and
79 physiological changes may affect the species' contribution to processes such as bioturbation, but
80 species-specific sensitivities to hypoxia vary greatly (Vaquer-Sunyer and Duarte 2008). Thus,
81 hypoxia results in a non-random species loss and the remaining community types may be adapted to
82 low-oxygen environments. It is, however, unclear how well they perform.

83
84 Much of our understanding of biodiversity-ecosystem functioning (BEF) relationships stems from
85 mechanistic, small-scale laboratory studies with only a few species and limited regard of changing
86 environmental conditions, such as increasing hypoxia (Snelgrove et al. 2014). While the potential
87 indirect effects of hypoxia via fauna on biogeochemical processes have been indicated in several
88 high-profile papers (e.g., Levin et al. 2009, Middelburg and Levin 2009, Friedrich et al. 2014),
89 actual measurements still appear to be virtually non-existent. To our knowledge, the effects of
90 hypoxia on BEF in terms of nutrient cycling have not been empirically tested in a meaningful way
91 in the laboratory or under natural field conditions. While it is challenging to assign causality in field
92 studies, relevant field measurements involving natural communities in a range of different
93 environments and geographical areas are imperative for developing a realistic understanding of the
94 effects of disturbance on BEF relationships. Using correlative field surveys and incubations of non-
95 manipulated sediment cores for measurement of benthic nutrient fluxes in a range of contrasting
96 environments, we have started to understand the importance of benthic macrofauna for nutrient
97 cycling and the concurrent effects of increasing hypoxia. This body of work includes a large-scale
98 study across the entire open Baltic Sea, spanning a salinity and corresponding diversity gradient as
99 well as areas that are more or less permanently hypoxic (Norkko et al. 2015) and a coastal study in
100 a brackish, seasonally hypoxic, low-diversity system in the northern Baltic Sea (Gammal et al.

101 2017). In order to pinpoint the mechanisms involved, the same research team has conducted coastal
102 field experiments where hypoxic events of different intensity were simulated *in situ* and the
103 responses assessed (Villnäs et al. 2012, Norkko et al. 2013, Villnäs et al. 2013). In all of these
104 studies, macrofauna was important for explaining the variability in nutrient fluxes across the
105 sediment-water interfaces, but this effect decreased as oxygen conditions deteriorated. It is now
106 imperative to investigate whether the patterns in higher-diversity systems are comparable to the
107 ones found in the low-diversity Baltic Sea.

108

109 Missing from these studies is also the consideration of several size classes and trophic levels
110 simultaneously, i.e. meiofauna and microbes. It is known that the influence of macrofauna on
111 ecosystem function, sediment biogeochemistry and nutrient cycling, is modulated by meiofauna
112 (Bonaglia et al. 2014, Piot et al. 2014) and by microbes (Yazdani Foshtomi et al. 2015), but studies
113 that examine these relationships in relation to hypoxia and nutrient cycling are rare. Also, the Baltic
114 Sea where our previous studies have been conducted is a low-diversity system (Villnäs and Norkko
115 2011), calling for comparative studies in fully marine systems that experience hypoxia but have
116 higher macrofaunal diversity (and potentially greater redundancy), such as the Swedish west coast.

117

118 Hypoxic or anoxic dead zones in deeper waters (e.g. the Baltic Sea, the Gulf of Mexico) have been
119 known for decades (e.g., Rabalais et al. 2002, Conley et al. 2009a), but the problem of near-shore,
120 coastal hypoxia is now receiving ever more attention (Conley et al. 2011). The coastal ecosystems
121 are heterogeneous and diverse, with enclosed inlets, and steep gradients in physical, chemical and
122 biological properties. They are hotspots of diversity and productivity, but at the same time human
123 impacts are very pronounced along the coasts (Levin et al. 2001, Halpern et al. 2008), highlighting
124 the need for management actions based on sound understanding of the links between hypoxic
125 disturbance and the functioning of these systems.

126

127 To investigate the links between benthic communities, hypoxic stress and nutrient fluxes across the
128 sediment-water interface, we conducted a field study in the seasonally hypoxic Havstensfjord and
129 Askeröfjord (Swedish west coast), at sites covering a gradient from oxic to anoxic bottom water,
130 with a corresponding gradient in diversity. The aim was to assess how macrofauna and meiofauna
131 communities change with increasing hypoxia and whether the effects of these changes on nutrient
132 cycling/fluxes could be quantified. We also investigated whether the different community types, or
133 successional stages, corresponded to different levels of sediment oxygen consumption (a proxy for
134 community metabolism). Given the higher species diversity and thus potentially higher functional
135 redundancy in Havstensfjord and Askeröfjord compared to our previous studies in the Baltic Sea
136 (Norkko et al. 2015, Gammal et al. 2017), we anticipated that the effect of hypoxia on the faunal
137 contribution to functioning would be smaller.

138

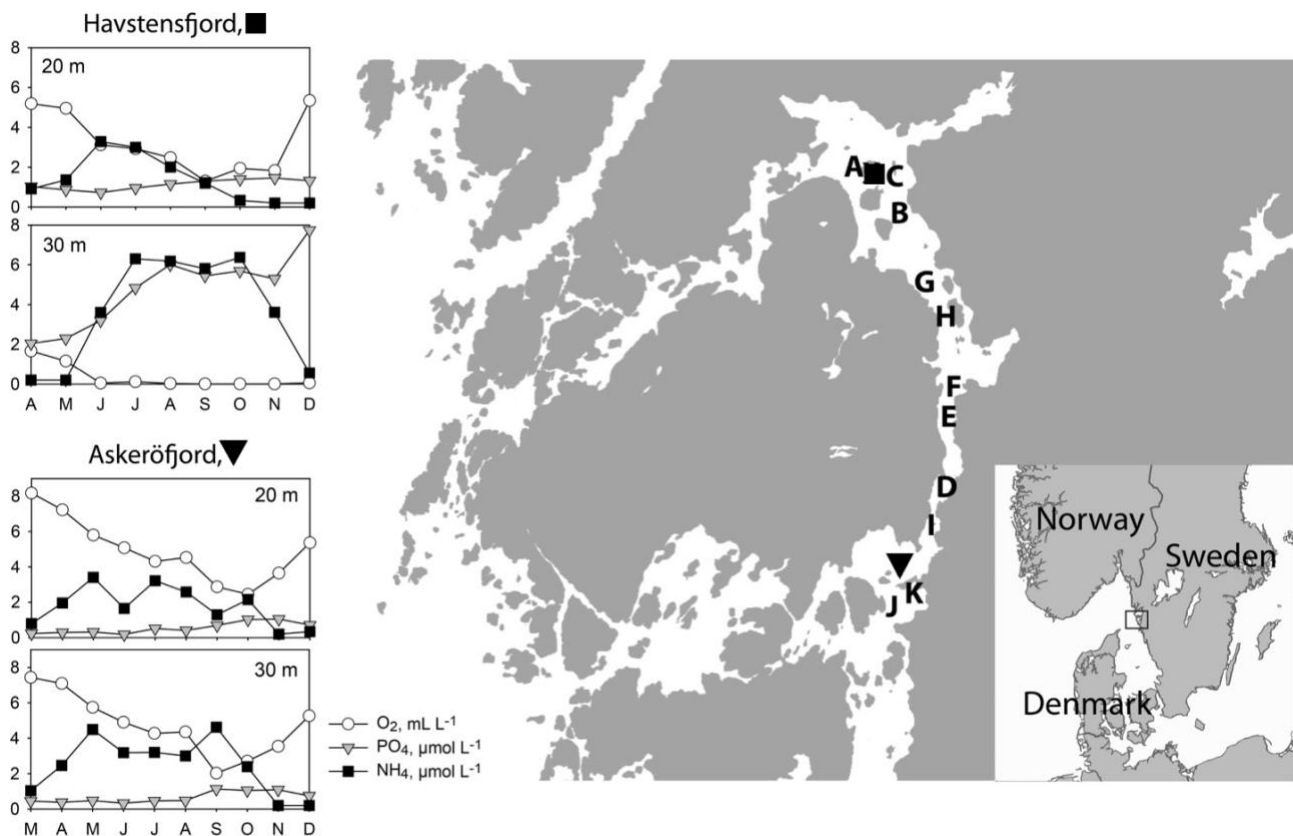
139 **MATERIAL AND METHODS**

140 **Study area, sampling and sample processing**

141 The Havstensfjord is a narrow fjord on the Swedish west coast and part of the Orust fjord system
142 (Fig. 1). The fjord extends about 25 km from north to south with its main connection to the sea
143 further south through Askeröfjord. The fjord system has been monitored since the 1950s and
144 bottom-water oxygen concentrations have steadily declined since then (Nilsson and Rosenberg
145 1997). The fjords suffer from seasonal hypoxia and particularly in the northern parts, deeper waters
146 may be anoxic for extended periods of the year (Hansson et al. 2013). The deeper parts of
147 Havstensfjord are usually ventilated once per year in late winter or early spring. The periods of
148 lower oxygen concentrations also correspond to higher concentrations of phosphate and ammonium
149 in the bottom waterers (Fig. 1).

150

151 During the peak of seasonal hypoxia, in early September 2011 we sampled 9 sites in the
 152 Havstensfjord and 2 outside the entrance in the Askeröfjord, covering a gradient from oxic sites
 153 outside the sill to Havstensfjord to hypoxic sites inside the sill, and then almost anoxic bottom water
 154 at the deepest site in the north (Fig. 1, Table 1). Although long-term monitoring data does not exist
 155 for all 11 sites, bottom-water oxygen concentrations in the fjord are strongly related to depth and
 156 therefore the assumption is that all sites follow a general pattern of lowest oxygen conditions at the
 157 end of summer. The choice of sampling sites was based on Nilsson and Rosenberg (1997) and the
 158 sampling was conducted on-board *R/V Skagerak*. All sites had muddy sediments, similar organic
 159 content and were 23-39 m deep.
 160



161
 162
 163 **Figure 1.** Map of the sampling area in the Orust fjord system on the Swedish west coast. Letters A-
 164 K indicate sites sampled during this study in September 2011. Inserted graphs present
 165 concentrations of oxygen (O₂, mL L⁻¹), phosphate (PO₄³⁻, µmol L⁻¹), and ammonium (NH₄⁺, µmol

166 L-1) measured at 20 and 30 m depth at monitoring sites in Havstensfjord (filled square; station name
167 Havstensfjord) and Askeröfjord (filled triangle; station name Galterö) from March or April to
168 December 2011, as part of the national Swedish coastal monitoring program (data obtained from the
169 SMHI database SHARKweb).

170

171 To characterise environmental conditions at each site, bottom-water salinity and temperature were
172 determined from CTD casts (Sea-Bird). Intact sediment cores were collected with a Gemax
173 twincoiler (ID = 90 mm) and the surface sediment (0-1 cm) analysed for organic content (OC, %
174 loss on ignition, 3 h at 500°C) and sediment silt/clay content (% <63 µm, determined by wet
175 sieving). To illustrate differences in the sedimentary environment between anoxic, hypoxic and oxic
176 sites, four sediment profile images (SPI) were obtained from each site and digitally analysed in
177 PhotoShop for sediment surface/subsurface structures and mean depth of the apparent redox
178 potential discontinuity (aRPD). Based on these variables a benthic habitat quality (BHQ) index was
179 calculated (see details in Nilsson and Rosenberg 1997).

180

181 Oxygen and nutrient fluxes across the sediment-water interface were estimated by on-board
182 incubation of undisturbed, intact sediment cores (n=5 per site). The upper parts of Gemax split
183 tubes were sealed and used as flux chambers (30 cm sediment + 10 cm bottom water). The core lid
184 contained a Teflon-coated magnetic stirring bar, which provided continuous gentle stirring by an
185 external magnet. Core incubations (in the dark at 11°C) started immediately after collection and
186 water samples for O₂ and nutrient concentrations (NO₃⁻, NO₂⁻, NH₄⁺, PO₄³⁻, SiO₄) were obtained at
187 the start and the end of incubation (4 h later). The differences in concentration were used to
188 calculate solute fluxes (µmol m⁻² d⁻¹). At the end of incubation, all cores were sieved to quantify
189 benthic macrofaunal species richness, abundance and biomass (0.5 mm sieve, preserved in 70%
190 ethanol, biomass estimated as blotted wwt). Dissolved oxygen was determined by Winkler titration,
191 while the nutrient samples were filtered (GF/F) and then frozen (-20°C) until analysed
192 spectrophotometrically with an autoanalyser (Lachat QuickChem 8000).

193

194 To provide an additional and more robust estimate of benthic macrofaunal species richness,
195 abundance and biomass, than that gained from the small flux cores (0.006 m²), we also sampled
196 with a Smith-McIntyre grab (0.1 m², 3 replicates per site, 1 mm sieve, preserved in 70% ethanol).

197

198 From a grab sample at each site, three subsamples (40 g ww_t) of the top 2 cm sediment were taken
199 for analyses of meiofauna community composition. These samples were stored in an
200 ethanol:glycerol (95:5 %) solution and meiofauna extracted using Ludox (Burgess 2001). Sediment
201 was rinsed with tap water on a 63 µm sieve to remove ethanol, salt and organic matter. Remaining
202 sediment was transferred to a 13-ml centrifuge tube, centrifuged at 800 G for 5 minutes and the
203 water was decanted. 10 ml of Ludox AS-40 was added to the tube and the tube was then vortexed at
204 1800 rpm for 30 s and then at 1400 rpm for 4.5 min. Afterwards the sample was centrifuged at 800
205 G for 5 minutes and approximately 2 ml of the top sample was transferred to a new tube. The old
206 tube was topped up with fresh Ludox and the procedure was repeated. The retained material was
207 sieved with milliQ water through a 63-µm sieve to remove the Ludox from the extracted meiofauna
208 and preserved in ethanol:glycerol solution. The extracted meiofauna were put on petri dishes and
209 diluted with milliQ water. The petri dishes were then digitally scanned on an Epson Perfection v500
210 (6400 dpi, 16-bit grey scale in positive film mode). The meiofauna were analysed using the image
211 analysis software ZooImage (Lindgren et al. 2013) to the following major taxonomic groups:
212 Nematoda, Harpacticoida, Rotaliina, Reophax, Allogromiina, Nonionella, Tanaidacea, Polychaeta
213 and Ostracoda.

214

215 **Statistical analyses**

216 Multivariate analyses in PRIMER 6 (Clarke and Gorley 2006) based on taxon abundance was used
217 to assess inter-site variations in faunal community composition. Macrofauna community data from

218 grab samples were transformed (square root) to lessen the influence of dominant taxa before Bray-
219 Curtis similarity calculations. Meiofauna data were not transformed because of the relatively coarse
220 taxonomic resolution and even distribution of taxa. Resulting linkages were visualised in a multi-
221 dimensional scaling (MDS) plot based on replicate data from which site clusters were identified.
222 The statistical validity of the site clusters ($p < 0.05$) was tested using the similarity profile test
223 (SIMPROF; Clarke 1993).

224

225 We used a correlation-based principal components analysis (PCA) to identify environmental
226 variables (Table 1) responsible for differences among sites. The analysis was conducted on
227 normalised environmental parameters (site average) using Euclidean distance to generate the
228 resemblance matrix. The purpose of this analysis was to identify a reduced set of independent
229 environmental predictors that could be used to explain variation in faunal composition and
230 ecosystem function (here focussing on sediment oxygen consumption (SOC) and nutrient
231 regeneration (NH_4^+ and PO_4^{3-} flux)).

232

233 The correlation between the site-averaged multivariate assemblage composition (grab samples) and
234 environmental variables was examined using distance based linear models (DistLMs) in
235 PERMANOVA+ (Anderson et al. 2008). The same approach was used for measures of ecosystem
236 function except similarity matrices were based on Euclidean distance rather than Bray-Curtis
237 similarity and we also included univariate measures of macrofaunal community composition
238 (abundance, diversity and biomass from flux cores) to assess their contribution to ecosystem
239 function. Because we had macrofaunal composition from each core, solute flux data was not site-
240 averaged prior to analysis. Site A was omitted from the DistLM analyses because it was almost
241 anoxic, contained no macrofauna and nutrient fluxes were much larger than the remaining
242 hypoxic/oxic sites (see results). The only exception to this was the analysis involving the

243 relationship between meiofauna community structure and environmental variables as meiofauna
244 were present at Site A. Although DistLM is a semi-parametric, permutation-based method that does
245 not rely on normally distributed data, we checked the normality of the environmental data with
246 Shapiro-Wilks tests and no transformations were necessary. We first performed marginal tests to
247 identify strong, significant predictors, irrespective of other variables, then partial tests to assess the
248 explanatory value of a predictor variable after all other significant predictors had been accounted
249 for. Finally, DistLMs were run using the step-wise selection procedure and r^2 selection criterion to
250 identify the (linear) combinations of significant predictor variables that explained the greatest
251 proportion of variation. P values were obtained for predictor variables by 9999 permutations.
252

253 **Table 1.** Environmental variables at the study sites in the Havstensfjord and Askeröfjord, sampled in September 2011. Depth, bottom-water
 254 temperature, salinity and oxygen concentration, surface sediment (0-1 cm) silt/clay and organic content (OC), and depth of the apparent redox
 255 potential discontinuity layer (aRPD; from SPI analyses). These factors were included in the PCA analysis. In addition, the benthic habitat quality
 256 (BHQ), the corresponding BHQ stage and site groupings based on macrofaunal abundance (see Table 2) are listed.

257
 258

Site	Longitude	Latitude	Depth (m)	Temp (°C)	Salinity	O ₂ (ml L ⁻¹)	Silt/clay (%)	OC (%)	aRPD (cm)	BHQ	BHQ stage	Macrofaunal group
A	58.31482	11.77350	39.1	6.8	32.2	0.11	66.7	9.9	0.1	1.50	0	
B	58.29382	11.80400	26.3	10.6	31.0	0.89	95.0	10.4	0.3	5.00	2	1
C	58.31382	11.80182	27.0	8.5	31.5	0.99	82.4	7.9	0.2	6.75	2	1
D	58.16616	11.85200	24.2	11.3	29.4	1.02	73.7	9.8	1.7	7.50	2	2
E	58.19532	11.85000	27.5	11.0	30.0	1.20	70.7	9.1	3.7	10.75	3	3
F	58.20620	11.85116	25.8	10.4	30.1	1.23	77.7	8.8	2.7	10.25	3	3
G	58.26146	11.80730	25.6	10.2	30.6	1.28	80.7	12.0	3.2	10.25	3	3
H	58.23506	11.84782	25.2	10.4	30.4	1.34	82.4	9.4	2.3	10.75	3	2
I	58.14816	11.84116	26.9	12.4	29.8	1.36	85.1	8.6	2.3	10.50	3	4
J	58.11682	11.83050	26.7	13.8	28.6	2.11	82.9	7.7	0.7	7.00	2	4
K	58.11250	11.81950	23.2	15.1	25.1	2.91	78.6	8.4	0.7	7.25	2	4

259

260 **RESULTS**261 **Environmental variables**

262 All sites had muddy sediments with a silt/clay content > 67% and OC between 8 and 12% (Table
263 1). Bottom-water temperature varied between 7°C at the deepest site and 15°C at the shallowest site.
264 Corresponding values for salinity and O₂ concentrations were 32 and 25, and 0.1 and 2.9 ml L⁻¹,
265 respectively. Thus, even at the shallowest site, the O₂ concentration was relatively low at the time of
266 sampling, although all sites except the innermost sites (A, B, C) likely experience relatively good
267 O₂ conditions during the rest of the year (Fig. 1). The 2-dimensional PCA ordination of the
268 environmental variables (Table 1) accounted for a large fraction (72%) of the total variance and
269 revealed sites primarily dispersed across two gradients (Fig. 2, PCA analysis). PCA1 alone
270 accounted for 52% of the variance and was most strongly correlated ($|r| = 0.5-0.6$) with
271 temperature, salinity and bottom-water O₂ concentration and to a lesser extent depth ($r = 0.4$).
272 PCA2 accounted for only 20% of the inter-site variation and was driven by differences in the aRPD
273 depth and OC ($r = 0.7$ and 0.5 , respectively). Salinity and temperature were strongly correlated with
274 oxygen concentration ($|r| > 0.9$, $p < 0.001$), consequently we used a reduced set of weakly
275 correlated ($|r| < 0.6$) environmental variables (depth, O₂ concentration, OC, silt/clay, aRPD depth)
276 in subsequent DistLM analyses. Variables excluded because of co-correlation explained less of the
277 variation in response measures than the variables retained.

278

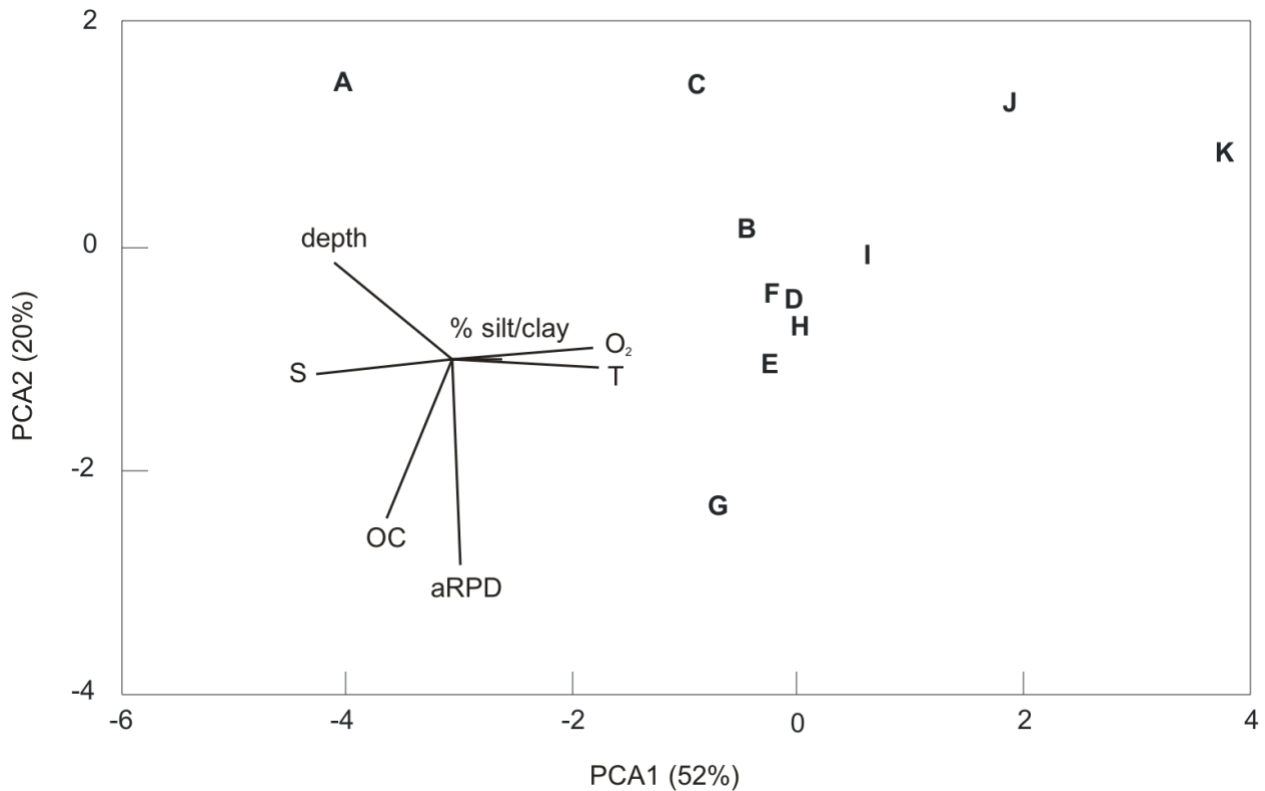
279

280

281

282

283

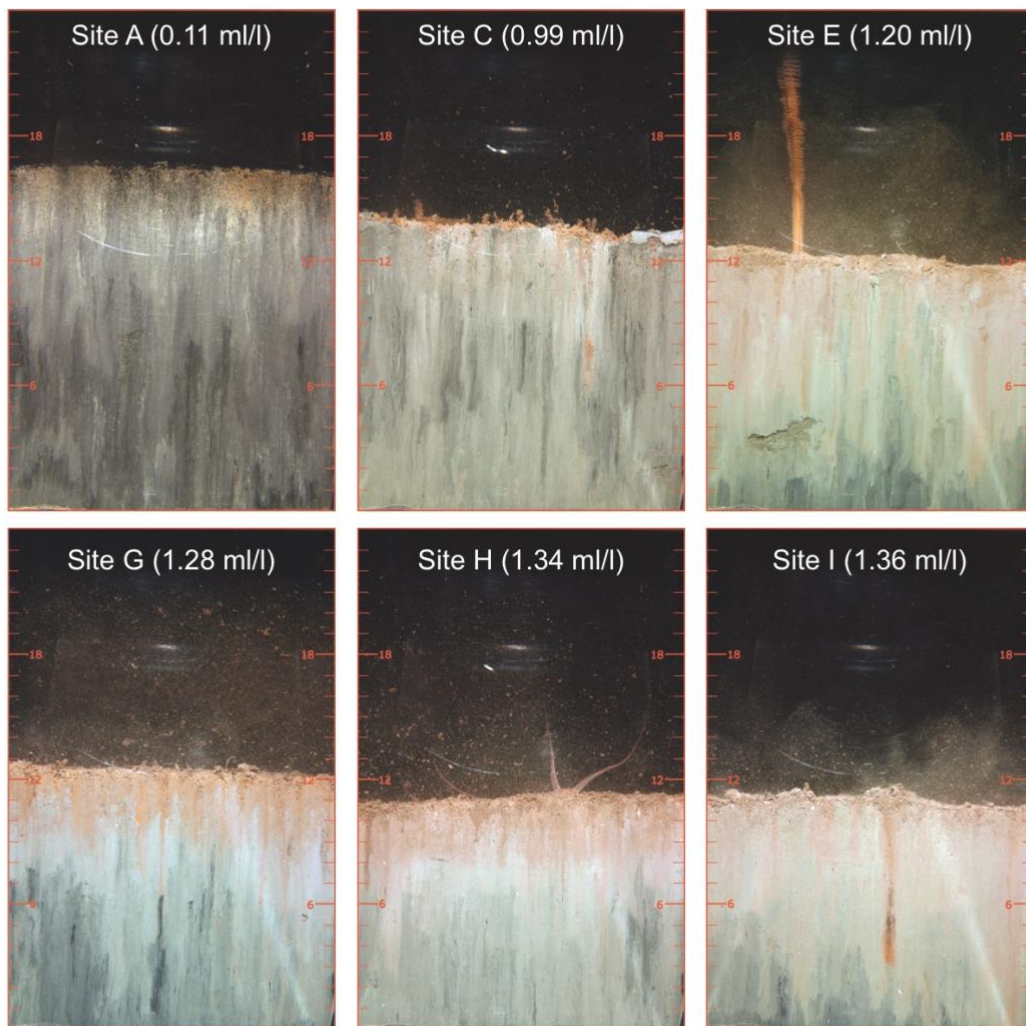


284
285
286
287
288
289
290

Figure 2. A two-dimensional ordination of a principal components analysis of site environmental variables, which collectively explained 72% of the variation between sites. Also shown is the correlation between the component axes and the environmental variables.

291 **Sediment Profile Images.** The O₂ gradient was also visible in the sediment profile images, with
292 clear differences between sites with different near-bottom O₂ concentrations, here exemplified by
293 images from almost anoxic (site A), hypoxic (sites C, E, G, H) and oxic areas (site I; Fig. 3). At the
294 almost anoxic site A, the whole sediment was reduced and large amounts of fecal pellets were seen
295 at the sediment surface. The image from site C, at about 1 ml L⁻¹ of near-bottom O₂ concentration,
296 showed no bioturbation activity and a reduced sediment with a marginally oxidised sediment
297 surface, but several small tubes were at the surface. These tubes are most likely inhabited by
298 spionids. BHQ was 5. Site E had an anthozoan, *Virgularia mirabilis*, at the well bioturbated
299 sediment surface and several vertical burrows. A feeding void was present at a depth between 8 and
300 10 cm. The mean aRPD was 2.6 cm and the BHQ index was 10. Site G demonstrated great
301 bioturbation activity at the sediment surface with some protruding tubes and several oxic burrows

302 going down into the surrounding reduced sediment. Some vertical black patches could indicate
 303 presence of dead animals. The mean aRPD was 2.0 cm and the BHQ index was 9. Similarly, site H
 304 had an ophiuroid at the sediment surface and great bioturbation activity. Many vertical tubes were
 305 stretching down to several centimetres in the sediment surrounded by reduced sediment. Mean
 306 depth of the aRPD was 2.8 cm and the BHQ index was 11. The image from site I showed some
 307 tubes at the sediment surface and some vertical oxidised burrows, where infauna is visible in one
 308 burrow. The sediment surface showed signs of great bioturbation activity, and the mean depth of the
 309 aRPD was 3.1 cm and the BHQ index was 12.



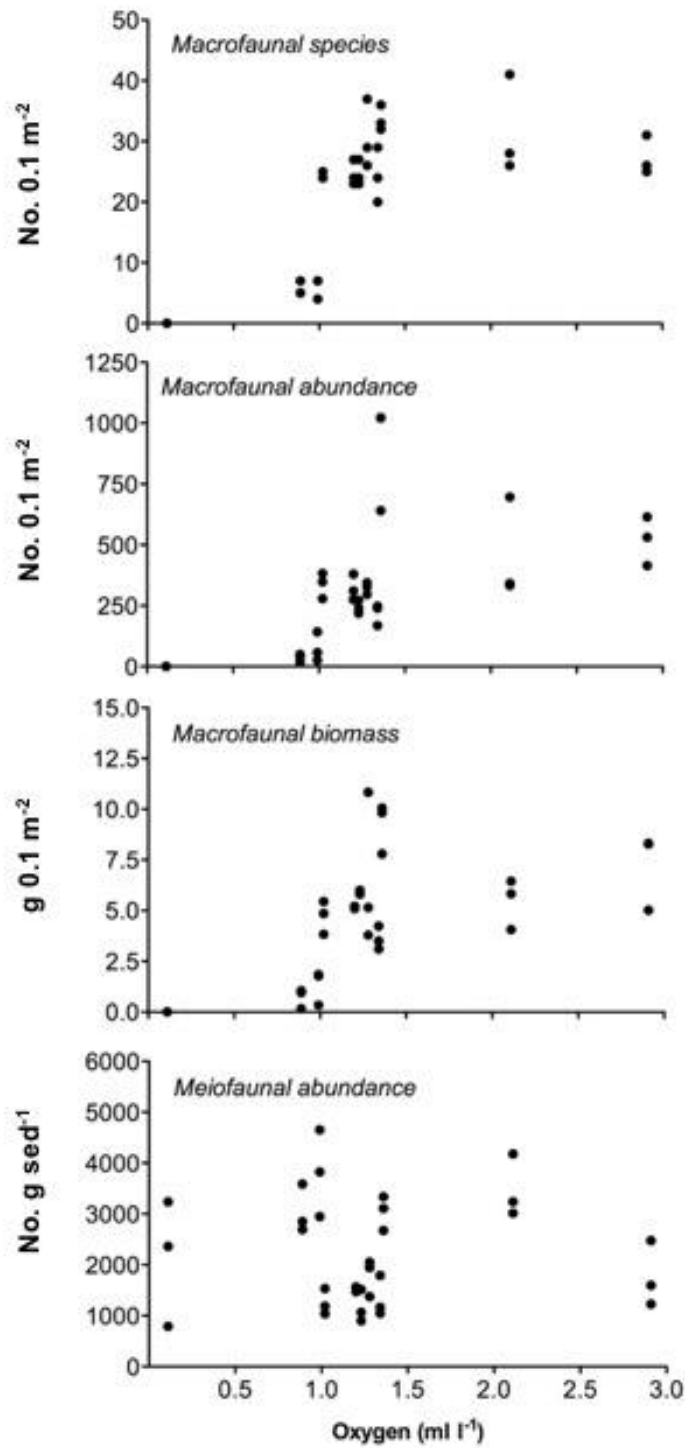
310

311 **Figure 3.** Selected sediment profile images with overlaying water from sites with different bottom-
 312 water O₂ concentrations. The colours have been digitally enhanced to facilitate interpretation. The
 313 vertical scale is in centimetres.

314
 315
 316 **Macrofauna.** In general, macrofauna species richness, abundance and biomass was greatly reduced
 317 at the sampling sites with bottom-water O₂ concentrations < 1.3 ml L⁻¹, while the highest abundance
 318 and biomass, and variability, was observed at the sites with intermediate O₂ concentration (Fig. 4).
 319 Multivariate analyses of macrofaunal abundance revealed four distinctive site clusters (confirmed
 320 by SIMPROF $p < 0.05$) that separated at 50-65% similarity (Fig. 5a). These groupings reflected
 321 changes in macrofaunal assemblage structure associated with the gradient in O₂ concentration
 322 (Table 1): severely hypoxic (0.9-1.0 ml L⁻¹, sites B, C), two hypoxic groups (1.0-1.3 ml L⁻¹, sites E,
 323 F, G and sites D, H), and oxic (1.4-2.9 ml L⁻¹, sites I, J, K). The dissimilarity between the oxic and
 324 the two hypoxic groups was approximately the same, likely driven by other co-varying spatial or
 325 environmental factors, which cannot be determined this from this dataset. The almost anoxic site A
 326 (0.1 ml L⁻¹) had no macrofauna and was excluded from this analysis. Among dominants in the
 327 hypoxic as well as oxic areas were the surface deposit-feeding bivalve *Abra nitida*, the
 328 chemosymbiotic bivalve *Thyasira flexuosa* (which can also suspension feed), the sub-surface
 329 deposit-feeding polychaete *Scalibragma inflatum* and the facultative suspension-feeding and
 330 surface deposit-feeding brittle star *Amphiura filiformis*, indicating a high level of tolerance to low
 331 O₂ concentrations in these species (Table 2). *Thyasira flexuosa* was additionally dominant even in
 332 the severely hypoxic areas. Conspicuous species in severe hypoxia were the tube building
 333 polychaetes *Maldane sarsi* and *Polydora caulleryi*, which have minor effects on the depth of the
 334 aRPD, the polychaete *Chaetozone setosa*, and the burrowing bivalve *Thyasira flexuosa*. Total
 335 abundance as well as total biomass decreased from the oxic to the severely hypoxic groups (Table
 336 2). Notable is that that the suspension-feeding bivalve *Arctica islandica* dominated the biomass,
 337 with only a few large individuals, at sites in the two hypoxic clusters (sites D, E, F, G).

338

339 In marginal tests multivariate macrofaunal assemblage structure (abundance) was most strongly
340 correlated with aRPD depth followed by silt/clay and O₂ concentration (Table 3). The other
341 variables (depth, OC) were not significantly correlated ($p > 0.2$). After correcting for the effect of
342 the other variables (i.e. in partial tests), aRPD depth, silt/clay as well as O₂ concentration remained
343 significant predictors and in linear combination collectively accounted for 60% of the variation in
344 assemblage structure.



345

346 **Figure 4.** Macrofauna species richness, abundance and biomass (per Smith-McIntyre grab, 0.1 m²),
 347 and meiofauna abundance (per gram sediment) as a function of bottom-water O₂ concentration at
 348 the sampling sites.

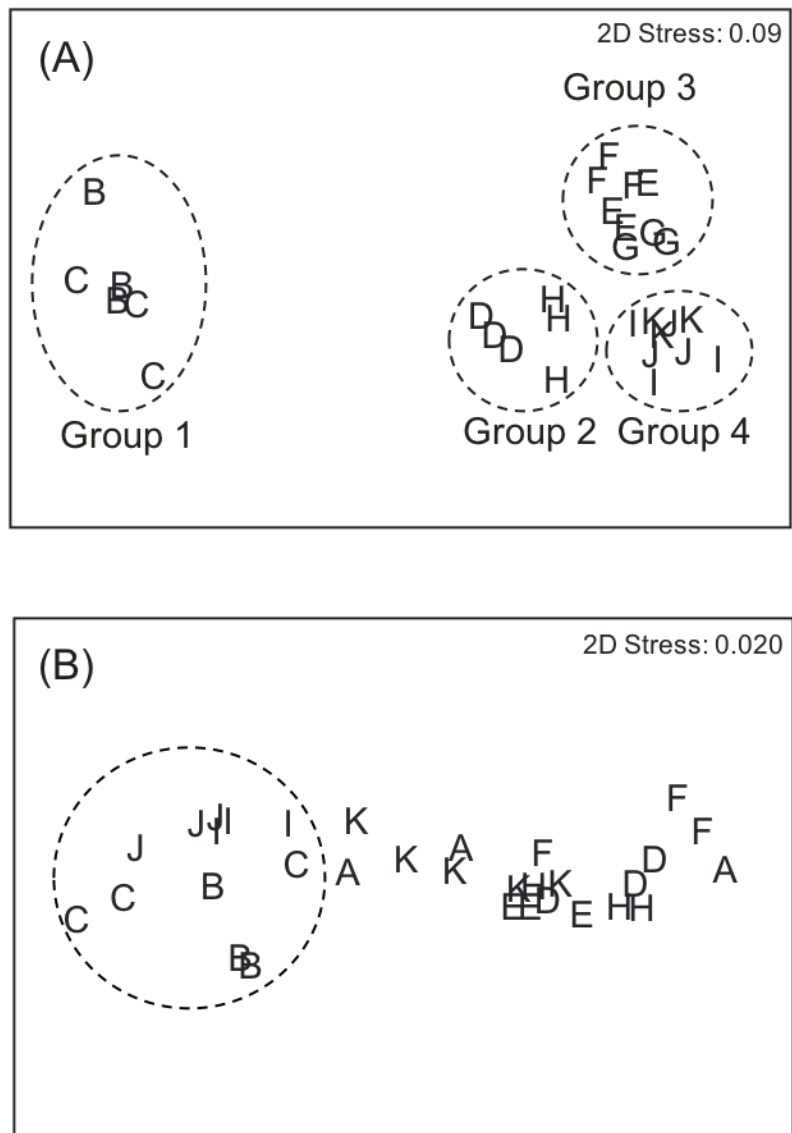
349

350

351 **Table 2.** Abundance and biomass (per 0.1 m²) of macrofaunal dominant taxa and community
 352 groupings identified as statistically distinctive site clusters using SIMPROF (based on abundance
 353 data, Fig. 5). The large bivalve *Arctica islandica* was excluded from the SIMPROF analysis, but
 354 dominated biomass, with only a few large individuals, at sites in the two hypoxic groups.
 355

Macrofaunal groupings	Group 1 Severely hypoxic	Group 2 Hypoxic	Group 3 Hypoxic	Group 4 Oxic
Abundance of dominant taxa				
<i>Scalibregma inflatum</i>		88	52	233
<i>Abra nitida</i>		22	49	188
<i>Nucula nitidosa</i>		10	9	61
<i>Thyasira flexuosa</i>	30	63	15	46
<i>Trochochaeta multisetosa</i>				17
<i>Amphiura filiformis</i>			91	10
<i>Amphiura chiajei</i>		4	20	
<i>Corbula gibba</i>		21	9	
<i>Anobothrus gracilis</i>			8	
<i>Chaetozone setosa</i>	13	16		
<i>Terebellides stroemi</i>		6		
<i>Hyala vitrea</i>		8		
<i>Heteromastus filiformis</i>		4		
<i>Maldane sarsi</i>	7			
<i>Polydora caulleryi</i>	4			
Total	54	242	253	555
Biomass of dominant taxa				
<i>Nucula nitidosa</i>		0.1		1.8
<i>Scalibregma inflatum</i>		0.6	0.3	1.5
<i>Ophiura ophiura</i>				0.9
<i>Abra nitida</i>			0.1	0.8
<i>Priapulius caudatus</i>			0.2	0.6
<i>Tubulanus polymorphus</i>				0.4
<i>Leptopentacta elongata</i>		1.3		0.2
<i>Amphiura sp.</i>			1.8	
<i>Amphiura filiformis</i>			1.3	
<i>Amphiura chiajei</i>			0.3	
<i>Thyasira flexuosa</i>	0.6	0.2		
<i>Corbula gibba</i>		0.2	0.3	
<i>Anobothrus gracilis</i>			0.1	
<i>Glycera alba</i>		0.4		
<i>Tubulanus polymorphus</i>		0.1		
<i>Phyllodoce groenlandica</i>		0.1		
<i>Maldane sarsi</i>	0.3			
Total	0.9	3	4.4	6.2

356
 357



358

359 **Figure 5.** Two dimensional MDS ordination of (A) macrofaunal (grab samples, square root
 360 transformed) and (B) meiofaunal (raw counts) community composition based on taxa abundance.
 361 Circles denote statistically distinctive site clusters determined in SIMPROF ($p < 0.05$).
 362

363 **Meiofauna.** In contrast to the macrofauna, meiofaunal total abundance changed less along the
 364 oxygen gradient (Fig. 4). Furthermore, multivariate analysis of meiofauna abundance data also did
 365 not identify distinct clusters of sites related to bottom-water O₂ concentration (Fig. 5b). For
 366 example, cluster analysis identified a group containing sites B, C, I and J (confirmed by
 367 SIMPROF), which contained in equal measures sites from the most oxic and severely hypoxic sites.
 368 Significant differences in community structure among sites was in most cases (97%) due to density

369 differences of nematodes (which was the most abundant group in all samples) and harpacticoids,
 370 however the foraminiferan group Rotaliina was also responsible in a few cases.

371

372 Multivariate analyses revealed that aRPD depth and silt/clay, but not O₂ concentration, were
 373 significant predictors also of meiofaunal assemblage structure (Table 3). The only other significant
 374 predictor in marginal tests was OC. Collectively, aRPD depth, silt/clay content and OC explained
 375 48% of the variation in meiofaunal assemblage structure. High fractions of silt/clay were found at
 376 sites B, C, H, I and J. This coincided with the highest densities of nematodes, with the exception of
 377 site H. Low levels of OC were found at site C and J, which corresponded with the highest
 378 abundances of harpacticoids. Low aRPD values were found at sites A, B and C, which matched
 379 high abundances of nematodes, with the exception of site A.

380 **Table 3.** Proportion of variation in infaunal multivariate assemblage composition (based on
 381 replicate grab sample abundance) explained by significant correlations with (site averaged)
 382 environmental variables derived from DistLMs. Marginal tests examine a single predictor
 383 separately, while partial tests take into account the effect of the remaining predictors. There was no
 384 macrofauna at site A so it was excluded from the analysis.

385

	Macrofauna		Meiofauna	
	Marginal	Partial	Marginal	Partial
Oxygen	0.26***	0.18***		
OC			0.11**	0.05*
Silt/clay	0.19***	0.05***	0.30***	0.22***
aRPD	0.28***	0.24***	0.22**	0.21***

386 * $p < 0.1$, ** $p < 0.05$, *** $p < 0.01$

387

388 Ecosystem function

389 Sediment oxygen consumption and nutrient fluxes varied strongly between sites (Table 4). For
 390 further analyses, we focussed on SOC, NH₄⁺ and PO₄³⁻. SOC increased while the PO₄³⁻ flux showed
 391 a general decrease with increasing bottom-water O₂ concentration, also when averaged between the
 392 sites corresponding to the groups identified based on the macrofaunal communities (Fig. 6). The

393 NH_4^+ flux was highly variable with no clear relationship with O_2 concentration, but similar to PO_4^{3-} ,
 394 there was a large efflux at the almost anoxic site A.

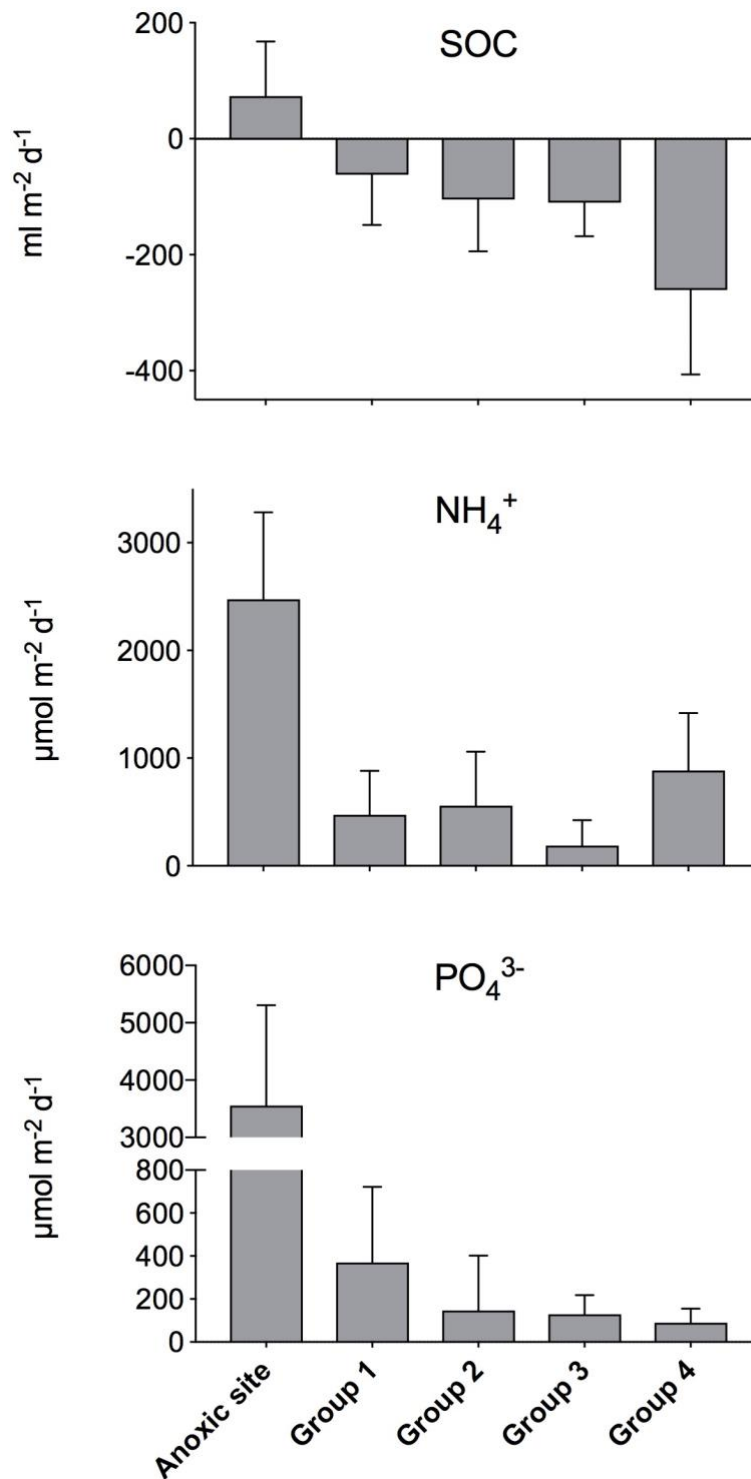
395

396 **Table 4.** Sediment O_2 consumption ($\text{ml m}^{-2} \text{d}^{-1}$) and nutrient fluxes ($\mu\text{mol m}^{-2} \text{d}^{-1}$) across the
 397 sediment-water interface at the study sites in the Havstensfjord and Askeröfjord, sampled in
 398 September (average \pm SD, five replicate flux cores per site). Negative values denote an influx into
 399 the sediment. Unfortunately, the NH_4^+ samples from sites B, E and J went missing during analysis.
 400

Site	O_2	NO_3^-	NO_2^-	NH_4^+	PO_4^{3-}	SiO_4
A	73 ± 94	8 ± 65	1 ± 20	2473 ± 809	3554 ± 1756	3128 ± 2544
B	-48 ± 66	-1138 ± 128	-50 ± 65	-	298 ± 162	2092 ± 908
C	-79 ± 117	752 ± 2219	57 ± 116	473 ± 409	442 ± 490	6323 ± 6034
D	-109 ± 37	-737 ± 110	-98 ± 19	910 ± 425	268 ± 108	3173 ± 688
E	-124 ± 21	-440 ± 241	-130 ± 17	-	199 ± 89	2495 ± 514
F	-61 ± 66	-571 ± 228	-110 ± 14	132 ± 304	91 ± 52	1789 ± 769
G	-149 ± 34	-1179 ± 208	-176 ± 31	244 ± 164	112 ± 100	3130 ± 254
H	-100 ± 128	-911 ± 2521	-86 ± 154	208 ± 278	25 ± 314	1022 ± 6574
I	-227 ± 76	-1323 ± 235	-178 ± 8	1274 ± 401	130 ± 82	5188 ± 1143
J	-124 ± 45	-116 ± 63	-76 ± 28	-	97 ± 51	2273 ± 899
K	-431 ± 74	80 ± 178	-13 ± 14	494 ± 315	43 ± 35	3103 ± 1842

401

402 In the DistLM we used the same four independent site environmental variables as above for the
 403 community analyses (depth, O_2 concentration (core specific, i.e. concentration at start of
 404 incubation), aRPD, OC and silt/clay), and the following faunal variables: core specific macrofauna
 405 abundance, biomass (less *Arctica*) and Shannon diversity and the site-averaged meiofaunal
 406 abundance and Shannon diversity. We did not include the number of taxa as a predictor variable
 407 because for macrofauna it was highly correlated with abundance (Pearson's $r > 0.8$) and the number
 408 of meiofauna taxa did not vary among sites. Site A was omitted from analyses because of the
 409 absence of macrofauna and predominantly chemically driven changes to measured fluxes,
 410 particularly for NH_4^+ and PO_4^{3-} . Unfortunately, the NH_4^+ samples from sites B, E and J went
 411 missing during analysis and so the DistLM analysis were conducted on a reduced set of sites.
 412 Fortunately, the missing sites were scattered across the oxygen gradient.



413

414 **Figure 6.** Sediment oxygen consumption (SOC) and effluxes of NH_4^+ and PO_4^{3-} (average \pm SD) in
 415 the site groupings identified based on multivariate analyses of macrofaunal abundances (Fig. 5),
 416 where Group 1 = severely hypoxic, Group 4 = Oxic. In addition, data from the almost anoxic site
 417 (site A) is included.

418

419 O₂ concentration was the single best predictor of SOC closely followed by macrofaunal abundance,
420 both being positively correlated to SOC (Table 5). Depth, macrofaunal biomass and meiofaunal
421 diversity were also correlated with SOC in marginal tests however in partial tests, after fitting other
422 significant predictors first, they did not explain a significant amount of the residual variation. The
423 single best linear combination of variables included O₂ concentration and macrofaunal abundance
424 which collectively explained 63% of the variation in SOC. Inclusion of the remaining predictors
425 only explained an additional 3% of the variability.

426

427 The reduced number of sites included in the analysis of NH₄⁺ flux restricted the environmental
428 gradient and as a consequence no water or sediment environmental variables were significant
429 predictors. Macrofaunal biomass was the single best predictor of NH₄⁺ flux followed by
430 macrofaunal abundance then meiofaunal abundance, all being positively correlated to the NH₄⁺ flux
431 (Table 5). In partial tests both macrofaunal biomass and meiofaunal abundance were still
432 significant, but macrofaunal abundance was not due it being correlated (albeit weakly) with
433 biomass. Macrofaunal biomass and meiofaunal abundance collectively explained 30% of the
434 variation in NH₄⁺ flux.

435

436 In marginal tests the PO₄³⁻ flux was best correlated with O₂ concentration with higher fluxes
437 associated with lower O₂ concentrations (Table 5). Phosphate flux was weakly correlated with most
438 faunal predictors. Interestingly, while the flux was negatively correlated with measures of
439 macrofaunal community composition it was positively correlated with meiofaunal indices. This
440 relationship is explained in part by decline in macrofaunal abundance/biomass with decreasing O₂
441 concentration. In partial tests only O₂ concentration, macrofaunal biomass and meiofaunal
442 abundance were significant predictors of phosphate flux and combined explained 28% of the
443 variability in the data.

444 **Table 5.** Proportion of variation in sediment oxygen consumption (SOC), NH₄₊ and PO₄₃₋ flux
 445 explained by significant correlations with faunal/environmental variables (the direction is given in
 446 brackets) derived from DistLMs. Marginal tests examine a single predictor separately, while partial
 447 tests take into account the effect of the remaining predictors. There was no macrofauna at site A so
 448 it was excluded from the analysis.
 449

	SOC		NH ₄₊		PO ₄₃₋	
	Marginal	Partial	Marginal	Partial	Marginal	Partial
Depth	0.23***(-)	0.02				
Oxygen	0.57***(+)	0.14***			0.13**(-)	0.07**
Macro-d	0.41***(+)	0.05**	0.13**(+)	0.04	0.10**(-)	<0.01
Macro-b	0.07*(+)	0.02	0.23***(+)	0.06*	0.06* (-)	0.05*
Macro-H'					0.11**(-)	<0.01
Meio-d			0.11*(+)	0.09*	0.11**(+)	0.07**
Meio-H'	0.13**(+)	0.01			0.13**(+)	0.02

450 Macro-d = abundance of macrofauna core-1, Macro-b = g ww core-1, Macro-H' the Shannon diversity, Meio-d =
 451 meiofaunal abundance g sediment-1
 452 **p* < 0.1, ***p* < 0.05, ****p* < 0.01
 453
 454

455 DISCUSSION

456 The deleterious effects of hypoxia on macrofaunal communities are well known and our results
 457 from the seasonally hypoxic Havstensfjord and Askeröfjord corroborate this pattern, with a general
 458 decrease in macrofaunal species richness, abundance and biomass with decreasing bottom-water O₂
 459 concentration. Meiofaunal communities on the other hand did not appear to be similarly affected by
 460 hypoxia (Table 3). Bottom-water O₂ concentration was the most important factor explaining
 461 variation in sediment oxygen consumption (SOC) and fluxes of NH₄₊ and PO₄₃₋. Nevertheless, after
 462 the key gradient in oxygen had been accounted for, macrofauna and meiofauna explained a small
 463 but significant fraction of the variation in ecosystem function. This implies that faunal burrowing
 464 and feeding indeed affect nutrient cycling, also when bottom-water concentrations are low, i.e. this
 465 is not a purely geochemically driven process. For example, the burrowing enhances the oxygenation
 466 of the sediment, thereby reducing the PO₄₃₋ efflux. The nearly anoxic site A was markedly different
 467 to the other sites, with no macrofauna and with massive effluxes of PO₄₃₋ and NH₄₊ from the

468 sediment (up to an order of magnitude higher fluxes than at other sites), indicating a complete shift
469 to geochemically-driven functioning.

470

471 The patterns of structural changes in macrofaunal communities following organic enrichment and
472 ensuing hypoxia have been well known for several decades (Pearson and Rosenberg 1978), but the
473 link to changes in ecosystem functioning remains elusive. The SPI analysis showed that the depth
474 of the aRPD was shallowest at the almost anoxic site and generally deeper at more oxic sites, but
475 not consistently (Table 1). The aRPD can indeed be used as a proxy for the transition between
476 redox states (Rosenberg et al. 2001, Simone and Grant 2017), but while it was a significant
477 predictor of both macrofauna and meiofauna (Table 3; although macrofauna activity is likely also
478 affecting the depth of the aRPD, so the interaction works both ways), it was not an important
479 predictor for SOC or fluxes of NH_4^+ or PO_4^{3-} (Table 5). Neither did the BHQ index correspond to the
480 nutrient fluxes, as it was a too coarse measure, with all our sites (except A) classified to the two
481 highest successional stages (BHQ 5-10 and >10). The multivariate analysis of macrofauna
482 community abundance, on the other hand, split these sites into four groups (in addition to site A).
483 There were indications that the four site clusters identified had a relationship with SOC and PO_4^{3-}
484 fluxes (Fig. 6), with increasing SOC and decreasing PO_4^{3-} flux towards the more oxic groups. The
485 four most abundant species were common in three of the four groups identified, indicating a high
486 level of tolerance to hypoxia. These species live buried in the sediment and have a generation time
487 of one year or more, and none of them are among the first colonizers in a succession pattern like
488 *Capitella* and *Polydora*. It is possible that such tolerant species might uphold bioturbation
489 throughout the year in seasonally hypoxic areas, although seasonal patterns in species-specific
490 bioturbation activity are not known. The number of sites (11 in total, 2-3 sites per group) did not,
491 however, allow analysing in detail any between-group differences in the different factors explaining
492 the variability of the fluxes. Future studies are needed to address this.

493

494 The Havstensfjord has suffered from seasonal hypoxia since the 1950s and the most likely cause is
495 eutrophication (Nilsson and Rosenberg 1997). Given the long history of hypoxic stress, the benthic
496 communities are probably adapted to this disturbance, but also likely permanently somewhat
497 degraded compared with communities that have not been exposed to hypoxia. Undisturbed
498 communities would likely be characterised by higher species richness, and large-bodied and long-
499 lived species (Pearson and Rosenberg 1978, Diaz and Rosenberg 1995, Gray et al. 2002), with a
500 potential for a stronger, more direct impact on nutrient cycling. The bottom-water O₂ concentration
501 was indeed relatively low at all sites at the time of sampling. The benthic habitat quality in relation
502 to hypoxia has previously (in 1994) been studied in the fjord using SPI (Nilsson and Rosenberg
503 1997), and no major changes in aRPD or further degradation of the BHQ index were observed in
504 the current study. Thus, the Havstensfjord appears to have been in a stable state of seasonal hypoxia
505 for decades already, with periods of good conditions every year. The big individuals of the bivalve
506 *Arctica islandica* found in the two hypoxic groups indicate that there had not been long periods of
507 complete anoxia at these sites in the last few years before the sampling, although these bivalves can
508 reduce metabolism and survive several months in hypoxic conditions. Given the proportionately
509 larger influence on nutrient cycling of such large individuals (Norkko et al. 2013), these few
510 bivalves may have a significant effect on ecosystem functioning in-between the hypoxic periods.

511

512 Fjords and other enclosed inlets and seas are prone to hypoxia because of limited water exchange.
513 Once a system has passed the threshold to hypoxia, the reversal might be difficult (Conley et al.
514 2009b), and there is evidence that once an ecosystem experiences hypoxia, it might be more
515 susceptible to hypoxia in the future (Conley et al. 2007, Diaz and Rosenberg 2008). This poses
516 challenges for the management of these systems. Another methodological challenge is often
517 introduced as part of the monitoring of hypoxia. We used the O₂ concentration measured 2-3 cm

518 above the sediment surface in flux chambers immediately upon core retrieval for the statistical
519 analyses. It is important to consider the difference between these concentrations and the ones
520 measured 1 m (or even higher with big CTD rosettes) above the sediment surface in most
521 monitoring programmes (Rosenberg 1977). Due to the benthic boundary layer processes, where
522 decreasing flow closer to the sediment surface usually corresponds to decreasing O₂ concentrations
523 (e.g., Jørgensen and Des Marais 1990), the monitoring data does not necessarily reflect the real
524 near-bottom O₂ concentrations, which hampers modelling efforts to predict species distribution
525 patterns in relation to O₂ conditions (Virtanen et al. 2019).

526
527 Temperature, salinity, and near-bottom O₂ concentrations were strongly correlated and therefore
528 only O₂ was included in the statistical analysis. Since all three variables can affect community
529 structure as well as ecosystem function, we cannot rule out the potential impact of co-variables, but
530 we cannot disentangle these effects from our field sampling because the hypoxic water was deep,
531 beneath the thermocline and therefore salty and/or cold. However, the lack of oxygen will most
532 likely override the effects of the other two on nutrient cycling, as it stresses the fauna as well as
533 alters biogeochemical pathways. While oxygen dropped below detrimental levels in this study,
534 neither the range in salinity (25-32) nor temperature (7-15) is likely to be a major driver of the
535 differences in function observed, given that this is an estuarine environment with organisms adapted
536 to fluctuations. It should also be noted, that in this relatively cool fjord system organisms are more
537 likely to oxyregulate and the effects of hypoxia may be therefore be less stressful compared to
538 warmer systems, where organisms are more likely to experience metabolic depression, i.e.
539 oxyconformation (Pörtner et al. 2005). Temperature does, however, affect nutrient fluxes, with
540 higher reaction rates at warmer temperatures. In our study, the bottom-water temperature varied
541 from 7 to 15 degrees, and the incubations were done at a standard 11 degrees, so it is possible that

542 the fluxes at the shallowest and the deepest site were slightly underestimated and overestimated,
543 respectively. This would, however, not have affected the main conclusions.

544

545 The importance of macrofauna for explaining variability in SOC and nutrient fluxes in the current
546 study was similar to that found across gradients of increasing hypoxia by Gammal et al. (2017) and
547 (Norkko et al. 2015), in the coastal and open Baltic Sea, respectively. This indicates that the number
548 of species is not crucial for functioning; the Baltic is very species poor (open sea often only 5-7
549 species) compared with the Swedish west coast (up to 40 species recorded per grab in the current
550 study). This is contrary to our prediction that the effect of hypoxia on the faunal contribution to
551 functioning would be smaller in a system with a higher background diversity. The number of
552 macrofaunal species was highly correlated with the macrofaunal abundance, further indicating that
553 the number of species is not the sole driving factor for functioning. Indeed, in previous studies
554 under normoxic conditions it has been suggested that species-specific traits of some particularly
555 important individual species, for example, the sediment reworking by large and abundant burrowing
556 urchins or bivalves, may override both species richness and functional diversity in terms of
557 influencing benthic nutrient cycling (Lohrer et al. 2004, Norling et al. 2007, Norkko et al. 2013).

558

559 Of particular interest is then whether the fauna can still contribute to ecosystem functioning when
560 conditions deteriorate, i.e. before the fauna is decimated, but when physiological and behavioural
561 changes are already likely to have been initiated. Hypoxia tolerance is highly species specific
562 (Vaquer-Sunyer and Duarte 2008), with potential for changes in behaviour already at relatively high
563 O₂ concentrations. For example, urchin grazing rates dropped already at 5.5 mg/L (3.85 ml L⁻¹),
564 with potential consequences on kelp recruitment (Low and Micheli 2018). Macrofauna can still
565 influence nutrient cycling in low O₂ conditions, although not as much as under good O₂ conditions
566 (Norkko et al. 2015). This points to the increasing importance of tolerant species when conditions

567 deteriorate, e.g. the invasive polychaete *Marenzelleria* spp. in the Baltic Sea (Norkko et al. 2015).
568 Such deep-burrowing species may also affect the propensity for rapid oxygen consumption by
569 burying organic matter deeper into the sediment, which slows down the oxidation of OM (Josefson
570 et al. 2012). Other studies have also found that larger-bodied, tolerant species may mediate
571 ecosystem functioning during periods of low-oxygen conditions (Rakocinski and Menke 2016). In
572 the present study the large bivalve *Arctica islandica* may have performed this role. Nevertheless,
573 the hypoxia-induced changes in species behaviour that is influencing nutrient cycling are not
574 sufficiently known and this requires further mechanistic studies.

575

576 **Meiofauna, microbes and altered energy pathways**

577 The differences in meiofaunal communities were due to the silt/clay fraction, OC and aRPD, not
578 differences in O₂ concentrations, which was the major factor explaining differences in the
579 macrofauna. Various taxa of meiofauna can use aerobic metabolism at O₂ concentrations as low as
580 0.1 µmol/l (Giere 2009). In addition, the non-correlation to O₂ concentration has been noted before,
581 by e.g. Josefson and Widbom (1988), who recorded no major decrease of any meiofaunal taxa
582 during a several month long period of anoxia in the Gullmar Fjord, Sweden. In the Baltic Sea,
583 Elmgren (1975) noted that meiofauna did decrease with decreasing oxygen conditions, but that the
584 meiofauna extended deeper into the hypoxic zone compared with the macrofauna. In the current
585 study the aRPD, which approximates the depth of the oxygenated sediment layer, also significantly
586 affected the differences in meiofaunal community structure, and thus O₂ is probably still involved in
587 the vertical distribution of the meiofauna, possibly through indirect effects on macrofaunal
588 bioturbation. Sediments with high silt/clay fractions have a lower permeability compared to coarser
589 sediments, which directly influences the amount of oxygenated water that can penetrate into the
590 sediment. Large abundances of nematodes were found at the sites with a high silt/clay fraction (B,
591 C, H, I, J), which is a common representation. The nematode community is then often made up of

592 non-selective deposit-feeding nematodes grazing on bacteria and detritus (Heip et al. 1985,
593 Boeckner et al. 2009, Delgado et al. 2009). Larger grain size tends to lead to more harpacticoids;
594 however they can thrive in various conditions (Giere 2009).

595

596 At site B and J high levels of NH_4^+ were registered. It is also at these sites the highest abundances of
597 meiofauna were found. This relationship can be stressed further using site H, which had low NH_4^+
598 while at the same time holding one of the lowest total meiofaunal abundances of the analyzed sites.
599 The high ammonium at site B and C can furthermore be explained by a reduced bioturbation under
600 the anoxic conditions at those sites. This can in its turn affect the nitrification and denitrification;
601 instead of nitrogen being removed as N_2 in denitrification processes, ammonia, ammonium and
602 phosphate are the main fluxes out of the sediment (Diaz and Rosenberg 2008). In addition,
603 meiofauna is known not to control the abundance of the microbial community with their grazing,
604 but to enhance the growth rate (Piot et al. 2014). In general, high OC leads to higher abundances of
605 meiofauna but at the sites in this survey the relationship was reversed. Nevertheless, no site had
606 very low OC (range 7.7 – 12%) and OC was likely not a limiting factor for the meiofauna.

607

608 Prolonged hypoxia will have larger-scale functional consequences, with miniaturization of benthic
609 food webs and altered energy pathways (the last successional stages 1 and 0) (Diaz and Rosenberg
610 2008). For example, deep-burrowing species which are important for organic matter processing,
611 priming it for further processing by the microbes, will be lost (Table 2), translating also into a loss
612 of functional spaces and a reduction of surface area. While our study design did not allow for
613 quantifying this, it is possible that meiofaunal and microbial communities will have a
614 proportionately larger influence on ecosystem function in stressed systems compared with
615 macrofauna. The resistance of sediment microbial communities to hypoxia is however poorly
616 known, but field experiments with *in situ* -induced hypoxia indicated that microbial diversity

617 decreased with increasing duration of hypoxic stress, concurrently with the deteriorating
618 macrofaunal community, likely due to the reduced macrofaunal bioturbation activity (Sinkko et al.
619 submitted).

620

621 **Conclusions and management implications**

622 Many studies on BEF have failed to demonstrate real-world relevance, highlighting the need to
623 combine insights derived from theory with detailed experiments and broad-scale monitoring and
624 well-designed field surveys (Snelgrove et al. 2014). While causality cannot be assigned in
625 correlative field studies, they are imperative for understanding the generality of the BEF
626 relationships. They need to be conducted in a range of different environments and geographical
627 areas, using the same methodology and focussing on the relative similarities or differences. There
628 are however inherent logistic constraints to field work, as the number of sites that it is possible to
629 include in a field study is almost always limited.

630

631 Ecosystem management decisions are based on model predictions of key ecosystem processes.
632 While our understanding of the links between biodiversity and ecosystem functioning, also under
633 increasing hypoxia, is growing, there are still large gaps in our knowledge about these processes
634 and how they can be included in the models, with significant effects on model performance
635 (Carstensen et al. 2014). For example, while there are sophisticated models used for nutrient
636 management around the entire Baltic Sea (BALTSEM; Savchuk et al. 2012), macrofauna data is
637 currently lacking altogether from these models, which is a serious drawback for being able to
638 predict, for example, the time required for recovery of the sea from eutrophication. Indeed our
639 failure to account for biodiversity in models of biogeochemical cycling is severely impeding our
640 ability to understand, quantify and predict, the consequences of changes in eutrophication status and
641 climate as expressed through altered carbon pathways (Snelgrove et al. 2018). This work thus needs

642 to continue with field studies targeting macrofauna, meiofauna as well as microbes linked with
643 controlled laboratory studies where specific mechanisms can be quantified, conducted in parallel
644 with model development. This will become increasingly important as we grapple with trying to
645 protect the dwindling biodiversity and trying to predict future changes in ecosystem functioning.

ACKNOWLEDGEMENTS

The study was supported by an ASSEMBLE grant (European Community, ASSEMBLE grant agreement 227799), the BONUS+ HYPER project (FP7/2007-2013, grant agreement 217246), the Walter and Andrée de Nottbeck Foundation, Victoriastiftelsen and the Academy of Finland (project ID 294853). We thank the crew on-board R/V Skagerak for help with sampling, the Sven Lovén Centre for Marine Sciences, Kristineberg, for laboratory facilities, and Pia Engström for help with preparations.

Role of the funding source

The funders had no influence on the execution of the study or the preparation of this article.

REFERENCES

- Aller, RC, JY Aller. 1998. The effect of biogenic irrigation intensity and solute exchange on diagenetic reaction rates in marine sediments. *J. Mar. Res.* **56**:905-936.
- Anderson, MJ, RN Gorley, KR Clarke. 2008. PERMANOVA+ for PRIMER. Guide to software and statistical methods.
- Boeckner, MJ, J Sharma, HC Proctor. 2009. Revisiting the meiofauna paradox: dispersal and colonization of nematodes and other meiofaunal organisms in low- and high-energy environments. *Hydrobiologia* **624**:91-106.
- Bonaglia, S, FJA Nascimento, M Bartoli, I Klawonn, V Bruchert. 2014. Meiofauna increases bacterial denitrification in marine sediments. *Nature Communications* **5**.
- Breitburg, D, LA Levin, A Oschlies, M Gregoire, FP Chavez, DJ Conley, V Garcon, D Gilbert, D Gutierrez, K Isensee, GS Jacinto, KE Limburg, I Montes, SWA Naqvi, GC Pitcher, NN Rabalais, MR Roman, KA Rose, BA Seibel, M Telszewski, M Yasuhara, J Zhang. 2018. Declining oxygen in the global ocean and coastal waters. *Science* **359**:eaam7240.

- Burgess, R. 2001. An improved protocol for separating meiofauna from sediments using colloidal silica sols. *Mar. Ecol. Prog. Ser.* **214**:161-165.
- Carstensen, J, DJ Conley, E Bonsdorff, BG Gustafsson, S Hietanen, U Janas, T Jilbert, A Maximov, A Norkko, J Norkko, DC Reed, CP Slomp, K Timmermann, M Voss. 2014. Hypoxia in the Baltic Sea: Biogeochemical cycles, benthic fauna and management. *Ambio* **43**:26-36.
- Chapin III, FS, BH Walker, RJ Hobbs, DU Hooper, JH Lawton, OE Sala, D Tilman. 1997. Biotic control over the functioning of ecosystems. *Science* **277**:500-504.
- Clarke, KR. 1993. Non-parametric multivariate analyses of changes in community structure. *Aust. J. Ecol.* **18**:117-143.
- Clarke, KR, RN Gorley. 2006. PRIMER v6: User Manual / Tutorial. PRIMER-E, Plymouth.
- Conley, DJ, S Björck, E Bonsdorff, J Carstensen, G Destouni, BG Gustafsson, S Hietanen, M Kortekaas, H Kuosa, HEM Meier, B Müller-Karulis, K Nordberg, A Norkko, G Nürnberg, H Pitkänen, NN Rabalais, R Rosenberg, OP Savchuk, CP Slomp, M Voss, F Wulff, L Zillén. 2009a. Hypoxia-related processes in the Baltic Sea. *Environ. Sci. Technol.* **43**:3412-3420.
- Conley, DJ, E Bonsdorff, J Carstensen, G Destouni, BG Gustafsson, L-A Hansson, NN Rabalais, M Voss, L Zillén. 2009b. Tackling hypoxia in the Baltic Sea: Is engineering a solution? *Environ. Sci. Technol.* **43**:3407-3411.
- Conley, DJ, J Carstensen, G Ærtebjerg, PB Christensen, T Dalsgaard, JLS Hansen, AB Josefson. 2007. Long-term changes and impacts of hypoxia in Danish coastal waters. *Ecol. Appl.* **17**:S165-S184.
- Conley, DJ, J Carstensen, J Aigars, P Axe, E Bonsdorff, T Eremina, B-M Haahti, C Humborg, P Jonsson, J Kotta, C Lännergren, U Larsson, A Maximov, M Rodriguez Medina, E Lysiak-Pastuszak, N Remeikaite-Nikiene, J Walve, S Wilhelms, L Zillén. 2011. Hypoxia is increasing in the coastal zone of the Baltic Sea. *Environ. Sci. Technol.* **45**:6777-6783.

- Cowan, JLW, WR Boynton. 1996. Sediment-water oxygen and nutrient exchanges along the longitudinal axis of Chesapeake Bay: Seasonal patterns, controlling factors and ecological significance. *Estuaries* **19**:562-580.
- Delgado, JD, R Riera, Ó Monterroso, J Núñez. 2009. Distribution and abundance of meiofauna in intertidal sand substrata around Iceland. *Aquat. Ecol.* **43**:221-233.
- Diaz, RJ, R Rosenberg. 1995. Marine benthic hypoxia: a review of its ecological effects and the behavioural responses of benthic macrofauna. *Oceanogr. Mar. Biol. Annu. Rev.* **33**:245-303.
- Diaz, RJ, R Rosenberg. 2008. Spreading dead zones and consequences for marine ecosystems. *Science* **321**:926-929.
- Elmgren, R. 1975. Benthic meiofauna as indicator of oxygen conditions in the northern Baltic Proper. *Merentutkimuslaitoksen julkaisuja* **239**:265-271.
- Friedrich J, F Janssen, D Aleynik, HW Bange, NBoltacheva, MN Çagatay, AW Dale, G Etiope, Z Erdem, M Geraga, A Gilli, MT Gomoiu, POJ Hall, D Hansson, Y He, M Holtappels, MK Kirf, M Kononets, S Konovalov, A Lichtschlag, DM Livingstone, G Marinaro, S Mazlumyan, S Naeher, RP North, G Papatheodorou, O Pfannkuche, R Prien, G Rehder, CJ Schubert, T Soltwedel, S Sommer, H Stahl, EV Stanev, A Teaca, A Tengberg, C Waldmann, B Wehrli, F Wenzhöfer. 2014. Investigating hypoxia in aquatic environments: diverse approaches to addressing a complex phenomenon. *Biogeosciences* **11**:1215-1259.
- Gammal, J, J Norkko, CA Pilditch, A Norkko. 2017. Coastal hypoxia and the importance of benthic macrofauna communities for ecosystem functioning. *Estuar. Coast.* **40**:457-468.
- Giere, O. 2009. *Meiobenthology: The microscopic motile fauna of aquatic sediments*. 2nd edition. Springer-Verlag, Berlin.
- Glud, RN. 2008. Oxygen dynamics of marine sediments. *Mar. Biol. Res.* **4**:243-289.
- Gray, JS, RSS Wu, YY Or. 2002. Effects of hypoxia and organic enrichment on the coastal marine environment. *Mar. Ecol. Prog. Ser.* **238**:249-279.

- Halpern, BS, S Walbridge, KA Selkoe, CV Kappel, F Micheli, C D'Agrosa, JF Bruno, KS Casey, C Ebert, HE Fox, R Fujita, D Heinemann, HS Lenihan, EMP Madin, MT Perry, ER Selig, M Spalding, R Steneck, R Watson. 2008. A global map of human impact on marine ecosystems. *Science* **319**:948-952.
- Hansson, D, A Stigebrandt, B Liljebadh. 2013. Modelling the Orust fjord system on the Swedish west coast. *J. Mar. Syst.* **113**:29-41.
- Heip, C, M Vincx, G Vranken. 1985. The ecology of marine nematodes. *Oceanogr. Mar. Biol. Annu. Rev.* **23**:399-489.
- Ingall, ED, RM Bustin, P Van Cappellen. 1993. Influence of water column anoxia on the burial and preservation of carbon and phosphorus in marine shales. *Geochim. Cosmochim. Acta* **57**:303-316.
- Jäntti, H, S Hietanen. 2012. The effects of hypoxia on sediment nitrogen cycling in the Baltic Sea. *Ambio* **41**:161-169.
- Jørgensen, BB, DJ Des Marais. 1990. The diffusive boundary layer of sediments: Oxygen microgradients over a microbial mat. *Limnol. Oceanogr.* **35**:1343-1355.
- Josefson, AB, J Norkko, A Norkko. 2012. Burial and decomposition of plant pigments in surface sediments of the Baltic Sea – role of oxygen and benthic fauna. *Mar. Ecol. Prog. Ser.* **455**:33-49.
- Josefson, AB, B Widbom. 1988. Differential response of benthic macrofauna and meiofauna to hypoxia in the Gullmar Fjord basin. *Mar. Biol.* **100**:31-40.
- Levin, LA, DF Boesch, A Covich, C Dahm, C Erséus, KC Ewel, RT Kneib, A Moldenke, MA Palmer, P Snelgrove, D Strayer, JM Weslawski. 2001. The function of marine critical transition zones and the importance of sediment biodiversity. *Ecosystems* **4**:430-451.
- Levin, LA, W Ekau, AJ Gooday, F Jorissen, JJ Middelburg, SWA Naqvi, C Neira, NN Rabalais, J Zhang. 2009. Effects of natural and human-induced hypoxia on coastal benthos. *Biogeosciences* **6**:2063-2098.

- Levinton, J. 1995. Bioturbators as ecosystem engineers: Control of the sediment fabric, inter-individual interactions, and material fluxes. Pages 29-38 in C. G. Jones and J. H. Lawton, editors. Linking species and ecosystems. Chapman & Hall, New York.
- Lindgren, JF, IM Hasselov, I Dahllof. 2013. Analyzing changes in sediment meiofauna communities using the image analysis software ZooImage. *J. Exp. Mar. Biol. Ecol.* **440**:74-80.
- Lohrer, AM, SF Thrush, MM Gibbs. 2004. Bioturbators enhance ecosystem function through complex biogeochemical interactions. *Nature* **431**:1092-1095.
- Low, NHN, F Micheli. 2018. Lethal and functional thresholds of hypoxia in two key benthic grazers. *Mar. Ecol. Prog. Ser.* **594**:165-173.
- McCarthy, MJ, KS McNeal, JW Morse, WS Gardner. 2008. Bottom-water hypoxia effects on sediment-water interface nitrogen transformations in a seasonally hypoxic, shallow bay (Corpus christi bay, TX, USA). *Estuar. Coast.* **31**:521-531.
- Meysman, FJR, JJ Middelburg, CHR Heip. 2006. Bioturbation: a fresh look at Darwin's last idea. *Trends Ecol. Evol.* **21**:688-695.
- Middelburg, JJ, LA Levin. 2009. Coastal hypoxia and sediment biogeochemistry. *Biogeosciences* **6**:1273-1293.
- Mortimer, CH. 1941. The exchange of dissolved substances between mud and water in lakes. *J. Ecol.* **29**:280-329.
- Nilsson, HC, R Rosenberg. 1997. Benthic habitat quality assessment of an oxygen stressed fjord by surface and sediment profile images. *J. Mar. Syst.* **11**:249-264.
- Norkko, A, A Villnäs, J Norkko, S Valanko, CA Pilditch. 2013. Size matters: implications of the loss of large individuals for ecosystem function. *Sci. Rep.* **3**:2646.
- Norkko, J, J Gammal, JE Hewitt, AB Josefson, J Carstensen, A Norkko. 2015. Seafloor ecosystem function relationships: In situ patterns of change across gradients of increasing hypoxic stress. *Ecosystems*:1-16.

- Norling, K, R Rosenberg, S Hulth, A Grémare, E Bonsdorff. 2007. Importance of functional biodiversity and species-specific traits of benthic fauna for ecosystem functions in marine sediment. *Mar. Ecol. Prog. Ser.* **332**:11-23.
- Pearson, TH, R Rosenberg. 1978. Macrobenthic succession in relation to organic enrichment and pollution of the marine environment. *Oceanogr. Mar. Biol. Annu. Rev.* **16**:229-311.
- Piot, A, C Nozais, P Archambault. 2014. Meiofauna affect the macrobenthic biodiversity-ecosystem functioning relationship. *Oikos* **123**:203-213.
- Pörtner, HO, M Langenbuch, B Michaelidis. 2005. Synergistic effects of temperature extremes, hypoxia, and increases in CO₂ on marine animals: From Earth history to global change. *J. Geophys. Res.* **110**:C09S10.
- Rabalais, NN, W-J Cai, J Carstensen, DJ Conley, B Fry, X Hu, Z Quiñones-Rivera, R Rosenberg, CP Slomp, RE Turner, M Voss, B Wissel, J Zhang. 2014. Eutrophication-driven deoxygenation in the coastal ocean. *Oceanography* **27**:172-183.
- Rabalais, NN, RE Turner, WJ Wiseman. 2002. Gulf of Mexico hypoxia, aka "The dead zone". *Annu. Rev. Ecol. Syst.* **33**:235-263.
- Rakocinski, CF, DP Menke. 2016. Seasonal hypoxia regulates macrobenthic function and structure in the Mississippi Bight. *Mar. Poll. Bull.* **105**:299-309.
- Reed, DC, CP Slomp, BG Gustafsson. 2011. Sedimentary phosphorus dynamics and the evolution of bottom water hypoxia: A coupled benthic-pelagic model of a coastal system. *Limnol. Oceanogr.* **56**:1075-1092.
- Rosenberg, R. 1977. Benthic macrofaunal dynamics, production, and dispersion in an oxygen-deficient estuary of west Sweden. *J. Exp. Mar. Biol. Ecol.* **26**:107-133.
- Rosenberg, R, S Agrenius, B Hellman, HC Nilsson, K Norling. 2002. Recovery of marine benthic habitats and fauna in a Swedish fjord following improved oxygen conditions. *Mar. Ecol. Prog. Ser.* **234**:43-53.

- Rosenberg, R, HC Nilsson, RJ Diaz. 2001. Response of benthic fauna and changing sediment redox profiles over a hypoxic gradient. *Estuar. Coast. Shelf Sci.* **53**:343-350.
- Savchuk, OP, BG Gustafsson, B Müller-Karulis. 2012. BALTSEM - a marine model for decision support within the Baltic Sea Region.
- Simone, M, J Grant. 2017. Visual assessment of redoxcline compared to electron potential in coastal marine sediments. *Estuar. Coast. Shelf Sci.* **188**:156-162.
- Slomp, CP, J Thomson, GJ de Lange. 2002. Enhanced regeneration of phosphorus during formation of the most recent eastern Mediterranean sapropel (S1). *Geochim. Cosmochim. Acta* **66**:1171-1184.
- Snelgrove, PVR, K Soetaert, M Solan, S Thrush, C-L Wei, R Danovaro, RW Fulweiler, H Kitazato, B Ingole, A Norkko, RJ Parkes, N Volkenborn. 2018. Global carbon cycling on a heterogeneous seafloor. *Trends Ecol. Evol.* **33**:96-105.
- Snelgrove, PVR, SF Thrush, DH Wall, A Norkko. 2014. Real world biodiversity–ecosystem functioning: a seafloor perspective. *Trends Ecol. Evol.* **29**:398-405.
- Vaquer-Sunyer, R, CM Duarte. 2008. Thresholds of hypoxia for marine biodiversity. *Proceedings of the National Academy of Sciences* **105**:15452-15457.
- Villnäs, A, A Norkko. 2011. Benthic diversity gradients and shifting baselines: implications for assessing environmental status *Ecol. Appl.* **21**:2172-2186.
- Villnäs, A, A Norkko, J Norkko, K Lukkari, J Hewitt. 2012. Impacts of increasing hypoxic disturbance on benthic biodiversity and ecosystem functioning. *PLoS ONE* **7**:e44920.
- Villnäs, A, J Norkko, S Hietanen, AB Josefson, K Lukkari, A Norkko. 2013. The role of recurrent disturbances for ecosystem multifunctionality. *Ecology* **94**:2275-2287.
- Virtanen, EA, A Norkko, A Nyström Sandman, M Viitasalo. 2019. Identifying areas prone to coastal hypoxia – the role of topography. *Biogeosciences*, accepted for publication.

Yachi, S, M Loreau. 1999. Biodiversity and ecosystem productivity in a fluctuating environment:

The insurance hypothesis. *Proceedings of the National Academy of Sciences* **96**:1463-1468.

Yazdani Foshtomi, M, U Braeckman, S Derycke, M Sapp, D Van Gansbeke, K Sabbe, A Willems,

M Vincx, J Vanaverbeke. 2015. The link between microbial diversity and nitrogen cycling in

marine sediments is modulated by macrofaunal bioturbation. *PLOS ONE* **10**:e0130116.