Effects of bottom trawling and hypoxia on benthic invertebrate communities

van Denderen, P Daniel; Tornroos, Anna; Sciberras, Marija; Hinz, Hilmar; Friedland, Rene; Lasota, Rafal; Mangano, Maria Cristina; Robertson, Craig; Valanko, Sebastian; Hiddink, Jan Geert

Marine Ecology Progress Series

DOI: 10.3354/meps14094

Published: 11/08/2022

Peer reviewed version

Cyswllt i'r cyhoeddiad / Link to publication

Dyfyniad o'r fersiwn a gyhoeddwyd / Citation for published version (APA): van Denderen, P. D., Tornroos, A., Sciberras, M., Hinz, H., Friedland, R., Lasota, R., Mangano, M. C., Robertson, C., Valanko, S., & Hiddink, J. G. (2022). Effects of bottom trawling and hypoxia on benthic invertebrate communities. *Marine Ecology Progress Series*, 694, 13-27. https://doi.org/10.3354/meps14094

Hawliau Cyffredinol / General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

· Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
 You may freely distribute the URL identifying the publication in the public portal ?

Take down policy If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Effects of bottom trawling and hypoxia on benthic invertebrate communities.

3

4	P. Daniël van Denderen ^{1,2*} , Anna Törnroos ³ , Marija Sciberras ⁴ , Hilmar Hinz ⁵ , Rafal Lasota ⁶ ,
5	Maria Cristina Mangano ^{7,8} , Craig Robertson ⁹ , Sebastian Valanko ¹⁰ , Jan Geert Hiddink ⁹
6	
7	¹ Centre for Ocean Life, DTU Aqua, Technical University of Denmark, Lyngby, Denmark
8	² Graduate School of Oceanography, University of Rhode Island, Narragansett, USA
9	³ Environmental and Marine Biology, Faculty of Science and Engineering, The Sea research
10	profile, Åbo Akademi University, 20520 Turku, Finland.
11	⁴ The Lyell Centre, Heriot-Watt University, Edinburgh, UK, EH14 4AP
12	⁵ Mediterranean Institute for Advanced Studies, IMEDEA (UIB-CSIC), Marine Ecosystem
13	Dynamics, Esporles, Balearic Islands, Spain
14	⁶ University of Gdansk, Faculty of Oceanography and Geography, Institute of Oceanography,
15	Department of Marine Ecosystems Functioning, 81-378 Gdynia, Poland
16	⁷ Department of Integrative Marine Ecology (EMI), Stazione Zoologica Anton Dohrn, Sicily
17	⁸ Marine Centre, Lungomare Cristoforo Colombo (complesso Roosevelt) 90149, Palermo, Italy
18	⁹ Bangor University, School of Ocean Sciences, Menai Bridge, UK, LL59 5AB
19	¹⁰ International Council for the Exploration of the Sea, Copenhagen, Denmark
20	
21	Running title (max 6 words): Benthic impacts from trawling and hypoxia

22 *Corresponding author: e-mail: <u>pdvd@aqua.dtu.dk</u>

23 Abstract

Marine benthic habitats in continental shelf regions are increasingly impacted by hypoxia 24 25 caused by the combination of eutrophication and climate warming. Many regions that have the potential for hypoxic conditions are being fished by mobile bottom-contacting fishing gears. 26 The combined effects of trawling and hypoxia may be synergistic and disproportionally impact 27 28 benthic fauna, or they may act antagonistically, leading to smaller trawl impacts in hypoxic areas. Yet, few studies have quantified how bottom trawling and hypoxia interact to effect 29 benthic communities. Here we examine these combined effects on benthic community biomass 30 and abundance, the number of large organisms, the longevity distribution of the community 31 and the vertical position of fauna in the sediment in the southern Baltic Sea. Our results show 32 large declines in benthic biomass and abundance with declines in near-bed oxygen 33 concentrations from 5.8 to 0.8 ml O₂ l⁻¹. Conversely, no effects and weak positive effects of 34 bottom trawl disturbances are found on the benthic community. No interacting effects between 35 hypoxia and trawling are detected. Our findings therefore highlight a low likelihood of 36 synergistic impacts of bottom trawling and hypoxia on the benthic communities studied. These 37 results suggest that management may prioritize benthic protection from fishing in regions that 38 are not in a state of oxygen stress. 39

40

41 Keywords: bottom-trawling, hypoxia, Baltic Sea, macrofauna, body size, longevity, vertical
42 position in sediment

43 **1. INTRODUCTION**

Marine benthic habitats in continental shelf regions are increasingly impacted by 44 eutrophication-induced hypoxia (Diaz & Rosenberg 2008). Hypoxic regions are predicted to 45 expand due to the warming of shelf waters with climate change, which increases both the 46 oxygen demand and the strength of stratification of the water column (Breitburg et al. 2018). 47 48 Many periodic hypoxic regions, such as the Baltic and Adriatic Seas, are currently also being fished by mobile bottom-contacting fishing gears, and these bottom trawl fisheries are the most 49 widespread source of anthropogenic physical disturbance to seabed habitats (Amoroso et al. 50 2018). 51

Since management of nutrient loads and subsequent declines of hypoxic regions will take 52 53 decades to centuries to occur, strategic management of bottom fisheries may be used to 54 alleviate some of the impacts on benthic fauna and the ecosystem. Such strategic management requires quantifying the cumulative impacts of the two pressures on benthic fauna. The 55 56 cumulative impacts of trawling and hypoxia may be synergistic when hypoxic conditions make organisms more sensitive to bottom fishing disturbance. Such effects may occur in areas where 57 hypoxic and anoxic bottoms result in upward migration of burrowing organisms to the sediment 58 surface (Pihl et al. 1992, Nilsson & Rosenberg 1994), thereby making these organisms more 59 60 vulnerable to damage by bottom gears dragged along the seabed surface. Synergistic effects 61 have also been proposed for brittle stars that became more sensitive to trawl disturbance due to arm-tipping behaviour, i.e. the elevation of the central disk to escape the low oxygen 62 concentrations closest to the seafloor (Baden et al. 1990, Diaz & Rosenberg 1995). Under this 63 64 scenario, management measures that protect hypoxic or hypoxia-prone areas from fishing may thus disproportionally benefit benthic fauna by reducing fishing-derived mortality of hypoxia 65 66 sensitive fauna, and thus lowering the risk of benthic habitats being pushed into a permanently altered state (Riedel et al. 2016). Alternatively, when hypoxia leads to asphyxiation of sessile 67

fauna and/or migration of mobile fauna, leading to a depauperate state or even absence of benthic fauna, trawling may have a negligible additional impact on the benthos. Animals that can survive hypoxia events may also have physical characteristics that make them less vulnerable to trawling and trawling may not affect these species significantly. Fisheries management measures in such areas will have only limited benefits to the benthic ecosystem.

The above highlights that depending on the cumulative effects of bottom trawling and hypoxia, 73 management of human impacts on benthic habitats may either prioritize protection from 74 fisheries of hypoxic areas or prioritize the protection of areas that are non-hypoxic. So far, few 75 studies have quantified the interactive effects of bottom trawling and hypoxia on benthic 76 communities, limiting the development of management plans that consider the cumulative 77 effects. Recent work on a continental slope off Vancouver Island found that bottom trawling 78 continues to impact benthic communities in hypoxic areas, suggesting cumulative impacts that 79 are either additive or synergistic (De Leo et al. 2017). 80

81 Here we study the effects of bottom trawling and hypoxia on benthic communities in the Baltic Sea. Throughout its historic past the Baltic Sea has seen hypoxic events on the seafloor (natural 82 and human induced), mainly due to limited water exchange with the Atlantic and in recent 83 decades excess nutrient input (Carstensen et al. 2014, Andersen et al. 2017). The Baltic Sea's 84 85 benthic communities lack large habitat forming epifauna, in particular in the deeper areas. This 86 is due to a low salinity, that in combination with its historic hypoxic past, has created the characteristically low species diverse, but highly abundant benthic communities (Bonsdorff 87 2006). Most of these benthic communities consist of short-lived fauna (Törnroos et al. 2015, 88 89 2019, van Denderen et al. 2019), which are predicted to be largely resilient to bottom trawling disturbance (Hiddink et al. 2018). Yet, approximately two thirds of the Baltic Sea bottom trawl 90 fishing footprint occurs in areas with seasonal oxygen concentrations <3.2 ml O₂ l⁻¹ that may 91 impact benthic organisms (van Denderen et al. 2019). It is therefore an area where trawling 92

93 impacts may be severely underestimated if synergistic effects are found to occur. The objective of this study is to estimate interactive effects of bottom trawling and hypoxia on macrofaunal 94 biomass and abundance, large-bodied macrofauna, benthic community longevity and the 95 vertical position of fauna in the sediment. The studied stations ranged over a gradient in near-96 bed oxygen concentrations from 0.8 to 5.8 ml $O_2 l^{-1}$ and in fishing intensity from little trawl 97 disturbance to up to 7 bottom trawl events per year. Our findings highlight a low likelihood of 98 synergistic effects between bottom trawling and hypoxia on the Baltic Sea benthic communities 99 studied. 100

101 **2. MATERIAL AND METHODS**

102 **2.1 Study area**

The effects of bottom trawling and hypoxia were examined over a gradient of oxygen and bottom trawling intensity in the Southern Baltic Sea in Polish waters (Fig. 1). The area is an active fishing ground with bottom trawling by otter trawls that target cod and flounder. The area was selected as earlier work showed it has gradients in oxygen concentration and bottom trawling intensity (van Denderen et al. 2019).

108 We selected sampling stations that covered a wide range of expected oxygen concentrations and trawling intensities on muddy sediment in a limited depth range (61 - 102 metres, Table 109 S1-1), and aimed to cover all crossed combinations of trawling and oxygen to end up with a 110 111 design where the effects of trawling and oxygen could be disentangled. Unsurprisingly, we 112 could not identify any stations that had a high fishing effort at very low oxygen, as no fish can survive in those conditions, but we did sample several stations with fishing and relatively low 113 oxygen concentrations (Fig. 2 and Table S1-1). All sampling stations were found in an area of 114 30 by 62 km, and the distance to the nearest station varied between 3 and 18 km. 115

116 **2.2 Sampling of benthic fauna**

117 Sampling was carried from on board the research vessel RV Oceanograf between 3 and 8 September 2018. Five replicate box corers and two replicate dredge samples were collected 118 from each station. Box corer samples (catching area 0.06 m² per core) were taken in a star 119 pattern, one at the selected position and four 100 m in each direction, using the vessel's 120 Dynamic Positioning System. At times, the box corer came up empty or lost most of the sample 121 122 before retrieval because the jaws did not close fully when pebbles or stones jammed the mechanism. When sampling was unsuccessful, the ship was moved 10s of meters and the box 123 core redeployed. Three 4.5 cm diameter sub-cores up to a maximum depth of 17 cm were 124

collected from the first two intact box core samples collected at each station to assess the 125 vertical distribution of fauna in the sediment in areas of different oxygen and fishing levels. 126 127 Each sub-core was sliced into four sections (0-2 cm, 2-4 cm, 4-8 cm, >8 cm) and sieved to obtain the fauna at different sediment depths. The rest of the box core sample was sieved over 128 0.5 mm sieve. Most samples had a solid clay layer below a surficial mud, sand and/or gravel 129 layer (surficial layer was ca. 8-18 cm deep); the clay layer was not sieved as no macrofauna 130 131 can and did occur in this solid layer. To separate the smaller fauna from the remaining gravel and stones, samples were back-washed at least 7 times to obtain the fauna, by adding water, 132 133 stirring the sample and draining the water over a sieve to capture suspended animals. The remaining sediment was checked for any further large fauna. 134

Semi-quantitative dredge samples were obtained using an Agassiz trawl type dredge (56 wide 135 x 31 cm high, outer net mesh size 1 x 1 cm, inner net mesh size 0.5 x 0.5 cm), except for Station 136 1 (see below). The dredge was towed for 2 minutes at 1 knot and was generally full on retrieval. 137 138 Samples were searched visually for fauna. Station 1 was sampled with a Naturalist dredge (60 cm wide x 30 cm high) that was towed for 10 and 5 minutes at 1 knot. We expect that the 139 alternative methodology did not affect the resulting data in Station 1 as, in all cases, the dredge 140 filled up quickly and stopped sampling afterwards. Inspection of the data does not suggest the 141 dredge data in Station 1 is an obvious outlier. 142

All biota were identified to the lowest taxonomic level possible and their individual wet weight was measured. For dredge samples, where species abundance was greater than approximately 100 individuals, a subsample was taken per species. In the subsample, length was measured for all individuals whereas weight was measured for 50 individuals of a range of lengths. For the remaining individuals, we converted length to weight using the established length-weight relationship. We afterwards calculated abundance and biomass of the total sample based on the relative size of the subsample.

150 **2.3 Environmental conditions**

A sub-core measuring 4 cm in diameter core by 4 cm in depth was collected from the first box 151 core sample collected at each station to determine sediment grain size composition. Particle 152 size distributions were determined using a Malvern laser diffraction particle sizer (Blott & Pye 153 2001). Sediment particle analysis showed that most of the stations had a high percentage mud 154 (Table S1-1). Visual inspection of the box cores showed that the muddy sediment was 155 156 sometimes mixed with pebbles, gravel or sand and typically with a deeper layer of glacial clay. Some stations had ferromanganese nodules on the seafloor surface. A principal component 157 158 analysis on depth and sediment conditions shows that stations 5 and 16 could be outliers due to a relatively high sand / gravel fraction, respectively (Fig. S1-1). Results showed similar 159 patterns and trends when these stations were excluded from the analysis. It was therefore 160 decided to maintain these stations within the analysis, despite small differences in sediment 161 composition. 162

163 Bottom water was collected at 20 cm above the seabed with a Niskin bottle and oxygen saturation (YSI model 58) and temperature and salinity (WTW Multiset 340i with TetraCon 164 325) were measured from three replicate water samples collected at each station (with the 165 exception of the first few stations where only a single measurement was taken) (Table S1-1). 166 Temperature and salinity information was used to convert oxygen saturation into oxygen 167 168 concentration in ml per litre following Benson and Krause (1984). A large fraction of stations was azoic, with concentrations below 1 ml O₂ l⁻¹ and with a strong smell of H₂S and a clear 169 black surface layer. In addition to the oxygen measurements taken at the sampling stations, we 170 analysed a time series of bottom oxygen concentration measurements collected from a nearby 171 site at 90 m depth (red asterisk in Fig. 1) between 2012 and 2018. Monthly bottom oxygen 172 concentration was calculated by averaging oxygen measurements at depths less than five 173

meters from the seafloor per month (where available). Data used were obtained from HELCOM
secretariat and based on ICES dataset on Ocean Hydrography.

Bottom trawling intensity data were obtained at a 0.05° x 0.05° grid resolution (ICES 2019), and expressed as the ratio of the area that is swept each year by trawl gears relative to the total surface area (hereafter termed swept-area-ratio, SAR). We used average SAR values per grid cell for the period 2013-2017, as we were interested in chronic effects of fishing and since recovery following trawling disturbance may last more than one year.

181 **2.4 Data analysis**

We analysed the effect of trawling and oxygen concentrations on community biomass and 182 183 abundance, the number of large organisms, the biomass-longevity distribution of the benthic 184 community and the vertical position of fauna in the sediment. In all analyses, we estimated 185 biomass and abundance as the sum of all box core or dredge samples per station. All results show biomass of bivalves with shell. Since the biomass patterns are strongly driven by bivalve 186 weights, we verified that similar results are obtained when shell-free wet weight is used (based 187 on a wet weight to shell-free wet weight conversions as shown in Table S1-2). The dataset, R 188 scripts and output are available on Github with DOI: 10.5281/zenodo.5579600. 189

190 <u>Community biomass and abundance</u>

We tested the effect of trawling and oxygen and their interaction on community biomass and abundance with a Tobit regression model and selected the best model using the Akaike Information Criterion (AIC); when models differ less than 2 AIC-units, the model with fewest parameters is selected. Tobit regression is designed to estimate relationships assuming a normal distribution for the response variable with left/right censoring, i.e. where a threshold value is set to the response variable above/below the response variable data are censored (Tobin 1958).We used Tobit regression as it was expected that a range of sampling stations at low 198oxygen concentrations had zero biomass and abundance and we therefore set the threshold for199censoring our data at zero biomass and abundance. The community biomass and abundance200analyses were done considering all stations, to examine the combined effect of trawling and201oxygen concentration, as well as for a subset of stations with oxygen concentrations > 3 ml O2202 l^{-1} to examine the effect of bottom trawling in isolation (Figure 2). The oxygen threshold value203was chosen since benthic biomass and abundance no longer correlate with oxygen above this204value.

205 <u>Large organisms</u>

Since previous work showed that the impact of bottom trawling is larger on larger benthic organisms (>4 mm) (McLaverty et al. 2020), we included a separate analysis to study the effect of oxygen and trawling on the number of large organisms in the benthic community. The analysis used the same Tobit regression model as described above. We used two length thresholds: >4 mm and >15 mm. We estimated individual length using length-weight relationships from the literature (Table S1-3).

212 <u>Community longevity</u>

Previous work has further shown that long-lived fauna are more impacted by bottom trawling 213 (Rijnsdorp et al. 2018, Hiddink et al. 2018). We therefore examined the effect of trawling and 214 oxygen on the longevity biomass distribution of the benthic community. We used the box core 215 216 data to study changes in the biomass-longevity composition as these data provide a more controlled estimate of biomass (the dredge filled up quickly, as noted above). Using 217 information on species longevity of benthic fauna in the Baltic Sea (Törnroos & Bonsdorff 218 219 2012, van Denderen et al. 2019), we classified biota in four longevity groupings: maximum lifespan of 0-1 years, 1-3 years, 3-10 years and > 10 years. The resulting taxon-by-trait matrix 220 was combined with the taxon biomass-by-station (gram wet weight per 0.3 m^2) matrix to create 221 the final station-by-trait matrix. We used a PerMANOVA analysis to examine if changes in 222

trawling intensity and/or oxygen concentrations had a significant effect on the distribution of benthic biomass in these four longevity groupings. This analysis was done on absolute and fractional biomass values per longevity grouping and sampled station.

226 Vertical position

We also used a PerMANOVA analysis to examine if changes in oxygen concentrations and trawling intensity affected the vertical distribution of fauna in the sediment. Since the number of observations was low in the vertical sub-cores, we limited our analysis to examining community changes in the vertical position using the abundance and biomass data, whereas we did not examine species-specific vertical changes. The analyses were done on absolute and fractional abundance/biomass data per vertical section and sampled station.

3. RESULTS

3.1 Temporal and spatial stability in fishing and near-bed oxygen

We used average fishing intensities per grid cell for the period 2013-2017 to examine the 235 chronic effects of fishing. A cross-comparison of fishing intensities between all years shows 236 clear correlations, with an average Pearson product-moment correlation of 0.89. These strong 237 238 correlations show there is temporal stability in the spatial fishing patterns studied. The studied gradient in fishing intensities is conserved in 2018, the year of sampling (Pearson correlation 239 of 0.85 when compared with the annual average for the period 2013-2017), although the 240 average level of fishing in the area in 2018 is marginally lower (average SAR in 2018 for all 241 stations is 1.0, whereas it varies between 1.3 and 2.9 in the other years). 242

The time series of near-bed oxygen concentrations (Fig. 3a) shows large fluctuations between months, with maximum concentrations close to 5 and minimum less than 0.5 ml O_2 l⁻¹. The median near-bed oxygen concentrations are lower in summer and autumn (Q3 and Q4, Fig. 3b), although very low oxygen concentrations have been found in all seasonal quarters. The time series highlights that there are significant uncertainties in the temporal history of near-bed oxygen in our sampled stations (see further *Discussion* section).

249 **3.2** Community biomass and abundance

The biomass of the community is dominated by the bivalves *Astarte elliptica*, *Mytilus trossulus* and *Limecola balthica* (formerly known as *Macoma balthica*). Together these species comprised 90% of total biomass in the box core data and 99% in the dredge. Abundance is dominated by the bristleworm *Scoloplos armiger* (27% of total abundance) in the box core data and by *A. elliptica* (81% of total abundance) in the dredge. No biota is found at stations with oxygen concentrations below 2 ml 1^{-1} . All stations with oxygen concentrations above 3 ml 1^{-1} had some fauna.

We find a significant positive relationship between oxygen and both community biomass and 257 abundance for both box core and dredge data (Fig. 4). In two of the four datasets (biomass and 258 259 abundance from the dredge), trawling intensity has a significant positive effect (Table 1). An interaction term between oxygen and trawling intensity does not provide a better fit to any 260 model (note that for box core abundance the interaction term has a p-value of 0.05, but the AIC 261 value differs less than 2 units from a model without the interaction, Table 1). When analysing 262 263 a subset of sampled stations with relatively high oxygen concentrations (> 3 ml l^{-1}), trawling has no relationship with community biomass or abundance (Fig. 5). 264

To explore further the community response to trawling, we examined changes in biomass for 265 the six most dominant species (in terms of biomass) in the box core samples and the five most 266 dominant in the dredge samples collected at stations with relatively high oxygen concentrations 267 $(> 3 \text{ ml } 1^{-1})$. It is worth noting that all species have a non-significant relationship with oxygen 268 in this subset (not shown). We find no significant relationship between biomass and trawling 269 270 intensity for most dominant species, except for the priapulid worm *Halicryptus spinulosus* where a significant positive relationship is observed between its biomass and trawling in both 271 dredge and box core data (Fig. 6). 272

273 **3.3 Large organisms**

Individuals larger than 4 mm comprised >99% of total biomass and 85% of total abundance. 274 275 For these individuals (n = 1587), we find a significant positive relationship between oxygen 276 and abundance and between trawling intensity and abundance (Table 1). Individuals larger than 15 mm comprised 30% of total biomass and 6% of total abundance. For these individuals (n =277 278 110), there is again a significant positive relationship between oxygen and abundance, whereas tobit regression indicates no support for a model that includes trawling (Table 1). The dominant 279 280 species contributing to the 'larger than 15 mm' group are the bivalve A. *elliptica* (n = 34), the isopod Saduria entomon (n=19) and the bristleworm Terebellides stroemii (n=16). The number 281

of individuals larger than 15 mm varies among stations between 0 and 21 per 0.3 m² (the sum of the five box cores); high numbers are only found in areas with relatively high oxygen concentrations (> $3.5 \text{ ml } l^{-1}$) (Fig. 7).

285 **3.4 Community longevity**

Almost 90% of the total biomass is grouped as fauna with longevities between 3 and 10 years and no species are found with lifespans more than 10 years. PerMANOVA analysis shows no support for a change in the biomass longevity composition of the community with changes in oxygen and/or trawling intensity (Table S1-4).

290 **3.5 Vertical position**

In total, 143 individuals are found in the vertical sub-cores across the stations. Most fauna (102 291 292 individuals, representing 90% of total biomass) are in the upper 0-2 cm of the sediment. Different polychaete species (Pygospio elegans, Aricidea catherinae, Nereis diversicolor and 293 Streblospio benedicti), nematodes and ostracods are found in low abundance at depths deeper 294 than 4 cm (in total 11 individuals, representing 1% of total biomass). PerMANOVA analysis 295 shows no support for a change in the vertical distribution of fauna with changes in oxygen 296 297 and/or trawling intensity (Table S1-5). The vertical biomass and abundance distribution is for each species shown in Figure S1-2 and S1-3. 298

299 **4. DISCUSSSION**

We examined the effects of bottom trawl disturbance and oxygen on benthic community 300 301 biomass and abundance, the number of large-bodied individuals, community longevity and the vertical position of fauna in the sediment. Our results show that changes in near-bed oxygen 302 concentrations have a large impact on benthic fauna, whereas trawling had a weak positive 303 304 effect on some species and community metrics and no effect on others. The lack of negative effects of trawling on most parameters, and a positive effect on some community metrics and 305 the priapulid worm Halicryptus spinulosus, are somewhat unexpected given the high annual 306 trawling frequencies at some sampling stations. We discuss below whether these non-negative 307 effects may be a true effect or the result of different confounding factors. Whatever the reason, 308 our findings highlight that bottom trawling appears to have limited impacts on the benthic 309 communities studied here. 310

311 4.1 Non-negative trawl effects on benthic fauna

Bottom trawl impacts depend on the frequency of trawl disturbance, the amount of mortality 312 (depletion) that bottom gears cause and the recoverability of the benthic community (Pitcher 313 314 et al. 2017). The fisheries data shows that all trawling in the area occurs with a demersal otter trawl. This is a relatively light gear that is associated with a low depletion rate of 0.03, i.e. 315 causing a biomass decline of 3% each time a benthic community is swept by this trawl (Hiddink 316 et al. 2017, Rijnsdorp et al. 2020). All fauna in the studied region are further relatively short 317 lived (<10 years), potentially the result of selection by environmental filtering due to low 318 salinity and variable oxygen levels (Bonsdorff 2006). Short-lived species have relatively high 319 320 recovery rates (Hiddink et al. 2018). Therefore, low trawl impacts may be expected in this area. 321 Nonetheless, model predictions based on the observed longevities and the above depletion rate suggest a 20% decline of community biomass relative to carrying capacity in a location with a 322 chronic fishing frequency of SAR = 7 a year (see SI 2 for the calculation). Although this is a 323

substantial effect, the variation in the recorded benthic communities is large due to naturalvariation and sampling error, and the power to detect such an effect may have been limited.

326 Biomass comparisons with the Kattegat and the more northern Gotland Basin reveal that box core biomass in our study area is on average 6 to 10 times lower than in those areas (Sköld et 327 al. 2018, van Denderen et al. 2019). Since low oxygen conditions tend to affect relatively large 328 329 areas homogenously, the low biomass in our study area may be the result of low and variable 330 oxygen levels that have hindered recruitment and recolonization from neighbouring regions (Whitlatch et al. 1998). Recruitment in the Baltic Sea is also more stochastic and dependent on 331 wind-induced waves since it is a non-tidal system (Valanko et al. 2010). It may therefore be 332 hypothesized that any survivors and/or new arrivals in the area will have little competition for 333 space and food and high growth rates, thus resulting in lower trawl impacts than expected. 334

335 The large fluctuations in oxygen concentration may also have limited our ability to detect negative bottom trawl effects given that we sampled at only one point in time. Some sampled 336 337 stations with high oxygen measurements during our survey may have experienced lower levels in earlier months that would have killed a large fraction of the biota, causing a large amount of 338 noise on the trawl impact signal. Nonetheless, all areas with relatively high oxygen 339 concentrations were found to have some large-bodied (> 15 mm) individuals in the box core 340 341 (Fig. 7). This suggests that the historic oxygen conditions were sufficient to allow the benthos 342 to survive and grow to a large size. Moreover, most species sampled have minimal mobility so are unlikely to have moved from neighbouring regions. 343

We observed a positive relation between trawling and some community metrics and the biomass of the priapulid worm *H. spinulosus*. These increases may be linked to changes in the biomass and abundance of *Saduria entomon*, an important benthic predator in the Baltic Sea, with trawling. In field enclosure experiments, *S. entomon* has been found to have a negative effect on priapulid abundances (Bonsdorff et al. 1995). *S. entomon* shows a negative trend with
trawling intensity that albeit non-significant may have lowered predation pressure on infaunal
prey in fished areas. Although speculative, the declining trend of *S. entomon* with trawling can
itself also be driven by predation, since cod, an important predator on *S. entomon* (Neuenfeldt
et al. 2020), may be found at higher abundances in areas of high fishing.

4.2 Low oxygen conditions and the absence of fauna

We found no benthic fauna in locations with oxygen concentrations less than 2 ml l⁻¹. Mass-354 mortality of benthic fauna typically occurs around 0.5 ml O₂ l⁻¹ (Diaz & Rosenberg 1995) and 355 it is therefore likely that these sampling locations had lower oxygen levels earlier in the year. 356 We also found no benthic fauna in Station 9, where an oxygen concentration of 1.3 ml l⁻¹ was 357 measured. Station 9 is only 3.7 km away from the location with a time series of near-bed 358 oxygen (Fig. 1, Fig. 3), making it likely that the time series reflect the temporal oxygen history 359 of Station 9. The time series shows that oxygen concentrations dropped below 0.5 ml $O_2 l^{-1}$ in 360 May 2018, which is four months before sampling. This drop is likely the moment where the 361 mass-mortality occurred that led to the defaunation of Station 9. Importantly, in June 2018 362 oxygen concentrations were again 2 ml $O_2 l^{-1}$, showing the fine-scale temporal fluctuations in 363 oxygen conditions. These results highlight that models predicting mortality on benthic fauna 364 from hypoxia need near-bed oxygen data that captures these fine-scale temporal fluctuations 365 366 in oxygen, as predictions will otherwise be at risk of significantly underestimating the effect. For example, all seasonal averages of the time series are well above 1 ml O₂ l⁻¹ and these values 367 would not suggest a benthic defaunation of the area. A conclusion that warrants a reassessment 368 of oxygen condition monitoring for assessing and disentangling seabed disturbance activities 369 in the future. 370

The fishing intensity at Station 9 declined most severely in comparison to all stations, over the period 2013-2018. The station exhibited values close to SAR = 10 in 2013 and 1 in 2018. Fig.

S1-4 shows there is a strong and positive relation between the annual fishing intensities at 373 Station 9 and the maximum yearly oxygen concentrations from the time series (linear model 374 shows a p-value of 0.01 and an adjusted R^2 of 0.78), whereas no relation is found for minimum 375 or average yearly oxygen conditions. This finding may imply that peaks in oxygen, which are 376 still below 5 ml O₂ l⁻¹, attract fish and subsequently fisheries in this area. Catch rates of cod, an 377 important target species, have been shown to peak at these intermediate oxygen conditions in 378 379 the neighbouring Bornholm Basin (Neuenfeldt & Beyer 2003). Since cod feeds on pelagic sprat as well as benthic animals (especially motile ones), cod abundance may be high in areas where 380 381 the state of the seafloor is low. This could explain why there is no relationship between trawling intensity and the average or minimum yearly oxygen conditions. If correct, there is an 382 important temporal aspect to the interactive impacts of trawling and hypoxia on benthic fauna, 383 where trawl disturbance will be highest in periods with relatively healthy oxygen conditions. 384

385 **4.3 No synergistic effects**

Despite significant uncertainties in the impact of trawling on benthic fauna and the temporal 386 history of oxygen conditions, it is clear from our study that trawling impacts are not 387 underestimated for this area because of clear synergistic effects between low oxygen conditions 388 and trawling. Compared to other regions, two characteristics of the benthic community and the 389 environment may have reduced the potential for synergistic effects in this area. Firstly, most 390 391 samples had a deeper layer of glacial clay and few deep-bioturbating species; 99% of total biomass was found in the upper 4 cm. This limits the potential of increasing trawl impacts with 392 hypoxia due to upward migration of burying organisms to the sediment surface as observed in 393 other regions (Pihl et al. 1992, Nilsson & Rosenberg 1994), because all fauna is already found 394 in the upper surface layers. Secondly, there were no ophiuroids and asteroids present in the 395 area, limiting the potential of high trawl impacts with hypoxia due to arm-tipping behaviour. 396

This highlights that other regions with a different benthic fauna and sediment composition mayshow a different response, as is also observed by De Leo et al., (2017).

399 4.4 Management implications

Our results suggest that significant long-term improvements of benthic communities in this 400 area are not to be expected when the area is solely protected from bottom trawl disturbance 401 402 (even though significant negative effects of trawling in certain months of the year cannot be ruled out). Management of bottom trawl fisheries could therefore prioritize the protection of 403 neighbouring non-hypoxic regions, under the assumption that trawl impacts do take place in 404 these regions. Bottom fisheries management targeting non-hypoxic regions can thus be used to 405 improve the overall status of benthic fauna in these specific areas within a relatively short 406 period of time, which could have positive spill-over effects for neighbouring seasonal or more 407 permanent hypoxic areas. Continued reductions of nutrient loads and subsequent declines of 408 hypoxic regions would also benefit the state of the benthic ecosystem, but such management 409 410 actions targeting eutrophication and hypoxia take decades before they yield positive results on the benthic ecosystem (Carstensen et al. 2014). 411

412

5. ACKNOWLEDGEMENT

The field study was funded by the Assemble+ program. We thank the crew of RV Oceanograf for their help in collecting the data. PDvD was funded by the Centre for Ocean Life, a Villum Kann Rasmussen center of excellence supported by the Villum Foundation. PDvD was further funded by the European Union's Horizon 2020 research and innovation programme under the Marie Sklodowska-Curie grant agreement No 101024886. AT was funded by the research profile "The Sea" at Åbo Akademi Univeristy, supported by the Academy of Finland.

419 **6. REFERENCES**

Amoroso RO, Pitcher CR, Rijnsdorp AD, McConnaughey RA, Parma AM, Suuronen P, 420 421 Eigaard OR, Bastardie F, Hintzen NT, Althaus F, Baird SJ, Black J, Buhl-Mortensen L, Campbell AB, Catarino R, Collie J, Cowan JH, Durholtz D, Engstrom N, Fairweather TP, 422 Fock HO, Ford R, Gálvez PA, Gerritsen H, Góngora ME, González JA, Hiddink JG, 423 424 Hughes KM, Intelmann SS, Jenkins C, Jonsson P, Kainge P, Kangas M, Kathena JN, Kavadas S, Leslie RW, Lewis SG, Lundy M, Makin D, Martin J, Mazor T, Gonzalez-425 Mirelis G, Newman SJ, Papadopoulou N, Posen PE, Rochester W, Russo T, Sala A, 426 Semmens JM, Silva C, Tsolos A, Vanelslander B, Wakefield CB, Wood BA, Hilborn R, 427 Kaiser MJ, Jennings S (2018) Bottom trawl fishing footprints on the world's continental 428 shelves. Proc Natl Acad Sci 115:E10275–E10282. 429 430 Andersen JH, Carstensen J, Conley DJ, Dromph K, Fleming-Lehtinen V, Gustafsson BG, Josefson AB, Norkko A, Villnäs A, Murray C (2017) Long-term temporal and spatial 431 trends in eutrophication status of the Baltic Sea. Biol Rev 92:135–149. 432 Baden SP, Loo L-O, Pihl L, Rosenberg R (1990) Effects of eutrophication on benthic 433 434 communities including fish: Swedish west coast. AMBIO A J Hum Environ 19:113–122. Baltic Sea Hydrographic Commission (2013) Baltic Sea Bathymetry Database version 0.9.3. 435 Downloaded from http://data.bshc.pro/ on 21-02-2017 436 Benson BB, Krause D (1984) The concentration and isotopic fractionation of oxygen dissolved 437 in freshwater and seawater in equilibrium with the atmosphere. Limnol Oceanogr 29:620-438 439 632. Blott SJ, Pye K (2001) GRADISTAT: a grain size distribution and statistics package for the 440

441 analysis of unconsolidated sediments. Earth Surf Process Landforms 26:1237–1248.

- Bonsdorff E (2006) Zoobenthic diversity-gradients in the Baltic Sea: Continuous post-glacial
 succession in a stressed ecosystem. J Exp Mar Bio Ecol 330:383–391.
- Bonsdorff E, Norkko A, Sandberg E (1995) Structuring zoobenthos: the importance of
 predation, siphon cropping and physical disturbance. J Exp Mar Bio Ecol 192:125–144.
- 446 Breitburg D, Levin LA, Oschlies A, Grégoire M, Chavez FP, Conley DJ, Garçon V, Gilbert D,
- 447 Gutiérrez D, Isensee K, Jacinto GS, Limburg KE, Montes I, Naqvi SWA, Pitcher GC,
- 448 Rabalais NN, Roman MR, Rose KA, Seibel BA, Telszewski M, Yasuhara M, Zhang J
- 449 (2018) Declining oxygen in the global ocean and coastal waters. Science 359:eaam7240.
- 450 Carstensen J, Andersen JH, Gustafsson BG, Conley DJ (2014) Deoxygenation of the Baltic
 451 Sea during the last century. Proc Natl Acad Sci 111:5628–5633.
- van Denderen PD, Bolam SG, Friedland R, Hiddink JG, Norén K, Rijnsdorp AD, Sköld M,
 Törnroos A, Virtanen EA, Valanko S (2019) Evaluating impacts of bottom trawling and
 hypoxia on benthic communities at the local, habitat, and regional scale using a modelling
- 455 approach. ICES J Mar Sci 77:278–289.
- 456 Diaz RJ, Rosenberg R (1995) Marine benthic hypoxia: a review of its ecological effects and
 457 the behavioural responses of benthic macrofauna. Oceanogr Mar Biol An Annu Rev
 458 33:203–245.
- 459 Diaz RJ, Rosenberg R (2008) Spreading dead zones and consequences for marine ecosystems.
 460 Science 321:926–929.
- 461 Hiddink JG, Jennings S, Sciberras M, Bolam SG, Cambiè G, McConnaughey RA, Mazor T,
 462 Hilborn R, Collie JS, Pitcher R, Parma AM, Suuronen P, Kaiser MJ, Rijnsdorp AD (2018)
 463 Assessing bottom-trawling impacts based on the longevity of benthic invertebrates. J Appl
 464 Ecol 56:1075–1084.

465	Hiddink JG, Jennings S, Sciberras M, Szostek CL, Hughes KM, Ellis N, Rijnsdorp AD,
466	McConnaughey RA, Mazor T, Hilborn R, Collie JS, Pitcher CR, Amoroso RO, Parma
467	AM, Suuronen P, Kaiser MJ (2017) Global analysis of depletion and recovery of seabed
468	biota after bottom trawling disturbance. Proc Natl Acad Sci 114:8301-8306.

- 469 ICES (2019) Working Group on Spatial Fisheries Data (WGSFD). ICES Scientific Reports.
 470 1:52. 144 pp.
- 471 De Leo FC, Gauthier M, Nephin J, Mihály S, Juniper SK (2017) Bottom trawling and oxygen
 472 minimum zone influences on continental slope benthic community structure off
 473 Vancouver Island (NE Pacific). Deep Sea Res Part II Top Stud Oceanogr 137:404–419.
- McLaverty C, Eigaard OR, Gislason H, Bastardie F, Brooks ME, Jonsson P, Lehmann A,
 Dinesen GE (2020) Using large benthic macrofauna to refine and improve ecological
 indicators of bottom trawling disturbance. Ecol Indic 110:105811.
- 477 Neuenfeldt S, Bartolino V, Orio A, Andersen KH, Andersen NG, Niiranen S, Bergström U,
 478 Ustups D, Kulatska N, Casini M (2020) Feeding and growth of Atlantic cod (*Gadus*479 *morhua* L.) in the eastern Baltic Sea under environmental change. ICES J Mar Sci 77:624–
 480 632.
- 481 Neuenfeldt S, Beyer JE (2003) Oxygen and salinity characteristics of predator-prey
 482 distributional overlaps shown by predatory Baltic cod during spawning. J Fish Biol
 483 62:168–183.
- Nilsson HC, Rosenberg R (1994) Hypoxic response of two marine benthic communities. Mar
 Ecol Prog Ser 115:209–217.
- Pihl L, Baden SP, Diaz RJ, Schaffner LC (1992) Hypoxia-induced structural changes in the
 diet of bottom-feeding fish and Crustacea. Mar Biol 112:349–361.

488	Pitcher CR, Ellis N, Jennings S, Hiddink JG, Mazor T, Kaiser MJ, Kangas MI, McConnaughey
489	RA, Parma AM, Rijnsdorp AD, Suuronen P, Collie JS, Amoroso R, Hughes KM, Hilborn
490	R (2017) Estimating the sustainability of towed fishing-gear impacts on seabed habitats:
491	a simple quantitative risk assessment method applicable to data-limited fisheries. Methods
492	Ecol Evol 8:472–480.
493	Riedel B, Diaz R, Rosenberg R, Stachowitsch M (2016) The ecological consequences of
494	marine hypoxia: from behavioural to ecosystem responses. Stress Mar Environ Oxford
495	Univ Press Oxford:175–194.
496	Rijnsdorp AD, Bolam SG, Garcia C, Hiddink JG, Hintzen NT, van Denderen DP, Van Kooten
497	T (2018) Estimating sensitivity of seabed habitats to disturbance by bottom trawling based
498	on the longevity of benthic fauna. Ecol Appl 28:1302–1312.
499	Rijnsdorp AD, Hiddink JG, van Denderen PD, Hintzen NT, Eigaard OR, Valanko S, Bastardie
500	F, Bolam SG, Boulcott P, Egekvist J (2020) Different bottom trawl fisheries have a

- differential impact on the status of the North Sea seafloor habitats. ICES J Mar Sci
 77:1772–1786.
- Schernewski G, Friedland R, Carstens M, Hirt U, Leujak W, Nausch G, Neumann T, Petenati
 T, Sagert S, Wasmund N, Weber M von (2015) Implementation of European marine
 policy: New water quality targets for German Baltic waters. Mar Policy 51:305–321.
- 506 Sköld M, Göransson P, Jonsson P, Bastardie F, Blomqvist M, Agrenius S, JG H, HC N,
 507 Bartolino V (2018) Effects of chronic bottom trawling on soft-seafloor macrofauna in the
 508 Kattegat . Mar Ecol Prog Ser 586:41–55.
- Tobin J (1958) Estimation of relationships for limited dependent variables. Econom J Econom
 Soc:24–36.

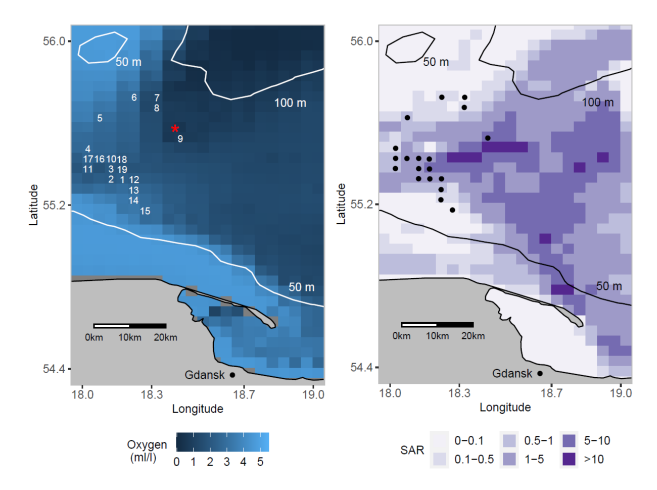
511	Törnroos A, Bonsdorff E (2012) Developing the multitrait concept for functional diversity:
512	lessons from a system rich in functions but poor in species. Ecol Appl 22:2221–2236.
513	Törnroos A, Bonsdorff E, Bremner J, Blomqvist M, Josefson AB, Garcia C, Warzocha J (2015)
514	Marine benthic ecological functioning over decreasing taxonomic richness. J Sea Res
515	98:49–56.
516	Törnroos A, Pecuchet L, Olsson J, Gårdmark A, Blomqvist M, Lindegren M, Bonsdorff E

interlinkage across trophic groups in a large marine ecosystem. Glob Chang Biol 25:1235–
1246.

517

(2019) Four decades of functional community change reveals gradual trends and low

- Valanko S, Norkko A, Norkko J (2010) Strategies of post-larval dispersal in non-tidal softsediment communities. J Exp Mar Bio Ecol 384:51–60.
- Whitlatch RB, Lohrer AM, Thrush SF, Pridmore RD, Hewitt JE, Cummings VJ, Zajac RN
 (1998) Scale-dependent benthic recolonization dynamics: life stage-based dispersal and
 demographic consequences. Hydrobiologia 375:217–226.



525

Figure 1. Maps of the sampling station locations in the southern Baltic Sea (numbers 526 correspond to Table S1-1). The oxygen concentrations (left) show the lowest seasonal 527 concentration in the year based on modelled bottom oxygen concentrations from an ecosystem 528 model representative of 2006-2010 (Schernewski et al. 2015). The red asterisk shows a nearby 529 530 location with time-series of near-bed oxygen measurements (see further Fig. 3). The fishing intensity data (right) show average SAR (Swept Area Ratio per year) values for the period 531 2013-2017. Depth contours were extracted from the Baltic Sea Hydrographic Commission 532 (2013). The oxygen and depth data were solely used to select sampling stations as depth and 533 oxygen data used in the analysis were measured in-situ at each station. 534

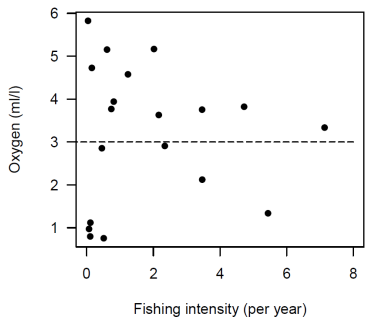


Figure 2. Fishing intensity (average SAR, per year) and oxygen concentration of the sampling 536 stations. We analysed all sampling stations together to examine the combined effect of trawling 537 and oxygen concentration, as well as a subset of stations with oxygen concentrations $> 3 \text{ ml } O_2$ 538 1⁻¹ (stations above dashed line) to explore trawling effects in isolation. This threshold was 539 chosen as no effect of oxygen on benthic biomass or abundance is observed above this 540 concentration. 541

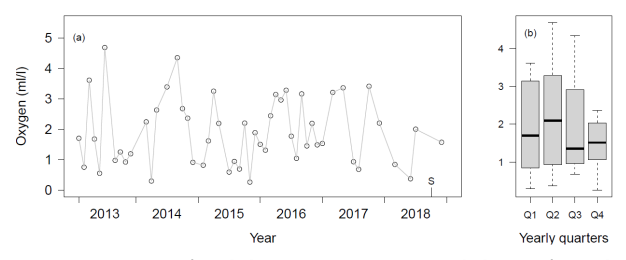
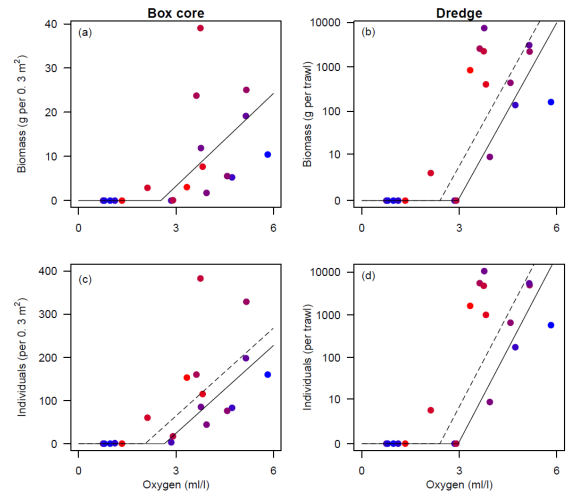


Figure 3. (a) Time series of near-bed oxygen concentrations at a nearby location of our study area (red asterisk in Fig. 1) from 2013 to 2018. The letter "S" indicates when we sampled the study area. (b) Boxplot of oxygen concentrations for each seasonal quarter based on all data points in (a), indicating lower median oxygen concentrations in quarter 3 and 4. Data used were obtained from HELCOM secretariat and based on ICES dataset on <u>Ocean Hydrography</u>.



548

Figure 4. Relationships between oxygen and total faunal biomass and abundance collected using the box corer (a, c) and dredge (b, d). Lines are fitted with a Tobit-regression. In (b-d) where trawling intensity is part of the best model (Table 1), the solid lines show the prediction without fishing (SAR = 0) and the dashed lines with SAR = 2. Points are coloured using a linear colour gradient from the station with the lowest (blue, SAR = 0.04) to highest (red, SAR = 7.14) fishing intensity. The dredge data is $log_{10}(x+1)$ transformed because there are four orders of magnitude variation between stations.

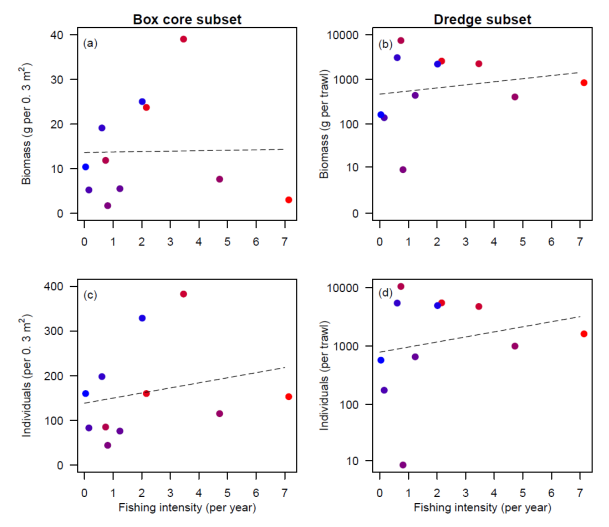
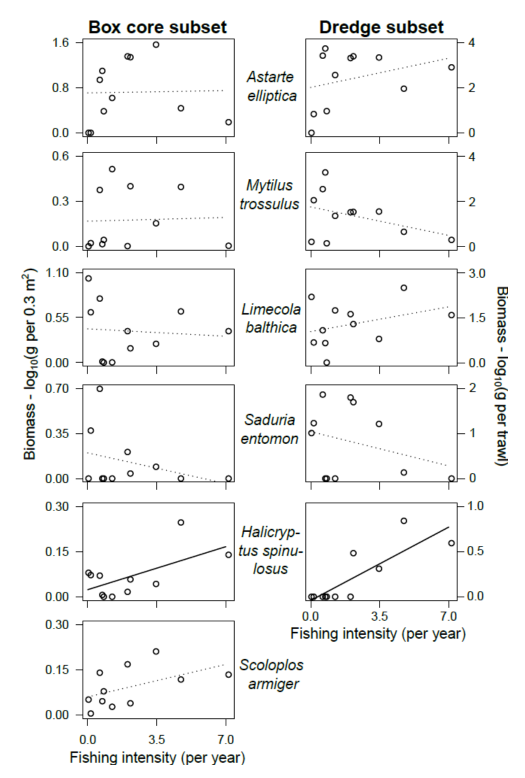


Figure 5. Relationship between fishing intensity (average SAR, per year) and faunal biomass and abundance of box corer and dredge for stations with oxygen concentrations above 3 ml l⁻ ¹. The relationships are all non-significant based on linear regression. Points are coloured using a linear colour gradient from the station with the lowest oxygen concentration (red, 3.3 ml l⁻¹) to the highest (blue, 5.8 ml l⁻¹) in this data subset.



562

Figure 6. Relationship between fishing intensity (average SAR, per year) and the biomass of different dominant species in box corer and dredge for all stations with oxygen concentrations above 3 ml 1^{-1} . Solid lines show significant relationships (p-value <0.05), dashed lines nonsignificant. Note that all species in this subset have a non-significant relationship with oxygen concentration.

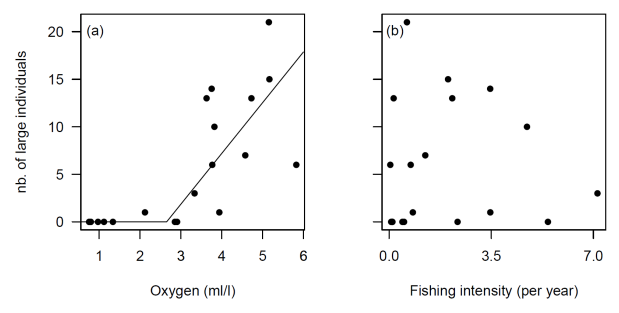


Figure 7. Relationships between the number of large individuals (>15 mm) and oxygen (a) and
fishing intensity (b) for all sampling stations. The solid line is based on Tobit regression. There
is no support for a model with trawling (Table 1).

572	Table 1. Model selection and coefficients. B is benthic biomass, A is benthic abundance, T is
573	trawling intensity (year-1) and O is measured oxygen concentration (ml l-1). Tobit regression
574	lower limit was set at zero biomass/abundance. The Tobit model includes an ancillary statistic,
575	termed σ , which describes the log-standard deviation of the latent variable. Asterisks show best
576	model based on AIC (when models differ less than 2 AIC-units, the model with fewest
577	parameters is selected). Since Hauck-Donner effects were observed in some p-value estimates
578	using Wald statistics, p-values were obtained by conducting a likelihood ratio test.

Model	σ	p-values	AIC
Box core biomass			
$B = -17.71 + 7.00 \cdot O$	2.35	$P_{\rm O} < 0.001$	114.7*
$B = -23.37 + 7.58 \cdot O + 1.72 \cdot T$	2.32	$P_0 < 0.001; P_T = 0.20$	115.1
$B = -16.11 + 5.54 \cdot O - 4.00 \cdot T + 1.73 \cdot T \cdot O$	2.27	$P_O=0.01;P_T=0.41;P_{T+O}=0.21$	115.5
Box core abundance			
$A = -132.65 + 65.44 \cdot O$	4.53	$P_{O} < 0.001$	187.7
$A = -179.66 + 67.86 \cdot O + 20.12 \cdot T$	4.43	$P_0 < 0.001; P_T = 0.06$	186.3*±
$A = -110.58 + 46.60 \cdot O - 45.77 \cdot T + 20.71 \cdot T \cdot O$	4.32	$P_O = 0.005; P_T = 0.20; P_{T+O} = 0.05$	184.4
Dredge biomass			
$Log_{10}(B+1) = -2.57 + 1.16 \cdot O$	0.24	$P_0 < 0.001$	52.4
$Log_{10}(B+1) = -3.87 + 1.31 \cdot O + 0.36 \cdot T$	0.10	$P_0 < 0.001; P_T = 0.03$	49.5*
$Log_{10}(B+1) = -2.70 + 1.01 \cdot O - 0.27 \cdot T + 0.19 \cdot T \cdot O$	0.03	$P_O\!<\!0.001;P_T\!=\!0.60;P_{T+O}\!=\!0.21$	49.9
Dredge abundance			
$Log_{10}(A+1) = -2.78 + 1.26 \cdot O$	0.30	$P_{O} < 0.001$	53.8
$Log_{10}(A+1) = -4.28 + 1.44 \cdot O + 0.41 \cdot T$	0.14	$P_0 < 0.001; P_T = 0.02$	50.2*
$Log_{10}(A+1) = -3.02 + 1.12 \cdot O - 0.24 \cdot T + 0.19 \cdot T \cdot O$	0.07	$P_0 < 0.001; P_T = 0.66; P_{T \cdot 0} = 0.22$	50.7
Box core abundance (individuals >4 mm)			
$A = -161.08 + 66.60 \cdot O$	4.56	$P_{\rm O} < 0.001$	176.6
$A = -252.91 + 76.43 \cdot O + 27.27 \cdot T$	4.42	$P_0 < 0.001; P_T = 0.02$	173.3*
$A = -162.55 + 52.31 \cdot O - 29.86 \cdot T + 17.09 \cdot T \cdot O$	4.34	$P_{O} = 0.005; P_{T} = 0.45; P_{T \cdot O} = 0.12$	172.9
Box core abundance (individuals >15 mm)			
$A = -14.16 + 5.33 \cdot O$	1.71	$P_{O} < 0.001$	85.8*
$A = -18.72 + 5.98 \cdot O + 0.99 \cdot T$	1.69	$P_0 < 0.001; P_T = 0.21$	86.3
$A = -14.34 + 4.86 \cdot O - 1.74 \cdot T + 0.78 \cdot T \cdot O$	1.65	$P_0 < 0.001; P_T = 0.58; P_{T \cdot 0} = 0.37$	87.5

 \pm note that a model without trawling (model 1) differs less than 2 AIC values from the selected model (model 2). Yet, a model with interaction (model 3) is more than 3 AIC units lower than model 1. Model 2 is therefore selected as best intermediate model.