1 The effects of bottom trawling and primary production on the biological

2 traits composition of benthic assemblages

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8 Running page head: Trawling and production on benthic functioning

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- 11

12 Abstract

Although many studies have investigated the effects of disturbance and environmental drivers on 13 14 marine ecosystems, comparatively few have studied their interactions. Using fuzzy coded biological 15 traits, we compared the functional composition, diversity and evenness of benthic communities in the English Channel, and Celtic and Irish Seas, across interacting gradients of bottom trawling and 16 17 primary production. Fuzzy correspondence analysis indicated greater similarity in traits composition at sites of high trawling pressure than those of low trawling. In contrast, the analysis revealed no 18 19 relationship between traits composition and primary production. Trawling and primary production had no effect on the traits "longevity", "sediment position" and "feeding mode". However, trawling 20 had negative effects on all modalities within the trait "living habit", which were strongest for 21 22 attached and epifaunal organisms, but weakest for burrow- and tube-dwelling species. Trawling also 23 negatively affected most modalities within the trait "maximum weight", which were strongest for 24 organisms weighing between < 0.1 g and 1 kg. Conversely, trawling positively affected organisms 25 weighing > 10 kg. For the trait "bioturbation", upward conveyors were positively related with primary production, whilst other modalities exhibited no clear pattern. Because trawling affected 26 some traits more than others, community biomass was less evenly distributed across traits in highly 27 28 trawled areas, which resulted in lower levels of functional diversity and evenness. Overall, the 29 effects of bottom trawling were greater in areas of high primary production.

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30 Introduction

31 Marine ecosystems are subject to a range of anthropogenic stressors, many of which are increasing 32 in intensity and occurrence (Lotze et al. 2006, Poloczanska et al. 2013). The use of bottom trawls has 33 greatly increased over the last century (Watson et al. 2013, Howarth et al. 2014). In contacting the 34 seafloor, bottom trawling can damage benthic habitats (Kaiser et al. 2000), reduce the abundance of 35 target and non-target species (Hiddink et al. 2017), and truncate age and size distributions (Beamish 36 et al. 2006, Jørgensen et al. 2007). In parallel, ocean acidification, eutrophication, and climate 37 change are altering global levels of primary production (Frederiksen et al. 2006, Tait & Schiel 2013), 38 which could significantly alter food web dynamics, fisheries production and ocean biogeochemistry 39 (Gregg et al. 2003, Brander 2007, Blanchard et al. 2012). In areas where primary production 40 increases, benthic communities should receive a greater input of energy, increasing their growth, 41 reproduction and resilience to bottom trawling (Blanchard et al. 2009, Hiddink et al. 2017). 42 Investigating the interactive effects of bottom trawling and primary production could therefore help 43 to inform management and improve understanding of how multiple drivers affect marine 44 ecosystems.

45 Most studies quantify the ecological effects of disturbance by measuring changes in species 46 abundances (Mouillot et al. 2013, Stuart-Smith et al. 2013). However, species-based approaches 47 convey little direct information on how disturbances affect ecosystem processes, such as nutrient 48 cycling and habitat provisioning (Bremner & Frid 2005), which play a greater role in maintaining 49 ecosystem integrity than the abundance of a particular species (Loreau et al. 2001). Hence, biological 50 traits are increasingly being used to describe the physical, behavioural and life-history characteristics of species to evaluate their potential vulnerability to disturbance ('response traits') and contribution 51 52 towards ecosystem function ('effect traits' – see Bolam et al. 2016). For example, bioturbation can be 53 considered an 'effect trait' as the burrowing of benthic infauna can transport nutrients and oxygen 54 from the sediment surface to deeper layers (Olsgard et al. 2008, Sciberras et al. 2016). In contrast, 55 bottom trawls are more likely to damage sessile organisms than pelagic organisms (Kaiser et al.

2000, Tillin et al. 2006), meaning living habit can be considered a 'response trait'. Given that some
disturbances affect life histories and functions more than others, biological traits can help
disentangle the ecological effects of multiple stressors (Bremner et al. 2006)

59 Numerous studies show bottom trawling can change benthic traits composition. By reducing the 60 abundance of large predators, trawling can increase the abundance of small and fast-growing 61 species, which can recover quickly from disturbance and benefit from reduced predation (Bremner & 62 Frid 2005, Tillin et al. 2006). Additionally, trawling can increase the availability of organic matter and 63 dead / injured animals, boosting the local abundance of mobile scavengers (Tillin et al. 2006, Kaiser 64 & Hiddink 2007, Craven et al. 2013). Trawling can also plough and re-suspend sediments which can 65 reduce the feeding efficiency of suspension feeders (Collie et al. 2000, Bradshaw et al. 2003). 66 Compared to bottom trawling, little is known about the effects of primary production on traits 67 composition. Nonetheless, areas of high primary production often support greater abundances of small-bodied organisms that can quickly incorporate pulses of energy into growth and reproduction 68 69 (Macpherson et al. 2002, Jennings & Blanchard 2004, Gómez-Canchong et al. 2013). High levels of 70 primary production should also result in greater quantities of phytoplankton and other organic 71 matter sinking to the seafloor, which would benefit suspension and deposit feeders (Blanchard et al. 72 2009).

73 Given that trawling and primary production likely have contrasting consequences, their interaction 74 may produce different effects to them acting alone (Crain et al. 2008). These interactions could be 75 synergistic or additive, where the effects of one driver equal or exceed the sum of each in isolation 76 (see Gunderson et al. 2016). For example, the positive relationship between primary production and 77 small body size might be intensified by trawling, because fishing removes the largest individuals 78 (Beamish et al. 2006, Jørgensen et al. 2007, Fu et al. 2018). Alternatively, interactions may be 79 antagonistic, where the cumulative effect of several drivers is less than them acting alone. For 80 instance, the increased detritus and organic matter caused by high primary production levels may

81 increase population growth rates of suspension and deposit feeders, reducing recovery times and
82 their sensitivity to trawling effects.

83 Persistent and intense disturbances can reduce species abundances to levels that eliminate their 84 influence on ecosystem function (Howarth et al. 2014). Given that multiple species can perform the 85 same functional role (Fonseca & Ganade 2001), high species diversity presumably increases ecosystem resilience because of the low probability of losing all species capable of performing a 86 87 specific function (Loreau et al. 2001, Cardinale et al. 2002). Biological traits can therefore measure 88 the distribution (functional evenness) and range (functional diversity) of functional roles present 89 within an ecosystem (Díaz & Cabido 2001, Mooney et al. 2009). Given the greater effects of bottom 90 trawling on some groups, trawling is expected to reduce the functional diversity and evenness of 91 marine communities, and reduce their resilience to further disturbance (Schleuter et al. 2010, 92 Howarth et al. 2014).

93 In order to evaluate how primary production and fishing influence the functioning of benthic 94 ecosystems, we compare the functional composition, diversity and evenness of benthic communities in the English Channel, and Celtic and Irish Sea, across interacting gradients of bottom trawling and 95 96 primary production. First, we hypothesize that bottom trawling will reduce the functional diversity 97 and evenness of benthic communities by reducing the abundance of long-lived predatory fish and 98 attached and suspension feeding invertebrates. Second, we hypothesize that both trawling and 99 primary production increase abundances of scavengers and small, short-lived species. Last, we 100 hypothesize reduced effects of trawling on suspension feeders in areas of high primary production, 101 but intensified effects of trawling on body size.

102 Methods

103 Outline

104 Benthic organisms were sampled over gradients of trawling intensity and primary production at 105 sampling stations with otherwise homogeneous environmental conditions. Their biological trait 106 composition, functional diversity and functional evenness were then related to trawling intensity107 and primary production.

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109 Study area

110 This work was carried out in the Irish Sea, Celtic Sea, and western English Channel. These areas are 111 characterised by moderate levels of wave energy, depths of 20 - 100 m, and similar circalittoral 112 sediments of muddy sand and gravel (Cooper et al. 2010). However, they also differ slightly in that 113 sea surface temperatures are often lower, and primary production levels higher, within the Irish Sea. 114 Between March and May, the Celtic Sea undergoes a distinct algal bloom that originates south of Ireland and temporarily increases chlorophyll a concentrations in the Celtic Sea, southern Irish Sea 115 116 and western English Channel (Fasham et al. 1983, Garcia-Soto & Pingree 2009). This algal bloom can 117 increase chlorophyll a levels to such an extent that it can homogenise any spatial differences that 118 typically exist during the rest of the year. European beam and otter trawlers operate throughout 119 these regions and target a wide range of species such as hake (Merluccius merluccius), plaice 120 (Pleuronectes platessa), sole (Solea solea), monkfish (Lophius piscatorius) and whiting (Merlangius 121 merlangus). However, the Irish Sea is characterised by a greater presence of otter trawlers targeting Nephrops prawns (Nephrops norvegicus), the Celtic Sea is characterised by a greater presence of 122 123 beam trawlers targeting flat fish, and the western English Channel is the focus of a large king scallop 124 (Pecten maximus) dredge fishery (Guénette & Gascuel 2012, Campbell et al. 2014, Howarth & 125 Stewart 2014).

126 Data sources

Water column depths (m) were extracted from GEBCO (<u>www.gebco.net</u>) at a resolution of halfminute intervals. Information on seabed substrate type and bed shear stress was obtained from the UkSeaMap 2016 (<u>http://jncc.defra.gov.uk/ukseamap</u>). Annual levels of primary production (mg C m⁻² yr⁻¹) between 2009 and 2013 (estimated by the MODIS satellite sensor) were provided by NEODAAS (<u>www.neodaas.ac.uk</u>) at a resolution of 1.1 x 1.1 km and averaged across all years. These data used 132 Morel & Berthon's (1989) algorithm to convert depth, surface chlorophyll, and photosynthetically 133 active radiation (PAR) into estimates of net primary production. However, like most satellite 134 chlorophyll algorithms, it can over-estimate primary production in waters of high sediment loads or 135 coloured dissolved organic matter. Fishing effort (hours spent bottom trawling and dredging) 136 between 2011 2013 MMO and was provided by the 137 (www.gov.uk/government/organisations/marine-management-organisation) at a resolution of 3.5 x 3.5 km, which was averaged by year. Trawling intensity (swept area ratio yr⁻¹) averaged between the 138 139 years 2010 and 2012 was supplied by Eigaard et al. (2016) at a resolution of 1.8 x 1.8 km. Their data 140 take into account differences in trawl size and trawl type, which can strongly influence the level of 141 disturbance trawling gears have on the benthos (Hiddink et al. 2007). Swept area ratio can be 142 interpreted as the mean number of times fishing gear impacts an area in a year. A swept area ratio 143 of one indicates that the swept area equals the cell area (Gerritsen et al. 2013). Because swept area 144 ratio is more informative than the number of hours spent fishing, the MMO measures of fishing 145 effort were converted to swept area ratio as described by Gerritsen et al., (2013). For this, we 146 assumed that the average fishing vessel towed gears 24 m in width at a speed of two knots. Our 147 study solely used fishing effort provided by (Eigaard et al. 2017) except for two stations (O and P) 148 where we used MMO data instead. Beam trawlers in North Devon intensively target this area 149 (Campbell et al. 2014) which was not highlighted in the Eigaard dataset, likely because of a lack of 150 French and Spanish VMS data for their trawl fleets operating in the Celtic Sea.

151 Experimental design

Trawling intensity and primary productivity were both divided into four categorical levels (Table 1) and 1.8 x 1.1 km sampling stations were chosen to cover all combinations within the Irish Sea, Celtic Sea and English Channel. To ensure comparability and to avoid confounding effects, stations were restricted to "sand" and "muddy sand" substrates, between 40-100 m in depth, and of "moderate" bed shear stress as defined by the UkSeaMap. Final sampling stations were selected based on their proximity to one another, and their distance from restricted (military and fishing) zones and offshore structures; greater distances were preferred in all cases. Stations were also selected based on their similarity in depth and sediment composition. Some treatment combinations were rarer than others, meaning there were instances where we had to sample stations coarser in substrate or located at greater depth. Hence, there is some spatial clustering of sampling stations, and differences in environmental characteristics may provide some scope for confounding effects which we test for in our analyses.

164 Sample collection

A total of 20 stations were sampled both in September 2015 and April 2016 (Figure 1 and Table S1) and data were pooled across both sampling events. Most benthic recovery processes are seasonal and rely on distinct pulses in recruitment, which require time for larvae to settle and grow (Rodríguez et al. 1993, Mann 2009). Hence, by sampling during and after the spring algal bloom, we aimed to account for these seasonal differences.

170 At each station, seabed water samples were taken to quantify the chlorophyll a (Chl-a) content at 171 time of sampling, and to allow comparisons with the satellite-generated measures of primary 172 production. At each sampling station, a one-litre Niskin bottle fitted with a digital reversing 173 thermometer was deployed to one metre above the seafloor and triggered with a brass messenger 174 three times. Upon retrieval, the mean of 15 temperature measurements were recorded along with 175 their standard deviation (SD). For each station, we filtered between 900 and 1800 ml of seawater 176 over 47 mm GF/F filter papers in triplicate and stored these filters at -20°C in the dark until further 177 processing.

Five 0.1 m² Day grabs were taken within each sampling station. These samples were later used to quantify sediment size, Chl-a, and infauna. To quantify sediment Chl-a, we subcored using a cut-off syringe 14.5 mm in diameter to a depth of 8 mm which was preserved at -20°C in the dark. To sample meiofauna, we took two subcores 25 mm in diameter to a depth of 40 mm which were preserved in a 4% buffered formalin seawater solution. To sample infauna, we washed whatever

sediment remained of the Day grab samples over a 1-mm sieve before preserving the remainingmaterial in 4% formalin.

185 Epifaunal organisms were sampled using two 2 m beam trawls (fitted with a 10 mm mesh and 2 mm 186 cod-end liner) which were towed for five minutes along the seafloor at a speed of 1.5 knots. Benthic 187 megafauna and demersal fish were sampled using two 4 m beam trawls (fitted with a chain matrix 188 and an 82 mm diamond cod-end) which were towed for 30 minutes at a speed of 3 knots. All 189 organisms caught by the trawls were identified to species level where possible, counted, and 190 weighed using a motion compensating balance. Subsampling was used for species comprising of 191 more than 50 individuals. Small animals < 1 g were weighed in groups and their mean individual 192 weight calculated.

193 Particle size analysis (PSA)

Sediment samples collected by the Day grab were analysed for particle size using both dry sievingand a laser particle analyser. Both datasets were then combined as described in Mason (2011).

196 Chl-a analysis

Water samples were analysed for Chl-a content using the methods and equations described in Lorenzen (1966) and JGOFS (1994). The same method was applied to the sediment cores, however, these were centrifuged for 20 minutes at 2400 rpm before analysis in order to separate sediments from the supernatant.

201 Invertebrate sorting and identification

Approximately 10 ml of 0.1% Rose Bengal was added to the formalin-preserved grab samples, gently mixed, and left to stain for at least one hour before sorting. Samples were then washed over a 1-mm sieve to remove all traces of formalin and moved to a white tray. All organisms were separated from the sediment and preserved in 70% industrial methylated spirit (IMS). These organisms were sorted into broad taxonomic groups using a dissection microscope and methyl blue dye. All individuals were then identified to at least family level, counted, blotted dry, and weighed to the nearest 1 mg. Body 208 parts were reassembled to make whole organisms but were discarded if less than 20% of the 209 individual remained. Individuals were not counted if they did not possess a head. Mollusc shells 210 were smashed and discarded if empty. Tube-dwelling animals were separated from their tubes 211 before weighing, and hermit crabs were removed from their shells or epibionts and weighed 212 separately.

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215 Data analysis

216 Comparisons of environmental characteristics

217 Multivariate tests were used to test if differences in measured environmental characteristics existed 218 between sampling sites, and whether they could confound the effects trawling and primary 219 production. These environmental data were mean particle size, percentage gravel, percentage mud, 220 percentage sand, and water depth. A resemblance matrix using a Euclidean Distance measure was 221 created and visualised using non-Metric Dimensional Scaling (nMDS) with clusters generated from a 222 Similarity Profile (Simprof) routine. This analysis indicated two sampling stations (D and P) were 223 dissimilar to the others (S1). As station D had a very low outlying mean particle size (69 μ m ± 2 SE) 224 and station P had a very high outlying mean particle size (937 μ m ± 203 SE), these two stations were 225 excluded from further analysis.

226 Verification of experimental treatments

Trawling intensity, sediment and water Chl-a content were compared between the categorical levels of fishing effort and primary productivity. Trawling intensity increased exponentially across levels of fishing effort (S2a). The Chl-a content of the sediment samples displayed no relationship with levels of primary production (S2b). In contrast, water Chl-a correlated tightly with levels of primary production in September but not in April (S2c).

232 Gear calibrations

233 Because we used multiple types of sampling gears to capture several components of the benthic 234 ecosystem, each gear partly overlapped in the size of organisms they captured, but differed in 235 sampling area and catch efficiency. For each sampling gear, small animals were undersampled 236 because they passed through the sieves or nets of the gear, and large animals were rarely found 237 because their abundance was too low relative to the area sampled. This meant that the abundance 238 and biomass values obtained by the different sampling gears could not be combined until they were 239 corrected to represent the same sampling area and sampling efficiency, and were restricted to the 240 size ranges that were sampled effectively. Hence, we scaled the data from the different sampling 241 gears based on the assumption that the abundance of size ranges that were sampled by two 242 sampling gears should be the same after correction. We carried out the following correctional 243 procedure on the whole dataset for all stations combined to get the most robust correction factors.

244 Normalised biomass size spectra were created as described in Sprules & Barth (2015). The size 245 spectrum of each individual sampling gear showed an optimum relationship with different minimum 246 and maximum size ranges of organisms sampled (S3a). The size range of organisms sampled 247 effectively by each gear was identified as the range where the size spectrum exhibited a monotonic 248 decline (S3b). Abundance values from the Day grab samples were used as the base for the 249 corrections because the area sampled by the grab was known to be exactly 0.1 m². The abundance 250 of organisms sampled by the 2-m beam trawl were then corrected by performing a linear regression 251 on the normalised biomass per size class captured by the Day grab and 2-m beam trawl for the size 252 ranges in which they overlapped (S4a). This procedure was then repeated for the combination of the 253 2-m and the 4-m beam trawl (S4b). To correct those size classes that were sampled by multiple 254 gears, the total biomass within each size class was then divided by the number of gears contributing 255 to each size class.

256 Constructing a fuzzy coded database

257 Many traits-based studies assign species to discrete functional roles, such as 'predator' or 258 'detritivore' (e.g. Friedlander & DeMartini 2002; Micheli & Halpern 2005; Williams et al. 2015). 259 However, this approach is overly simplistic because most aquatic species exhibit multiple traits (e.g. 260 part-predator, part-detritivore) and express ontogenetic shifts in their ecology as they develop (e.g. 261 from planktivorous larvae, to piscivorous adults). Hence, we used a method known as 'fuzzy coding' 262 to help overcome these issues. Fuzzy coding divides traits (e.g. maximum weight) into categories, or 263 'modalities,' that cover the full range of possible values for that trait (e.g. < 10 g; 10 - 100 g; > 100 g). 264 We then assigned scores to each trait indicating the affinity of species to those modalities; where 265 low scores represented no affinity, and high scores represented total and exclusive affinity 266 (Chevenet et al. 1994; Tillin et al. 2006). These scores were then converted to proportions totalling 267 to one, and multiplied by species biomass or abundance, effectively spreading out their abundance 268 across multiple traits.

269 We modified an existing fuzzy coded traits database of species, genus's and families compiled by 270 Bolam et al. (2017) to include maximum biomass and demersal fish (Table 2). These additional 271 information came from FishBase (<u>www.fishbase.org</u>), the BIOTIC database (<u>www.marlin.ac.uk/biotic</u>) 272 and Jennings et al. (2001). For taxa identified at a resolution higher than species (e.g. genus and 273 family), the database assigned scores based on their most closely-related taxa using the Best 274 Professional Judgement (BPJ) approach (Bolam et al. 2014). This strategy resulted in a reasonably 275 accurate completion of modalities wherever the entries across closely related taxa were fairly 276 consistent, but we were less confident where traits were variable across closely related taxa, making 277 it necessary to spread the fuzzy-scores across a wider number of modalities.

Overall, our functional traits database included information on six biological traits spanning across modalities (Table 2). There is currently no accepted methodology for selecting the most appropriate traits for a given study. However, the traits used in our study were intended to cover a combination of response and effect traits with the potential to reflect an organism's life history (e.g.

life span), ecology (e.g. living habit), vulnerability (e.g. sediment position), and contribution towards ecosystem processes (e.g. bioturbation). We also aimed to ensure our selected traits did not overlap in the information they conveyed. For instance, both maximum weight (g) and maximum length (cm) can describe an organism's size. However, maximum weight is more informative about how much energy an organism has invested into its own growth (Brose et al. 2005, Gómez-Canchong et al. 2012), and can more accurately reflect the size of an organism that does not exhibit bilateral symmetry. Hence, we chose maximum weight over maximum length as a biological trait.

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291 Relating functional indices to trawling and primary production

Functional diversity and evenness were estimated from the Shannon-Wiener Diversity Index and Pielou's Evenness using the biomass of each modality as described in Schleuter *et al.* (2010). To test whether functional diversity and evenness were significantly related to primary production and trawling intensity, two linear models were constructed as follows:

296 Functional diversity $\sim \log_{10}(\text{trawling intensity}) * \log_{10}(\text{primary production})$

297 Functional evenness ~ log₁₀(trawling intensity) * log₁₀(primary production)

298 For visualisation purposes, fitted values were plotted against continuous levels of trawling and 299 compared between categorical levels of primary production, and vice a versa. These visualisations 300 therefore differed from the models, as the models fitted these as continuous variables. This was 301 done using the predict function in R (Team 2017) across trawling gradients and the mean of each 302 level of primary production, and vice a versa. Linear models are used throughout our analyses 303 because previous studies show the log biomass of communities decrease linearly with the log of 304 trawling pressure (Hiddink et al. 2006). This pattern occurs because a fixed fraction of benthic 305 biomass is typically removed with every pass of a trawl.

306 Relating functional composition to trawling and primary production

307 To visualise the influence of trawling pressure and primary production on overall functional 308 composition, a fuzzy correspondence analysis (FCA) was performed using the R package "ade4" 309 (Dray et al. 2017). FCA is a type of correspondence analysis that can describe relationships among 310 objects of interest (i.e. sampling stations) based on the abundances of fuzzy coded variables (i.e. 311 modalities). To aid interpretation, the total biomass of each modality was summed across both 312 survey periods. The scores generated by the first two FCA axes were then plotted for each station 313 and modality. This meant each bubble represented an individual sampling station, the size of which 314 was based on its trawling intensity or primary production. Stations that are plotted closer together in 315 the ordination have similar patterns of biomass distributions across modalities. Hence, distances 316 between the stations reflect differences in the proportional biomass of modalities. For example, if 317 the trawling ordinations clearly separated the larger bubbles from the small, then trawling intensity 318 could be interpreted as having a strong influence on trait composition. In addition, modalities 319 labelled closely to the larger bubbles would be more associated with areas of high trawling pressure than areas of low trawling pressure. 320

321 Relating modality biomass to trawling and primary production

To determine whether modality biomasses were significantly related with trawling pressure and primary production, linear models were created as:

324 Log₁₀(biomass) ~ modality * log₁₀(trawling intensity) * log₁₀(primary production)

and repeated for each trait, creating a total of six models. We based this approach assuming that a significant interaction between modality and trawling and / or primary production would indicate unequal response in the modalities within a trait to these pressures. Significant interactions were then plotted using the visualisation methods described earlier.

329 Results

330 Dataset description

331 This study identified 332 different taxa, 52 (or 16%) of which were identified to species level, 221 were to genus (63%) and 59 (21%) were to family. The modalities with the fewest taxa (< 20) were 332 333 organisms weighing > 10 kg, organisms buried > 10 cm deep within the sediment, and upward and 334 downward bioturbators (Table 2). In contrast, surface-depositors, free-living organisms and animals 335 living on the sediment surface contained the most taxa (> 200). The modalities with the lowest biomass (< 100 g m⁻²) were organisms with a life span of < 1 year, organisms weighing > 10 kg, 336 337 crevice-dwelling organisms, and pelagic species. In contrast, the most abundant modalities (> 3000 g m^{-2}) were free-living animals, organisms with a life span between 3 – 10 years, and animals buried 338 339 down to 5 cm in the sediment.

340 Effect of trawling and primary production on functional indices

Functional diversity and functional evenness related positively to primary production and negatively to trawling (Table 3); the highest levels of functional evenness and diversity therefore coincided with low trawling and high primary production (Figure 2). The interaction between trawling and primary production was significantly negative, with the strongest trawling effect at high primary production. At low primary production, trawling did not affect functional evenness and diversity.

346 Effect of trawling and primary production on traits composition

347 FCA ordinations showed greater similarity in traits composition at stations subjected to high levels of 348 trawling pressure than stations subjected to lower levels of trawling (Figure 3A). Intensively trawled 349 stations were characterised by a greater biomass of organisms that weighed over 1 kg (g1001 and 350 g10000), lived for over 10 years (l10), resided deep within the sediment (sp10) or were pelagic 351 (spPelagic), did not contribute to bioturbation (bNone), and were predatory (fPred) and / or crevice-352 dwelling (IhCrev). In contrast, stations subjected to low levels of trawling were characterised by 353 animals that attached to the seabed (lhAtt) or other animals (lhEpi), lived high up (sp0.5 and sp6.10) 354 or on top of the sediment (spSurf), were diffusive (bDiff) bioturbators, weighed between 11 g - 1 kg (g11 and g101), lived < 1 year to 3 years (l1 and l1.3), and were suspension (fSusp) and / or surface-
feeders (fSurf). Unlike these comparisons in trawling pressure, the FCA ordinations did not clearly
separate stations of low and high primary production (Figure 3B).

358 Effect of trawling and primary production on modality biomass

359 Linear models relating modality biomass with primary production and trawling pressure for each 360 biological trait showed significant interactions between modality and trawling and / or primary 361 production, thus indicating different reactions by modalities to these two stressors (Table 4). 362 Trawling and primary production did not interact significantly with modality biomass for the traits "longevity", "sediment position" and "feeding mode". For "living habit", we observed significant 363 364 interactions between modality and primary production, and between modality and trawling 365 pressure. Trawling had negative effects on all modalities within this trait, and these effects were 366 more pronounced under high levels of primary production (Figure 4). This negative relationship was 367 strongest for attached and epifaunal organisms, and weakest for burrow- and tube-dwelling species. 368 For "maximum weight", the three-way interaction between modality, trawling, and primary 369 production was significant. This meant trawling had negative effects on most modalities within this 370 trait, with the negative effect of trawling strengthening under high levels of primary production 371 (Figure 5). These negative relationships were strongest in organisms weighing between < 0.1 g and 372 1000 g. In contrast, organisms weighing > 10 kg exhibited a broadly positive relationship with 373 trawling, which strengthened with increasing primary production. Primary production had positive 374 effects on the biomass of most modalities when plotting primary production as a continuous 375 variable (Figures S5 and S6), and that the positive effect of primary production weakened under high 376 levels of trawling. For "bioturbation", only the interaction between modality and primary production 377 was significant. Of the modalities within this trait, upward conveyors related positively to primary 378 production (Figure 6), whilst the other modalities exhibited no clear pattern.

379 Discussion

Our study is the first to measure the joint effects of trawling and primary production on the functional diversity and traits composition of benthic communities. Overall, bottom trawling had negative effects on the functional diversity, evenness, and biomass of 12 of the 32 modalities investigated, whilst primary production had positive effects. We also observed strong interactive effects between the two, in that the degree to which one influenced the benthic community depended on the strength of the other. This meant greater effects of bottom trawling on benthic communities in areas of high primary production.

387 Community biomass was less evenly distributed across modalities in highly trawled areas, resulting 388 in lower levels of functional diversity and functional evenness. Whilst some studies suggest fishing 389 can reduce functional diversity and evenness (Worm et al. 2006, Martins et al. 2012), ours is the first 390 to directly relate reductions to incremental increases in trawling pressure. We also found stronger 391 effects of trawling on functional diversity and evenness in areas of high primary production. The 392 cause of this pattern remains unclear, especially considering recent evidence which suggests that 393 high primary production should buffer benthic communities from trawling impacts (Hiddink et al. 394 2017). Then again, highly productive freshwater systems often coincide with high community 395 biomass but low species diversity (McQueen et al. 1989, Rudstam et al. 1993), and should therefore 396 be less resilient (Macpherson et al. 2002, Jennings & Blanchard 2004, Gómez-Canchong et al. 2013). 397 Given the effects of climate change, acidification, and eutrophication on global levels of primary production, such a relationship could make marine ecosystems and the fisheries they support less 398 399 resilient to the impacts of fishing, particularly in areas where primary production increases 400 (Blanchard et al. 2012, Tait & Schiel 2013).

Intensely trawled areas were characterised by lower biomasses of attached and epiphytic organisms such as sea squirts (Tunicata), sponges (Porifera), bryozoans (e.g. *Flustra foliacea*, Crisiidae and *Cellaria* spp), and soft (e.g. *Alcyonium digitatum*) and hard corals (e.g. *Caryophyllia smithii*). Many studies document strong impacts on these organisms by mobile gears through physical disruption of

405 sediments (Kaiser et al. 2000, 2006, Bradshaw et al. 2003, Howarth, Pickup, et al. 2015) and 406 associated negative effects on suspension feeders (Collie et al. 2000, Bradshaw et al. 2003). 407 However, these organisms are functionally important to temperate marine ecosystems because they 408 add three dimensional structure to the seabed (Howarth, Roberts, et al. 2015). In doing so, they can 409 provide nursery habitats to a wide range of fish and invertebrates, supporting local levels of 410 biodiversity and the recruitment of commercially important species (Beck et al. 2001, Kamenos et al. 411 2004, Gibb et al. 2007, Howarth, Roberts, et al. 2015). Consequently, negative effects of bottom 412 trawling could potentially reduce their nursery habitat function (Kaiser et al. 2000, Bradshaw et al. 413 2001, Howarth et al. 2011).

414 High levels of fishing pressure are expected to reduce the abundance of large predators, and favour 415 smaller competitor and prey species with shorter life histories (Bremner & Frid 2005, Tillin et al. 416 2006). Alternatively, all body sizes could be affected given that bottom trawling is non-selective, 417 causing mortality in a wide range of non-target and target species ranging from nematodes to large 418 sharks (Fennessy 1994, Hiddink et al. 2006, Hinz et al. 2008, 2009). However, we found negative 419 effects of bottom trawling on a variety of body sizes (< 0.1 g to 10 kg) but no relationship with 420 feeding mode. Hence, our results indicate that bottom trawling removes biomass from most of the 421 benthic community, and not just large predators. Our results therefore suggest that fishing mortality 422 is more important for small-bodied organisms than the release from predation pressure caused by 423 the removal of targeted fish species.

Surprisingly, trawling had a broadly positive effect on the biomass of large organisms weighing > 10 kg such as rays (e.g. *Amblyraja radiata* and *Dipturus batis*), sharks (e.g. *Mustelus Asterias*), gadoids (e.g. *Gadus morhua* and *Molva molva*) turbot (*Scophthalmus maximus*) and monkfish (*Lophius piscatorius*). This unexpected result may reflect the low number of species (i.e. 11) represented in this modality. Hence, any differences in the spatial distribution of these species across our sampling stations would have had disproportionally strong effects on our results. Alternatively, this finding

may reflect the confounding effects of using fishing effort as an indicator of fishing disturbance.
Logic suggests fishing pressure should concentrate in areas with larger, more commercially viable
species. If true, this would reduce our ability to detect a negative relationship between fishing effort
and the abundance of large predatory fish.

434 The abundance of mobile scavengers can increase by up to 200 times in trawled areas as a result of 435 increased availability of carrion and other organic matter (Tillin et al. 2006, Kaiser & Hiddink 2007). 436 However, we detected no relationship between bottom trawling and the biomass of scavenging 437 organisms such as hermit crabs (e.g. Pagurus spp). Then again, given the dispersion of odour plumes, 438 resettlement of sediment and predation of damaged / injured organisms, high scavenger densities 439 on fishing grounds are likely to be relatively short-lived events (Howarth, Pickup, et al. 2015). Hence, 440 our approach, which was designed to detect long-term and large-scale changes, is unlikely to detect 441 such an event. However, our results support two other studies that suggest the benefits of an 442 increased food supply to scavengers cannot compensate for the direct mortality caused by bottom 443 trawling (Bolam 2014, Bolam et al. 2017). These two studies also observed recovery of bioturbating 444 species in 3-5 months following bottom fishing. In our study, this modality was composed primarily 445 of burrowing polychaetes (e.g. Pectinariidae, Maldanidae and Eunicidae) which recover quickly from 446 disturbance (reviewed in Jennings, Kaiser, et al. 2001). This capacity may explain why we observed 447 no relationship between bottom trawling and the biomass of bioturbating organisms.

High levels of primary production should provide more energy to benthic deposit and suspension feeders (Blanchard et al. 2009). However, we detected no relationship between primary production and the biomass of deposit and suspension feeding organisms, such as bivalves (e.g. Cardiidae and *Abra* spp), polychaetes (e.g. *Lagis koreni*), and anemones (Actiniaria). This may reflect the depth of our sampling stations. Detritus is a poor energy resource, subject to degradation through microbial action and consumption as it sinks through the water column towards the seabed (Gerlach et al. 1985). Given that our sampling sites were located at depths between 40 and 90 m, the detritus

reaching the seabed may have been too low in energy to offer measurable benefit to deposit and suspension feeders. We also expected a positive relationship between primary production and small body size, and for this relationship to intensify under high trawling pressure. Whilst we found evidence of primary production increasing the biomass of small-bodied organisms, in reality, this relationship weakened with increasing trawling pressure. Hence, this pattern suggests the fishing mortality inflicted on small organisms outweighs any benefits they receive from enhanced growth rates under high primary production (Posey et al. 2002).

462 In summary, our evidence suggests that bottom trawling and primary production can cause 463 functional changes to benthic communities. Bottom trawling had negative effects on functional 464 diversity, functional evenness, and the maximum weight and living habit of the benthic community, 465 and no effects on their feeding mode, longevity, bioturbation, and sediment position. We also found 466 greater effects of trawling on benthic ecosystem functioning in areas of high primary production. 467 Noting changing levels of primary production globally, this interaction may reduce resiliency of 468 ecosystems and fish stocks to future fishing impacts. We therefore suggest further study of the 469 interactions between fishing disturbance and environmental perturbations, which could have strong 470 implications for conservation and fisheries management.

471 Data accessibility

- 472 Data used in this paper are archived in the British Oceanographic Data Centre (<u>www.bodc.ac.uk</u>)
- 473 under doi:10.5285/674d4224-7cc5-4080-e053-6c86abc0626e.

474 Acknowledgements

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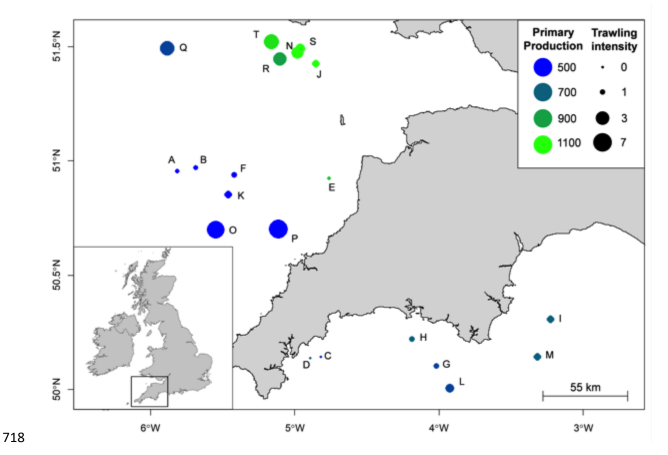


Figure 1. Sampling stations. Each point represents a 1 x 0.6 nautical mile box, the shade and size of which signifies the level of primary production (mg C $m^{-2} yr^{-1}$) and trawling intensity (yr⁻¹).

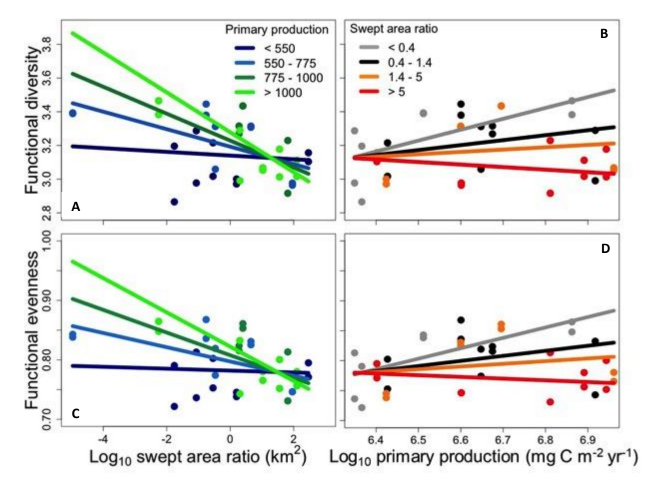


Figure 2: Functional diversity (A) and functional evenness (B) plotted against trawling intensity and
 categorical levels of primary production. Functional diversity (C) and functional evenness (D) plotted
 against primary production and categorical levels of trawling intensity. Trend lines fitted by linear
 models.

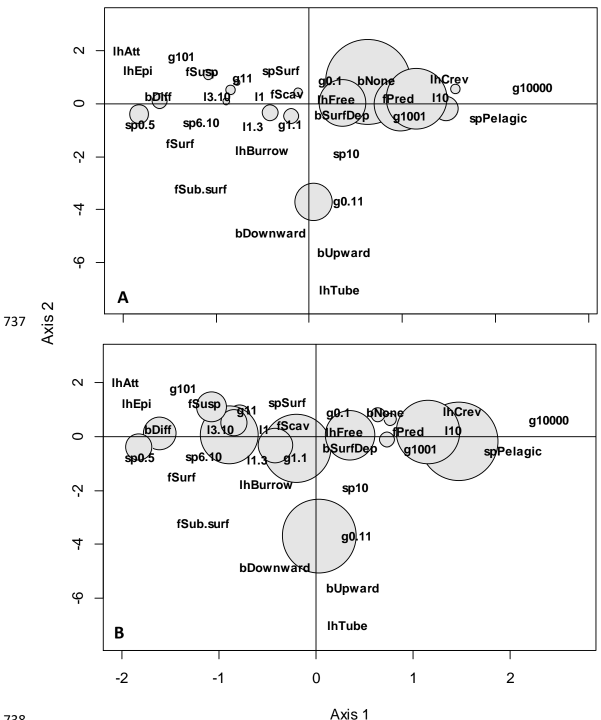
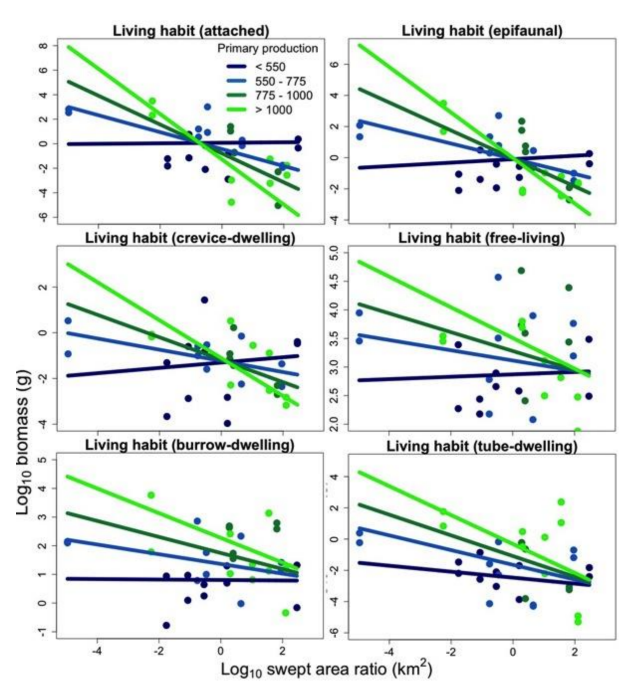
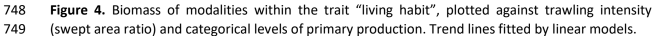


Figure 3. Plots of ordination scores from fuzzy correspondence analysis of modality biomass. Each bubble indicates a sampling station. Plots are identical except for the area of circles: (A) trawling intensity (yr-1); (B) primary production (mg C m-2 yr-1). The abbreviations give the names of the modalities. Stations that appear further apart have greater differences in overall trait composition. The location of the modalities indicates the types of stations with which they are associated.







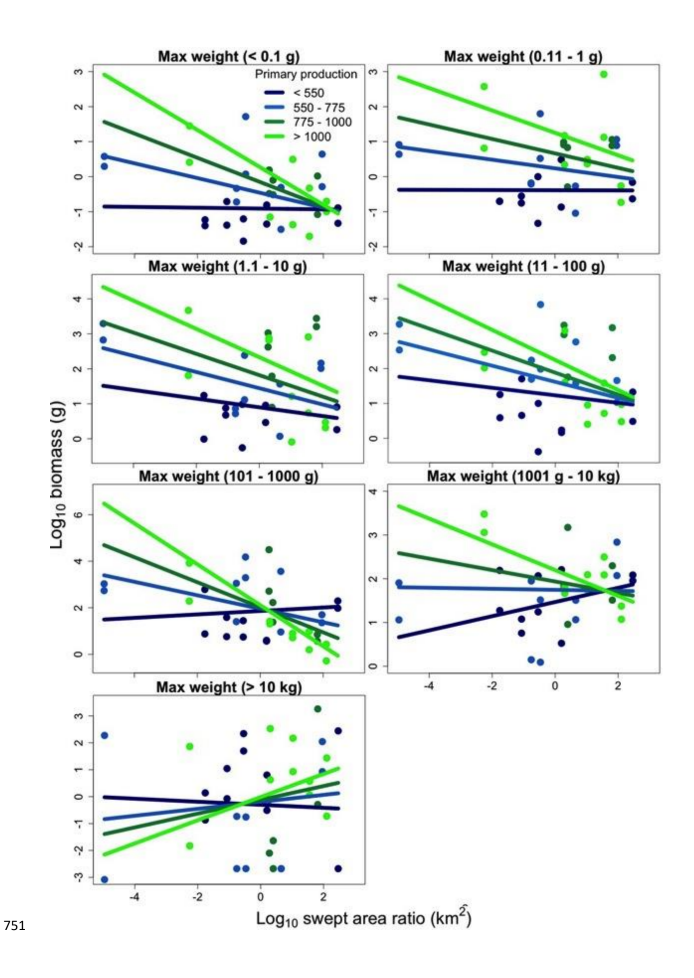


Figure 5. Biomass of modalities within the trait "maximum weight", plotted against trawling
 intensity (swept area ratio) and categorical levels of primary production. Trend lines fitted by linear
 models.

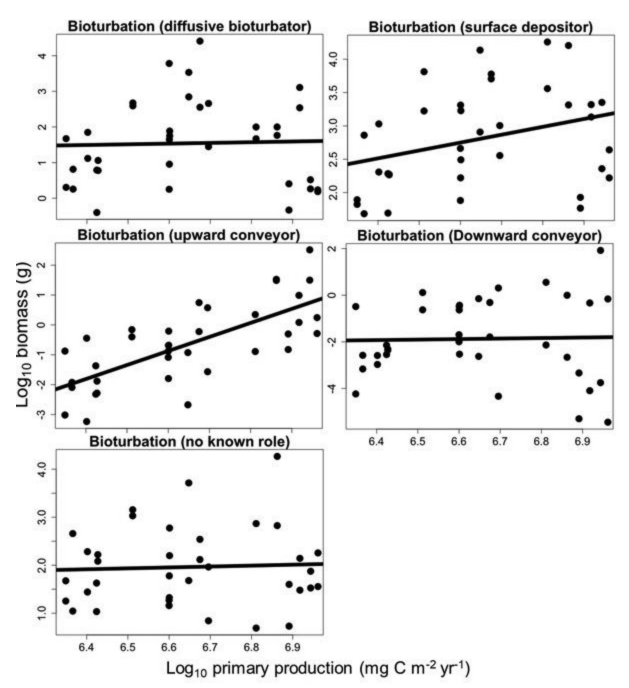


Figure 6. Biomass of modalities within the trait "bioturbation", plotted against primary production.

757 Trend lines fitted by linear models.

764	Table 1. The range of values that defined the four experimental treatments of trawling intensity and
765	primary production sampled in this study.

	Treatment	Swept area ratio (km ²)	Primary production (mg C m ⁻² yr ⁻¹)
	1 - Low	<u><</u> 0.4	<u><</u> 550
	2 - Medium low 3 - Medium high	<u>></u> 0.4 ≤ 1.4 ≥ 1.4 ≤ 5	> 550 <u><</u> 775 > 775 <u><</u> 1000
	4 - High	<u>≥ 1.4 ≤ 5</u> > 5	> 1000
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Trait	Code	Description	No. of taxa	Biomass (g m⁻²)	Relevance
Maximum weight	g0.1	< 0.1 g	108	298.1	Reflects trophic position, metabolic rate, energy
	g0.11	0.11 - 1 g	71	801.8	requirements, generation time, commercial value,
	g1.1	1.1 - 10 g	105	2305.8	and catchability (Jennings et al. 2002, Brose et al.
	g11	11 - 100 g	84	936.5	2005, Sprules & Barth 2015).
	g101	101 - 1000 g	46	1421.8	
	g1001	1001 - 10000 g	33	256.5	
	g10000	> 10000 g	11	33	
Longevity	1	< 1 yrs	45	15.3	Reflects age at maturity, somatic growth, and
	1-3	1 - 3 yrs	125	1152.2	disturbance frequency / severity (Pearson &
	3-10	3 - 10 yrs	171	3557.9	Rosenberg 1978, Musick 1999).
	110	> 10 yrs	83	1328.1	
Living habit	lhTube	Tube-dwelling	31	329	Indicates potential to evade, or to be exposed to
	lhBurrow	Burrow-dwelling	89	2015.8	disturbance and predation pressure (Kaiser et al.
	lhFree	Free-living	222	3266.5	2000).
	lhCrev	Crevice-dwelling	48	62.1	
	lhEpi	Epiphytic	34	140.6	
	lhAtt	Attached	29	239.5	
Location within	spPelagic	Pelagic	63	79.9	Implications for sediment-water biogeochemistry
sediment	spSurf	Atop of sediment	233	1366.5	and oxygen availability, and susceptibility to
	sp0-5	0 <i>-</i> 5 cm deep	124	3731.3	disturbance (Aller 1982, Collie et al. 2000).
	sp6-10	6 - 10 cm deep	53	729.2	
	sp10	> 10 cm deep	26	146.5	
Feeding mode	fSusp	Suspension feeder	127	1847.6	Implications for energy flow and susceptibility to
	fSurf	Surface deposit feeder	107	1434	disturbance (Rosenberg 1995, Bergmann et al. 2002
	fSub-surf	Sub-surface deposit feeder	49	1559.7	Craven et al. 2013).
	fScav	Scavenger	60	355.8	
	fPred	Predator	163	854.9	
Bioturbation	bDiff	Diffusive bioturbator	112	2304.8	Implications for sediment-water biogeochemistry
	bSurfDep	Surface depositor	249	2897.3	and oxygen availability (Mermillod-Blondin 2011).
	bUpward	Upward conveyor	12	303.5	
	bDownward	Downward conveyor	17	186.4	
	bNone	None	88	361.4	

Table 2. Description of biological traits used in this study, and the total biomass (pre-gear calibration) and number of taxa that fell within each modality.

Table 3. Linear model outputs modelling functional diversity and functional evenness to trawlingintensity and primary production.

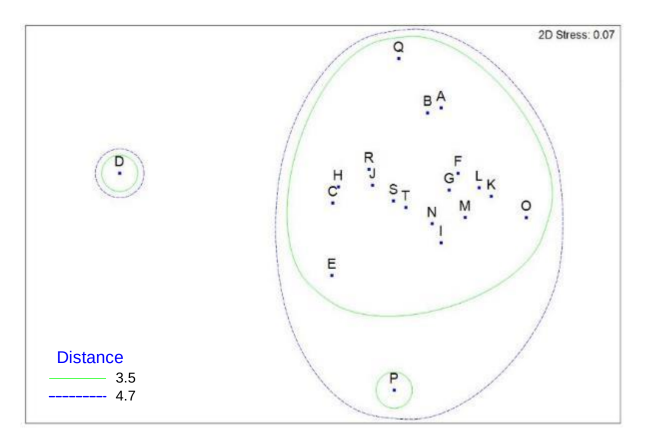
	Model	Predictor	Value	SE	t	Р	R^2
		Intercept	1.23	0.99	1.25	0.233	
	Functional diversity	log(trawling)	1.6	0.61	2.61	0.020	0.27
	Functional diversity	log(primary production)	0.3	0.15	2.1	0.044	0.37
	_	log(trawling):log(primary production)	-0.25	0.09	-2.7	0.018	
		Intercept	0.26	0.24	1.07	0.305	
	Functional evenness	log(trawling)	0.4	0.15	2.68	0.018	0.38
	runctional eveniness	log(primary production)	0.08	0.04	2.25	0.041	0.50
		log(trawling): log(primary production)	-0.06	0.02	-2.77	0.015	
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Table 4. Linear model outputs modelling trait biomass to modality, trawling intensity, and primary

818 production.

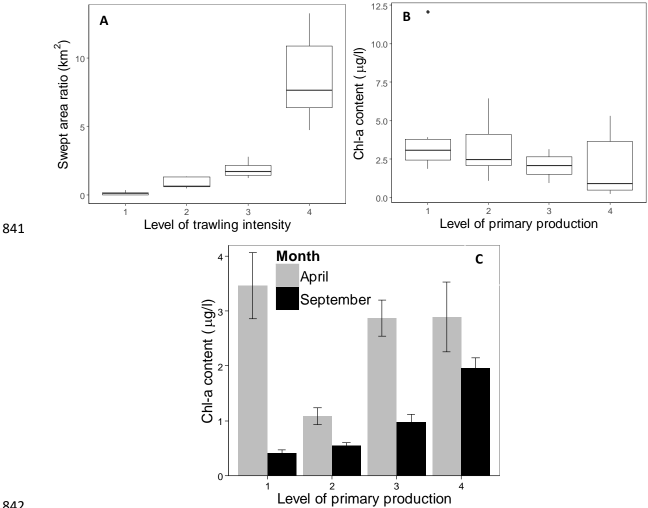
Trait	Predictor	df	MSE	F	Р	R^2			
	Modality	5	126.2	59.9	<0.001				
	log(trawling)	1	75.6	35.9	< 0.001				
	log(production)	1	1.5	0.7	0.401				
Living habit	Modality : log(trawling)	5	9.1	4.3	<0.001	0.6			
	Modality : log(production)	5	8.6	4.1	0.002				
	log(trawling) : log(production)	1	67.6	32.1	< 0.001				
	Modality : log(trawling) : log(production)	5	4.6	2.2	0.057				
	Modality	6	52.3	17.4	< 0.001				
	log(trawling)	1	7.9	2.7	0.104				
Maximum	log(production)	1	33.2	11.1	0.001				
weight	Modality : log(trawling)	6	1.4	0.5	0.822	0.3			
Weight	Modality : log(production)	6	1.7	0.6	0.734				
	log(trawling) : log(production)	1	2.7	0.9	0.339				
	Modality : log(trawling) : log(production)	6	8.1	2.7	0.015				
	Modality	3	351.7	274.9	<0.001				
	log(trawling)	1	13.6	10.6	0.001				
	log(production)	1	7	5.5	0.02				
Longevity	Modality : log(trawling)	3	0.8	0.6	0.605	0.8			
	Modality : log(production)	3	0.6	0.5	0.711				
	log(trawling) : log(production)	1	16.2	12.7	<0.001				
	Modality : log(trawling) : log(production)	3	0.4	0.3	0.799				
	Modality	4	136.1	110.6	<0.001				
	log(trawling)	1	7.8	6.5	0.011				
	log(production)	22.1	18	< 0.001					
Bioturbation	Modality : log(trawling)	4	1.2	1	0.432				
	Modality : log(production)	4	4.7	3.8	0.005				
	log(trawling) : log(production)	1	10.5	8.6	0.004				
	Modality : log(trawling) : log(production)	4	0.4	0.3	0.863				
	Modality	4	101.4	89	<0.001				
	log(trawling)	1	5	4.4	0.037				
	log(production)	1	9.5	8.3	0.004				
Sediment position	Modality : log(trawling)	4	2.3	2	0.091	0.7			
position	Modality : log(production)	4	1.1	1.1	0.376				
	log(trawling) : log(production)	1	9.2	9.2	0.002				
	Modality : log(trawling) : log(production)	4	1.2	1.2	0.319				
	Modality	4	28.7	28.7	<0.001				
	log(trawling)	1	6.3	6.3	0.013				
E I'	log(production)	1	16.2	16.2	<0.001				
Feeding	Modality : log(trawling)	4	1.21	1.2	0.305	0.5			
mode	Modality : log(production)	4	0.45	0.4	0.769				
	log(trawling) : log(production)	1	19.2	19.2	<0.001				
	Modality : log(trawling) : log(production)	4	1.2	1.2	0.315				

819 Electronic supplementary material

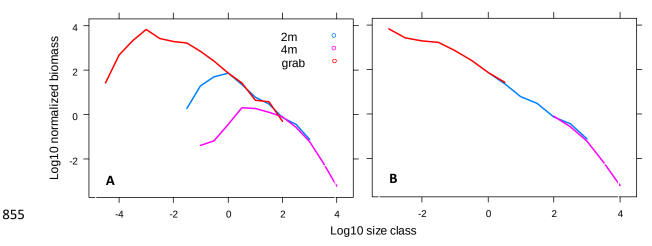




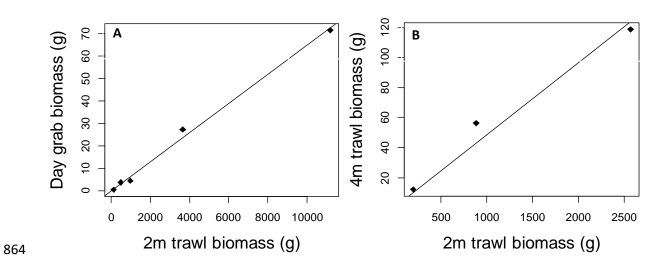
823 S1. nMDS plots of the different stations and their environmental characteristics before removing824 outliers.

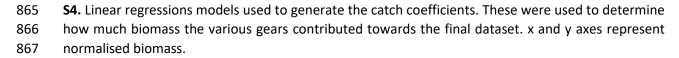


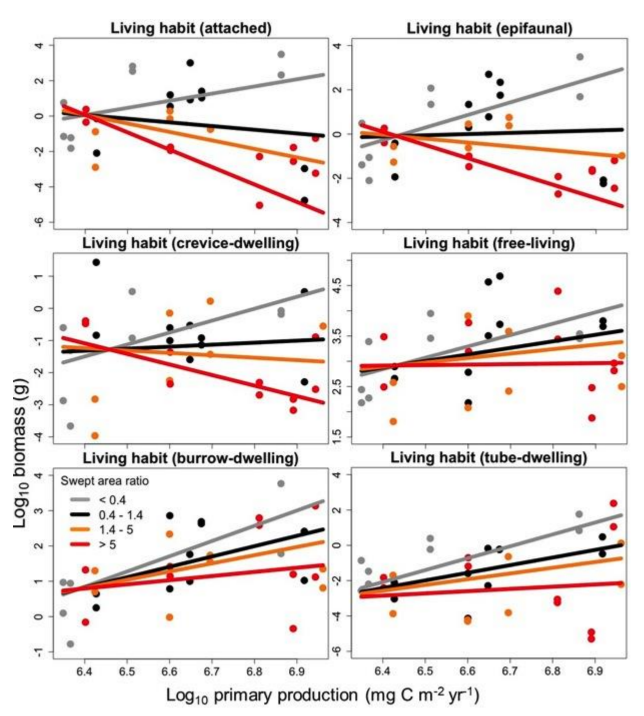
S2. The mean swept area ratio of the sampling stations plotted against their *a priori* experimental level of trawling intensity (A). The mean total Chl-a content of the sediment samples collected in April plotted against their a priori experimental level of primary production (B). The mean total Chl-a content of the water samples collected in September and April plotted against their a priori experimental level of primary production (C). Error bars represent ±1 SE.



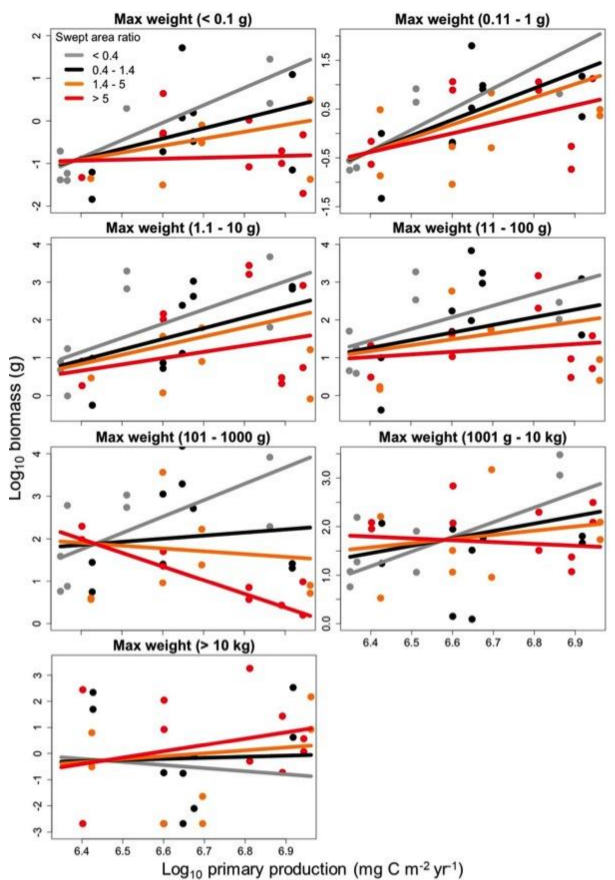
S3. The sampling gears captured organisms of markedly different size ranges (A). The amount of biomass each gear contributed was corrected to represent the same sampling area and were restricted to the size ranges that were sampled effectively. The extreme size classes inefficiently captured by each gear were then removed, resulting in a clear negative slope (B).

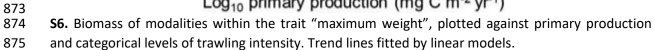






870 871 S5. Biomass of modalities within the trait "living habit", plotted against primary production and 872 categorical levels of trawling intensity. Trend lines fitted by linear models.





- **Table S1.** The latitude, longitude, mean annual primary productivity, mean annual trawling pressure,
- 877 depth and mean particle size of the 1 x 0.6 nautical mile stations sampled in this study.

Ctation	Longitudo	Latituda	Drimony production	Swont area ratio	Donth
Station	Longitude	Latitude	Primary production	Swept area ratio $(1 m^{-1})$	Depth
			(mg C m ⁻² yr ⁻¹)	(yr⁻¹)	(m)
А	50.95762	-5.81468	581	0.17	86.5
В	50.96952	-5.68377	572	0.34	83.4
С	50.14108	-4.81793	672	0.01	62.0
D	50.13779	-4.89094	777	0.01	47.7
E	50.92419	-4.76168	955	0.11	41.6
F	50.94027	-5.42183	618	0.58	76.3
G	50.10285	-4.01841	735	0.47	64.0
Н	50.2217	-4.19	770	0.63	41.3
I	50.3062	-3.22847	792	1.32	53.5
J	51.42451	-4.85315	1009	1.36	55.3
К	50.8516	-5.46147	616	1.22	77.0
L	50.00633	-3.9267	735	1.92	70.0
Μ	50.14212	-3.31622	808	1.49	62.0
Ν	51.49193	-4.96128	1054	2.80	51.4
0	50.69918	-5.5469	603	11.76	75.3
Р	50.7029	-5.11145	613	13.23	59.4
Q	51.49384	-5.8869	735	7.10	96.0
R	51.44706	-5.10275	907	6.14	57.0
S	51.47474	-4.98225	1036	4.74	50.8
Т	51.52256	-5.15893	983	8.18	43.7