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Similar effects of bottom trawling and natural disturbance on composition and function of benthic communities across habitats

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ABSTRACT: Bottom trawl fishing has widespread impacts on benthic habitats and communities. The benthic response to trawling seems to be smaller or absent in areas exposed to high natural disturbance, leading to the hypothesis that natural and trawl disturbance affect benthic communities in a similar way. However, systematic tests of this hypothesis at large spatial scales and with data from sites spanning a large range of natural disturbance do not exist. Here, we examine the effects of trawl and natural (tidal-bed shear stress) disturbance on benthic communities over gradients of commercial bottom trawling effort in 8 areas in the North and Irish Seas. Using a traitbased approach, that classified species by life-history strategies or by characteristics that provide a proxy for their role in community function, we found support for the hypothesis that trawl and natural disturbance affect benthic communities in similar ways. Both sources of disturbance caused declines in long-living, hard-bodied (exoskeleton) and suspension-feeding organisms. Given these similar impacts, there was no detectable trawling effect on communities exposed to high natural disturbance. Conversely, in 3 out of 5 areas with low bed shear stress, responses to trawling were detected and resulted in community compositions comparable with those in areas subject to high natural disturbance, with communities being composed of either small-sized, deposit-feeding animals or mobile scavengers and predators. The findings highlight that knowledge of the interacting effects of trawl and natural disturbance will help to identify areas that are more or less resilient to trawling and support the development of management plans that account for the environmental effects of fishing.

KEY WORDS: Bottom trawling \cdot Benthic community \cdot Biological trait approach \cdot Bed shear stress \cdot Ecosystem function \cdot Disturbance \cdot Beam trawling \cdot Otter trawling

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INTRODUCTION

Bottom trawl fisheries account for around $23\,\%$ of global fisheries yield (FAO 2009) and are among the most widespread sources of human disturbance

affecting benthic communities in shallow shelf seas (Eastwood et al. 2007, Foden et al. 2011). The fishery physically disturbs the seabed by dragging the fishing gear over the seabed to catch bottom-dwelling fish and benthic invertebrates. This disturbance

modifies benthic habitats and leads to mortality of benthic invertebrates in the path of the gear (Kaiser et al. 2002).

The effects of trawling vary markedly among benthic species (Collie et al. 2000a, Kaiser et al. 2006), as a result of their different vulnerability to a trawl pass (e.g. Bergman & van Santbrink 2000) and different recovery rates following impact, varying from months to many years (e.g. Lambert et al. 2014). Generally, studies have found that long-living, sessile and suspension-feeding organisms show the greatest declines in response to a given type and frequency of trawl disturbance (Tillin et al. 2006, de Juan et al. 2007, Kenchington et al. 2007), while opportunistic species, e.g. short-living polychaetes, are less affected (Kaiser et al. 2006).

The response of a benthic community to trawling will also depend on the pre-fished composition of the community (Kaiser et al. 2002). This composition is largely affected by the degree of natural disturbance, due to currents, waves or storms (Thistle 1981, Probert 1984). Natural disturbance may erode seabed sediment, cause resuspension of organic matter (Morris & Howarth 1998) and may affect settlement of new recruits (Thistle 1981, Hunt & Scheibling 1997). Such effects promote species that are adapted to natural disturbance; species that usually have opportunistic life-history strategies and may also be resistant to trawl disturbance (Jennings & Kaiser 1998, Kaiser 1998). Indeed, changes in response to trawling seem to be smaller or undetectable in communities exposed to high natural disturbance (e.g. Kaiser & Spencer 1996, Collie et al. 2000b, Tillin et al. 2006), leading to the hypothesis that natural and trawl disturbance affect benthic communities in a similar way (Kaiser 1998).

Support for this hypothesis can be inferred from Hiddink et al.'s (2006) study of the relationships between trawling intensity, benthic biomass and richness in 4 areas in the North Sea subject to varying levels of natural disturbance. However, systematic tests of the hypothesis in many areas with data spanning many levels of natural disturbance have not been conducted. One approach that allows the hypothesis to be tested in a systematic way in many areas where communities will differ in species composition is to group benthic species in each area by traits that provide proxies for their role in community function (for review see Pearson 2001, Bremner 2008). Trait-based approaches have already been used successfully to describe the impacts of bottom trawling on benthic communities (Bremner et al. 2003, Tillin et al. 2006, de Juan et al. 2007, Kenchington et al. 2007, Bolam et al. 2014).

Here, we use a biological trait approach to assess the effects of trawling and natural disturbance on benthic community composition and function. We combine data from 8 studies of trawling impacts at different sites throughout the North and Irish Seas. Our results confirm the hypothesis that bottom trawling and natural disturbance have comparable effects on benthic communities. These findings are expected to be relevant to management in that they may help to identify areas that are more or less resilient to trawl impact (sensu Diesing et al. 2013) and support the development of management plans that take into account the environmental effects of fishing.

METHODS

Study area

The effects of trawling were assessed in 8 areas where soft-sediment benthic communities were sampled across a gradient in trawling disturbance. Of these, 7 were located in the North Sea and 1 in the Irish Sea (Fig. 1). Sampling sites were selected to cover the trawling intensity gradient in each area, while keeping the environmental conditions as homogenous as possible (Table 1). The 8 areas differed in terms of habitat type (expressed as depth, sediment type and primary productivity) and as such in their degree of natural disturbance, as predicted by calculating the force per unit area exerted on the seabed by the tidal currents (i.e. tidal-bed shear stress). Areas were categorised and named on the basis of their mean tidal-bed shear stress, assigning A to the area with the lowest shear stress and H to the highest. For 6 areas (A, B, D, E, F, G), homogeneity of other habitat characteristics was maximised by limiting the distance between sampling sites in the area and by selecting sites with similar habitat conditions (based on depth and sediment maps and/or habitat information from previous field studies; see Table 1: Ref. to area). For the 2 other areas (C, H), sampling sites were selected from monitoring sites in the Dutch Exclusive Economic Zone based on similarity of sediment grain size conditions, and so these covered a much larger area (Table 1: Ref. to area). Sample data from 4 areas (B, C, D, H) have been published previously (Table 1: Ref. to dataset), but have not been used to investigate benthic community composition and function. Temporal differences between areas,

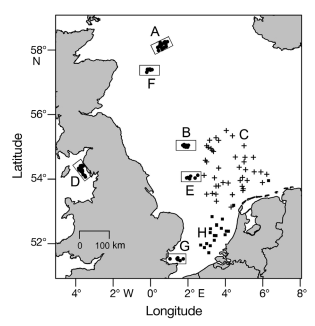


Fig. 1. The 8 different study areas (see Table 1 for area codes) and associated sampling sites

due to timing of sampling, were minimised as most of the selected areas were sampled between June and August (Table 1). In addition, we expected that temporal differences were of minor importance for explaining the variation in community composition due to the large differences in habitat conditions between areas.

Benthic sampling and trait classification

The number of benthic sampling sites ranged from 6 to 230 among areas, while the number of samples taken at each site ranged from 1 to 5 (Table 1). When there was more than 1 replicate per site, samples were pooled to provide an integrated description of the benthic community at each sampling site. The benthos was sampled using a 0.1 m² Day grab (Areas A and D), a 0.1 m² Hamon grab (Areas B and F) or a 0.078 m² Reineck box corer (Areas C, E, G and H). The different gears have a different penetration depth and sample a different surface area but they were selected because no single gear can operate effectively on all substrate types. However, all these gears do sample the smaller epi- and infaunal component of the benthic fauna and provide a quantitative estimate of their abundance and biomass (Eleftheriou & Moore 2005). Samples from all areas were sieved over a 1 mm mesh sieve and biota were identified to the lowest taxonomic level possible. Biomass

A: Fladen Ground; B: Dogger Bank; C: Dutch Exclusive Economic Zone (EEZ) fine sediment; D: Sellafield; E: Silver Pit; F: Long Forties; G: Thames; H: Dutch EEZ coarse Table 1. Overview of the different study areas and their characteristics. Areas are ranked on the basis of their mean tidal-bed shear stress (lowest in A, highest in H) (5) Tillin et al. (2006), (6) Hinz et al. (2009) (9) Jennings et al. (2002) (2011), (4) (7) Jennings et al. (2001a), (8) Jennings et al. (2001b), et al. ((2014), (3) Hiddink Denderen et sediment. (1) Queirós et al. (2006), (2) van

Area	Area No. of stations Sampling code (samples date per station) (mo/yr)	Sampling date (mo/yr)	Sampling device (surf. area in m^2)	Max. dist. between stations (km)	Depth (m)	Sediment type	Silt (%)	Prim. Tre product int (gr C m ⁻² yr ⁻¹) ($\begin{array}{c} {\rm Trawling} \\ {\rm intensity} \\ {\rm (Yr^{-1})} \end{array}$	$\begin{array}{c} \text{Shear} \\ \text{stress} \\ \text{(N m}^{-2)} \end{array}$	Ref. to dataset	Ref. to area
4	14 (5)	6/04	Day grab (0.1)	41	143-153	Mud	93-97	125-141	0.1-3.0	0.1-0.1		(5)
В	7 (5)	9/03	Hamon grab (0.1)	20	25-30	Sand	1-3	171 - 192	0.1-1.5	0.1 - 0.2	(1)	(1)
U	230 (1)	3-6/02-07	Box corer (0.078)	264	26-56	Sand	0 - 21	149-417	0.0 - 4.0	0.0 - 0.4	(2)	(2)
О	15 (5)	60/9	Day grab (0.1)	42	21 - 42	Muddy sand	42 - 89	ı	0.5 - 11.9	0.2 - 0.3	(3,4)	(3, 4, 6)
Ш	6 (4)	7/02		40	68-78	Muddy sand	9 - 27	173-184	$1.8-99^{a}$	0.2 - 0.3		(4,8,9)
Ц	7 (2–5)	9/03	Hamon grab (0.1)	19	74–83	Gravelly sand	2–6	183-198	0.1 - 1.6	0.3 - 0.4		(2)
U	6 (4)	7/02	Box corer (0.078)	49	16 - 40	Sand	3-22	130 - 340	$0-100^{a}$	0.5 - 1.1		
H	84 (1)	3-6/02-07		329	22–36	Sand	0-1	258-436	0.0-4.8	0.4 - 1.4	(2)	(2)
aAre	^a Areas E and G have different trawl disturbance	different traw	vl disturbance metric	s compared wi	th the other	with the other areas. These va	dnes rep	resent relative fre	equencies of c	disturban	disturbance on a linear scale	ear scale

per taxonomic group was estimated in grams ashfree dry weight (Areas C and H) or wet weight (other areas). Some large and low-density individuals are not effectively sampled by the gears and were removed from the data. We removed individuals when their biomass was larger than the mean biomass (excluding these individuals) of all samples in the area. A total of 29 large individuals were removed from the entire data set (e.g. a masked crab *Corystes cassivelaunus*, a common otter shell *Lutraria lutraria* and a heart urchin *Echinocardium cordatum*), representing <0.001% of all individuals and 17.8% of the total biomass.

Besides these few large individuals, most areas had 1 or 2 genera (heart urchins from the genera *Echinocardium* in 4 areas and *Brissopsis* in 2 areas, and a razor clam *Ensis* and a brittle star *Amphiura* in 1 area) that dominated the biomass of the sampled fauna (28 to 74% of total biomass). Fishing effects on these dominant taxa were expected to overshadow the wider community responses that are the focus of this study (see Supplement 1 at www.int-res.com/articles/suppl/m541p031_supp.pdf) and so we assessed the responses of these taxa to trawling separately.

We used a suite of 10 biological traits to describe changes in the function and resilience of benthic communities in response to bottom trawling and shear stress. In total, trait information was obtained for 222 different genera and 59 unique higher taxonomic groupings (mostly 'family') for which biomass data were available (Bolam et al. 2014). Each trait was subdivided into multiple modalities (Table 2). For each genera-trait combination, a single trait modality was assigned a score of 1 when the genus showed total affinity for that particular modality. When the genus could not be assigned unequivocally to a single trait modality, multiple modalities were assigned fractional scores that summed to 1, depending on the affinity of that genus for that modality (fuzzy coding, see Chevene et al. 1994). When genera could not be identified, traits were defined for higher taxonomic levels. From this genera-by-trait matrix (including the higher taxonomic levels), we calculated a table of sampling sites by biomass-weighted modalities. This was done for each sampling site by multiplying the total biomass per taxonomic grouping by the score for each trait modality. These were summed by modality to produce a biomass-weighted trait modality table for all sampling sites (Tillin et al. 2006, Bolam et al. 2014). Four taxonomic groups, representing 0.5% of the biomass, were excluded from the analysis as no trait data were available.

Trawl disturbance, natural disturbance and habitat conditions

To assess the intensity of trawling and natural disturbance, and to describe environmental factors that may affect community composition, we combined depth and sediment data collected during the benthic sampling with estimates of trawl disturbance, tidal-bed shear stress and primary production for the same sites.

Type of sediment, silt percentage and depth were site-specific data collected during the benthic sampling. The distinction in sediment type was based on the classification diagram of Folk (Folk 1954). Silt percentage was obtained from particle size analysis. Except for Areas C and H, depth was directly measured at the benthic sampling location. Depths for Areas C and H were extracted from bathymetric data (see van Denderen et al. 2014).

Table 2. Benthic traits, their modalities and corresponding abbreviations

Traits	Modalities	Abbreviation
Size (mm)	<21 21–100 101–200 >200	S<21 S21-100 S101-200 S>200
Morphology	Soft Exoskeleton	M_soft M_exo
Longevity (yr)	<3 3-10 >10	L<3 L3-10 L>10
Larval develop- ment	Planktotrophic Lecithotrophic/direct	LD_plank LD_le/di
Egg development	Pelagic Benthic Brooded	ED_pela ED_bent ED_brood
Living habit	Tube-dwelling Burrow-dwelling Free-living	LH_tube LH_burrow LH_free
Sediment position	Surface Shallow (0–5 cm) Mid-depth (6–10 cm) Deep (>10 cm)	SP_surf SP0-5 SP6-10 SP>10
Feeding type	Suspension-feeder Deposit-feeder Scavenger Predator	F_susp F_dep F_scav F_pred
Mobility	Sessile Swimmer/crawler Burrower	M_sessile M_swi/cr M_bur
Bioturbation activity	Diffusive mixing Surface deposition Others	BT_dif BT_dep BT_others

Estimates of the amount of trawl disturbance were based on quantifying the fishing activities of both beam and/or otter trawls. Both types of trawling disturb seabed sediment and impact benthic communities (Kaiser et al. 2006). Trawl disturbance for Areas A, B, C, D, F and H was estimated using satellite Vessel Monitoring System (VMS) data and expressed as the ratio between the area of the site that is trawled each year and the total area of the site (the data sources and exact calculations of trawl disturbance have been explained in previous articles, see Table 1: Ref. to area). Trawl disturbance for Areas E and G was estimated from aerial survey data collected by fisheries inspection services (Jennings et al. 2001a).

Tidal-bed shear stress was estimated using a 2-dimensional hydrographic model. This model predicts shear stress (the force per unit area exerted on the seabed by the tidal currents: $N\ m^{-2}$) per sampled station on a 1/8° longitude by 1/12° latitude spatial scale. The shear stress calculations are explained in more detail in Hiddink et al. (2006).

Primary productivity was obtained through predictions from GETM-ERSEM (General Estuarine Transport Model-European Regional Seas Ecosystem Model) (Baretta et al. 1995). GETM-ERSEM describes the temporal and spatial patterns of the biogeochemistry of the water column and sediment using 2 coupled hydrodynamic models. These models predicted total production of new phytoplankton biomass for each year (g C m $^{-2}$ yr $^{-1}$) on a 10 \times 10 km spatial scale. Total production was estimated for each sampling site, except for Area D, over a period of 1 yr prior to the sampling date. These modelled productivities approximate measured primary productivity (Ebenhoh et al. 1997).

Statistical analysis

We first analysed the effect of trawling on trait composition for all areas together by aggregating the sampling sites for each area into 'low', 'intermediate' and 'high' trawl disturbance treatments. We then analysed the effects of trawling on trait composition for each area individually using the gradient in trawling disturbance rather than the 3 categories. Finally, we analysed the effects of trawling for the few dominant genera separately (see Supplement 1).

For the first analysis, we examined the proportion of biomass per modality within trait categories, as this allowed us to compare areas that may vary

greatly in their total biomass and that were sampled with different gears. We defined low trawl disturbance as an intensity $\leq 0.2 \text{ yr}^{-1}$ as this means that there will be, on average, a trawl pass once every 5 yr. We defined intermediate trawl disturbance at an intensity of >0.2 to ≤ 0.5 yr⁻¹. All other sampling sites were grouped into the high trawl disturbance group. Since trawl disturbance of both Areas E and G was based on different metrics, we rescaled the overflight effort data of these areas to trawling intensity based on the maximum trawling intensity estimates found in Area E by Hiddink et al. (2006). Differences in trait composition between areas and 'low', 'intermediate' and 'high' trawl disturbance treatments were examined with a correspondence analysis that included bootstrapped p-values from a hierarchical cluster analysis (Suzuki & Shimodaira 2006).

In the second analysis, we described the effects of trawling on trait composition for each area separately using the trawling intensity gradient and biomass per modality (instead of proportion of biomass per modality). This was done with a redundancy analysis where we used the trawling intensity gradient as a predictor variable. With only 1 predictor variable present, the redundancy analysis is the multivariate analogue of linear regression (Legendre & Legendre 2012) and may be used to determine which trait modalities are positively or negatively correlated to trawl disturbance. The approach will show whether the observed shifts in response to trawling are relative, indicating that some organisms are less (negatively) affected by trawl disturbance than others, or absolute, indicating an increase in the biomass abundance of certain trait modalities at high trawl disturbance. The redundancy analysis assumes there is a linear relationship between the predictor variable and its response. For that reason trawl disturbance was log transformed as we expected the trait modalities sensitive to trawl disturbance to decline exponentially (Tillin et al. 2006, Hiddink et al. 2011). Whether trawl disturbance had a significant effect on community composition for each area was tested using a permutation test. Since sampling sites in Areas C and H covered a large spatial scale and were only selected on the basis of similar sediment grain size conditions, the effects of trawling in these 2 areas were examined using a partial redundancy analysis that controlled for the environmental conditions. All multivariate analyses were done using the package 'vegan' in program R (Oksanen et al. 2013).

Finally, we analysed the effects of trawling on a few dominant genera separately as their responses overshadow the community response to trawling (Fig. S1 in Supplement 1). This was achieved by investigating the relationship between trawling intensity and biomass which was $\log(x+1)$ transformed to improve model fit.

RESULTS

Effects of trawling on trait composition for all areas combined

When sites were grouped into 'low', 'intermediate' and 'high' trawl disturbance treatments, the analysis of trawling effects on trait composition showed that high trawling disturbance led to trait compositions similar to those in areas subject to high shear stress. The correspondence analysis based on trait composition produced 5 different clusters (Fig. 2). Five of the areas (A, C, F, G and H) remained within the same cluster at low, intermediate and high trawl disturbance, and this suggests that their trait composition does not change with trawling. The other 3 areas (B, D and E) had a similar trait composition at low and/or intermediate trawling (Figs. 2a & 3, Area D had no low trawl disturbance treatment), which is most associated with the modalities exoskeleton, a maximum longevity of >10 yr and suspension-feeder (Fig. 2b). Trawling caused significant changes in trait composition in Areas B, D and E, and these changes led to community compositions comparable with those in areas subject to high shear stress (Figs. 2 & 4). Thus, Area B, at intermediate and high trawl disturbance, and D, at high trawl disturbance, clustered with Area H. This group is most associated with the modalities swimmer/crawler, scavenger, predator and diffusive mixing activity. Area E clustered, at high trawl disturbance, with Areas F and G (Fig. 2a), and this group is most strongly associated with the modalities small-sized (<21 mm), a maximum longevity of <3 yr, surface-living, benthic or brood egg development, tube-dwelling and deposit-feeder.

No detectable effects of trawling were found in the area with the lowest shear stress (Area A) or in Area C. The trait composition for Area A, at low and high trawl disturbance, is most strongly associated with the modalities direct or lecithotrophic larval development, large-sized (>200 mm), a maximum longevity of < 3 yr, and soft-bodied (Figs. 2 & 3). Area C was not strongly associated with any specific trait modalities.

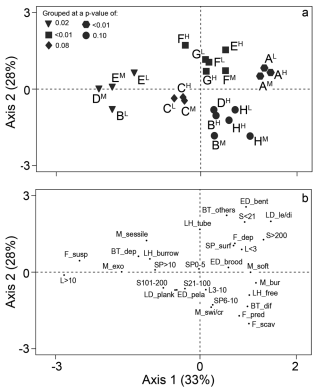
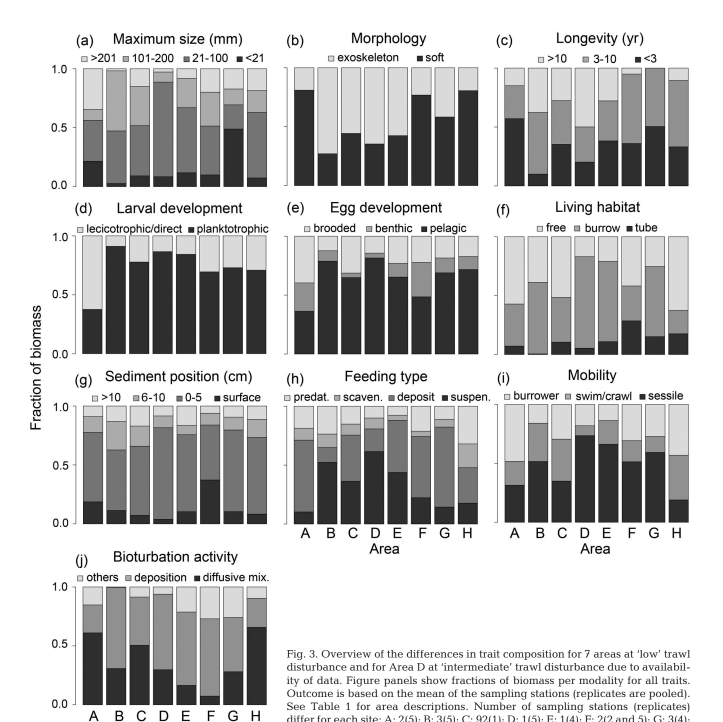


Fig. 2. Outcome of the correspondence analysis based on trait composition for (a) the different sampling areas, split into a 'low' (L), 'intermediate' (M) and 'high' (H) trawl disturbance treatment (see Table 1 for area descriptions), and (b) trait modalities. The correspondence analysis shows 5 different clusters in (a), grouped at a p-value between <0.01 and 0.10 (represented by different symbols). The abbreviations in (b) correspond to the trait modalities in Table 2

Effects of trawling on trait composition tested for each area separately

The 3 areas that show shifts in trait composition (Areas B, D and E) were also significantly affected by trawling when they were treated independently (Fig. 5). Trawling explained 52% of the variation in trait composition in B, 63% in D and 55% in E. A number of modalities were consistently and negatively correlated with trawl disturbance (Fig. 5) in all 3 areas. These were the modalities exoskeleton, sessile, suspension-feeder, planktotrophic larval development, pelagic egg development, burrowdwelling, positioned 0-5 cm in the sediment, surfacedeposition activity and a maximum longevity of 3-10 or >10 yr. A few abundant bivalve genera in these areas have all these modalities combined (such as Dosinia, Spisula, Acanthocardia, Ensis, Phaxas and Abra).



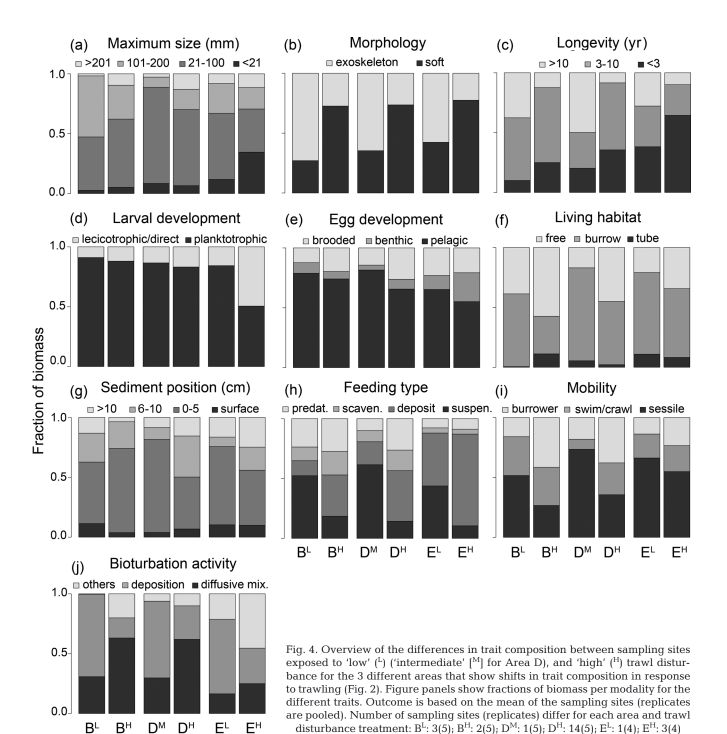
Trawling in Areas B and D was also negatively correlated with all other trait modalities and this means that the shift in trait composition is only based on relative increases. In contrast, trawl disturbance in Area E was positively correlated with the modalities smallsized (<21 mm), direct or lecithotrophic larval devel-

Area

opment and the bioturbation activity 'other'. This indicates absolute increases in biomass of fauna with these modalities in response to trawling, but these effects were not significant when tested using univariate statistics (see Table S2 in Supplement 2 at www.int-res.com/articles/suppl/m541p031_supp.pdf).

differ for each site: A: 2(5); B: 3(5); C: 92(1); D: 1(5); E: 1(4); F: 2(2 and 5); G: 3(4);

H: 9(1)



No significant effects of trawl disturbance on trait composition were observed in the other areas (all p-values in permutation tests > 0.24). Trawling explained only $9\,\%$ of the variation in trait composition in A, $0.3\,\%$ in C, $19.5\,\%$ in F, $26\,\%$ in G and $1.4\,\%$ in H.

Effects of trawling on dominant genera

Biomass in each study area was dominated by only 1 or 2 genera. The relationships between trawling intensity and log biomass of each of these dominant genera is shown in Table 3. The biomass of the dominant genera in Area D, *Amphiura* and *Echinocardium*, was negatively related to trawl disturbance, while no significant effects of trawling were observed in the other dominant genera.

DISCUSSION

We found no effects of trawling on benthic invertebrate communities at locations with high natural disturbance (Areas F, G and H), while in 3 out of 5 areas with more stable natural conditions, clear shifts were observed in trait composition in relation to trawling disturbance (Areas B, D and E). In these areas, trawling resulted in community compositions comparable with those in areas subject to high natural disturbance. Hence, our results provide support for the hypothesis that trawl and natural disturbance affect benthic communities in similar ways. Both sources of disturbance cause declines in long-living, hard-bodied (exoskeleton) and suspension-feeding animals and these effects are likely to affect community function.

The comparable effects of trawl and natural disturbance may help to identify areas that are particularly susceptible or resistant to trawl disturbance. Methods to identify such areas have already been proposed by Diesing et al. (2013), who estimated the probability that fishing disturbance exceeded natural disturbance by comparing fishing intensity and bed shear stress. They identified areas that are expected to be particularly vulnerable or resilient to bottom fishing on the basis of the bed shear stress in these areas without an understanding of the associated benthic communities. Our results broadly confirm the applicability of their proposed method, even though there will be some complexity of response that reflects local site characteristics.

Trawl disturbance reduced the proportion of 10 modalities of 9 different traits in 3 areas subject to low shear stress. The same type of trait modalities have been observed to decline in previous studies (see Kaiser et al. 2006 for a meta-analysis, Tillin et al. 2006, de Juan et al. 2007, Kenchington et al. 2007). Most of these studies focused on the epifaunal benthic component, while our results clearly reveal that similar effects of bottom trawling may be expected when infaunal data are used. Trawling had most adverse effects on infaunal organisms positioned between 0 and 5 cm in the seabed, and this may be expected as trawl gears penetrate at least a few cm into most soft sediments (Eigaard et al. 2015). Species positioned deeper in the sediment and species living

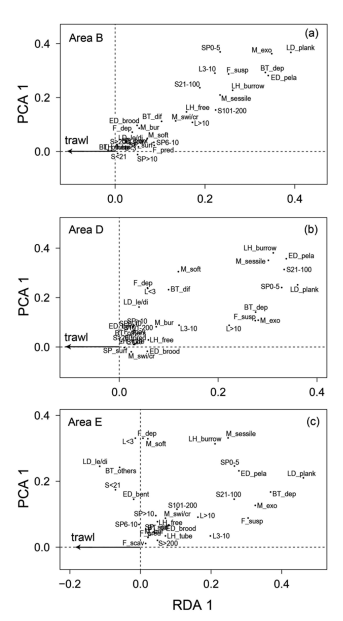


Fig. 5. Outcome of the redundancy analysis for all areas (B, D, E; see Table 1) where trawl disturbance had a significant effect on community composition (all p-values <0.05). Abbreviations correspond to the trait modalities in Table 2

on the seabed surface were less affected by trawling in our study. The latter is unexpected but could be explained by the fact that most surface-living animals in our dataset were mobile swimmers or crawlers, which may repopulate trawled grounds more easily after trawling disturbance (de Juan et al. 2007). The effects of trawling led, in our study, to community compositions comparable with those in areas subject to high natural disturbance, being composed of either small-sized, deposit-feeding animals or mobile scavengers and predators.

Table 3. Trawl effects on 4 different genera that were not included in the trait-based analysis as their responses overshadow the community response (see Supplement 1). Analysis was done using a linear regression model, genera biomass is $\log(x+1)$ transformed. The regression model for Area C includes the effects of productivity and percentage silt, which are both significantly related to *Echinocardium* biomass (not shown). See the Table 1 legend for area codes

Genera	Area	Intercept	Slope	\mathbb{R}^2	р
Brissopsis	Α	1.07	0.32	0.08	0.17
Echinocardium	В	1.48	-0.52	0.08	0.48
Echinocardium	С	-0.06	0.04	0.09	0.31
Echinocardium	D	2.45	-0.26	0.48	0.003
Amphiura	D	2.35	-0.26	0.57	0.001
Brissopsis	E	2.09	0.02	0.11	0.51
Echinocardium	Н	0.27	-0.02	0.00	0.50
Ensis	Н	0.14	0.00	0.01	0.92

The clearest indication of changes in community function in response to trawling is the strongly negative association of surface deposition (a modality of bioturbation) with trawl disturbance. This is mainly the result of a decline in suspension-feeding organisms. A decline in the biomass of this functional group means that less organic material is captured in the water column and deposited onto the seabed (Gili & Coma 1998, Snelgrove 1999, Pearson 2001, Lohrer et al. 2004, Thrush & Dayton 2010), which potentially reduces benthic secondary production by suspension feeders in the trawled areas.

We detected no effects of trawling in the area characterised by the lowest shear stress (Area A). The community composition of this area, at both low and high trawl disturbance, is most similar to a naturally perturbed community, but differs as many organisms are large-sized and have direct or lecithotrophic larval development. These types of development are often observed in deep areas with limited amounts of planktonic food (Vance 1973). Indeed, Area A is located deepest and has the lowest primary productivity of our study areas. Food limitation has also been observed in the area for one long-living suspension-feeder, Arctica islandica, which had relatively low growth rates (Witbaard et al. 1999). The changes in the benthic community in response to a low benthic production in the area could have interfered with the response of the benthic community to the effect of trawl disturbance.

We sampled the smaller and more abundant epiand infaunal component of the benthic ecosystem. This resulted in a relatively low power to detect the effects of trawling on larger epifauna (species like shrimps, starfish and sea pens). In 2 of the areas where we detected no effects of trawling (A and F), trawl effects have been found on trait composition for larger epifauna sampled using a small beam trawl (Tillin et al. 2006). In these datasets, long-living and suspension-feeding trait modalities were particularly negatively affected by trawl disturbance. Furthermore, we found no effects of trawling in Area C, while trawl effects on benthic species richness have been previously detected in this area (van Denderen et al. 2014). Conversely, we observed fishing effects in Area B, whereas no effects of trawling on larger epifauna have been detected in this area (Tillin et al. 2006). These comparisons show that trawling can have differential effects on different components of the benthos, with the result that impacts may be overlooked unless several sampling gears and community indicators (e.g. diversity, biomass and trait composition) are used.

The effects of trawling were examined in all areas over a gradient of commercial bottom trawling intensity. Such a comparative analysis can result in differences in community composition along the trawling gradient that seem to be related to fishing impact, while in fact these patterns result from the fishery selecting areas with a particular community composition where they catch the most fish (see also Tillin et al. 2006). Such effects may be especially relevant at large spatial scales (scales at which the fishery fleet operates), where a large part of the variation in fishing effort can be explained by gradients in environmental conditions (van Denderen et al. 2014). However, others have shown that unfished habitats are not necessarily unsuitable for fishing (Dinmore et al. 2003) and it has been suggested that fisheries often return to areas that are known to be free from obstructions that could damage the gear (Holland & Sutinen 2000). In addition, Tillin et al. (2006) postulated that trawl effects on benthic communities can have a much larger impact than is expected from small changes in environmental conditions. This is also true for our study areas, where even large differences in silt content of the seabed sediments in Areas B and E (Table 1) were associated with similar trait compositions (Figs. 2 & 4).

Since the North and Irish Seas have been trawled for a long time (Brander 1980, de Groot 1984), the sampling stations least exposed to trawl disturbance are unlikely to be in a pristine state. For those areas where there were no detectable effects of trawling, we cannot be certain whether the areas had not recovered from historic trawl disturbance or whether the effects of trawl disturbance were relatively low in relation to natural disturbance. However, it is clear

from our results that trawl disturbance has a limited additional effect on the benthic ecosystem in areas exposed to high shear stress compared to areas exposed to low shear stress.

In a previous study conducted in Area E, Jennings et al. (2001b) examined whether there was an increase in biomass of some benthic species in response to trawling. In their study, they investigated the benthic infauna as an aggregated group and this approach revealed a lack of a significant positive response to trawl impact. They recommended that future trawl studies should focus on the smallest macrofauna (and meiofauna) as these have sufficiently fast life cycles to benefit from trawl disturbance. These faunal groups were positively correlated with bottom-trawling intensity in our study in Area E, although their increase was not statistically significant when trait modalities were individually analysed (see Supplement 2). The increase was mostly related to a high abundance of polychaetes from the family Scalibregmatidae and, to a lesser extent, the Sipunculidae at the trawled stations. Such an increase may be expected when the species that are relatively less sensitive to trawl disturbance benefit from an increase in available food, due to a decline of their more sensitive competitors (Jennings et al. 2001b, Hiddink et al. 2008, van Denderen et al. 2013).

The biomass of fauna in most of the study areas was dominated by 1 or 2 genera. The responses of these genera to trawling were assessed separately to avoid confounding responses of the community (see Supplement 1). Except for Area D, where the biomass of Amphiura and Echinocardium decreased significantly, none of the dominant genera showed a response, despite their assumed sensitivity to trawl gears (Bergman & van Santbrink 2000, Callaway et al. 2007). Although individuals are heavy in relation to other fauna, the small sample sizes taken by grabs and cores may not provide a good indication of mean density and thus the power to detect any responses to trawling disturbance will be low. Samples from small beam trawls and dredges that sample larger areas and integrate some of the expected small scale patchiness in the distribution of these genera would likely provide better data for testing whether these genera are affected by trawling disturbance. These genera contain species that are important habitat facilitators, e.g. the brittle star Amphiura filiformis and the heart urchins Brissopsis lyrifera and Echinocardium cordatum (Hollertz & Duchêne 2001, Lohrer et al. 2004, 2013, van Nes et al. 2007), and these may facilitate other benthos by providing resources and shelter (Thrush et al. 1992, Stachowicz

2001). Effects of trawling on these facilitators may, hence, indirectly affect the benthic component that is the focus of this study.

We conclude that high levels of natural disturbance that affect soft-sediment habitats will lead to community compositions and functions that are more resilient to a given level of trawling disturbance than those found in areas with less natural disturbance. Such asymmetric impacts of bottom fishing will help to identify areas that are particularly susceptible or resilient to trawling and thereby support the development of spatial management plans that deal with the ongoing process of balancing fisheries exploitation and conservation of marine benthic ecosystems.

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