Influence of Benthic Macrofauna as a Spatial Structuring Agent for Juvenile Haddock (Melanogrammus aeglefinus) on the Eastern Scotian Shelf, Atlantic Canada

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Abstract

We examined the habitat of juvenile haddock on the eastern Scotian Shelf (off Nova Scotia, Canada) in relation to grab-sampled benthic macrofaunal invertebrate species assemblages in order to determine whether there were significant differences in benthic macrofauna between areas of historically persistent high and low juvenile haddock abundance. Our analyses were conducted over two spatial scales in each of two years: among banks (Emerald, Western and Sable Island), approximately 60 km distant from each other, and between areas of high and low juvenile haddock abundance at distances of 10 to 30 km—all in an area that had not experienced groundfishing in the decade prior to sampling. We also examined fine-scale (10s of metres) within-site variability in the macrofauna and used surficial sediment characteristics, along with hydrographic variables, to identify environmental correlates. PERMANOVA identified statistically significant differences in biomass, density and composition of the benthos associated with juvenile haddock abundance; however it was difficult to determine whether the results had biological relevance. Post hoc tests showed that these differences occurred only on Sable Island Bank where both fish and benthos may have been independently responding to sediment type which was most different there (100% sand in the area of low haddock abundance vs. 22% gravel in the area of high haddock abundance). In total, 383 benthic taxa representing 13 phyla were identified. Annelida was the most specious phylum (36.29% of taxa, representing 33 families), followed by Arthropoda (with Crustaceans, mostly Amphipoda, accounting for 25.07% of the total number of taxa). The strongest pattern in the macrofauna was expressed at the largest scale, between banks, accounting for approximately 25% of the variation in the data. Emerald Bank, deeper, warmer and saltier than the Western and Sable Island Banks, had a distinctive fauna.
Introduction

As in other vertebrates, the spatial structure of fish populations is shaped both by factors endogenous to the populations and by habitat heterogeneity [1]. In marine ecosystems, currents and water masses play major roles in determining fish distributions at large spatial scales (100s of kms), while smaller-scale structure is often attributed to endogenous behavioral responses [2–9]. Within populations, heterogeneities of seabed habitat, both biotic and abiotic, have generally been under-studied as a spatial-structuring agent for boreal fishes living on continental shelves. Exceptions include those species that clearly utilize particular habitats for spawning (e.g., herring Clupea harengus) or burrowing (e.g., sand eels Ammodytes spp.). Yet, it is well known that benthic species and habitats play a critical role in the population dynamics of some marine fish [10,11], especially the juveniles. Benthic invertebrates are known to provide food [10–13], while biotic and abiotic benthic habitat can provide refuge [10,14].

Establishing a relationship between fish density and the spatial structuring of benthic species and habitats requires joint collection of data over the range of spatial scales relevant to the fish populations. A number of studies have related fish distribution to the presence of structure-forming benthic species such as corals and sponges [15,16], or to particular physical habitats [17–19], but few studies have examined the spatial structuring of benthic species and habitats over the range of spatial scales relevant to fish stock units. Recently, Sell and Kröncke [20] found a correlation between benthic species distributions and demersal fish assemblages on the Dogger Bank (North Sea), which is one of the few temperate areas where the benthos and the fishery resources have been surveyed over a similar spatial extent. There, the two assemblages showed similar spatial structuring, likely arising through common responses to depth and sediment type. In some instances, individual fish species showed spatial correlation with invertebrate prey species.

If benthic invertebrate species distributions impose spatial structure within continental-shelf fish populations, then the relationship is most likely to be identified in those demersal fish assemblages which live most intimately with the sea floor [20], and especially in those species which have benthic life-history stages. In 2000, Canada’s Department of Fisheries and Oceans (DFO) commenced a multiyear, multidisciplinary project on the Scotian Shelf to investigate the role of seabed habitat for a demersal fish species and to test remotely-sensed seabed classification systems [19,21]. Fish on the Scotian Shelf are distributed independently along environmental gradients, rather than comprised of highly co-evolved, inter-dependent species [22]. Consequently, a single key species, haddock (Melanogrammus aeglefinus), rather than a fish assemblage was selected for study. 

Haddock is a commercially harvested gadoid that is intimately associated with the seabed. It is found in the North Atlantic at depths from less than 50 m to about 350 m and temperatures ranging from 4–8°C [23]. On the Scotian Shelf, haddock spawn on gravel bottoms in April/May. The young have a 4-to-5 month pelagic phase, before moving to the seabed for the remainder of their juvenile stage [24,25]. That shift from a pelagic to a benthic existence occurs at approximately 8 cm length and is reflected in their diet [26]. Haddock populations typically exhibit highly temporally variable recruitment [22], which is commonly attributed to the effects of the environment and food supply during the pelagic phase [12], although the period of transition to the seabed and the following months as benthic juveniles have been cited as a determinant of year-class strength [3].

The haddock on the eastern Scotian Shelf are considered as a single management unit, with a range that formerly included the southern Gulf of St. Lawrence (encompassing Northwest Atlantic Fisheries Organization (NAFO) Divisions 4TVW), and as distinct from adjacent stocks on the western Scotian Shelf and north of the Laurentian Channel. In recent decades,
the majority of eastern Scotian Shelf haddock have lived, and spawned, on the offshore banks from Emerald Bank in the west to Banquereau Bank in the east. In 1984, DFO closed a large area (within NAFO Division 4W) to trawlers in an attempt to reduce discarding of undersized haddock [27,28]. In 1987, a larger area on Western and Emerald Banks, reaching \(12,776 \text{ km}^2\) or 13% of the area occupied by the population [28], was closed year round to most groundfish fishing (though not to long-lining with large hooks for part of this time, nor to scallop dragging which was very light in this region and outside of the study areas). In September 1993, the fisheries for cod and haddock on the eastern Scotian Shelf were closed, to protect the depleted stocks, and remain so today. Those closures allowed a rare opportunity to sample benthic species and physical habitats on the fishing grounds of the eastern Scotian Shelf in a relatively undisturbed state.

We used that opportunity to examine the role of benthic macrofaunal communities as a spatial-structuring agent for juvenile haddock on the eastern Scotian Shelf. While other studies have focused on benthic habitat features [15–18] or on broad correlation of species distributions [20], we took a different approach. Areas (100 km\(^2\)) of persistent high and low juvenile abundance were identified on each of three offshore banks within the area occupied by the stock unit, based on the probability of encountering juvenile haddock determined through the analyses of 32 years of data from the DFO summer groundfish surveys [21]. We then sampled the benthic macrofaunal invertebrate communities in those areas in each of two years to address the question: Do areas of preferred juvenile haddock habitat (areas of persistent high abundance) differ in benthic macrofaunal species composition from those that are not utilized as much (areas of persistent low abundance)?

Our intensive, temporally replicated grab-sampling also provided new data on the benthic macrofaunal communities of the eastern Scotian Shelf. In general, the benthic macrofauna of the offshore Scotian Shelf are poorly studied, and our study is also the first to compare those communities over large spatial scales. The grab-sampled macrofauna on Western and Banquereau Banks have been examined previously over small spatial scales in relation to fishing impacts [29–31], while others [32,33] minimally extended their own studies to the north and south respectively onto the Scotian Shelf. We additionally examined surficial sediment distribution, combined with dynamic bed-form characteristics (e.g., sandwaves) interpreted from sidescan sonar mosaics [34], along with hydrographic characteristics, to interpret the fine-scale (10s of metres) within-site and larger-scale (between sites within banks, between banks) macrofaunal patterns.

**Materials and Methods**

**Study Area**

The Scotian Shelf is the portion of northwest Atlantic continental shelf lying off Nova Scotia, Canada, between the Laurentian and Northeast Channels. It is approximately 700 km long and between 125 and 230 km wide, and characterized by highly productive marine ecosystems and complex bottom topography [35]. The Laurentian Channel delimits the Scotian Shelf in the northeast, while to the southwest the Northeast Channel separates it from Georges Bank. The outer part of the Scotian Shelf is characterized by a number of large, shallow banks separated by transverse troughs; Sable Island Bank rising above the surface as its eponymous island [35]. The present study focused on three outer Scotian Shelf banks; Emerald, Western and Sable Island Banks (Fig 1).

Historically these banks were heavily exploited, having a fishing history dating back to the 17th century. Groundfish, particularly Atlantic cod (\textit{Gadus morhua}) and haddock, were the mainstay of the 20th century fisheries [36]. Overfishing and environmental variability led to
population collapses, which affected both fisheries and trophic structure. Most groundfish fisheries were closed in September 1993 as noted previously, but recovery has not followed expected trajectories [37].

**Sampling Design**

Catches of juvenile haddock (including all younger than the age of sexual maturity) taken by summer research-vessel bottom-trawl surveys on Emerald, Western and Sable Island Banks from 1970 to 2001 had previously been mapped and areas of persistent high and low density identified [21]. While spatially variable within each bank, juvenile densities were similar on Emerald and Western Banks, and only slightly lower on Sable Island Bank [19]. Habitat utilization changes over time, however. In particular, when abundance is high, haddock expand over a larger area, apparently occupying less preferred habitats [27] and obscuring the relationship between optimal habitat types and fish distributions. We therefore focused on longer-term relationships, captured in the 32-year span of the survey data. In the event, a very strong 1999 year class of haddock was broadly spread across the Scotian Shelf during the period of our field program [21, 38]. Hence, spatial variations in the short-term responses of the benthos to the fish and vice versa may have been blurred.
Three pairs of 100 km$^2$ study sites, one pair on each bank (Table 1, Fig 1), were selected as representative of areas with persistently the highest and lowest juvenile haddock densities, respectively within each bank [21]. We designated the members of each pair as “High” or “preferred” and as “Low” or “non-preferred” areas. Survey trawling for fish prior to benthic sampling in 2002 and in the last year of the study in 2005 found that the densities of juvenile haddock in the High sites on Western and Sable Island Banks were approximately an order of magnitude greater than those in the paired Low sites, consistent with the differential densities seen in the 1970–2001 survey data, but densities on the two Emerald Bank sites were equal [38]. On Emerald and Western Banks the paired sites were approximately 10 km distant from one another, while on Sable Island Bank they were approximately 30 km apart. Within each 100 km$^2$ study site a 1 km x 5 km swath was randomly selected for detailed study.

**Benthos Sampling**

Between 6 and 12 grab samples were taken from each swath in each of 2003 and 2005 (Table 1, Fig 2). Benthic invertebrates were sampled with a video-grab, an electro-hydraulically actuated grab fitted with video cameras and halogen lights, which samples a 0.5 m$^2$ area of seafloor [39,40]. The cameras were used to increase sampling efficiency by ensuring that the bottom was suitable for sampling and that the grab closed properly before being recovered. An ORE Trackpoint II ultra-short baseline acoustic tracking system was used to determine the position of the video-grab relative to the ship [41]. No permits were required to undertake this sampling and no endangered or protected species were collected in the samples.

The video-grab contents were washed over a 1 mm screen, with retained material being preserved in buffered formalin on board ship. After return to the laboratory, all organisms were identified to species level, where possible. Poor condition of specimens, lack of information about juvenile forms or gaps in taxonomic knowledge prevented species-level identifications for some specimens. Abundance and biomass (formalin wet weight, including both mantle cavity liquid and shells for molluscs) were determined for each taxon.

**Environmental Data**

Depth and sediment type were determined for each grab sample, as were the estimates of mean bottom salinity, temperature and current (S1 Table, Fig 2). Depth was recorded from the ship's
sounder. Five sand and nine gravel sediment types had previously been interpreted from sidescan sonar mosaics and towed video observations on each swath. The categories were defined by sediment grain size and superimposed dynamic bedforms [34,42]. Sediment types were: Sand (S); Sand with Megaripples (SM); Sand to Sandy Gravel (SG); Sand with Scattered Boulders (SB); Sand Ribbons (SR); Gravel (G); Gravel to Gravelly Sand (GS); Gravel Ripples (GR); Gravel Ripples, Short Wave Length (GRS); Gravel Ripples, Long Wave Length (GRL); Gravel Ripples, Incised (GRI); Gravel Lag (GL); Gravel, Hummocky (GH); and Gravel with Small Sand Patches (GSP), although not all categories occurred in patches large enough to be visible on the maps presented here (Fig 2). Additionally, zones of boulders and sand ribbons, individual boulders, and the orientations of bedform crests were mapped [34]. Each grab sample was associated with one of these categories, based on its mapped position (S1 Table). The Sable Island Bank Low swath was the most homogeneous, with only categories S, SM, and SG present; the matching High site being mostly sand (78%), though with 4 gravel categories present (Fig 2). The Emerald Bank sites contained both sand and gravel categories (Fig 2), the Low site being 60% sand and the High site, 71%. The Western Bank High site was 60% sand, and the Low site there was 66% sand. Overall, Western Bank had the greatest diversity of sediment types (S1 Table, Fig 2).

Due to the absence of direct measurements, data for mean bottom temperature, salinity, and current speed were extracted from interpolated surfaces extracted from the Global Ocean Reanalysis and Simulations (GLORYS) model by Dr. Z. Wang, Oceans and Ecosystem Science Division, Bedford Institute of Oceanography. GLORYS is a numerical general circulation model with 1/4° resolution. The model used observational data collected from 1992 to 2011. Values for each sample were drawn from the interpolated surfaces for each variable (created with ordinary kriging) in ArcGIS v. 10.1 (Environmental Systems Research Institute, Redlands, CA).

Statistical Analyses

Species accumulation curves were constructed with PRIMER version 6.1.6 [43] separately for each study site and year to ensure that the benthic assemblages were adequately sampled prior to analysis. The number of observations (Sobs) was permuted 999 times to produce standard deviations.

The Total Number of Taxa (S), Total Abundance of organisms (individuals m\(^{-2}\)), Total Biomass of organisms (g m\(^{-2}\)), Pielou’s Evenness Index (J’) and Shannon-Wiener’s Diversity Index (H’) were determined. Levene’s statistic was used to test the null hypothesis that the group variances were equal. Most variables had homogeneous variances (P ≥ 0.01) without transformation, however total biomass required ln (x+1)-transformation to achieve homogeneity of variances. Their variations were tested using full factorial univariate ANOVAs (performed with IBM SPSS Statistics, version 20.0.0, IBM Corporation, Somers, New York), with three fixed factors: (Juvenile) Haddock Abundance (2 levels: High, Low); Bank (3 levels: Emerald, Sable Island, Western); and Year (2 levels: 2003, 2005). Sums of squares were calculated taking the interaction terms into consideration. Tukey’s HSD was used to test for equality of group means in post hoc tests of significant factors and interactions. Statistical significance was evaluated after Bonferroni correction for multiple tests (α ≤ 0.001).
Bray-Curtis similarities were calculated on the ln (x+1)-transformed species abundance and biomass data and on the untransformed presence/absence of species (including colonial taxa). Prior to statistical analyses, taxa which contributed ≤ 1% of total abundance in each sample year were removed to reduce the effect of rarities on the analyses [44]. The effects of setting the removal criterion at ≤ 3% or ≤ 5% were examined but results did not differ from those which arose with a ≤ 1% cut off. Using the same design as for univariate ANOVA, permutation multivariate analyses of variances (PERMANOVAs) were conducted on each matrix with 999 permutations [45]. Permutation of residuals was performed under a reduced model [46] and permuted pairwise tests of significant factors were conducted. Statistical significance was evaluated after Bonferroni correction for multiple tests (α ≤ 0.001).

nMDS plots were used to visualize variations related to significant factors. Similarity percentages tests (SIMPER) were used to determine the macrofaunal taxa that contributed most to significant dissimilarities among factors. PERMANOVA, nMDS and SIMPER routines were implemented in PRIMER-E (Plymouth Routines in Multivariate Ecological Research; Primer-E Ltd., 3 Meadow View, Lutton, Ivybridge, UK).

The relationships between the environmental parameters (depth, sediment type (categorical variable), mean bottom temperature, salinity and current) and the abundance and biomass of the benthic communities were examined using a distance-based linear model (DISTLM) and distance-based redundancy analysis ordination (dbRDA) performed on the Bray-Curtis similarity matrices. The best selection procedure was run with 9999 permutations and with the adjusted R² selection criterion implemented in PRIMER-E.

Results

Description of the Benthos

A total of 383 benthic taxa representing 13 phyla were identified (S2 Table). Of those, 52 species were only observed once and 30 were only observed twice. The samples collected in 2005 showed greater abundance and biomass and presented 52 more species than those collected in 2003. The Annelida was the most speciose phylum (36.3% of taxa, representing 33 families), followed by Arthropoda (with Crustaceans, mostly Amphipoda, accounting for 25.1% of the total number of taxa), Mollusca (19.1%—mostly Bivalvia and Gastropoda), Cnidaria (7.8%), Echinodermata (5.0%), Bryozoa (2.9%), Chordata (1.0%—mostly genera of Ascidians), and seven other phyla accounting for 2.9% of the total. Polychaetes and amphipods prevailed in abundance with 10 to 13 species accounting for 50% of the total, whereas bivalves and echinoderms prevailed in biomass, with 4 to 7 species accounting for 90% of the total. The three most frequently sampled species, based on presence/absence data, were the amphipod Unciola irrorata, and the polychaetes Ampharete finmarchica and Clymenura borealis.

The average Number of Taxa (S) found in each of the six study sites ranged from 44 to 69 (2003) and from 44 to 73 (2005). The highest numbers of taxa were found within Western Bank samples; two grab samples from the Low Haddock Abundance site sampled in 2003 yielded the highest number of taxa (126 and 109 species). Average Total Abundance ranged from 779 individuals m⁻² (Emerald Bank, High Haddock Abundance) to 2,578 individuals m⁻² (Western Bank, High Haddock Abundance) in 2003 and from 1,759 individuals m⁻² (Sable Island Bank, Low Haddock Abundance) to 2,406 individuals m⁻² (Western Bank, Low Haddock Abundance) in 2005. The top four most abundant taxa in both years were the bamboo worm Clymenella zonalis, the amphipod Unciola irrorata, and the polychaetes Polygordius sp. and Chone sp. Average Total Biomass ranged from 0.0176 kg m⁻² (Emerald Bank, High Haddock Abundance) to 12.116 kg m⁻² (Sable Island Bank, Low Haddock Abundance) in 2003 and from 0.212 kg m⁻² (Emerald Bank, High Haddock Abundance) to 44.538 kg m⁻² (Western
Bank, High Haddock Abundance) in 2005. The highest biomass was found for echinoderms and bivalves, with the sea cucumber *Cucumaria frondosa*, the propeller clam *Cyrtodaria siliqua*, and the sand dollar *Echinarchnnius parma* showing the highest biomass per species.

**Effects of Haddock Abundance, Bank and Year on Diversity Indices**

Species-accumulation curves for each bank and area within bank approached the asymptote suggesting that sampling was adequate to compare species richness among stations (S1 Fig). Shannon’s Diversity (H’), Pielou’s Evenness Index (J’) and the Total Abundance of organisms (individuals m⁻²) showed no significant differences among factors (S2 Fig) or their interactions in the univariate ANOVAs. A significant model effect was found for the Total Number of Taxa (S) but none of the individual effect tests were significant. Total Biomass of organisms (g m⁻²) was significantly different among Banks, among levels of Haddock Abundance and in the interaction of those factors (S3 Table). Post hoc tests revealed significantly lower transformed macrofaunal biomass on Emerald Bank than on the other banks and in the areas where there was High Haddock Abundance versus Low (S3 Table). The significant interaction between these factors resulted from the High Haddock Abundance site on Sable Island Bank grouping with the Emerald Bank sites, and the Low Haddock Abundance site with the Western Bank sites as indicated by Tukey’s HSD post hoc test; the trends between High and Low Haddock Abundance within each Bank were all in the same direction.

**Community Analyses**

**Macrofaunal Species Abundance.** PERMANOVA of the transformed abundance of the macrofauna showed an interaction between the factors Haddock Abundance and Bank, and one between Bank and Year; all other interactions terms were non-significant (Table 2). Each of the three factors was significant, with Bank explaining the largest proportion of the variance, followed by Year and Haddock Abundance (Table 2). Post hoc pairwise tests identified significant differences between every combination of main-effect factor levels. The Haddock Abundance x Bank interaction identified significant differences between High and Low Haddock Abundance within Sable Island Bank but not in the other banks. The Bank x Year interaction had significant differences between all combinations.

nMDS plots confirmed the greater explanatory power of the Bank factor over Haddock Abundance (Fig 3). Labelling of the samples by Year showed no change to the observed pattern with large overlap between years within banks (Fig 3). SIMPER identified a high degree of variability among grab samples within each Bank, with average similarity ranging from 36.10% on

Table 2. PERMANOVA of the transformed abundance of macrofaunal taxa based on Bray-Curtis similarity.

| Source              | Degrees of Freedom | Sums of Squares (SS) | Mean Square (MS) | Pseudo-F | P(Perm) | Est. of Variance Component | Sq. Root of Variance Component |
|---------------------|--------------------|----------------------|------------------|----------|---------|---------------------------|-------------------------------|
| Haddock Abundance   | 1                  | 7421.3               | 7421.3           | 4.571    | 0.001   | 100.15                    | 10.01                         |
| Bank (BA)           | 2                  | 50231                | 25116.0          | 15.469   | 0.001   | 606.97                    | 24.64                         |
| Year (YR)           | 1                  | 8271.1               | 8271.1           | 5.094    | 0.001   | 114.82                    | 10.72                         |
| HAxBA               | 2                  | 19322.0              | 9661.1           | 5.951    | 0.001   | 415.33                    | 20.38                         |
| HAxYR               | 1                  | 2892.2               | 2892.2           | 1.781    | 0.043   | 43.83                     | 6.62                          |
| BAxYR               | 2                  | 9053.7               | 4526.9           | 2.788    | 0.001   | 150.03                    | 12.25                         |
| HAxBAxYR            | 2                  | 3305.0               | 1652.5           | 1.018    | 0.401   | 2.99                      | 1.73                          |
| Residual            | 108                | 1.75 x10⁵            | 1623.6           |          |         | 1623.60                   | 40.29                         |
| Total               | 119                | 2.78 x10⁵            |                  |          |         |                           |                               |

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Emerald Bank to 43.5% on Western Bank (Table 3). Ten taxa contributed to 50% of that variability on Emerald Bank, with 14 taxa accounting for that level on each of Sable Island and Western Banks. In the associated taxon lists (Table 3), only Emerald Bank had no molluscs or echinoderms present in that percentage, and variability was influenced by small polychaetes and crustaceans. The small amphipod *E. fasciatus* was not found in Sable Island Bank samples and accounted for 2% of the variation between Emerald and Sable Island Bank (Table 4). These two areas were the most dissimilar (75.84%; Table 4) and the dissimilarity was drawn from a large number of species, with 12 taxa contributing to just 20% of the total. Emerald and Western Banks had dissimilar community composition (71.39%) with 11 taxa contributing to 20% of the total. A maldanid polychaete *Clymenella zonalis* and the small amphipod *E. fasciatus* contributed to 4% of the total dissimilarity, with *E. fasciatus* present in the Western Bank samples but at lower abundance than in the Emerald Bank samples (Table 4). Western and Sable Island Banks were less dissimilar (65.10%) and the single taxon contributing most to that difference was again the maldanid polychaete *C. zonalis* (Table 4). SIMPER analysis of the two levels of Haddock Abundance showed that they were 68% dissimilar with 127 taxa contributing to 90% of that variation and all differences between those due to proportional abundance differences between the two groups and not absence of taxa. The maldanid

![Fig 3. Non-metric multidimensional scaling (nMDS) of transformed abundance-based Bray Curtis similarities.](image-url)
polychaete *C. zonalis* explained the highest proportion of the variation of the individual taxa, although that was only 2%. This species was present in greater abundance in the samples from areas where Haddock Abundance was high.

| Bank (%Sim.) | Taxon                  | Av. Abund. | Av. Sim. | Sim./SD | Cont.% | Cum.% |
|--------------|------------------------|------------|----------|---------|--------|-------|
| Emerald      | *Unciola irrorata*     | 2.75       | 2.81     | 1.71    | 7.78   | 7.78  |
| (36.10%)     | *Clymenura borealis*   | 2.39       | 2.62     | 1.00    | 7.26   | 15.04 |
|              | *Ericthonius fasciatus* | 2.67       | 2.11     | 1.06    | 5.85   | 20.90 |
|              | *Chone sp.*            | 2.77       | 1.99     | 1.15    | 5.50   | 26.40 |
|              | *Aglaothamnus circinata* | 1.63       | 1.86     | 1.69    | 5.16   | 31.55 |
|              | *Glyceria capitata*    | 1.88       | 1.74     | 1.50    | 4.82   | 36.38 |
|              | *Lumbrinerides acuta*  | 1.54       | 1.18     | 0.92    | 3.27   | 39.65 |
|              | *Ampharete finmarchica*| 1.32       | 1.12     | 1.17    | 3.11   | 42.75 |
|              | *Exogone sp.*          | 1.72       | 0.98     | 0.63    | 2.71   | 45.46 |
|              | *Syllidae*             | 1.69       | 0.96     | 0.66    | 2.66   | 48.13 |
| Sable Island | *Ampharete finmarchica*| 2.86       | 2.14     | 1.74    | 5.71   | 5.71  |
| (37.55%)     | *Nemertea*             | 2.01       | 1.77     | 1.49    | 4.73   | 10.43 |
|              | *Lumbrinerides acuta*  | 2.13       | 1.66     | 0.78    | 4.43   | 14.86 |
|              | *Unciola irrorata*     | 2.75       | 1.60     | 1.05    | 4.27   | 19.13 |
|              | *Clymenura borealis*   | 1.94       | 1.37     | 1.25    | 3.65   | 22.79 |
|              | *Spiophanes bombyx*    | 2.07       | 1.32     | 1.16    | 3.52   | 26.30 |
|              | *Echinarchnius parma*  | 2.03       | 1.29     | 1.07    | 3.43   | 29.74 |
|              | *Ophelia limacina*     | 1.51       | 1.23     | 1.05    | 3.28   | 33.02 |
|              | *Ascidiaea (solitary)* | 1.93       | 1.22     | 0.83    | 3.24   | 36.26 |
|              | *Scoloplos armiger*    | 1.63       | 1.08     | 0.81    | 2.87   | 39.14 |
|              | *Tharyx sp.*           | 1.70       | 1.08     | 0.98    | 2.87   | 42.01 |
|              | *Clymenella zonalis*   | 2.34       | 0.98     | 0.71    | 2.60   | 44.61 |
|              | *Nereis sp.*           | 1.70       | 0.96     | 1.15    | 2.55   | 47.17 |
| Western      | *Aglaothamnus circinata* | 1.72       | 0.89     | 0.82    | 2.37   | 49.54 |
| (43.53%)     | *Unciola irrorata*     | 3.98       | 2.79     | 2.07    | 6.41   | 6.41  |
|              | *Clymenella zonalis*   | 3.55       | 2.01     | 1.57    | 4.63   | 11.04 |
|              | *Edwardsia elegans*    | 2.59       | 1.86     | 1.70    | 4.26   | 15.30 |
|              | *Ampharete finmarchica*| 2.49       | 1.74     | 1.59    | 4.00   | 19.30 |
|              | *Lumbrinerides acuta*  | 2.37       | 1.67     | 1.15    | 3.84   | 23.14 |
|              | *Nemertea*             | 2.38       | 1.50     | 1.95    | 3.44   | 26.57 |
|              | *Echinarchnius parma*  | 1.94       | 1.35     | 0.99    | 3.09   | 29.67 |
|              | *Clymenura borealis*   | 2.08       | 1.33     | 1.02    | 3.05   | 32.71 |
|              | *Oligochaeta*          | 2.33       | 1.26     | 1.15    | 2.89   | 35.60 |
|              | *Polygordius sp.*      | 2.60       | 1.20     | 0.92    | 2.77   | 38.37 |
|              | *Paraonis sp.*         | 2.05       | 1.18     | 1.60    | 2.72   | 41.09 |
|              | *Cyrtodaria siliqua*   | 1.90       | 1.18     | 1.40    | 2.71   | 43.80 |
|              | *Exogone sp.*          | 2.16       | 1.14     | 1.26    | 2.63   | 46.43 |
|              | *Tharyx sp.*           | 2.03       | 1.06     | 1.11    | 2.44   | 48.86 |

Table limited to the taxa contributing to 50% of the similarity among samples within banks.
Abbreviations: (% Sim.), total similarity among samples expressed as a percentage; Av.Abund., average abundance; Av. Sim., average contribution to the total similarity; Sim./SD, average contribution to the total similarity divided by standard deviation; Cont%, percentage contribution to similarity; Cum%, cumulative percentage contribution of contribution to similarity.

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The interaction between Haddock Abundance and Bank also explained a high proportion of the variability in the data (Table 5, Fig 3), being second only to the Bank factor (Fig 3). The nMDS plot (Fig 3) showed spatial separation of the Haddock Abundance levels on Western and Sable Island Banks, but not on Emerald Bank, although the levels of dissimilarity between levels on each Bank were all greater than 57% (Table 5). The pairwise permutations noted above identified significant differences only on Sable Island Bank. SIMPER identified relative

| Bank (%Diss.) | Taxon | Av. Abund. (1) | Av. Abund. (2) | Av. Diss. | Cont.% | Cum.% |
|---------------|-------|----------------|----------------|-----------|--------|-------|
| Emerald (1)  | Ericthonius fasciatus | 2.67 | 0.00 | 1.67 | 2.20 | 2.20 |
| Sable Island (2) | Chone sp. | 2.77 | 0.43 | 1.54 | 2.03 | 4.23 |
| Emerald (1)  | Clymenella zonalis | 1.61 | 2.34 | 1.43 | 1.88 | 6.11 |
| Western (2)  | Unciola irrorata | 2.75 | 2.75 | 1.39 | 1.83 | 7.94 |
| Emerald (1)  | Ampharete finmarchica | 1.32 | 2.86 | 1.22 | 1.61 | 9.55 |
| Western (2)  | Echinarchnus parma | 0.18 | 2.03 | 1.20 | 1.59 | 11.14 |
| Emerald (1)  | Lumbrinerides acuta | 1.54 | 2.13 | 1.19 | 1.57 | 12.71 |
| Western (2)  | Spiophanes bombyx | 0.34 | 2.07 | 1.15 | 1.52 | 14.23 |
| Emerald (1)  | Clymenura borealis | 2.39 | 1.94 | 1.15 | 1.52 | 15.75 |
| Western (2)  | Asciadiaea (solitary) | 1.01 | 1.93 | 1.12 | 1.48 | 17.23 |
| Emerald (1)  | Glyceria capitata | 1.88 | 0.36 | 1.10 | 1.45 | 18.68 |
| Western (2)  | Exogone sp. | 1.72 | 0.17 | 1.06 | 1.40 | 20.08 |
| Emerald (1)  | Clymenella zonalis | 1.61 | 3.55 | 1.58 | 2.21 | 2.21 |
| Western (2)  | Ericthonius fasciatus | 2.67 | 0.39 | 1.44 | 2.01 | 4.22 |
| Emerald (1)  | Polygordius sp. | 1.18 | 2.60 | 1.31 | 1.84 | 7.92 |
| Western (2)  | Unciola irrorata | 2.75 | 3.98 | 1.25 | 1.75 | 9.67 |
| Emerald (1)  | Edwardsia elegans | 0.70 | 2.59 | 1.20 | 1.68 | 11.35 |
| Western (2)  | Echinarchnus parma | 0.18 | 1.94 | 1.17 | 1.63 | 12.99 |
| Emerald (1)  | Oligochaeta | 0.78 | 2.33 | 1.16 | 1.63 | 14.61 |
| Western (2)  | Exogone sp. | 1.72 | 2.16 | 1.10 | 1.54 | 16.15 |
| Emerald (1)  | Glyceria capitata | 1.88 | 2.31 | 1.08 | 1.51 | 17.66 |
| Western (2)  | Cyrtodaria siliqua | 0.07 | 1.90 | 1.07 | 1.50 | 19.16 |
| Sable Island (1) | Clymenella zonalis | 2.34 | 3.55 | 1.44 | 2.22 | 2.22 |
| Western (2)  | Unciola irrorata | 2.75 | 3.98 | 1.27 | 1.95 | 4.17 |
| Sable Island (1) | Polygordius sp. | 1.64 | 2.60 | 1.16 | 1.78 | 5.95 |
| Western (2)  | Edwardsia elegans | 1.77 | 2.59 | 1.09 | 1.67 | 7.62 |
| Sable Island (1) | Glyceria capitata | 0.36 | 2.31 | 1.04 | 1.61 | 9.22 |
| Western (2)  | Oligochaeta | 1.17 | 2.33 | 1.00 | 1.54 | 10.77 |
| Sable Island (1) | Exogone sp. | 0.17 | 2.16 | 1.00 | 1.54 | 12.30 |
| Western (2)  | Protomediela fasciata | 0.55 | 2.05 | 0.89 | 1.37 | 13.67 |
| Sable Island (1) | Lumbrinerides acuta | 2.13 | 2.37 | 0.87 | 1.34 | 15.01 |
| Western (2)  | Syllidae | 0.49 | 1.97 | 0.87 | 1.33 | 16.35 |
| Sable Island (1) | Spiophanes bombyx | 2.07 | 1.51 | 0.87 | 1.33 | 17.68 |
| Western (2)  | Chaetozoon sp. A | 0.70 | 1.86 | 0.86 | 1.32 | 18.99 |
| Sable Island (1) | Arctica islandica | 1.44 | 1.16 | 0.85 | 1.30 | 20.30 |

Table limited to the taxa contributing to 20% of the dissimilarity between banks.

Abbreviations: %Diss., total dissimilarity between banks expressed as a percentage; Av.Abund., average abundance; Av. Diss., average contribution to the total dissimilarity; Cont%, percentage contribution to dissimilarity; Cum%, cumulative percentage contribution of contribution to dissimilarity.

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proportions in the abundance of taxa as contributing to the differences between High and Low Haddock Abundance on each Bank as opposed to different species present (Table 5). The list of species contributing to 10% of the total dissimilarity between Haddock Abundance levels on each Bank is provided in Table 5. On Sable Island Bank, where the differences were most pronounced (average dissimilarity = 69.29%), the ocean quahog *Arctica islandica* and the sand dollar *Echinarachnius parma*, were much more abundant in the areas where Haddock Abundance was Low, while the polychaetes *Lumbrinerides acuta* and *Clymenella zonalis* were much more abundant in the areas of High Haddock Abundance (Table 5).

**Macrofaunal Biomass and Presence/Absence.** PERMANOVA of the transformed biomass of the macrofaunal taxa showed an interaction effect between Haddock Abundance and Bank; all other interactions were non-significant (Table 6). As for transformed abundance, all three factors were significant with Bank explaining the largest proportion of the variance, followed by Haddock Abundance with Year explaining the least proportion of the variance (Table 6). The PERMANOVA of the presence/absence data showed interaction effects between Haddock Abundance and Bank, and between Bank and Year; all other interactions were non-significant and all 3 factors were significant (Table 6). As for the other analyses, Bank had the largest variance component. The nMDS configurations for these analyses (not shown) were very similar to that of those generated from abundance data both with respect to Bank separating the samples from one another and with the two levels of Haddock Abundance showing overlapping distributions.

**Macrofauna Contributing to Differences in Areas of High and Low Haddock Abundance.** Haddock Abundance showed significant differences in PERMANOVA between High and Low levels in all 3 variables: transformed abundance and biomass, and presence/absence of taxa (Tables 2 and 6). A SIMPER analysis listing those taxa accounting for 20% of the variation between the two levels, for each variable, is provided in Table 7. Areas with High Haddock Abundance were between 58% and 68% different in benthic community composition from areas with Low Haddock Abundance and many species contributed to the differentiation (Table 7). Areas with High Haddock Abundance were characterized by larger numbers of the

| Bank (%Diss.) | Taxon      | Av.Abund. (Low HA) | Av.Abund. (High HA) | Av.Diss. | Cont.% | Cum.% |
|--------------|------------|-------------------|---------------------|---------|--------|-------|
| Emerald      | *Chone* sp.| 2.97              | 2.61                | 1.71    | 2.63   | 2.63  |
| (64.94%)     | *Ericthonius fasciatus* | 2.33              | 2.93                | 1.60    | 2.47   | 5.09  |
|              | *Unciola irrorata* | 3.82              | 1.92                | 1.56    | 2.40   | 7.49  |
|              | *Syllidae* | 2.49              | 1.07                | 1.52    | 2.35   | 9.84  |
| Sable Island | *Arctica islandica* | 3.05              | 0.28                | 1.50    | 2.16   | 2.16  |
| (69.29%)     | *Lumbrinerides acuta* | 0.53              | 3.28                | 1.50    | 2.16   | 4.33  |
|              | *Echinarchnius parma* | 3.54              | 0.94                | 1.46    | 2.11   | 6.44  |
|              | *Aglaophamus cinctinata* | 3.17              | 0.67                | 1.41    | 2.04   | 8.47  |
|              | *Clymenella zonalis* | 1.98              | 2.60                | 1.36    | 1.96   | 10.43 |
| Western      | *Clymenella zonalis* | 3.10              | 4.01                | 1.25    | 2.15   | 2.15  |
| (57.96%)     | *Polygordius sp.* | 2.74              | 2.45                | 1.17    | 2.02   | 4.17  |
|              | *Glycera capitata* | 1.88              | 2.76                | 1.05    | 1.81   | 5.99  |
|              | *Unciola irrorata* | 3.29              | 4.70                | 1.05    | 1.81   | 7.80  |
|              | *Tharyx sp.* | 1.22              | 2.89                | 0.99    | 1.70   | 9.50  |

Table limited to the taxa contributing to 10% of the dissimilarity between the High and Low Haddock Abundance (HA) sites on each bank. Abbreviations: % Diss., total dissimilarity between banks expressed as a percentage; Av.Abund., average abundance; Av. Diss., average contribution to the total dissimilarity; Cont%, percentage contribution to dissimilarity; Cum%, cumulative percentage contribution of contribution to dissimilarity.

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maldanid polychaete *C. zonalis*, the elegant burrowing anemone *Edwardsia elegans*, the polychaete *Lumbrinerides acuta*, the bristle worm *Glycera capitata*, and the amphipod *Erichthonius fasciatus*. Biomass of the elegant burrowing anemone *E. elegans*, the bristle worm *G. capitata*, the polychaete *Ophelia limacina*, and *Cerianthidae* was higher in areas with High Haddock Abundance. While the bristle worm *Scoloplos armiger*, the polychaete *O. limacina*, the oval spoonclam *Periploma leanum*, Gastropoda, the polychaete *Exogone* sp., the bristle worm *Noto- mastus latericeus*, Mytiloidea, the bristle worm *Orbinia swani*, the bubble snail *Cylichna alba*, and the moonsnail *Euspira* sp. occurred more frequently in areas with Low Haddock Abundance.

**Environmental Influence on Macrofauna**

DISTLM models constructed the best combination of environmental variables that accounted for the variation seen in the data. Sediment Type was the factor explaining the highest variability in all three analyses, with respect to abundance, biomass and presence/absence of macrofauna. The rest of the variables increased the value of Adjusted $R^2$ up to 43.8–45.1% of the explained variation (Table 8).

A total of 43.9% of the total variability in macrofaunal abundance was explained by all the variables analyzed; with Sediment Type explaining 27.6% and subsequent variables 16.2%. Sediment Type also explained the highest variability in the biomass and presence/absence data (28.1%), followed by Depth, and Mean Bottom Current, Salinity and Temperature. The variables combined explained 44 and 45.1% of total variation respectively.

On the abundance dbRDA plot (Fig 4) the first two axes explained 59.2% of the fitted variation and 25.9% of the total variation. The pattern of the macrofaunal samples on the plot suggested two gradients of variation. The first gradient was driven by the variable Depth, with

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Table 6. PERMANOVA of the transformed biomass and untransformed presence/absence of macrofaunal taxa based on Bray-Curtis similarity.

| Variable          | Source                    | Degrees of Freedom | Sums of Squares (SS) | Mean Square (MS) | Pseudo-$F$ | $P_{(perm)}$ | Est. of Variance Component | Sq. Root of Variance Component |
|-------------------|---------------------------|--------------------|----------------------|------------------|------------|-------------|-----------------------------|------------------------------|
| Biomass           | Haddock Abundance (HA)    | 1                  | 7625.5               | 7625.5           | 4.631      | 0.001       | 103.28                      | 10.16                        |
|                   | Bank (BA)                 | 2                  | 49606.0              | 24803.0          | 15.064     | 0.001       | 598.29                      | 24.46                        |
|                   | Year (YR)                 | 1                  | 5862.1               | 5862.1           | 3.560      | 0.001       | 72.82                       | 8.53                         |
|                   | HAxBA                     | 2                  | 21583.0              | 10791            | 6.554      | 0.001       | 472.55                      | 21.74                        |
|                   | HAxYR                     | 1                  | 2939.9               | 2939.9           | 1.786      | 0.040       | 44.68                       | 6.68                         |
|                   | BAxYR                     | 2                  | 7182.7               | 3591.4           | 2.181      | 0.003       | 100.50                      | 10.03                        |
|                   | HAxBAxYR                  | 2                  | 3013.3               | 1506.7           | 0.915      | 0.574       | -14.46                      | -3.80                        |
| Presence/ Absence| Haddock Abundance (HA)    | 1                  | 4727.0               | 4727.0           | 3.893      | 0.001       | 60.68                       | 7.79                         |
|                   | Bank (BA)                 | 2                  | 22520.0              | 18.5             | 18.992     | 0.001       | 550.48                      | 23.46                        |
|                   | Year (YR)                 | 1                  | 7182.1               | 5.9              | 6.135      | 0.001       | 103.09                      | 10.15                        |
|                   | HAxBA                     | 2                  | 5871.0               | 4.8              | 5.027      | 0.001       | 240.64                      | 15.51                        |
|                   | HAxYR                     | 1                  | 2379.1               | 1.9              | 1.987      | 0.025       | 40.24                       | 6.34                         |
|                   | BAxYR                     | 2                  | 4170.7               | 3.4              | 3.548      | 0.001       | 152.78                      | 12.36                        |
|                   | HAxBAxYR                  | 2                  | 2467.0               | 1233.5           | 1.016      | 0.401       | 2.00                        | 1.41                         |
|                   | Residual                  | 108                | 1.31x10^5            | 1214.2           |            |             | 1214.20                     | 34.85                        |
|                   | Total                     | 119                | 2.79x10^5            |                  |            |             |                             |                              |

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### Table 7. SIMPER analysis of transformed abundance, transformed biomass and presence/absence of macrofauna between High and Low Haddock Abundance sites across Banks.

| Variable (%Diss.) | Taxon               | Low HA Aver. | High HA Aver. | Av.Diss. | Cont.% | Cum.% |
|-------------------|---------------------|--------------|---------------|----------|--------|-------|
| **Abundance**     | Clymenella zonali s | 2.41         | 2.64          | 1.42     | 2.09   | 2.09  |
|                   | (68.14%)            |              |               |          |        |       |
|                   | Unciola irrorata    | 3.32         | 3.08          | 1.26     | 1.85   | 3.94  |
|                   | Polygordius sp.     | 1.86         | 1.81          | 1.15     | 1.68   | 5.62  |
|                   | Chone sp.           | 1.62         | 1.62          | 1.14     | 1.67   | 7.29  |
|                   | Echinarchnus parma  | 1.99         | 0.90          | 1.02     | 1.50   | 8.79  |
|                   | Edwardsia elegans   | 1.67         | 1.76          | 1.01     | 1.48   | 10.27 |
|                   | Lumbrinerides acuta | 1.71         | 2.29          | 0.99     | 1.45   | 11.72 |
|                   | Exogone sp.         | 1.59         | 1.22          | 0.99     | 1.45   | 13.17 |
|                   | Glyceria capitata   | 1.47         | 1.62          | 0.97     | 1.42   | 14.59 |
|                   | Syllidae            | 1.69         | 1.18          | 0.96     | 1.41   | 16.00 |
|                   | Clymenura borealis  | 2.31         | 1.99          | 0.96     | 1.41   | 17.41 |
|                   | Oligochaeta         | 1.54         | 1.39          | 0.95     | 1.40   | 18.81 |
|                   | Erichthonius fasciatus | 0.72      | 1.25          | 0.91     | 1.34   | 20.15 |
| **Biomass**       | Echinarchnus parma  | 5.86         | 2.56          | 1.74     | 2.55   | 2.55  |
|                   | (68.08%)            |              |               |          |        |       |
|                   | Cyrtodaria siliqua  | 4.88         | 2.92          | 1.60     | 2.34   | 4.89  |
|                   | Arctica islandica   | 4.49         | 2.88          | 1.46     | 2.14   | 7.04  |
|                   | Clymenura borealis  | 6.16         | 4.86          | 1.21     | 1.78   | 8.82  |
|                   | Clymenella zonali s | 4.06         | 3.91          | 1.19     | 1.74   | 10.56 |
|                   | Nephys caeca        | 3.41         | 2.91          | 1.17     | 1.73   | 12.29 |
|                   | Edwardsia elegans   | 3.69         | 3.90          | 1.12     | 1.64   | 13.92 |
|                   | Glyceria capitata   | 2.85         | 3.42          | 1.03     | 1.51   | 15.43 |
|                   | Aglaophamus cinctata| 4.11         | 3.19          | 1.03     | 1.51   | 16.94 |
|                   | Ophelia limacina    | 2.29         | 2.89          | 1.02     | 1.49   | 18.43 |
|                   | Cerianthidae        | 1.75         | 3.10          | 1.01     | 1.48   | 19.91 |
| **Occurrence**    | Chone sp.           | 0.47         | 0.33          | 0.54     | 0.92   | 0.92  |
|                   | Cyrtodaria siliqua  | 0.40         | 0.29          | 0.53     | 0.90   | 1.83  |
|                   | Scoloplos armiger   | 0.28         | 0.38          | 0.53     | 0.90   | 2.73  |
|                   | Polycrius sp.       | 0.47         | 0.35          | 0.53     | 0.90   | 3.63  |
|                   | Ophelia limacina    | 0.35         | 0.44          | 0.52     | 0.89   | 4.52  |
|                   | Chaetozone sp. A    | 0.37         | 0.35          | 0.52     | 0.88   | 5.40  |
|                   | Euchone papillosa   | 0.42         | 0.37          | 0.52     | 0.88   | 6.28  |
|                   | Aricidea catherinae | 0.39         | 0.37          | 0.52     | 0.88   | 7.16  |
|                   | Gastropoda          | 0.24         | 0.35          | 0.51     | 0.88   | 8.04  |
|                   | Hippomedon serratus | 0.30         | 0.31          | 0.51     | 0.88   | 8.92  |
|                   | Periploma leanum    | 0.29         | 0.32          | 0.51     | 0.88   | 9.79  |
|                   | Cistenides granulata| 0.37         | 0.35          | 0.51     | 0.88   | 10.67 |
|                   | Nephys caeca        | 0.34         | 0.30          | 0.51     | 0.87   | 11.54 |
|                   | Exogone sp.         | 0.39         | 0.41          | 0.51     | 0.87   | 12.42 |
|                   | Arctica islandica   | 0.42         | 0.38          | 0.51     | 0.87   | 13.29 |
|                   | Notomastus latericeus| 0.37       | 0.42          | 0.51     | 0.87   | 14.16 |
|                   | Mytiloidea          | 0.28         | 0.36          | 0.51     | 0.87   | 15.03 |
|                   | Syllidae            | 0.47         | 0.39          | 0.51     | 0.87   | 15.90 |
|                   | Oligochaeta         | 0.43         | 0.39          | 0.51     | 0.87   | 16.76 |
|                   | Orbinia swani       | 0.38         | 0.47          | 0.51     | 0.86   | 17.63 |
|                   | Clylichna alba      | 0.28         | 0.32          | 0.50     | 0.86   | 18.48 |

(Continued)
deeper samples on the lower right quadrant (Emerald Bank) and shallowest on the upper left and explaining 13.6% of the total variation. There was a clear separation between Emerald Bank samples and the rest, while Sable and Western Bank samples were not strongly differentiated. dbRDA ordination of biomass and presence/absence samples showed similar patterns, with samples plotted according to different sediment types and depth. The second gradient was driven by the variable Sediment Type, distinguishing samples of Sand-group bottom types in the lower left quadrant and gravel-types in the upper right and explaining 12.3% of the total variation. Samples of Sable Bank Low Haddock Abundance were displayed forming a tight group in the lower left quadrant, indicating a high proportion of sandy sediments.

Discussion
Link between Benthos and Haddock Spatial Structure
Haddock are an ecologically and commercially important fish of the continental shelves of the North Atlantic, with strong links to the benthos [47–50]. Physical parameters such as temperature and depth play a key role in the large-scale spatial structuring of haddock distribution

Table 7. (Continued)

| Variable (%Diss.) | Taxon                   | Low HA | High HA | Av.Diss. | Cont.% | Cum.% |
|------------------|-------------------------|--------|---------|----------|--------|-------|
|                  | Euspira sp.             | 0.26   | 0.30    | 0.50     | 0.85   | 19.34 |
|                  | Spio filicornis         | 0.53   | 0.39    | 0.50     | 0.85   | 20.19 |

Table limited to the taxa contributing to 20% of the dissimilarity between levels of Haddock Abundance for each variable. Abbreviations: %Diss., total dissimilarity between Haddock Abundance levels expressed as a percentage; HA, Haddock Abundance; Aver., average; Av. Diss., average contribution to the total dissimilarity; Cont%, percentage contribution to dissimilarity; Cum%, cumulative percentage contribution of contribution to dissimilarity.

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Table 8. Distance-based linear model (DistLM) of Bray-Curtis similarities between samples in abundance, biomass and presence/absence of macrofauna against environmental variables.

| Variable | Environmental Variable | Adj. $R^2$ | Pseudo-$F$ | $P$ | Prop. | Cum. |
|----------|------------------------|------------|------------|-----|-------|------|
| Abundance| Sediment Type          | 0.194      | 3.378      | 0.0001 | 0.276 | 0.276 |
|          | Mean Bottom Temperature | 0.271      | 12.117     | 0.0001 | 0.074 | 0.351 |
|          | Mean Bottom Current    | 0.303      | 5.940      | 0.0001 | 0.035 | 0.386 |
|          | Mean Bottom Salinity   | 0.341      | 6.942      | 0.0001 | 0.038 | 0.425 |
|          | Depth                  | 0.350      | 2.453      | 0.0008 | 0.013 | 0.439 |
| Biomass  | Sediment Type          | 0.200      | 3.465      | 0.0001 | 0.281 | 0.281 |
|          | Depth                  | 0.258      | 9.261      | 0.0001 | 0.058 | 0.340 |
|          | Mean Bottom Current    | 0.303      | 7.805      | 0.0001 | 0.046 | 0.386 |
|          | Mean Bottom Salinity   | 0.347      | 8.085      | 0.0001 | 0.044 | 0.430 |
|          | Mean Bottom Temperature| 0.352      | 1.713      | 0.0240 | 0.094 | 0.440 |
| Presence/| Sediment Type          | 0.200      | 3.461      | 0.0001 | 0.281 | 0.281 |
| Absence  | Depth                  | 0.284      | 13.465     | 0.0001 | 0.081 | 0.363 |
|          | Mean Bottom Current    | 0.323      | 7.108      | 0.0001 | 0.040 | 0.403 |
|          | Mean Bottom Salinity   | 0.361      | 7.071      | 0.0001 | 0.038 | 0.442 |
|          | Mean Bottom Temperature| 0.365      | 1.768      | 0.0310 | 0.009 | 0.451 |

Abbreviations: Adj. $R^2$, adjusted $R^2$; Prop., proportion of variance explained by each variable; Cum., cumulative proportion of variance explained by multiple variables.

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while small-scale spatial structure of haddock populations is linked to life processes such as growth and survival determined in part by predator avoidance, prey availability and protection by camouflage and cover [21].

Our study embedded knowledge of the long-term (32 years), persistent distribution of juvenile haddock abundance in its experimental design, in the form of a factor with two levels, High and Low, characterized by long-term catch rates differing by an order of magnitude [38]. We were interested in determining whether the benthic species composition of the macrofauna could be a factor in determining preferred juvenile habitat for this species. As a factor explaining patterns in benthic macrofaunal communities it could be interpreted in three ways. Significant effects could be due to habitat selection by the juvenile haddock actively choosing an area based on benthic species composition; to habitat modification produced by the fish selectively feeding on the benthos over a long period of time in a given area; and/or to fish and benthos responding separately to the same environmental drivers. Juvenile Haddock Abundance was a statistically significant factor in our multivariate analyses of macrofaunal community composition as assessed using data on abundance, biomass and species occurrence. Therefore one or more of these dynamics between fish and benthos could be operating to induce this effect.

Descriptively the benthic communities at High and Low sites, within Banks, were between 58% and 70% dissimilar. However, the biological relevance of those differences was not conspicuous and samples did not show clear separation in the MDS ordinations. Statistically significant pairwise tests between areas of High and Low juvenile Haddock Abundance were only identified on Sable Island Bank, creating a significant interaction effect that explained a large portion of the variability in the data. The statistical difference in the benthic communities based on abundance, biomass and species occurrence between the sites on Sable Island Bank was clearly explained by differences in the sediment types (abundance: Fig 2 and Fig 4). The Low Haddock Abundance samples were characterized entirely by fine sandy sediment bottoms with large and continuous areas of low bathymetric relief (Fig 2), whereas the High Haddock Abundance samples showed more diverse, rugged and spatially heterogeneous bottoms with both coarse gravel and sand sediments [21]. As a result, the Sable Island Bank benthic communities in the area with Low Haddock Abundance were characterized by infaunal species associated with mud or sand bottoms.

As observed in previous studies, physical habitat can have a strong impact on juvenile haddock spatial distribution due to its essential role in predator avoidance which is the primary source of mortality for demersal fish [21,51]. We note that species such as the sand dollar *Echinarchnium parma*, the bivalve *Arctica islandica*, and the clam *Cyrtodaria siliqua* are found in fine sands [52–54] and their association with the Low Haddock Abundance site on Sable Island Bank may reflect an avoidance of fine sands by the juvenile fish where they would have little camouflage. Abiotic and biotic habitat parameters including sediment composition and presence of emergent fauna or topography, mediate avoidance of predators [51,55–57]. Thus, structurally complex benthic habitats such as those containing coarse sediments, small-scale topographical variability, patched sediment distribution or emergent epibenthic fauna are stated to enhance predator avoidance and therefore increase juvenile fish survivorship [16,51,55,58–60]. These sorts of habitats were more widespread within High Haddock Abundance areas (Fig 2). Haddock are known to be associated with gravel bottoms [10,11,18,23,57,58], including in the study areas [60] and could take advantage of the higher availability of these types of bottoms within preferred areas to camouflage by mimicking the coloration and texture of the uneven seafloors to block visual recognition by predators. Similar behaviors have previously been documented on Georges Bank by Lough et al. [58]. Also, the higher number of crevices among coarser sediments might serve as refuge providing spaces inaccessible to larger-bodied predators [18,51] which has been observed in another demersal
fish species, Atlantic cod *Gadus morhua*, by Gotceitas and Brown [61]. They found that in the presence of predators, juvenile cod changed their substrate preference from sand or gravel-pebble to cobble. Finally, these hard substrates support diverse communities of sessile taxa such as Porifera, Hydrozoa or Bivalvia [62,63] that can provide microhabitat biogenic structure and resources [60] that are also used as shelter for juvenile fish [16,64]. Therefore, the extreme difference in sediment type and in the associated communities between preferred and non-preferred sites on Sable Island Bank may indicate that sediment type, rather than the benthic species composition that occupy it, is the key determinant of juvenile haddock abundance in that area, with both fish and benthos responding to it in different ways. Further, as the same trend was not seen on the other banks which also differed in the proportions of sand and gravel between areas of high and low haddock abundance, this relationship is likely non-linear, with some minimal patch size involved in rendering the area non-preferred by the fish. However, we

**Fig 4.** Distance-based RDA ordination of macrofauna based on transformed abundance data as predicted by a DISTLM model. The environmental variables that best explain the variation in infaunal abundance on the three banks are shown. Vectors indicate direction of the effect of quantitative variables in the ordination plot. The variable Sediment Type is illustrated using the end point of the vector (*) for each of the 14 categories.

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were unable to test that hypothesis with these data, not having data on fish abundance at the same fine spatial scale as the benthos. Integrating such data on haddock abundance in future, obtained through video observations, could help determine the relative importance of the relationship between juvenile haddock size and abundance and macrobenthic communities.

It was expected that our results would show a positive association between macrofaunal density and juvenile haddock abundance, as benthic prey density, a subset of the macrofauna, influences the opportunistic diet of haddock [12,13,20,48–50]. The main prey of haddock are small and slow moving benthic infauna [50, 65–67] typically including crustaceans, polychaetes, molluscs, and echinoderms [12, 23,26,50,67,68] ranking in different order according to haddock age [49], prey availability, location and year [48]. On the Scotian Shelf, several studies have analyzed the diet of haddock in the study area [12,13,65,66] although only Kohler and Fitzgerald [66] and Mahon and Neilson [12] focused on juveniles. Both of those studies ranked crustaceans, mainly amphipods and decapods, as the major prey group, followed by echinoderms. In our study we found that at the species level the link between juvenile haddock abundance and benthic prey distribution was not consistent. Some known prey [12,66], such as the amphipod *Unciola irrorata* or the shrimp *Crangon septempsinosa* among others, were more abundant within High Haddock Abundance areas as expected, while others such as the amphipod *Leptocheirus pinguins* or the echinoid *E. parma* were more abundant within Low Haddock Abundance areas. Thus, it was not possible to conclude that juvenile haddock were actively selecting the preferred areas according to benthic species composition.

**New Insights into Benthic Macrofaunal Communities on the Eastern Scotian Shelf**

Our study is the first to describe the benthic macrofaunal communities over broad as well as smaller spatial scales on the eastern Scotian Shelf. The regional benthic fauna off eastern Canada have previously been described using various sampling gears [32,33,69,70–73] although none of those studies were focused on the eastern Scotian Shelf. Comparable studies to ours were Davis and Gilhen [74], Davis [75], Gilkinson et al. [31], Henry et al. [30], and Kenchington et al. [30], but these were limited to small-spatial scales, latterly collected to examine the impacts of experimental bottom trawling on benthic communities.

Emerald, Western and Sable Island Banks are located in the outer part of the Scotian Shelf and this study has shown them to contain diverse benthic communities. The overall level of community dissimilarity in relative abundance of macrofaunal species between banks was high (Emerald Bank vs Sable Island Bank, 75.8%; Emerald Bank vs. Western Bank 71.4%; Western Bank vs. Sable Island Bank, 65.1%) and was based on differences in a large number of species, and paralleled in biomass and species occurrence. This large scale-variation was expressed as a clear differentiation between Emerald Bank macrofaunal communities and those in Sable Island and Western Banks in the nMDS ordinations, driven primarily by differences in depth, and to a lesser degree by bottom salinity and temperature, as indicated in the dbRDA ordinations. The macrofaunal communities on Sable Island and Western Banks were also distinctive from each other with respect to abundance, biomass and occurrence; however they presented themselves as a continuum rather than as a discontinuity in two dimensional space, as visualized by nMDS, and driven by Sediment Type (Fig 4, Table 8). Overall, Sediment Type was the dominant variable explaining the largest percentage of variation in the macrofaunal community data. Although statistically significant differences were found between the two years of study, there was no temporal change in this dominant among-bank pattern.

Average depth decreases from the southwest (Emerald Bank) to northeast (Sable Island Bank), with the samples from Emerald Bank being at about 80 m, or twice as deep as those
from Western Bank at about 43 m and from Sable Island Bank at about 38 m. Bottom temperature and salinity decrease in the same direction, with Emerald Bank being not only deeper, but warmer and saltier. The number of taxa observed and total biomass were both significantly lower on Emerald Bank due in part to the relative paucity of echinoderms and molluscs which were prevalent on Western and Sable Island Banks.

Macrofaunal communities from Western and Sable Island Banks, although significantly different, were more similar to one another than either was to Emerald Bank, which is in concordance with the findings of Courtney et al. [42] who stated that from a geological perspective, Western Bank was a continuum of Sable Island Bank, rather than a separate bank. This distinction of the Emerald Bank macrofauna was not previously known. Seabed topography and substrate type are known to be key structuring factors of benthic assemblages controlling the presence or absence of several sediment-dependent species [62,70,76–80] and they explained the differentiation among the Western and Sable Island Bank samples, as seen in the dbRDA ordination. Emerald Bank seabed showed a lesser variety in sediment types and benthic habitats in comparison to Western Bank seabed that hosted a larger variety of sediments and habitats, which is associated with higher number of organisms and high diversity [36,70,81] especially sessile epifauna [62]. Hence, a more species-rich and abundant community of diverse phyla was found there, which is in concordance with previous studies [29].

This strong statistical signal in the macrobenthos, separating Emerald Bank from the other banks does not correspond with the contemporary abundance of haddock which were generally similar across banks but lower on Sable Island Bank [38].

Conclusions

We were not able to detect compelling evidence that the temporally-persistent distribution of juvenile haddock density was spatially correlated with differences in benthic macrofaunal communities on the eastern Scotian Shelf. Only on Sable Island Bank were distinct macrofaunal assemblages associated with areas of preferred and non-preferred juvenile haddock habitat. There, both juvenile fish and benthos may be independently responding to the same environmental driver, namely sediment type—the area of non-preferred habitat being 100% sand, while approximately 22% of the habitat in the preferred area was gravel. On the other banks, the sediment types were much more similar in the preferred and non-preferred areas, which lacked statistically significant difference in associated macrofaunal communities. We hypothesize that selection of preferred habitats in the studied banks if present, occurs over fine spatial scales of less than 1 km and may be related to the availability of complex boundaries between gravel and sand areas and their greater topographic relief [82] that allows juvenile haddock to balance predator avoidance on gravel habitats with increased prey abundance/biomass. A future comparative study including stomach contents of juvenile haddock in the area would give insight into the role of benthic prey species as structuring drivers of haddock spatial distribution and abundance [83].

Supporting Information

S1 Fig. Species accumulation curves for each study area and year. The number of observations (Sobs) was used as the estimator and the bars represent standard deviations derived from 999 permutations of the data. A) Curves constructed by Bank (E = Emerald; S = Sable Island; W = Western) and Year (3 = 2003; 5 = 2005); B) Curves constructed with 2005 data only, by Bank and Haddock Abundance Level (High, Low); C) Curves constructed with 2003 data only, by Bank and Haddock Abundance Level (High, Low).
S2 Fig. Mean and standard deviation of Total Number of Taxa (S), Pielou’s Evenness Index (J’), Shannon-Wiener Diversity Index (H’), and Total Abundance by Bank, Juvenile Haddock Abundance, and Year.

TIFF

S1 Table. Location and associated environmental data for each video-grab sample.

DOCX

S2 Table. Abundance and biomass of macrofaunal taxa by video-grab sample.

DOCX

S3 Table. Summary of ANOVA of transformed Total Biomass (g m⁻²) testing for effect of Year, Bank and Haddock Abundance.

DOCX

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Author Contributions

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Supervision: ELK.
Validation: ELK BR.
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References

1. May RM. Unanswered questions in ecology. Philos T Roy Soc B. 1999; 354: 1951–1959.
2. Levin SA. The problem of pattern and scale in ecology. Ecology. 1992; 73 (6): 1943–1967.
3. Sissenwine MP. Why do fish populations vary? In: May RM editor. Exploitation of Marine Communities. Dahlem Konferenzen 1984. Berlin: Springer-Verlag; 1984. p. 59–94.
4. Houde ED. Emerging from Hjort’s shadow. J Northw Atl Fish Sci. 2008; 41: 53–70.
5. Cadrin SX, Secor DH. Accounting for spatial population structure in stock assessment: past, present and future. In: Beamish J and Rothschild BJ, editors. The Future of Fishery Science in North America. Dordrecht: Springer; 2009. p. 405–425.
6. Kerr LA, Cadrin SX, Secor DH. The role of spatial dynamics in the stability, resilience, and productivity of an estuarine fish population. Ecol Appl. 2010; 20:497–507. PMID: 20405802
7. Kerr LA, Cadrin SX, Secor DH. Simulation modelling as a tool for examining the consequences of spatial structure and connectivity on local and regional population dynamics. ICES J Mar Sci. 2010; 67: 1631–1639.
8. Christensen V, Coll M, Piroddi Ch, Steenbeek J, Buszowski J, Pauly D. A century of fish biomass decline in the ocean. Theme Section ‘Trophicdynamics in marine ecology’. Mar Ecol Prog Ser. 2014; 512: 155–166.
9. Young E, Belchier M, Hauser L, Horsburgh GJ, Meredith MP, Murphy EJ, et al. Oceanography and life history predict contrasting genetic population structure in two Antarctic fish species. Evol Appl. 2015; 8(5):486–509. doi: 10.1111/eva.12259 PMID: 26029262
10. Linehan J. Annotated bibliography of habitat associations of juvenile demersal fish in offshore shelf waters from Newfoundland, the Gulf of St. Lawrence, and the Maritimes region. Can Tech Rep Fish Aquat Sci; 2004. Report No.: 2526.
11. Gregory R, Simon J, Linehan J, Hurley P. Habitat associations and stock status of haddock and Atlantic cod on the eastern Scotian Shelf. In: Anderson JT and Gordon DC Jr. editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci; 2007. Report No.: 2770. p. 10–13.
12. Mahon R, Neilson JD. Diet changes in Scotian Shelf haddock during the pelagic and demersal phases of the first year of life. Mar Ecol Prog Ser. 1987; 37: 123–130.
13. Kenchington E, Gordon DC Jr, Bourbonnais C, Maclsaac K, Glikinson KD, McKeown DL, et al. Effects of experimental otter trawling on the gravel bottom ecosystem of Western Bank, N.S.: Feeding by demersal fish and analysis of trawl catch. Proceedings of the Symposium on Effects of Fishing Activities on Benthic Habitats: Linking Geology, Biology, Socioeconomics and Management. Am Fish S S. 2004; 41: 391–409.
14. Lindholm JB, Auster PJ, Kaufman LS. Habitat-mediated survivorship of juvenile (0-year) Atlantic cod Gadus morhua. Mar Ecol Prog Ser. 1999; 180: 247–255.
15. Kenchington E, Power D, Koen-Alonso M. Associations of demersal fish with sponge grounds on the continental slopes of the northwest Atlantic. Mar Ecol Prog Ser. 2013; 477: 217–230.
16. Kutti T, Fossā HJ, Bergstad OA. Influence of structurally complex benthic habitats on fish distribution. Mar Ecol Prog Ser. 2015; 520:175–190.
17. Uiblein F, Lorance P, Latrouite D. Behaviour and habitat utilisation of seven demersal fish species on the Bay of Biscay continental slope, NE Atlantic. Mar Ecol Prog Ser. 2003; 257:223–232.
18. Auster PJ, Lindholm J. The ecology of fishes on deep boulder reefs in the western Gulf of Maine (NW Atlantic). In: Godfrey JM, Shumway SE, editors. Diving for science. Proc Amer Acad Underw Sci. 24th Ann Symp. Connecticut Sea Grant, Groton, CT: 2005. p. 89–107.
19. Cuff A, Anderson JT, Devillers R. Comparing surficial sediments maps interpreted by experts with dual-frequency acoustic backscatter on the Scotian shelf, Canada. Cont Shelf Res. 2015; 110: 149–161.
20. Sell AF, Kröncke I. Correlations between benthic habitats and demersal fish assemblages—A case study on the Dogger Bank (North Sea). J Sea Res. 2013; 80: 12–24.
21. Anderson JT, Gordon DC, Jr. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci; 2007. Report No.:2770.
22. Mahon R, Smith RW. Demersal fish assemblages on the Scotian Shelf, northwest Atlantic: Spatial distribution and persistence. Can J Fish Aquat Sci. 1989; 46(S1): 134–152.
23. Cargnelli LM, Griesbach SJ, Berrien PL, Morse WW, Johnson DL. Essential fish habitat document: Haddock, Melanogrammus aeglefinus, life history and habitat characteristics. 1999. NOAA Technical Memorandum No.: NMFS-NE-128.
24. Miller DJ, Colton RR, Marak RR. A study of the vertical distribution of larval haddock. J Cons Int Explor Mer. 1963; 28: 37–49.
25. Koeller P. A., Hurley P., Perley P., Neilson J.D. Juvenile fish surveys on the Scotian Shelf: implications for year-class size assessments. J Cons Int Explor Mer. 1986; 43: 59–76.
26. Bowman RE. Feeding habits of then northwest Atlantic juvenile groundfish. U.S. Department of Commerce. National Oceanic and Atmospheric Administration, Northeast Marine Fisheries Centre. Woods Hole, Massachusetts 02543. No.: 79–43, 1979.
27. Frank KT, Shackell NL, Simon JE. An evaluation of the Emerald/Western Bank juvenile haddock closed area. ICES J Mar Sci. 2000; 57:1023–1034.
28. O’Boyle R. Benefits of marine protected areas and fisheries closures in the Northwest Atlantic. Can Tech Rep Fish Aquat Sci; 2001. Report No.: 2948.

29. Kenchington ELR, Gilkinson KD, MacIsaac KG, Bourbonnais-Boyce C, Kenchington TJ, Smith SJ, et al. Effects of experimental otter trawling on benthic assemblages on Western Bank, northwest Atlantic Ocean. J Sea Res. 2006; 56: 249–270.

30. Henry L-A, Kenchington E, MacIsaac K, Bourbonnais-Boyce C, Gordon D. Impacts of otter trawling on colonial epifaunal assemblages on a cobble bottom ecosystem on Western Bank (northwest Atlantic). Mar Ecol-Prog Ser. 2006; 6: 63–78.

31. Gilkinson KD, Gordon DC Jr, MacIsaac KG, McKeown DL, Kenchington ELR, Bourbonnais C, et al. Immediate impacts and recovery trajectories of macrofaunal communities following hydraulic clam dredging on Banquereau, eastern Canada. ICES J Mar Sci. 2005; 62 (5): 925–947.

32. Nesis KNN. Biocoenoses and biomass of benthos of the Newfoundland-Labrador region. Fisheries Research Board of Canada Translation Series No. 1375. Translated by the Translation Bureau Foreign Languages Division, Department of the Secretary of State of Canada. 1970.

33. Theroux, Roger B, Roland, Wigley L. Quantitative composition and distribution of the macrobenthic invertebrate fauna of the continental shelf ecosystems of the northeastern US. US Dep Commer; 1988. NOAA Tech. Rep. No.: NMFS 140.

34. Fader GBJ. Seabed sediment distributions, morphology, dynamics and features of detailed study areas on Emerald, Western and Sable Island Banks, outer Scotian Shelf. In: Anderson JT and Gordon DC Jr. editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci: 2007. Report No.: 2770. p. 29–31.

35. Shaw J, Todd BJ, Li MZ, Mosher DC, Kostylev VE. Continental Shelves of Atlantic Canada. Chapter 2. In: Chiocci FL, Chivas AR, editors. Continental Shelves of the World: Their Evolution During the Last Glacio-Eustatic Cycle. Geological Society, London, Memoirs; 2014. p. 7–20.

36. Breeze H, Fenton DG, Rutherford RJ, Silva MA. The Scotian Shelf: An Ecological Overview for Ocean Planning. 2002. Can. Tech. Rep. Fish. Rep. Aquat. Sci.; 2393. x + 259 p.

37. Frank KT, Petrie B, Fisher JAD, Leggett WC. Transient dynamics of an altered large marine ecosystem. Nature. 2011; 477: 86–89. doi: 10.1038/nature10285 PMID: 21796120

38. Dalley EL, Anderson JT, Davis DJ. Fish communities within the Scotian Shelf Habitat Study area: Observations from trawling. In: Anderson JT and Gordon DC Jr, editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci; 2007. Report No.: 2770. p. 18–23.

39. Rowell TW, Schwinghamer P, Gilkinson KD, Gordon DC Jr, Hartgers E, Hawryluk M, et al. Grand Banks otter trawling impact experiment: II. Experimental design and sampling equipment. Can Tech Rep Fish Aquat Sci; 1997. Report No.: 2190.

40. Gordon DC Jr, Kenchington ELR, Gilkinson KD, McKeown DL, Steeves G, Chin-Yee M, et al. Canadian imaging and sampling technology for studying marine benthic habitat and biological communities. ICES 2000 Annual Conference; 2000 Sep 27–30; Bruges, Belgium. ICES Paper CM 2000/T:07.

41. McKeown DL, Gordon DC Jr. Grand Banks otter trawling impact experiment: III. Experimental design and sampling equipment. Can Tech Rep Fish Aquat Sci; 1997. Report No.: 2190.

42. Courtney R, Anderson JT, Lang C, Fader G. Comparative seabed classification using sidescan and normal incidence sonar data at selected study sites on the Scotian shelf, Canada. Proceedings of the International Conference “Underwater Acoustic Measurements: Technologies & Results”; 2005 Jun 28–Jul 1; Heraklion, Crete, Greece. 2005.

43. Clarke KR, Gorley RN. PRIMER v6: User Manual/Tutorial. PRIMER-E, Plymouth; 2006.

44. Legendre P, Legendre L. Numerical ecology. 2nd English ed. Amsterdam: Elsevier; 1998.

45. Anderson MJ. A new method for nonparametric multivariate analysis of variance. Austral Ecol. 2001; 26: 32–46.

46. Anderson MJ. PERMANOVA: a FORTRAN computer program for permutation multivariate analysis of variance. Department of Statistics, University of Auckland, New Zealand; 2005.

47. Kenchington ELR, Gordon DC Jr, Bourbonnais-Boyce C, MacIsaac KG, Gilkinson KD, McKeown DL, et al. Effects of experimental otter trawling on the feeding of demersal fish on Western Bank, Nova Scotia. Am Fish S S. 2005; 41: 391–409.

48. Langton RW, Bowman RE. Food of fifteen northwest Atlantic Gadiform fishes. U.S. Department of Commerce; 1980. NOAA Tech Rep No.: NMFS SRF-740.

49. Methven DA. Annotated bibliography of demersal fish feeding with emphasis on Selected studies from the Scotian Shelf and Grand Banks of the Northwestern Atlantic. 1999. Can Tech Rep Fish Aquat Sci; 1999. Report No.: 2267.
50. Schückel S, Ehrich S, Kröncke I, Reiss H. Linking prey composition of haddock *Melanogrammus aeglefinus* to benthic prey availability in three different areas of the North Sea. J Fish Biol. 2010; 77, 98–118. doi: 10.1111/j.1095-8649.2010.02657.x PMID: 20646141

51. Scharf FS, Manderson JP, Fabrizio MC. The effects of seafloor habitat complexity on survival of juvenile fishes: Species-specific interactions with structural refuge. J Exp Mar Bio Ecol. 2006; 335: 167–176.

52. Stanley DJ, James NP. Distribution of *Echinarchus parma* (Lamarck) and associated fauna on Sable Island Bank, Southeast Canada. Smithsonian contribution to the Earth Sciences, No.: 6. Washington: Smithsonian Institution press; 1971.

53. Cargnelli LM, Griesbach SJ, Packer DB, Weissberger E. Ocean quahog, Arctica islandica, life history and habitat characteristics; 1999. NOAA Tech. Memo No.: NMFS-NE-148.

54. Kilada R, Campana SE, Roddick D. Growth and sexual maturity of the northern propeller clam (*Cyrtodaria siliqua*) in Eastern Canada, with bomb radiocarbon age validation. Mar Biol. 2009; 156:1029–1037.

55. Diaz RJ, Cutter GR, Able KW. The importance of physical and biogenic structure to juvenile fishes on the shallow inner continental shelf. Estuar Coast. 2003: 26: 12–20.

56. Auster PJ, Malatesta RJ, Donaldson CLS. Distributional responses to small-scale habitat variability by early juvenile silver hake, *Merluccius bilinearis*. Environ Biol Fishes. 1997; 50: 195–200.

57. Auster PJ, Joy K, Valentine PC. Fish species and community distributions as proxies for seafloor habitat distributions: the Stellwagen Bank National Marine Sanctuary example (Northwest Atlantic, Gulf of Maine). Environ Biol Fishes. 2001; 60: 331–346.

58. Lough RG, Valentine PC, Potter DC, Auditore PJ, Bolz GR, Neilson JD, et al. Ecology and distribution of juvenile cod and haddock in relation to sediment type and bottom currents on eastern Georges Bank. Mar Ecol Prog Ser. 1989; 56: 1–12.

59. Auster PJ, Lindholm J, Valentine P C. Variation in Habitat use by Juvenile Acadian Redfish, Sebastes fasciatus. Environ Biol Fishes. 2003; 68: 381–389.

60. Ollerhead LMK, Anderson JT. Habitat suitability criteria for juvenile Haddock and Atlantic Cod on the Scotian Shelf. In: Anderson JT and Gordon DC Jr, editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci; 2007. Report No.: 2770. p. 49–52.

61. Gotceitas V, Brown JA. Substrate selection by juvenile Atlantic cod (*Gadus morhua*): effects of predation risk. Oecologia. 1993; 93: 31–37.

62. Kostylev V, Todd B, Fader G, Courtney R, Cameron G, Pickrell R. Benthic habitat mapping on the Scotian Shelf based on multibeam bathymetry, surficial geology and sea floor photographs. Mar Ecol Prog Ser. 2001; 219: 121–137.

63. Gilkinson KD. Epifaunal communities on Western Bank. In: Anderson JT and Gordon DC Jr, editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci; 2007. Report No.: 2770. p. 46–48.

64. Langton R, Auster PJ, Schneider DC. A spatial and temporal perspective on research and management of groundfish in the northwest Atlantic. Rev Fish Sci. 1995; 3:201–229.

65. Homans RES, Needler AWH. Food of the haddock. Proc N S Inst Sci.1944; 21:1–35.

66. Kohler AC, Fitzgerald DN. Comparisons of Food of Cod and Haddock in the Gulf of St. Lawrence and on the Nova Scotia Banks. Can J Fish Aquat Sci. 1969; 26 (5): 1273–1287.

67. Mattson S. Food and feeding habits of fish species over a soft sublittoral bottom in the northeast Atlantic: 3. Haddock (*Melanogrammus aeglefinus* (L.)) (Gadidae). Sarsia. 1992; 77, 33–45.

68. Bowman RE, Michaels WL. Food of seventeen species of Northwest Atlantic fish. U.S. Department of Commerce; 1984. NOAA Tech Memo No.: NMFS-F/NEC-28.

69. Mills EL, Fournier RO. Fish Production and the Marine Ecosystems of the Scotian Shelf, Eastern Canada. Mar Biol. 1979; 54: 101–108.

70. Desrosiers G, Savenko C, Olivier M, Stora G, Juniper K, Caron A, et al. Trophic structure of macrobenthos in the Gulf of St. Lawrence and on the Scotian Shelf. Deep Sea Res Part 2. 2000; 47: 663–697.

71. Barrio Frojaño CR, Maclsaac KG, McMillan AK, Sacau Cuadrado M., Large PA, Kenny AJ, et al. An evaluation of benthic community structure in and around the Sackville Spur closed area (Northwest Atlantic) in relation to the protection of vulnerable marine ecosystems. ICES J Mar Sci. 2012; 69(2): 213–222.

72. Gilkinson KD. Recent DFO (Newfoundland & Labrador Region) studies of the Grand Banks benthos at small and large spatial scales. DFO Can Sci Advis Sec Res Doc; 2012. Report No.: 2012/114. 34 p.
73. Murillo FJ, Serrano A, Kenchington E, Mora J. Epibenthic assemblages of the Tail of the Grand Bank and Flemish Cap (northwest Atlantic) in relation to environmental drivers. Deep Sea Res Part 1. 2015. doi: 10.1016/j.dsr.2015.08.006

74. Davis SD, Gilhen J. Marine Animals Collected During FRB Cruise 104 1972. Nova Scotia Museum; 1973. Curatorial Report No.: 10.

75. Davis SD. Benthic animals from Cruises P.114 and P.139, 1973 and 1974. Nova Scotia Museum; 1976. Curatorial Report No.: 31.

76. Thorson G. Bottom Communities (Sublittoral or Shallow Shelf). In: Hedgpeth G, editor. Treatise on Marine Ecology and Palaeoecology. Ch. 17 Mem Geol Soc Amer; 1957; 67: 461–534.

77. Maciolek-Blake N, Grassle JF, Blake JA, Neff JM. Georges Bank benthic infauna monitoring program: final report for second year of sampling (July,1982–May,1983). Battelle New England Marine Research Laboratory. Woods Hole Oceanographic Institution; US. Minerals Management Service; 1984. Contract No.: 14-12-001-29192.

78. Schneider DC, Gagnon J-M, GIlkinson KD. Patchiness of epibenthic megafauna on the outer Grand Banks of Newfoundland. Mar Ecol Prog Ser. 1987; 39: 1–13.

79. Auster PJ, Langton RW. The effects of fishing on fish habitat. Am Fish Soc Symp. 1999; 22:150–187.

80. Hargrave BT, Kostylev VE, Hawkins CM. Benthic epifauna assemblages, biomass and respiration in The Gully region on the Scotian Shelf, NW Atlantic Ocean. Mar Ecol Prog Ser. 2004; 270: 55–70.

81. Kaiser MJ, Bergmann M, Hinz H, Galanidi M, Shucksmith R, Rees EIS, et al. Demersal fish and epifauna associated with sandbank habitats. Estuar Coast Shelf Sci. 2004; 60: 445–456.

82. Anderson JT, Simon JE, Gordon DC Jr, Hurley PC. Linking fisheries to benthic habitats at multiple scales: Eastern Scotian Shelf Haddock. Am Fish Soc Symp. 2005; 41: 251–264.

83. Gordon DC Jr. Project overview. In: Anderson JT and Gordon DC Jr. editors. Report of the Spatial Utilization of Benthic Habitats by Demersal Fish on the Scotian Shelf Synthesis Meeting 2007. Can Tech Rep Fish Aquat Sci. 2007. Report No.: 2770. p. 1–5.