Rapid biotic homogenization of marine fish assemblages

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The role human activities play in reshaping biodiversity is increasingly apparent in terrestrial ecosystems. However, the responses of entire marine assemblages are not well-understood, in part, because few monitoring programs incorporate both spatial and temporal replication. Here, we analyse an exceptionally comprehensive 29-year time series of North Atlantic groundfish assemblages monitored over 5\(^\circ\) latitude to the west of Scotland. These fish assemblages show no systematic change in species richness through time, but steady change in species composition, leading to an increase in spatial homogenization: the species identity of colder northern localities increasingly resembles that of warmer southern localities. This biotic homogenization mirrors the spatial pattern of unevenly rising ocean temperatures over the same time period suggesting that climate change is primarily responsible for the spatial homogenization we observe. In this and other ecosystems, apparent constancy in species richness may mask major changes in species composition driven by anthropogenic change.
The Anthropocene1,2 is characterized by marked changes in the diversity of natural assemblages across the Earth3. There is, however, increasing evidence that community responses to anthropogenic impacts are complex and scale dependent4. Although heavily transformed terrestrial habitats exhibit species loss3, recent meta-analyses have detected no systematic change in local a diversity5,6. In contrast, temporal turnover (temporal β diversity)7 in local assemblages is occurring at rates in excess of background levels predicted by null and neutral models8, β diversity measures change in community composition over time (or space), and will be influenced by a range of processes including species invasion and shifts in species ranges. Dornelas et al.6 hypothesized that elevated rates of turnover could be a result of increasing biotic homogenization, a process whereby previously differentiated assemblages increasingly resemble one another in species composition. This hypothesis of a linkage between untrending local species richness, high turnover in local species composition and declining spatial β diversity (increasing homogenization) has not yet been tested in any ecosystem.

To date, investigations of biotic homogenization have focused on direct human influences on freshwater and terrestrial assemblages8. For example, deliberate introductions of gamefish are largely responsible for the greater homogeneity of contemporary freshwater fish assemblages in the United States compared with the time of the European settlement9. However, open water marine communities—especially those in offshore localities—are arguably less vulnerable to biotic homogenization than terrestrial ones because they are initially less spatially differentiated10–12 and less subject to some of the direct human interventions driving terrestrial homogenization such as species introductions and habitat modification. Nonetheless, for many marine species13, recent shifts in geographic ranges and phenology have been linked to anthropogenic climate change14–17 and temporal turnover in marine assemblages has also increased relative to null expectations6. This makes offshore marine ecosystems a good candidate for testing the role of elevated community turnover in homogenization.

Using data collected during a systematic Scottish groundfish survey, we probe marine community shifts by quantifying temporal trends in a and β diversity over almost three decades. We show that biotic homogenization is driven by elevated species turnover, transforming marine assemblages in ways that resemble terrestrial assemblages.

**Figure 1** | Temporal trends in a diversity (rarefied species richness) and β diversity (Jaccard similarity, relative to the initial year), in each latitudinal band, over the duration of study. Trend lines (OLS regression) are colour coded red if significantly negative (P < 0.05, n = 28) and grey if not significant. (There were no significant positive slopes). See Supplementary Figure 2 for overall trend.
Results

\( \alpha \) and \( \beta \) diversity. With a total regional count of 131 species recorded, and an average local species richness of 21.5 (per 30 latitudinal band, per year), there was no systematic change in rarefied species richness across the 29 years of sampling (ordinary least squares (OLS) regression, \( r^2 = 0.005 \), \( P = 0.15 \), \( n = 252 \), see Supplementary Figure 2), although the nine 30° latitudinal bands exhibit heterogeneous temporal trends in species richness (\( \alpha \) diversity; left-column panels, Fig. 1). In contrast to species richness, community composition shows increasing change relative to the baseline level within each latitudinal band, reflecting the temporal turnover that is typical for other marine, terrestrial and freshwater assemblages\(^6\) (Jaccard similarity; right-column panels, Fig. 1).

Biotic homogenization. It is possible for temporal turnover to be independent at each site, with no spatial signature. Instead, we found the temporal turnover depicted in Fig. 1 is at least partly explained by the species composition in the different latitudinal bands becoming progressively more similar over time (Fig. 2). Moreover, the distance-decay relationship in composition is less steep in later years, indicating that the northern and southern assemblages are now less differentiated from one another than they were three decades ago (Fig. 3 and Supplementary Fig. 4). The result of these progressive changes is spatial biotic homogenization. These analyses confirm the hypothesis of Dornelas et al.\(^6\) that biotic homogenization can lead to a pattern of constant local species richness but increasing change in species composition. As further support, an analysis of spatial \( \beta \) diversity partitioning\(^18,19\) reveals that turnover in species identity is a consistently stronger driver of community reorganization than change in species richness (Supplementary Fig. 5 and Supplementary Fig. 6). This result is in contrast to a recent meta-analysis\(^20\) which argued that changes in community similarity are driven by changes in species richness rather than changes in species composition.

Water temperature. What forces are responsible for this widespread reorganization of marine fish assemblages? In this region of the North Atlantic, average ocean temperature has increased by c. 1 °C over the time frame of this study (Fig. 4a). Moreover, the south–north temperature gradient has decreased, so that the latitudinal bands in Fig. 1 are now thermally less differentiated than they used to be (Fig. 4d), meaning the northerly waters have increased in temperature more rapidly than the southerly waters. The mean annual pairwise difference in temperature between latitudinal bands (Fig. 4c) also declines through time. This decline mirrors the shift in community similarity indicating that, as water temperatures within the latitudinal bands converge over time, so too does the composition of the assemblages found there.

Discussion

This investigation has documented rapid biotic homogenization in marine fish assemblages and argued that these community shifts are linked to climate. Other recent studies also highlight the potential of anthropogenic climate change\(^21,22\) to restructure marine communities. Climate velocity has been identified as a predictor of biotic change\(^14\). Our data show that these transformations are manifested over relatively small geographic areas and short time periods. It is likely that climate change acts in synergy with other anthropogenic stressors, such as pollution and over-exploitation\(^23\). However, although fishing has heavily impacted the North Atlantic for over a century, fishing effort (as reported landings and trawling hours) has not increased appreciably over the time period of this study\(^24,25\). Moreover, coastal human population densities in this region have remained low over the last three decades (for example, in 2011 mean human population density was 5.1 km\(^{-2}\)).

Our perception of the natural state of marine communities is shaped by shifting baselines\(^26\). Contemporary accounts of Scottish waters in the eighteenth century attest to greater
abundances of fish and sea mammals present at that time while drawing attention to the impacts that commercial fishing were having over 200 years ago. The homogenization we document here is a continuation of the ongoing change in fish communities exposed to long-term anthropogenic pressures.

As we have shown, apparent constancy in widely used measures of α diversity (that is, lack of trends in local species richness) can mask enormous changes in an ecosystem. Local richness has, on average, been maintained over the duration of this study, with the same common species (including the Norway pout Trisopterus esmarkii, and Atlantic herring Clupea harengus) continuing to be overall dominants. However, our analyses reveal that groundfish communities off western Scotland are undergoing rapid reorganization. This change is driven by spatial homogenization, that is increasing similarity between locations 100 s of kms apart—possibly linked to climate change. Our data suggest that predicted changes in marine fish communities are already underway. Biogeographical shifts in zooplankton species in the North East (NE) Atlantic provide further evidence that the pace of change is marine communities exceeds that in the terrestrial realm. This collapse of spatial β diversity (that is, increasing homogenization) is arguably a far greater and more pressing crisis than the loss of local species richness. As long as species are not globally extinct this homogenization is potentially reversible. However, this crisis is largely unrecognized, and adds to the challenges already facing marine biodiversity.

Methods
Data. The Scottish groundfish survey takes place annually, in the seas to the north and west of Scotland (Supplementary Fig. 1). Sampling began in 1985. Trawling occurs in the first quarter of the year (January, February and March), and the same methodology has been used throughout. Each sample is a discrete trawl occurring at a precise geographical point within an International Council for the Exploration of the Seas (ICES) statistical rectangle (rectangles represent a 30’ latitude by 1’ longitude grid cell). Sample number varies between rectangles in a given year. Species abundances are reported as catch per unit effort (that is the number of individuals per species caught during a 1 h trawl). Body size (standard length) is also recorded. We focus on the 35 rectangles (Supplementary Fig. 1) for which there are good community time series. A total of 131 taxa have been sampled in this study area over the 29 years of the survey. 126 of these taxa are fish species, the remaining 5 macroinvertebrates (squid and macrocrustaceans). As 1992 and 1995 have substantially fewer samples than other years, we exclude them from our analyses. Species and temperature data were downloaded from the ICES portal (DATRAS Fish Survey Data ‘Scottish West Coast Survey For Commercial Fish Species 1985–2013’ (Available at https://datras.ices.dk, accessed on 2014)). Information on coastal human population density was obtained from National Records of Scotland, 2011 Census: Aggregate data (Scotland). UK Data Service Census Support. Downloaded from: http://infuse.mimas.ac.uk. This information is licensed under the terms of the Open Government Licence (http://www.nationalarchives.gov.uk/doc/open-government-licence/version/2) accessed on 2015.

Analysis. To understand how the composition of these groundfish communities is changing through time, relative to latitude, we first assign the rectangles to nine 30’ latitudinal bands. We then compile a community time series for each latitudinal band. Sample rarefaction (as in ref. 6) ensures equal sampling effort across bands and is used in the calculation of temporal β diversity (species richness) and temporal β diversity (jaccard similarity (presence/absence) and Bray–Curtis similarity (quantitative)). We first calculate similarity in relation to the start of the survey. Next, for each year, we compute the pairwise compositional similarity of these latitudinal bands. We also construct distance-decay plots (similarity plotted against geographic distance) for each year. Quantile regression is used to fit the median slope of these relationships as it is robust against outliers. β diversity partitioning, enables us to compute the relative contributions of turnover and nestedness to community change. To do this we recalculate Jaccard similarity as Jaccard dissimilarity because β diversity partitioning conventionally focuses on dissimilarity.

Geographical distances (in km) are calculated using the point distance tool in ArcGIS. Analyses use R statistical software. The vegdist function in the R package vegan is used to compute similarities. The quantreg package is used to fit median slopes, and the betapart package to partition β diversity into components of turnover and nestedness.

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Author contributions

A.E.M., M.D. and F.M. conceived the project in discussion with NJG and BMcG. F.M. was responsible for data handling and ran all the analyses. All authors contributed to the final manuscript.

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