Genomics of habitat choice and adaptive evolution in a deep-sea fish

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Intraspecific diversity promotes evolutionary change, and when partitioned among geographic regions or habitats can form the basis for speciation. Marine species live in an environment that can provide as much scope for diversification in the vertical as in the horizontal dimension. Understanding the relevant mechanisms will contribute significantly to our understanding of eco-evolutionary processes and effective biodiversity conservation. Here, we provide an annotated genome assembly for the deep-sea fish Coryphaenoides rupestris and re-sequencing data to show that differentiation at non-synonymous sites in functional loci distinguishes individuals living at different depths, independent of horizontal spatial distance. Our data indicate disruptive selection at these loci; however, we find no clear evidence for differentiation at neutral loci that may indicate assortative mating. We propose that individuals with distinct genotypes at relevant loci segregate by depth as they mature (supported by survey data), which may be associated with ecotype differentiation linked to distinct phenotypic requirements at different depths.

While longitudinal and latitudinal habitat transitions have been proposed to define marine communities and promote intraspecific differentiation1,2, little is known about the importance of transitions along ocean depth gradients3,6, although substantial changes in species assemblages with depth have been recorded (for example, ref. 8), and relatively narrow depth ranges may distinguish closely related species (for example, refs 9,10). Understanding the relevant mechanisms will contribute significantly to our understanding of eco-evolutionary processes and the origin of marine biodiversity. We chose the roundnose gendier (Coryphaenoides rupestris) as a model system because it is a widespread species that can inhabit a comparatively broad range of depths9 from ~180 m to 2,600 m. It is a batch spawner, producing up to 69,000 pelagic eggs per female10. It has a spawning season peaking in autumn11 (recent observations were in September at 1,500 m; ref. 12), and the depth gradient can retain polymorphisms within or among populations, this potential to promote incipient speciation through ecological processes in sympatry13. When environmental change exposes new habitats and niche potential, adaptive radiations may rapidly generate a new lineage of species18,19. To the extent that differential selection at these loci; however, our data indicate assortative mating. We propose that individuals with distinct genotypes at relevant loci segregate by depth as they mature (supported by survey data), which may be associated with ecotype differentiation linked to distinct phenotypic requirements at different depths.

Results and discussion

We produced an annotated reference genome for C. rupestris with a total length of 0.829 gigabase pairs, a mean depth of 104× and an N50 of 159,738 (see Supplementary Methods for details). We used this draft genome to map 60 additional genomes sequenced to a mean depth of ~6×, representing a transect from 750 m to 1,800 m over a horizontal sampling range of 25 km (Supplementary Fig. 1), collected on the same day. Manhattan plots using generalized linear model (GLM) and \( F_{ST} \) metrics (Fig. 1a and Supplementary Fig. 2) consistently showed the same pattern of outliers for comparisons between 1,800 m and shallower sites along the transect, although they were the most pronounced for the comparison between 1,000 m and 1,800 m. Among 25 genomic regions showing clusters of outlier single nucleotide polymorphisms (SNPs) exceeding the Manhattan plot threshold correction for multiple tests (a total of 346 outliers; Fig. 1a), 9 SNPs coded for non-synonymous changes within 6 genes, and genotypes were strongly correlated with habitat depth (Fig. 2). These 9 non-synonymous sites were on 5 contigs and surrounded the depth gradient, to experience differential selective pressures. We tested hypotheses about adaptation to these deep-sea habitats using genome sequence data together with data on the ecology and life history of the subject species. We found that juvenile fish of this species are found primarily in relatively shallow depths (near the transition between the mesopelagic and bathypelagic zones) and then migrate as they mature to different depths, this is strongly associated with their genotype at a set of functional loci. In particular, all adults below ~1,800 m share the same homozygous genotype at each locus. There is evidence for strong selection maintaining this difference, but no clear evidence for differentiation driven by assortative mating.
by multiple significant outliers on the same contig (for example, the non-synonymous sites on contig 1041 have 32 significant outliers nearby on the same contig). Principal component analysis (PCA) for all 346 outliers also showed a strong correlation with depth, while the remaining 5.9 million non-outlier SNPs did not (Fig. 3a), and pairwise \( F_{ST} \) values for a subset of 44,650 neutral loci were not significantly different from zero (see Methods and Supplementary Table 1). Simulations showed this number of loci to be sufficient to detect an \( F_{ST} \) of 0.0007 with a power of 0.86 (see Methods). An independent assessment of genomic regions associated with habitat depth (cacti plots; Fig. 1b and Supplementary Table 2) reinforced the evidence for selection. Although some catch data have previously been interpreted to suggest diurnal and seasonal vertical migrations\(^9\), these strong genomic associations instead suggest considerable adult fidelity to specific habitat depths. Temporary migrations during spawning to an intermediate depth (~1,500 m) are possible based on recent data\(^12\).

Evidence for linkage disequilibrium due to population mixture and a two-locus Wahlund effect\(^20\) further reinforces the pattern of differentiation at outlier loci for comparisons between 1,800 m and shallower depths (Supplementary Fig. 3). To determine whether the observed linkage disequilibrium was driven by concerted evolution or mixture linkage disequilibrium, we therefore needed to restrict our analyses to just one depth (which greatly attenuates the effects from mixture linkage disequilibrium; Supplementary Fig. 3). Comparing the decay profiles for linkage disequilibrium associated with physical linkage for the samples from 1,000 m, we find a consistent evidence for selection. Although some catch data have previously been interpreted to suggest diurnal and seasonal vertical migrations\(^9\), these strong genomic associations instead suggest considerable adult fidelity to specific habitat depths. Temporary migrations during spawning to an intermediate depth (~1,500 m) are possible based on recent data\(^12\).

DNA sequencing analyses (RAD-seq; see Methods) and distinguished putative neutral loci from outlier loci using the \( F_{Dist} \) method. Focusing on four sample sites near the Hebrides (see Supplementary Fig. 1), we found ordination clustering by depth (Fig. 2a). The six coding regions are indicated, and contig references are included putative neutral loci from outlier loci using the \( F_{Dist} \) method. Focusing on four sample sites near the Hebrides (see Supplementary Fig. 1), we found ordination clustering by depth (Fig. 2a). The six coding regions are indicated, and contig references are included.

## Supporting Information

- **Supplementary Fig. 1**: Variance explained by principal components (PCs) for each depth.
- **Supplementary Fig. 2**: Volcano plots showing the strongest outlier SNPs.
- **Supplementary Fig. 3**: Linkage disequilibrium decay profiles for all outlier loci.
- **Supplementary Table 1**: Summary of outlier SNPs.
- **Supplementary Table 2**: Genomic loci with positive selection signals.

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**Fig. 1** | Comparisons by habitat depth. **a**, Negative \( \log_{10} \) of \( P \) values based on the GLM for individuals at 750 m versus 1,500 m, and 1,000 m versus 1,800 m. The horizontal lines mark Bonferroni (red) and FDR (blue) corrections for multiple comparisons, with 346 outliers in the 1,000 m versus 1,800 m comparison above the red line. Alternating blue and red dots show the transition from one contig to another. **b**, Localized phylogenetic patterns within the genome ('cacti'). The majority of the genome exhibits neutral genetic divergence (as exemplified by cactus 4), while one cactus (cactus 5), which accounts for less than 1% of the genome, displays all but one 1,800 m sample clustering in the same lineage (see Supplementary Table 2).

**Fig. 2** | Genotypes of 60 re-sequenced genomes for nine non-synonymous changes. The six coding regions are indicated, and contig references are given. Each column represents an SNP, colour coded for homozygote (blue or yellow) and heterozygote (green). Each row represents an individual, arranged by habitat depth. N denotes an unresolved genotype at that position for that individual.
Comparisons among contigs are given for outliers showing non-synonymous changes within coding regions. The proportion of individuals at each depth with the ‘deep-adapted’ allele is shown (pooling heterozygotes and homozygotes).

For the 60 re-sequenced genomes along the depth transect (Supplementary Fig. 1), there was a highly significant pattern of homozygote excess at the nine outlier loci with non-synonymous changes (Supplementary Table 3). Homozygote excess was not more generally observed, as most of the 5.9 million non-outlier loci were consistent with Hardy–Weinberg equilibrium (HWE) (3.7% divergent after correction for false discovery), and non-outlier loci out of HWE were more likely to show heterozygous excess (while the 346 outliers mostly showed homozygote excess; Supplementary Table 3).

In the contig showing the most non-synonymous changes, with two outlier loci associated with three non-synonymous changes (contig 1041), we detected five RAD-seq outlier SNPs (out of 4,989 SNPs derived from 207 samples across 9 populations) and assessed their genotypes across the full geographic range. For samples taken at ≥1,800 m, one allele dominated at each of these loci, while the same allele was less frequent among samples from <1,800 m (Supplementary Table 4). In each case there was significant homozygote excess, consistent with under-dominance and disruptive selection (P < 0.00001; Supplementary Table 4).

We considered juvenile fish separately because trawl survey data demonstrate that juveniles of this species are predominantly found in waters at 1,200 m or less (Supplementary Fig. 4). We collected 96 juveniles from 1,000 m near the 60-genome transect location (transect 2; Supplementary Fig. 1) and genotyped these individuals at four non-synonymous outlier SNPs we genotyped in the adults. The allele frequencies of juveniles differed significantly from the adults captured at the same depth (Fig. 5, Supplementary Table 4 and Supplementary Fig. 5), showing roughly the same proportion of genotypes associated with adults from 1,800 m as with adults from 1,000 m (Fig. 5). Genotype frequencies in juveniles also showed significant deviation from HWE at all four loci in a pattern consistent with disruptive selection (Supplementary Table 4). Disruptive selection has been proposed as an important driver of ecological speciation when coupled with assortative mating21–23. However, as illustrated in Fig. 3, for non-outlier SNPs there was no clear evidence for differentiation among samples taken at different depths from the same geographic region, and differentiation would be expected if there was sustained assortative mating. This assumes that over time (sufficient time for concerted evolution in this case; see Fig. 4), assortative mating would result in differentiation at neutral loci through genetic drift. There is also no evidence of distinct breeding aggregations separated by depth12,19.

Cohorts tracked from samples collected by trawl (see Methods) in eight different years indicate an ontogenetic depth migration (Supplementary Fig. 4). A latent-class model (two-component mixture model; see Methods) applied to a subset of high-density age classes (>299 individuals per km²) revealed two ontogenetic
analyses found 29 gene ontology terms significantly over-represented in depth-correlated outlier SNPs detected from the Manhattan plot. A gene ontology analysis for all 69 genes associated with the 346 loci had up to 14 SNPs associated with it (one within 30 Kb, 15 within 10 Kb and the rest having SNPs within introns or exons). Of the 29 terms, 16 were associated with development or morphogenesis (Supplementary Table 5). This may be consistent with a more general trend for deep-demersal species to show distinct phenotypes compared with related species at shallower depths, although intraspecific morphological variation with depth has not yet been reported for this genus. There is also no well-studied, closely related species to serve as a reference for gene ontology terms, so proposed functions should be interpreted with caution.

A key difference between habitats at 1,000 m and below 1,800 m is access to the deep-scattering layer within the mesopelagic zone. The deep-scattering layer is a mid-water (200–1,000 m) mass of small fishes, cephalopods, crustaceans and zooplankton that provides a rich source and variety of prey items. It should be possible for C. rupestris to feed there at a relatively high trophic level (and some data are consistent with this); Supplementary Fig. 6), although the predation risk may also be relatively high. In deeper water, the benthic and pelagic systems become increasingly decoupled, resulting in low particulate organic carbon influx and lower food availability, so feeding may be more closely associated with the benthos and from a less diverse prey resource. We compared stable isotope data for carbon and nitrogen. Fish from ≥1,800 m were 13C enriched and 15N depleted compared with samples from 1,000 or 1,050 m (multivariate analysis of variance, F=6.43, P=0.003; Supplementary Fig. 6). These data support our genetic and life-history data, indicating two strategies: one in shallower water with access to the relatively abundant resources of the deep-scattering layer, but potentially greater predation risk, and another in relatively deep water feeding on distinct prey (suggested by carbon isotope data).

Our data reveal genomic differences associated with ecological specializations that are probably related to the anatomical and physiological requirements of the different environments and behavioural strategies, and indicate that fish adopt a habitat suited to their genotype as they mature. This type of vertical population structuring has important implications for fisheries management in general, and is not currently recognized for stock definition. For C. rupestris in particular, the Northeast Atlantic stock is currently assessed as single unit with a total allowable catch of around 2,000 tonnes. From 2017, the European Union has prohibited bottom trawling at depths greater than 800 m (http://data.consilium.europa.eu/doc/document/ST-11142-2016-INIT/en/pdf). While this management strategy protects the deep population of C. rupestris, it means that the entire total allowable catch must now be drawn solely from the shallow population, possibly leading to over-exploitation.

The evolutionary implications are that both temporal (developmental) and spatial factors (associated with habitat characteristics) affect the spatial distribution of diversity established by natural selection, and that this process is not necessarily associated with incipient speciation promoted by assortative mating. Species maintaining intraspecific diversity by strong selection (perhaps for partitioning resources with different phenotypic requirements) could be predisposed to rapid speciation in a conducive environment. Across the broader phylogeny for this genus, a number of species other than C. rupestris inhabit a similarly wide depth range, especially those species also found in the abyss, while other species show narrower, non-overlapping habitat depth ranges. The primary phylogenetic division in this genus is between abyssal and non-abyssal species, with species within those lineages radiating at similar times. This suggests that in some cases habitat depth has led to divisions in alpha taxonomy within this genus; however, our data for C. rupestris suggest instead a system whereby conspecific ecotypes associated with habitat depth are being maintained by disruptive selection at a set of loci in linkage disequilibrium.

Methods

Data reporting. Sample sizes for the detection of outliers were determined based on published simulation analyses that consider power in the context of the depth of sequence and the number of samples (for example, ref. 32). For the analysis of alternative ontogenetic strategies, to achieve model convergence, only data for which the estimated density of fish was greater than 299 individuals per km² were kept, corresponding to 81,549 out of 176,733 fish. There were no randomized experiments, and investigators were not blinded to allocation for outcome assessment.
Sampling. Samples were obtained by trawl fishing for this and earlier projects
(14, 15). Muscle tissue or fin clips were collected as soon after landing as possible and stored in 20% dimethyl sulfoxide saturated with salt or 95% ethanol, and long-term at –20°C. Details of specific sample sets are provided in the Supplementary Methods.

Genome sequencing. Illumina libraries were constructed for short-read, mate-
pair and genome sampling (RAD-seq) using genomic DNA extracted using a phenol chloroform protocol (37), and sequenced on Illumina HiSeq 2000 and 2500 machines. Additional sequencing was undertaken on a PacBio sequencer using single-molecule real-time (SMRT) chemistry. Genomes were assembled and annotated in conjunction with the genome re-sequencing protocols, strategy for RAD-seq sequencing and data analyses were performed as in refs 38–40 and are described in detail in the Supplementary Methods.

SNP calling and outlier detection. We aligned 60 re-sequenced genomes from transect 2 (Supplementary Fig. 1) to our newly assembled and annotated genome for C. rupestris, as described in the Supplementary Methods. SNPs were called using SAMtools version 1.3 (mpileup command with –q 20 –Q 10 –ug flags) and BCFTools (call command with –bvcf flags) (41). We subsequently removed indels using VCFTools (gvcf) and filtered low-quality SNPs using the filter command (e –QUAL < 30 –s ‘LOW.Qual’), this pipeline resulted in 23,883,346 biallelic SNPs.

Outlier detection. We tested for outliers using the GLM implemented in the software TASSEL version 5.0 (ref. 42). First, we ordered the SNPs in the vcf file, ‘LOW_QUAL’). This pipeline resulted in 23,883,346 biallelic SNPs. Data analyses were performed as in refs 37–55 and are described in detail in the Supplementary Methods.

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Comparison of putative populations at outlier and non-outlier loci. PCAs were conducted on both the 5,928,719 non-outlier SNPs from the 60 re-sequenced genomes and the 346 outlier SNPs using the Principal Components Plugin for TASSEL. The genotypes were converted to numeric scores using the Numeric Genotypic function and the missing data imputed to the mean score for each site. PCAs were then performed using eigenvalue decomposition of the covariate matrix, and eigenvectors were calculated from a singular value decomposition of the covariance matrix. The number of principal components was determined by default to five. The full set of SNPs from the re-sequencing analysis was randomly subsampled for 50,000 SNPs using the shuf command in Linux and 44,650 neutral loci identified in LocusScan (50,000 simulations, forced mean Fst confidence interval 0.95). The four depths were compared for Fst at these neutral loci using Arlequin version 3.5 (ref. 43) (results shown in Supplementary Table 1). We tested the power of this Fst analysis by running simulations in R (1,000 replicates). Briefly, we calculated Weir and Cockerham’s multi-locus Fst, for 44,650 loci and 15 individuals per population to generate two distributions—one for a specific Fst level and the other for panmixia—and calculated the power from the overlap between these distributions.

Outlier detection in Hebrides RAD-seq data. We used two methods to identify outlier loci among the four populations in the Hebrides region. First, we ran the FDist outlier approach as implemented in Lositan (52), which incorporates heterozygosity and simulates a distribution for neutrally distributed markers. This method has been shown to have lower rates of type 1 and 2 errors compared with the Fst outlier method implemented in Arlequin version 3.5 (ref. 43). We ran 50,000 simulations under the infinite alleles model with the option of neutral mean Fst and forced mean Fst. We employed a 95% confidence interval and a false discovery rate of 0.05. With this method, we detected 121 outlier loci. Next, we used the more conservative outlier loci detection methods of BayeScan version 2.1 (ref. 44) to detect loci under divergent selection. We ran the programme with the default sample size of 5,000 and the thinning interval of 10. We ran 20 pilot runs each of 5,000 iterations and an additional burn-in of 50,000. We conducted runs with prior odds for the neutral model set at 100 and a FDR of 0.05, which resulted in 122 outliers. There was a >90% overlap (110 loci) between the BayesScan and BayeScan outliers.

PCA of Hebrides RAD-seq data. Based on the results of Lositan for the four Hebrides populations, we divided the SNPs into outlier and non-outlier datasets.

Genotype function and the missing data imputed to the mean score for each site. PCAs were then performed using eigenvalue decomposition of the covariate matrix, and eigenvectors were calculated from a singular value decomposition of the covariance matrix. The number of principal components was determined by default to five. The full set of SNPs from the re-sequencing analysis was randomly subsampled for 50,000 SNPs using the shuf command in Linux and 44,650 neutral loci identified in LocusScan (50,000 simulations, forced mean Fst confidence interval 0.95). The four depths were compared for Fst at these neutral loci using Arlequin version 3.5 (ref. 43) (results shown in Supplementary Table 1). We tested the power of this Fst analysis by running simulations in R (1,000 replicates). Briefly, we calculated Weir and Cockerham’s multi-locus Fst, for 44,650 loci and 15 individuals per population to generate two distributions—one for a specific Fst level and the other for panmixia—and calculated the power from the overlap between these distributions.

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We evaluated population clustering among the four populations using the PCA method implemented in the R package Adegenet version 2.0 (ref. 70). Adegenet converts the diploid allele information from the Stacks output for each individual into a data frame and uses those data to perform the PCA.

Screening of nine populations for outlier SNPs on contig 1041. We conducted an outlier analysis using Loisian on the ‘nine populations’ file of 4,989 SNPs. We ran 50,000 simulations under the infinite alleles model with the option of neutral mean $F_T$ and forced mean $F_S$. We employed a 95% confidence interval and a false discovery rate of 0.05. With this method, we detected 341 outliers. Of these, five mapped to contig 1041 (which had the largest number of strong outliers from the comparison among 60 re-sequenced genomes). We extracted the genotypes for these five outlier loci and determined the allele frequencies across the dataset.

Gene ontology analysis. We searched for gene ontology terms using the FatGO tool on the Babelomics 5 platform (http://babelomics.bioinfo.cipf.es/). FatGO75 is a enrichment test whereby two lists of genes are compared to detect significant over-representation of functional annotations in the subject compared with the reference list. In this case, we compared the set of the 69 genes (Supplementary Table 4) with the SLAY database of 44,446 depth-correlated outlier SNPs detected in the Manhattan plot analyses and found within 30 kilobases of a coding gene, against the full list of 15,114 genes identified from our C. rupestris genome annotation that were associated with gene name, gene description and gene ontology terms using InterProScan. Gene ontology term functions were identified with reference to the blast2go database (chosen due to the relative completeness level of annotation and functional analysis and the lack of a suitable reference from a closely related species). A Fisher’s exact test for 2 X 2 contingency tables (testing for over-representation in list 1) was used to check for significance. Significance was corrected for multiple testing using Benjamini and Hochberg’s FDR-controlling procedure76. Gene ontology biological processes, molecular function, cellular components and GO slim annotation, InterPro and the Genome-Scale Metabolic Network database were searched, filtering terms by 5–500 annotated identifications in each database.

Stable isotope analysis. For analysis of carbon and nitrogen stable isotopic ratios (δ13C and δ15N denotes isotopic ratios of 13/12 C and 14/15 N relative to known standards), white muscle tissue that had been stored in 95% ethanol from 25 C. rupestris samples from shallower than 1,100 m (including 16 from 1,000 m sequenced for genomes and 9 from a nearby site) and another 25 from 1,800 m or deeper (including 16 from 1,800 m sequenced for genomes and 9 from a nearby site) were subjected to lipid extraction preparation. Briefly, tissue pieces of approximately 0.25 cm3 were (1) finely diced and sonicated in 1 ml of 3:1 dichloromethane:methanol solution for 15 min, then (2) centrifuged at 3,000 rpm for 10 min before excess solution was removed. Steps 1 and 2 were repeated three times. The remaining solid sample was then sonicated in 1 ml deionized water for 15 min before being centrifuged at 3,000 rpm for 10 min. Excess water was removed and samples were air-dried at 50°C for 48 h. Dried samples were then mechanically powdered and 0.4 mg of each sample was sealed into tin caps for analysis. Carbon and nitrogen isotope analyses of the samples was performed at the Stable Isotope Biogeochemistry Laboratory, Durham University, using a ECS 4010 Nitrogen / Protein Analyzer (Costech Analytical) connected to a Delta V Advantage Isotope Ratio Mass Spectrometer (Thermo Fisher Scientific). Carbon isotope ratios are corrected for δ13O contribution and reported in standard δ notation in per mil (%) relative to Vienna Pee Dee Belemnite. Isotopic accuracy was monitored through routine analyses of in-house standards, which were stringently calibrated against international standards (for example, United States Geological Survey (USGS) 40, USGS 24, International Atomic Energy Agency (IAEA) CH3, IAEA CH7, IAEA N1 and IAEA N2). Thus provided a total linear range in δ13C between −46‰ and +3‰, and between −4.5‰ and +20.4‰ for δ15N. Analytical uncertainty in δ13C and δ15N was typically ±0.1‰ or better for replicates analyses of the international standards and <0.2‰ for replicate sample analysis. Total organic carbon was obtained as part of the isotope analysis using an internal standard (glutamic acid, 40.82‰ and C, 9.52‰). δ13C was determined by using an underestimation of the depth at settlement, particularly for populations sharing a spawning site at 1,500 m (ref. 13). Samples were collected during the spawning period and so the distribution could reflect this and result in an underestimation of the depth at settlement, particularly for the deep strategy.

Life Sciences Reporting Summary. Further information on experimental design is available in the Life Sciences Reporting Summary.

Data availability. Sequence data have been deposited at GenBank under accession codes PRJNA417962 for the reference genome, SRP129631 for the 60 re-sequenced individuals and PRJNA430030 for the RAD-seq data. Figures 1–4 have associated source data, as do Supplementary Figs. 2–6. There are no restrictions on data availability.

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Modelling ontogenetic depth migration strategies. Size distributions were transformed into age distributions using the length at age table provided in ref. 76 and a multinomial logistic model as described by ref. 77.

Ontogenetic depth migration. To test whether C. rupestris shows an ontogenetic depth migration78,79 from shallow as juveniles to deeper as adults, we assigned individuals to age cohorts and modelled depth change over years. Cohorts of fish spawned between 1996 and 2003 were used to assess depth change over the first 12 years of life. Depth at capture was described using an asymptotic regression model to age cohorts and modelled depth change over years. Cohorts of fish that were sampled during the spawning period and so the distribution could reflect this and result in an underestimate of the depth at settlement, particularly for the deep strategy.

Alternative ontogenetic strategies. To assess the possibility of different ontogenetic depth migration strategies (and eventual adult settlement depth), depth at age data were modelled using a latent-class model (two-component mixture model) where each fish belongs to one of two unobserved (latent) classes or strategies; that is, one shallower and one deeper (Fig. 6). To achieve model convergence, it was necessary to reduce the asymptotic function to two parameters and use a subset of the data for which the estimated density of fish was greater than 299 individuals per km2 (this corresponded to 74 hauls out of 120, and 81,549 fish). As before, the Bayesian model was executed in R 3.2.3 using the R package rjags and formalized as:

$$Depth_{ij} = \text{dnorm}(mu_{ij}, tau_{ij})$$

The depth observed for individual ‘i’ from cohort C is drawn from a normal distribution with parameters $mu_{ij}$ and $tau_{ij}$, where:

$$mu_{ij} = \text{Asym} + \left( R_{ij} - \text{Asym} \right) \times \text{exp}^{-\text{exp}\left(\text{irc}_{ij} \times \text{Age}_j\right)}$$

Asym is the asymptotic depth, $R_i$ is the depth at age 0 and $irc_i$ is a cohort-specific parameter corresponding to the rate of depth change with age. A hierarchical structure was adopted for the cohort-specific parameters $irc_i$ and $tau_{ij}$, with priors assigned to a normal and a gamma distribution, respectively, and non-informative uniform hyperparameters. Non-informative uniform priors were used for all the other parameters.

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Author contributions
A.R.H. conceived the study and contributed to the analyses. A.R.H. and M.R.G. wrote the paper with input from all authors. M.R.G., G.A.G., M.d.J., F.S., F.N., T.R., D.M., D.R.G., N.H., X.L., J.K., A.L., M.H. and S.H. contributed to data generation and analyses.

Competing interests
The authors declare no competing interests.

Additional information
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Experimental design

1. Sample size

Describe how sample size was determined.

For genotype detection from the re-sequencing data we based sample size and coverage on published simulation data. For the cohort modeling, all individuals collected during the dedicated MSS (Marine Scotland Science) deep-water bottom trawl survey of the continental slope west of Scotland between the years 1998-2015 were included in the analysis (120 hauls yielding a total of 176,733 fish).

2. Data exclusions

Describe any data exclusions.

For the cohort modeling, in order to assess cohort-specific depth change over the first 12 years of life, cohorts of fish spawned between years 1996 and 2003 were selected (age 12 and below), corresponding to 33,741 fish. For the analysis of alternative ontogenetic strategies, in order to achieve model convergence, only data for which the estimated density of fish was greater than 299 individuals per km^2 were kept, corresponding to 81,549 fish.

3. Replication

Describe whether the experimental findings were reliably reproduced.

There were no experimental findings to be replicated.

4. Randomization

Describe how samples/organisms/participants were allocated into experimental groups.

No randomization was applied.

5. Blinding

Describe whether the investigators were blinded to group allocation during data collection and/or analysis.

Investigators were not blinded to allocation for outcome assessment for the genetic analyses. While no randomization or blinding was applied to the survey data, the modelling method used for the analysis of alternative ontogenetic strategies (latent class model) allows the assignment of fish to two strategies without investigator intervention and, by using non-informative priors, without use of prior knowledge.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.
6. **Statistical parameters**

For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).

| Item | Confirmed |
|------|-----------|
| n/a  |           |

- The exact sample size \((n)\) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
- A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- A statement indicating how many times each experiment was replicated
- The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
- A description of any assumptions or corrections, such as an adjustment for multiple comparisons
- The test results (e.g. \(P\) values) given as exact values whenever possible and with confidence intervals noted
- A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
- Clearly defined error bars

*See the web collection on statistics for biologists for further resources and guidance.*

7. **Software**

Describe the software used to analyze the data in this study.

For the genetic analyses a diversity of published software packages were applied and these are described in detail in the methods and supplementary texts. For the cohort modelling all models were developed in R 3.2.3 using the package rjags (Plummer et al. 2016).

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). *Nature Methods* guidance for providing algorithms and software for publication provides further information on this topic.

8. **Materials and reagents**

Materials are available on a collaboratory basis.

9. **Antibodies**

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

N/A

10. **Eukaryotic cell lines**

a. State the source of each eukaryotic cell line used.

N/A

b. Describe the method of cell line authentication used.

N/A
c. Report whether the cell lines were tested for mycoplasma contamination.

N/A
d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

N/A
Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals
   Provide details on animals and/or animal-derived materials used in the study.
   For genetic analyses samples were derived from materials collected during this study and earlier studies by deep water trawls, and details are provided in the supplementary methods. For the cohort analyses, data used in this analysis correspond to a re-utilisation of survey data collected from a total of 176,733 roundnose grenadier (Coryphaenoides rupestris) fish with pre-anal fin length ranging from 0.5 to 29.5cm over a period from 1998 to 2015. All fish were brought dead on board of the research vessel.

Policy information about studies involving human research participants

12. Description of human research participants
   Describe the covariate-relevant population characteristics of the human research participants.
   N/A