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Shared ancestral polymorphism and chromosomal rearrangements as potential drivers of local adaptation in a marine fish

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Abstract

Gene flow has tremendous importance on local adaptation, by influencing the fate of *de novo* mutations, maintaining standing genetic variation, and driving adaptive introgression. Furthermore, structural variation as chromosomal rearrangements may facilitate adaptation despite high gene flow. However, our understanding of evolutionary mechanisms impending or favoring local adaptation in the presence of gene flow is still limited to a restricted number of study systems. In this study, we examined how demographic history, shared ancestral polymorphism, and gene flow among glacial lineages contribute to local adaptation to sea conditions in a marine fish, the capelin (Mallotus villosus). We first assembled a 490 Mbp draft genome of M. villosus to map our RAD sequence reads. Then, we used a large dataset of genome-wide single nucleotide polymorphisms (25,904 filtered SNPs) genotyped in 1,310 individuals collected from 31 spawning sites in the northwest Atlantic. We reconstructed the history of divergence among three glacial lineages and showed that they likely diverged from 3.8 to 1.8 MyA and experienced secondary contacts. Within each lineage, our analyses provided evidence for large and high gene flow among spawning sites. Within the NWA lineage, we detected a polymorphic chromosomal rearrangement leading to the occurrence of three haplogroups. Genotypeenvironment associations revealed molecular signatures of local adaptation to environmental conditions prevailing at spawning sites. Our study also suggests that, both shared polymorphism among lineages, resulting from standing genetic variation or introgression, and chromosomal rearrangements may contribute to local adaptation in the presence of high gene flow.

Keywords: speciation, population genomics, RAD, Joint Site Frequency Spectrum, fish, *Mallotus villosus*, inversion, δaδi

Introduction

Local adaptation plays a critical role in organisms' response to the spatiotemporal variation of their environment (Kawecki & Ebert 2004, Blanquart et al. 2013, Savolainen et al. 2013). A population is considered as locally adapted when individuals have a higher fitness in their home environment than their counterparts from other populations experimenting different environmental conditions (Savolainen et al. 2013). Local adaptation relies on three forms of genetic variation: *de novo* mutations, standing genetic variation, and adaptive introgression (Savolainen et al. 2013, Tigano & Friesen 2016). Classic population genetics theory states that *de novo* mutations are the raw source of genetic adaptation (Smith & Haigh 1974, Kaplan et al. 1989, Barton 1998) for which empirical evidence has been previously reported (e.g., Linnen et al. 2009). Nevertheless, increasing evidences suggest that standing genetic variation play a central role in adaptation (Jones et al. 2012, Haenel et al. 2019, Bitter et al. 2019, Lai 2019) and might allow faster adaptation to a new environment than *de novo* mutation (Barret & Schluter 2008). Finally, adaptive introgression may also contribute to local adaptation when new mutants or variants from standing genetic variation with beneficial effect on fitness are introduced in the population through interbreeding with related taxa (Hedrick 2003, Racimo et al. 2015, Oziolor et al. 2019).

Gene flow, which is determined by organisms' dispersal capacity and land/seascape resistance, have tremendous effects on local adaptation (Lenormand 2002, Tigano & Friesen 2016). When selection is spatially heterogeneous but temporally constant, gene flow may erode local adaptation by swamping local alleles or impose a fitness cost to dispersers (Lenormand 2002, Blanquart & Gandon 2011). In the case of *de novo* mutations, simulation-based studies found that the interplay between selection s and rate of gene flow m is the strongest determinant of whether a beneficial de novo mutation establishes in a population. A *de novo* mutation is most likely to establish when s is greater than m (Wright 1931, Lenormand 2002, Yeaman & Otto 2011), or otherwise it will be swamped from the local genetic pool. By contrast, directed gene flow may favor local adaptation when organisms' dispersal decisions (i.e., context-dependent dispersal; Clobert et al. 2009) are adjusted according to local fitness prospects, leading to habitat matching choice (Edelaar et al. 2008), a process that have been reported in various taxa (e.g., Jacob et al. 2017, Lowe & Addis 2019, Camacho & Hendry 2020). Furthermore, gene flow can contribute to increase and/or maintain standing genetic variation on which selection can act, thus increasing the potential for local adaptation (Prezeworski et al. 2005, Monnahan et al. 2015, Tigano & Friesen 2016). In the case of polygenic adaptation in particular, high heritable genetic variation may provide a pool of potential combinations of small-effect alleles determining the expression of fitnessrelated traits (Pritchard & Di Rienzo, 2010; Hancock et al. 2010, Csillery et al. 2018). Lastly, gene flow may

also favor the introduction of beneficial variants from related species (i.e., adaptive introgression), the chance of fixation of introgressed alleles depending on the hybridization frequency and fitness of F1 hybrids and backcrosses (Hedrick 2013).

Three mechanisms allow maintaining local adaptation despite the presence of gene flow: (i) linkage with an already diverged locus, (ii) increased resistance to gene flow following secondary contact due to genetic incompatibilities, (iii) competition among genomic architectures including mechanisms that reduce or suppress recombination (Tigano & Friesen 2016). In the last few years, molecular ecologists paid an increasing attention to the latter mechanism and showed that chromosomal rearrangements, which involve structural changes such as inversion, fusion, fission, and translocation, may be a critical driver of local adaptation (Wellenreuther & Bernatchez 2018, Faria et al. 2019, Wellenreuther et al. 2019). Low recombination within chromosomal rearrangements may lead to independent evolution of the affected genomic regions despite high gene flow in the rest of the genome (Faria & Navarro 2010, Wellenreuther et al. 2017, Mérot et al. 2018, Westram et al. 2018, Wellband et al. 2019).

To date, empirical work focusing on the joint inference of local adaption and demographic history in the presence of high gene flow have been limited to a restricted number of study systems (e.g. in fishes: Tine et al. 2014, Le Moan et al. 2019, Barth et al. 2017, Petterson et al. 2019). Marine organisms are relevant biological models to address this issue since many marine species experience highly heterogeneous marine conditions potentially acting as selective agents, yet displaying very weak genetic differentiation due to large N_e and high connectivity associated with strong dispersal capacities (Palumbi 1992, Bradbury et al. 2008, Laporte et al. 2016, Selkoe et al. 2016, Xuereb et al. 2018). Capelin (Mallotus villosus) is a key-forage fish species (Buren et al. 2014) that displays such characteristics typical of marine organisms. Using mtDNA, microsatellite and AFLP markers, previous studies showed that capelin is characterised by high genetic diversity and weak local genetic structure, suggesting large Ne and high gene flow (Præbel et al. 2008, Colbeck et al. 2011). At a larger scale, three genetically-divergent, and parapatric glacial lineages have been reported in the North Atlantic and Arctic seas without apparent physical or geographic barriers separating them (Dodson et al. 2007). Within each lineage, capelin experiences spatiotemporally variable sea conditions that could be important selective drivers (Carscadden et al. 2001, Rose 2005). Following the pre-nuptial migration, adults may reproduce at sites located in a continuum spanning from the intertidal (i.e., beach spawning sites) to the benthic zone from 1 to 280 m (i.e., demersal-spawning sites) that strongly differ in terms of biotic and abiotic conditions (Nakashima & Wheeler 2002). Environmental variation is especially high in the intertidal zone where the survival and/or development of embryos and larvae depends on temperature, salinity, and trophic

productivity (Frank & Leggett 1981a, 1981b, 1982; Leggett et al. 1984, Præbel et al. 2013, Purchase 2018). The position along the gradient of water depth as well as the sea conditions prevailing in the intertidal zone, are expected to exert strong selective pressures on local populations. This offers a suitable system to investigate connectivity and adaptation at different scales in a marine ecosystem, between and within lineages. Studies that have previously focused on inter- and intra-lineage genetic variation in the capelin were undertaken before the 'genomic era', which hampered the investigation of complex scenarios of divergence history and the search of molecular signals (including chromosomal rearrangements) associated with adaptation to sea conditions.

In this study, we examined how shared polymorphism among the three aforementioned glacial lineages and chromosomal rearrangements underly local adaptation to prevailing sea conditions in capelin spawning sites. We used a large dataset of genome-wide single nucleotide polymorphisms (SNPs) from 1,310 fish collected at 31 spawning sites throughout the Northwest Atlantic and Arctic waters. First, we confirmed the existence of the three glacial lineages (Northwest Atlantic lineage, NWA; Greenland lineage, GRE; and Arctic lineage, ARC) identified by Dodson et al. (2007) by analyzing the pattern of genetic diversity and differentiation in the whole study area (Fig.1). Then, we inferred the demographic history of the three lineages using joint Site Frequency Spectrum (jSFS) for each pair of lineages. We took into account the confounding effect of barriers to gene flow, that affects the rate of migration (Barton & Bengsson 1986) and the confounding effect of linked selection, approximated as a reduction in local effective population size N_e (Charlesworth et al. 1993). Then, we analyzed in detail the genetic structure within the NWA lineage (Fig.1). In particular, we detected the molecular signature of putative chromosomal rearrangements (absent in the other glacial lineages) that best fit a pattern of chromosomal fusion. We examined the molecular signature of local adaption by identifying outlier loci putatively associated with the spawning site position along the water depth gradient (three categories of sites: beach spawning sites, demersal shallow-water sites, and demersal deepwater sites) and environmental variables in beach spawning sites including temperature and trophic production. We also evaluated the contribution of ancestral polymorphism to local adaptation by examining an excess of shared polymorphism among lineages at outlier loci associated with environmental factors. Finally, we explored the role of the chromosomal arrangement on local adaptive variation by determining the proportion of candidate loci captured by the putative arrangement.

Materials and methods

Genome sequencing and draft assembly

A first draft version of the capelin genome was constructed and used for the major purposes of mapping our RAD sequence reads. To estimate the chromosome-scale position of each SNP on the draft genome, we performed a synteny analysis with four other fish genomes. This allowed us to attribute each of the capelin contigs to one of the 24 orthologous chromosomes, i.e. to the chromosome with the longest total alignment across the four species. Capelin genome sequencing and assembly and syntheny analysis are described in details in the **Supplementary methods** section).

Sampling area and Genotyping-By-Sequencing approach

Sampling area and molecular analyses

A total of 1,359 capelins were sampled from 31 sites in the Northwest Atlantic, both in Canadian and Greenland waters (**Fig.1**, and **Table S6**), which were expected to include representatives from the three lineages according to a previous study using mitochondrial DNA and microsatellite genetic markers (Dodson et al. 2007). We sampled 25 spawning sites within the presumed NWA lineage including three demersal shallow-water sites, three demersal deep-water sites, and 18 beach spawning sites. These sites were located in four geographic regions (**Fig.1**): Kuururjuaq, Labrador, Newfoundland, and St. Lawrence. In parallel, we sampled two sites (beach-spawning fishes) within the presumed range of the ARC lineage and four offshore sampling sites in the range of the GRE lineage. The median sample size was 46.5 (range: 19 to 50) individuals per site. Fish were collected during the breeding period (captured by hand at beach spawning sites and using nets at demersal spawning sites), sexed, and a piece of fin was preserved in RNAlater.

Both DNA extractions and library preparation were performed following protocols fully described elsewhere (e.g., Moore et al. 2017; Rougemont et al. 2019). The methods are detailed in the **Supplementary methods** section. A scheme presenting all analyses performed in our study can found in **Fig.S13**.

DNA sequencing, genotyping, and diversity analysis

Barcodes were removed using cutadapt (Martin 2011) and trimmed to 80 bp, allowing for an error rate of 0.2. They were then demultiplexed using the 'process_radtags' module of Stacks v1.44 (Catchen et al. 2013) and aligned to the capelin draft genome assembly using bwa-mem (Li 2013) with default parameters, as detailed below. Then, aligned reads were processed with Stacks v.1.44 for SNP calling and genotyping. The 'pstacks'

module was used with a minimum depth of 3 and up to 3 mismatches were allowed in the catalog creation. We then ran the 'populations' module to produce a vcf file that was further filtered using python (https://github.com/enormandeau/stacks_workflow) and bash scripts. SNPs were kept if they displayed a read depth greater than 5 and less than 70 (based on the mean distribution of read depth computed with vcftools), the latter being set to control for paralogy and highly repetitive regions. Then, SNPs present in a least 70 % in each sampling location and with heterozygosity lower than 0.60 were kept to further control for paralogs and HWE disequilibrium (Dorant et al. 2019, Rougemont et al. 2019). The resulting vcf file comprised of 550,724 SNPs spread over 27,442 loci and was further filtered prior to being used for demographic inferences with $\partial a\partial i$ (see filtering details provided in the **Supplementary methods** section). This vcf file was then subsampled to meet the different assumptions (in particular, the use of unlinked SNPs) of the models underlying the different population genetic analyses performed as described below. We therefore kept one single SNP per locus (the one with the highest minor allele frequency) and further removed SNPs not present in at least 90 percent of the total dataset resulting in a final vcf of 25,904 high quality SNPs.

Hierarchical patterns of population genetic structure

Genetic differentiation between glacial lineages

We documented the extent of genetic differentiation among lineages using the whole dataset (i.e., the 31 sampling sites, Table 1). A principal component analysis (PCA) was performed to investigate the molecular variation among individuals using the R package *Adegenet* (Jombart et al. 2008). Then, we quantified ancestry and admixture proportion using the *snmf* function included in the R package LEA (Frichot & François 2015), with K-values from 1 to 10. We estimated ancestry coefficients for all levels of K and obtained cross-validations using the cross-entropy criterion with 5% masked genotypes. Next, we quantified the level of genetic differentiation among sampling sites using the *F*_{ST} estimator (Weir & Cockerham 1984) in the STAMPP R package along with 95% confidence intervals using 1000 bootstraps (Pembleton et al. 2013).

Geographical and habitat-dependent genetic structure within lineages

A second PCA was executed for each of the three lineages separately to examine within-lineage genetic variation. Admixture analyses were conducted only on the NWA lineage to examine intra-lineage genetic

structure. We did not analyze intra-lineage structure for ARC and GRE because of the limited number of sampling sites (2 and 4, respectively) and the absence of any genetic structure based on PCA results (**Fig. S4**). Moreover, we quantified F_{ST} estimates among all sampling sites, and between demersal and beach spawning sites more specifically.

In addition, we tested for patterns of Isolation-By-Distance (IBD) within the NWA lineage separately for neutral SNPs and SNPs putatively under divergent selection using a procedure detailed in the **Supplementary methods** section. IBD was tested using a linear model in which pairwise F_{ST} ($F_{ST}/(1-F_{ST})$) was included as the response variable and the Euclidean distance (z-score) was incorporated as an explanatory term. The significance of the effect of Euclidean distance was assessed using adjusted R^2 and ANOVA with *F*test.

Population structure as revealed by haplogroup distribution in the NWA lineage

Within the NWA lineage, we detected three genetic clusters using a PCA (hereafter called haplogroup1, haplogroup2, and haplogroup3), which occurred in all sampling sites across the range of the NWA lineage but were totally absent in the other two lineages. This pattern was also supported by a clustering analysis performed for the NWA lineage alone (see Results). This signature may be due to a chromosomal rearrangement (i.e., three haplogroups with higher heterozygosity for the heterokaryote and lower heterozygosity for the two homokaryotes; Berg et al. 2017, Wellband et al. 2019). We thus performed the following analyses to confirm this hypothesis. We identified a set of SNPs mainly located in Chr2 and Chr9 associated with these haplogroups based on their loadings along the PC1 axis (**Fig. S4** and **S5**). We tested whether the heterozygosity of those SNPs differed between haplogroups using a beta regression model; heterozygosity was included in the model as the dependent variable and the haplogroup as discrete explanatory variable. Furthermore, for each sampling site, we tested for Hardy-Weinberg equilibrium when considering the haplogroups as different karyotypes of a putative rearrangement using a chi-square test and a ternary plot implemented in the R package HardyWeinberg (Graffelman et al. 2015).

Next, we examined if the assignment probability of individuals to the haplogroups depended on their sex. We built a logistic regression model for each haplogroup in which individual assignment to the haplogroup was coded as a binary response variable (0 = not assigned to the haplogroup; 1 = assigned) and sex as a discrete explanatory variable. Moreover, we investigated how the frequency of the haplogroup within sampling sites was affected by the spawning habitat (deep-water and shallow-water demersal sites were merged into a single category to increase the power of the analysis). In addition, we examined how temperature and chlorophyll

concentration in beach spawning sites influenced haplogroup frequency. The marine data layers were downloaded from Bio-ORACLE (http://www.bio-oracle.org/) and the two environmental variables were extracted using the R package sdmpredictors (Bosch et al. 2017); we present the environmental data in **Table S5**. We used a linear model where the frequency of haplogroup 1 and 2 [$(2 \times N_{haplogroup1} + N_{haplogroup2}) / N_{total}$] was included as the response variable and the spawning habitat, temperature, and chlorophyll concentration as explanatory variables. Each explanatory variable was introduced separately in the model to avoid model over-fitting. The significance of explanatory variable effects was evaluated using adjusted R^2 and ANOVA with *F*-test.

Demographic history of divergence in $\delta a \delta i$ and identification of inter-lineage outliers

Divergence history of the three capelin lineages

We reconstructed the most likely demographic history using $\delta a \delta i$ (Gutenkunst et al. 2009). We compared four alternative models of historical divergence: (i) a model of Strict Isolation (SI), (ii) a model of divergence with continuous gene flow or Isolation with Migration (IM), (iii) a model of divergence with initial migration or Ancient Migration (AM) and (iv) a model of Secondary Contact (SC) (Fully described in Fig. S3). These models incorporate the confounding effects of selection at linked sites locally affecting Ne and the effect of differential introgression (i.e. due to barriers to gene flow accumulated during population divergence; Barton & Bengtson 1986, Le Roux et al. 2014) affecting the rate of migration (m) along the genome (Tine et al. 2014, Rougemont et al. 2017, Le Moan et al. 2016, Rougeux et al. 2017). Model choice was performed using the Akaike information criterion (AIC), Δ AIC, and AIC weights. We attempted to estimate biological parameters assuming a generation time of 3.8 years (Dodson et al. 2007) and a standard mutation rate value of 1e-8 mutation/bp/generation. Furthermore, we performed additional analyses of the possible confounding effect of the two chromosomal blocks (on separate Chr2 and Chr9) The modeling approach is detailed in the Supplementary methods section. Finally, to obtain an overview of the overall levels of divergence among the lineage we computed the net sequence divergence (Da) as well as level of absolute sequence divergence (Dxy), the former being informative of the position of a pair of populations along the speciation continuum (Roux et al. 2016). We used the vcf haplotype file obtained from stacks and computed these summary statistics using MScalc (Roux et al. 2011) to account for missing data and integrate the length of each rad loci.

Coalescent simulations and identification of inter-lineage outliers

We used the demographic parameter estimates along with their standard deviation under each best model for each pair of lineages to perform 4,000,000 coalescent simulations using the coalescent simulator ms (Hudson 2002). We assumed a strictly neutral model with homogeneous population size and homogeneous migration rate. For each SNP, we computed levels of expected heterozygosity (He) as well as Weir & Cockerham F_{ST} estimator between lineages. As proposed by Beaumont & Nichols (1996) we computed F_{ST} quantiles in heterozygosity intervals of 0.025. This method considers the uncertainty surrounding parameter estimates (Leroy et al. 2019). Loci departing from our neutral envelope (outside the 99.99th upper quantiles of the conditioned F_{ST} distribution) were inferred as candidate outliers. Finally, we used the sequences of the markers potentially under divergent selection between the three lineages to perform a BLAST (Altschul et al. 1990). More specifically, we extracted a 1kb length sequence on each side of the SNPs of interest. Then we performed blasts of those sequences against the NCBI NR database. To consider a SNP further we required a blast length of at least 80bp and an e-value of 1e-10. To further identify non-synonymous mutations, we blasted the fasta sequences against the reference transcriptome. All hits with more than 25 amino-acid similarity and more than 90% similarity with the transcriptome sequence were retained. Both variants of each SNP were used with the BLASTs against the transcriptome and translation results were compared pairwise. For each locus, results were only kept if the lowest e-value hit for both variants was the same (i.e. same protein name and length).

Searching for signals of local adaptation patterns within the NWA lineage

Genotype-environment associations and local adaptation to spawning habitat

We examined the molecular signature of local adaptation to spawning sites within the NWA lineage only, since not enough sampling sites were available within the other two lineages. More specifically, we aimed at detecting outlier loci associated with spawning habitats distributed along a water depth gradient: the beach spawning site located in the intertidal zone, the demersal spawning site in shallow water (from 2 to 5 m), and the demersal spawning site in deeper water (from to 10 to 20 meters). As recommended by Forester et al. (2018), we used a combination of latent factor mixed models (LFMM; Frichot et al. 2013, Frichot & François 2015) and partial redundancy analysis (pRDA; 'rda' function in 'vegan' R package; Lasky et al. 2012, Forester et al. 2018) to detect candidate loci.

We ran LFMMs in the R package LEA. Models were parameterized using the K-value (K = 2) obtained in our previous clustering analyses in LEA (**Fig.3c**). By doing so, we accounted for the genetic variation associated with the haplogroups in our analyses of genotype-environment associations. We ran each model 10 times with 5,000 iterations and a burnin of 2,500. The median of the squared z-scores was used to rank loci and to calculate a genomic inflation factor to evaluate model fit (François et al. 2016).

We used a partial redundancy analysis (pRDA) to detect multi-locus outliers that were associated with spawning habitats after controlling for the haplogroups (see Legendre & Legendre 2012, Forester et al. 2018 for details). Standard deviation of marker scores was multiplied by 3.5 to establish the multi-locus outlier detection p-value at 0.001 (Forester et al. 2018). We retained the outliers that were detected by both LFMM and pRDA to reduce the number of false positives as recommended by Forester et al. (2018) and represented the overlap with a Venn diagram. For each outlier locus associated with spawning habitats, we performed enrichment tests as detailed in the **Supplementary methods** section.

Genotype-environment associations and local adaptation to environmental conditions in beach spawning sites

We investigated the molecular signature of adaptation to local environmental conditions across beach spawning sites within the NWA lineage (i.e., 19 sampling sites). We considered two environmental variables: sea surface temperature and chlorophyll concentration as a proxy of trophic productivity, which may have dramatic effects on embryonic and larval performances (Frank & Leggett 1981a, 1981b, 1982; Leggett et al. 1984). The correlation between temperature and chlorophyll concentration (r = 0.47) was relatively low (for a discussion on multicollinearity in ecological studies, see Dormann et al. 2013). For this reason, these two variables were simultaneously introduced in the pRDA analyses presented below. We determined candidate loci associated with the two variables using a combination of LFMM and pRDA. In LFMM, the variables were introduced one by one in the models to identify the candidate loci associated with each variable. We then used the same procedure described for spawning site analyses for go_enrichment and blasting.

Results

Draft genome assembly and statistics

The resulting draft genome assembly was 490 Mbp with a N50 contig size of 230 Kbp. Running BUSCO with the *Actinopterygii* dataset on the assembly found 4070 complete (89%), 209 fragmented (4.5%), and 305

missing genes (6.5%) out of a total of 4584 genes searched. The genome assembly is available at DOI: 10.6084/m9.figshare.9752558.

Genetic differentiation between glacial lineages

The raw vcf from stacks contained 1,081,533 SNPs spread on 67,305 loci before filtration. After filtration we kept a final set of 25,904 SNPs spread on 25,904 loci (see details in methods). The mapping rate of our GBS data on the draft genome was 74%. The median number of reads per sample was 1,033,000 and the median depth was 21.

A high and heterogeneous level of differentiation was found along the genome for the comparison between lineages (**Fig. S1**). The first and second axes of the PCA explained 8.2% of the genetic variation (**Fig.2A**) and they supported the pattern found with F_{ST} . The ARC lineage was the most divergent among all three lineages. The clustering analysis also supported the existence of three lineages (K = 3) with an absence of admixture between lineages (**Fig.2C**).

Measures of net sequence divergence (d_a) which, unlike F_{ST} , are not affected by variation in the common ancestor, revealed a level of net divergence that was about twice as high between the ARC and the GRE lineages ($d_a = 0.00146$) and between the ARC and NWA lineages ($d_a = 0.00140$) compared to that between the GRE and NWA clades ($d_a = 0.00067$). The same was true for Dxy values with respective value of 0.0032, 0.0041 and 0.0041 between NWA and GRE, ARC and NWA, and ARC and GRE, respectively. Moreover, the mean F_{ST} was 0.2435 between NWA and GRE, 0.3783 between NWA and ARC, and 0.4106 between GRE and ARC (**Fig.2C**), thus revealing a high level of intraspecific genetic differentiation for a marine species (DeWoody & Avise 2005).

Geographical and habitat-dependent genetic structure within lineages

At the intra-lineage level, pairwise F_{ST} values between sampling sites were three orders of magnitude lower than observed between the three lineages (**Fig.2B**). The mean F_{ST} was 0.0033 within the GRE lineage (ranging from 0.0021 to 0.0054) and 0.0044 in the ARC lineage (only two sampling sites in this lineage). Within the NWA lineage, the mean F_{ST} was 0.0010 (ranging from 0 to 0.0090); 91% of the F_{ST} estimates had a *p-value* lower than 0.01 and a 95% CI that did not include 0 (**Table S13**).Within the NWA lineage, the mean F_{ST} among shallow-water spawning sites and among deep-water demersal sites was 0.0019 and 0.0017, respectively, whereas it was 0.0029 among beach spawning sites. The mean F_{ST} between shallow-water demersal sites and beach sites and between deep-water demersal sites and beach sites was 0.0031 and 0.0027, respectively.

We then tested for IBD within the NWA lineage, using neutral loci and loci under selection (detailed methods are given in the **Supplementary methods** section, and **Fig. S12**). Combining BAYESCAN and PCADAPT revealed a total of 12,600 putatively neutral SNPs while each method revealed respectively 20 and 42 SNPs putatively under divergent selection (additional results are given in the **Supplementary results** section). We detected a weak signal of IBD using the two sets of markers (with SNPs under divergent selection: F = 5.38, $R^2 = 0.01$, p = 0.02; with neutral SNPs: F = 3.89, $R^2 = 0.01$, p = 0.05) (**Fig. S12**).

A putative chromosomal rearrangement associated with population within the NWA lineage

The PCA performed for the GRE and ARC lineages did not resolve any genetic structure (**Fig. S4**). By contrast, the PCA revealed pronounced genetic structure within the NWA lineage (**Fig.3A**), with three genetic clusters along PC1 (explaining 0.3% of the variation). The distribution of the clusters did not show any geographic pattern (**Fig.3B**, but see below). A set of 248 SNPs displayed the highest loading along the first axis. Mapping of these 248 SNPs revealed that 85% of them were preferentially located on ancestral chromosomes Chr2 (n = 74 SNPs) and Chr9 (n = 139 SNPs), whereas an average of two SNPs was mapped on the remaining chromosomes. The PCA performed for Chr2 and Chr9 separately showed similar clustering patterns (**Fig. S10** and **S11**). Altogether, these observations raised the hypothesis that such genetic substructure may be due to a chromosomal inversion in Chr2 and Chr9 that could be fused in the capelin; alternatively, it could be two covarying chromosomal inversions or a polymorphic fusion. We thus considered the three clusters as three haplogroups. Based on the PCA, 57% of the individuals were assigned to haplogroup1, 38% to haplogroup2, and 5% to haplogroup3.

SNPs with the highest levels of differentiation between haplogroups were also predominantly located on Chr2 and Chr9 (**Fig.4**). The highest level of genetic differentiation was observed between haplogroups 1 and 3, with a genome-wide F_{ST} of 0.0047 but with a F_{ST} 50 times higher (0.29) when we only considered the SNPs located in the Chr2-Chr9 rearrangement (**Fig. S9**). This was higher than the differentiation observed between each haplogroup and the two other lineages for the same genomic region (F_{ST} haplogroup1 and GRE = 0.055; F_{ST} haplogroup1 and ARC = 0.012; F_{ST} haplogroup3 vs GRE = 0.072; F_{ST} haplogroup3 vs ARC = 0.148). The beta regression model also indicated that the heterozygosity of the SNPs from Chr2 and Chr9 differed among haplogroups (**Fig.3E**; **Table S2**): it was highest in haplogroup2 and similar in haplogroup1 and haplogroup3. The clustering approach also supported the existence of these haplogroups. We detected two genetic groups (K

= 2) across all sampling sites. All individuals of haplogroup1 are contained in one genetic group whereas all the individuals of haplogroup2 and haplogroup3 are admixed with the other genetic group (**Fig.3C**). Moreover, when considering haplogroup1 and haplogroup3 as the homokaryotes and haplogroup2 as the heterokaryote for the putative rearrangement, there was no deviation from Hardy-Weinberg equilibrium at any sampling site (**Table S12**), suggesting Mendelian inheritance of markers found in this rearrangement. Altogether, these observations suggest that haplogroup2 may represent a group of heterozygotes bearing two, perhaps partially non-recombining, versions of haploblocks that exist in the homozygote state in haplogroups 1 and 3.

Haplogroups 1 and 2 were present in 100% of the sampling sites, while haplogroup3 occurred in only 78% of the sites. The frequency of haplogroups varied among sampling sites (**Fig.3B**): the proportion of individuals in haplogroup1 ranged from 35 to 73%, from 24 to 52% for haplogroup2, and from 0 to 21% for haplogroup3. Logistic models indicated that the probability that an individual was assigned to haplogroup1 or haplogroup2 (encompassing 95% of the individuals) was not influenced by its sex (haplogroup1: F = 0.33, p-value = 0.33; haplogroup2: F = 0.69, p-value = 0.41). The probability that an individual was assigned to haplogroup3 was sex-dependent ($\beta_{males} = -0.97 \pm 0.34$, F = 8.73, p-value = 0.003), although this was likely due to the small number of individuals assigned to haplogroup3. Furthermore, linear models indicated that the frequency of the putative rearrangement differed between spawning habitats (**Fig.3D**). The frequency of haplogroups 1 and 2 combined was higher in beach spawning than in demersal sites (both deep-water and shallow-water sites combined) (df = 24, F = 6.74, p = 0.02). However, the rearrangement frequency was not influenced by temperature (df = 18, F = 0.67, p = 0.42) or chlorophyll concentration (df = 18, F = 0.08, p = 0.83). Additional PCA, performed for the three haplogroups separately, suggest the absence of underlying genetic structure in NWA lineage (**Fig. S6**).

Demographic history of divergence

History of divergence among glacial lineages

The divergence history inferred with $\partial a \partial i$ revealed the occurrence of ongoing gene flow between all three lineages. In two inter-lineage comparisons, GRE *vs* NWA lineages and GRE *vs* ARC lineages, the models of secondary contact including both linked selection and variable introgression rate among loci ("SC2N2m") received the highest statistical support (wAIC = 1; **Table S3, Fig.5**). In the comparison of the NWA *vs* ARC lineages, the model of isolation with migration including both linked selection and variable introgression rate among loci ("IM2N2m") received the highest statistical support (wAIC = 1; **Table S3, Fig.5**). In these results were

robust to the confounding effect of the putative chromosomal re-arrangement defined above (**Supplementary methods** section, **Table S3**, and **Fig. S8**).

Parameter estimates revealed that the NWA and ARC lineages diverged approximately 3.8 MyA, [95% CI: 3.6 - 3.8 MyA]. The GRE lineage was inferred to have diverged from the ARC lineage approximately 2.7 MyA, [95% CI: 2.1 - 3.2MyA). The NWA vs GRE comparison indicated a divergence time of approximately 1.8 MyA [95% CI: 0.8 - 2.8 MyA). These split times were older when we removed Chr2 and Chr9 and when we separated by haplogroup, with respective averaged values of 4.3 MyA for the NWA vs ARC split and 2.5 MyA for the NWA vs GRE split (**Table S4**). This suggests a more recent time to the most recent common ancestor (TMRCA) for Chr2 and Chr9. Gene flow was asymmetric and heterogeneous. Indeed, under SC, on average half of the genome displayed a reduced introgression rate, while our inference under IM2N2m indicated that 99% of the genome displayed reduced introgression. The rate of introgression under IM2N2m between the NWA and ARC lineages appears to be an order of magnitude lower than in other comparisons (**Table S4**). Finally, in all comparisons, our inference suggested that approximately half of the genome might be affected by selection at linked sites resulting in a reduced effective population size at such sites (**Table 1**). Our estimates of effective population sizes were high, varying from 220,000 (sd = 7,600) in the ARC lineage under IM2N2m to more than 1,330,000 (sd = 82,000) in the NWA lineage under the same model (**Table 1**; see also **Table S4** for details of parameter estimates when separating by haplogroups).

Candidate loci driving genetic divergence among lineages

Based on the parameters estimated with $\partial a \partial l$, we performed coalescent simulations to compute a neutral envelop of differentiation conditioned on expected heterozygosity. After excluding markers with reduced heterozygosity (He <0.10), a total of 2085, 622, and 330 markers departed from neutrality in the GRE vs ARC, NWA vs ARC and NWA vs GRE comparisons, respectively. These were all randomly distributed across the reconstructed chromosomes (**Table S7**). Of these, 276 were shared outliers between GRE *vs* ARC and NWA *vs* ARC, 130 were shared between NWA *vs* ARC and NWA *vs* GRE, and 114 were shared between GRE *vs* ARC and NWA *vs* GRE. A total of seven outliers were shared in all three pairwise comparisons, such that only a few outliers should be affected by background selection (see Discussion). A total of 118 outliers were fixed (i.e. *F*_{ST} = 1) between the GRE and ARC lineages, no outliers were fixed in the other two pairwise comparisons, and *F*_{ST} values were less pronounced in comparisons involving the NWA clades. Significant blasts (e-value <1e⁻¹⁰ and length >90) were obtained for 688 loci, with a total of 43 putative non-synonymous mutations. These significant hits were also randomly distributed in the genome (**Table S7**) and associated with various biological

processes (**Table S8**). The SNPs associated with the putative chromosomal rearrangement on Chr2 and Chr9 were significantly enriched with outliers when compared to the rest of the genome for the comparison between GRE and NWA (Fisher exact test, p < 0.0001) as well as between ARC and NWA (Fisher test, *p*-value = 0.0006), but not in the comparison between ARC and GRE (Fisher test, *p*-value = 0.7038) as expected given that this rearrangement is absent in these lineages.

Genotype-environment associations in the NWA clade

Outlier loci associated with spawning habitats

In the pRDA, genetic variance was significantly explained by spawning habitat (i.e., beach spawning, shallowwater demersal, and deeper-water demersal sites; overall significance, df = 2, F = 1.58, p < 0.001). We detected 407 and 1,737 outlier SNPs associated with spawning habitat using pRDA (overall significance, df = 2, F =1.12, p < 0.001) and LFMM, respectively (**Fig.6**). A total of 108 SNPs were detected by the two methods; 26% of the SNPs detected with pRDA were also detected using LFMM. Respectively, 5% and 9% of the 108 loci were located in the genomic regions of Chr2 and Chr9 associated with the haplogroups. Outlier loci were in excess in these genomic regions compared to the rest of the genome (see the results of the Fisher test in **Table** 2). Furthermore, 85% and 55% of the outlier SNPs were polymorphic (allelic frequency (AF) higher than 0) in the GRE (mean AF = 0.28) and ARC lineages (mean AF = 0.26), respectively. Fisher tests (**Table 2**) revealed an excess of shared polymorphism within both lineages at outlier loci associated with spawning habitat. We identified 112 transcripts in the 1Kbp window around the outlier loci enriched in various cellular and molecular processes (**Table S9**).

Outlier loci associated with temperature and chlorophyll concentration within beach-spawning sites

In the pRDA, the genetic variance was significantly explained by the two environmental variables (overall significance, df = 2, F = 1.09, p < 0.001). A total of 197 and 1,930 outlier SNPs were associated with temperature based on pRDA and LFMM, respectively (**Fig.6**). A total of 85 SNPs were identified by the two methods; 43% of the SNPs identified with pRDA were also detected using LFMM. Among those 85 SNPs, 7% and 2% respectively occurred within the genomic regions of Chr2 and Chr9 comprised in the haplogroups. Fisher tests indicated the outlier loci were in excess in these regions compared to the rest of the genome (**Table**

2). Furthermore, 76% and 47% of the SNPs were polymorphic in the GRE (mean AF = 0.220) and ARC lineages (mean AF = 0.22), respectively. Fisher tests showed an excess of shared polymorphism for both lineages at these outlier loci (**Table 2**). Within the 1Kbp window around the outlier loci, we detected 77 transcripts that were enriched in cellular and molecular processes (**Table S10**).

We also identified 186 and 2,079 outlier SNPs associated with the chlorophyll concentration with pRDA and LFMM, respectively (**Fig.6**). A total of 88 SNPs were identified using the two approaches; 47% of the SNPs detected with pRDA were also detected using LFMM. Among those 88 SNPs, 2% and 5% were present in the genomic regions Chr2 and Chr9 comprised in the haplogroups. However, outlier loci were not in excess in these genomic regions (**Table 2**). Furthermore, 91% and 55% of the SNPs were polymorphic in the GRE (mean AF = 0.26) and ARC lineages (mean AF = 0.23) respectively. Fisher tests showed an excess of shared polymorphism for both lineages in outlier loci (**Table 2**). We detected 93 transcripts within the 1Kbp window around the outlier loci that were enriched in cellular and molecular processes (**Table S11**).

Discussion

Our results revealed a complex demographic history of divergence among the three capelin glacial lineages. Models of secondary contact including both linked selection and variable introgression rates among loci received highest statistical support. They suggest that the lineages are old, having diverged from 3.8 to 1.8 MyA depending on the pair of clades considered. Using coalescent simulations, we also identified a set of candidate loci driving the genetic divergence among lineages and that might be involved in their reproductive isolation. These observations support the hypothesis that these lineages may in fact represent cryptic species of capelin or at the very least indicate that these lineages are part of an ongoing speciation process. Within lineages, our analyses revealed large N_e and weak genetic divergence among sampling sites, typical of other marine fishes. In the NWA lineage, a likely chromosomal rearrangement probably appeared during the interlineage divergence process resulted in the existence of three haplogroups spread across most spawning sites of the NWA lineage, but showing differences in frequencies between capelin from different spawning habitats. Genotype-environment associations revealed molecular signatures of local adaptation to sea conditions prevailing at spawning sites. Importantly, our study suggested that shared polymorphism among glacial lineages and chromosomal rearrangements occurring in NWA might have broadly contributed to the genetic polymorphisms involved in local adaptation in the presence of high gene flow.

An ongoing speciation process among lineages

Our modelling approach supports the hypotheses that i) the three lineages have diverged in allopatry, ii) gene flow during the secondary contact eroded past differentiation and resulted in heterogenous differentiation along the genome, and iii) that such differentiation is also probably driven by linked selection either in the form of background selection (Charlesworth et al. 1993) or hitchhiking (Smith & Haigh 1974). It was previously shown that neglecting these two effects can lead to bias in demographic inference in terms of model choice, false inference of ongoing gene-flow, or on the contrary, inference of the absence of gene flow while the genome might still be semi-permeable (Cruickshank & Hahn, 2014, Ewing & Jensens, 2016, Souza et al. 2013, Roux et al. 2016). The growing number of studies that have used this approach (e.g., Roux et al. 2013, 2014, 2016, Rougemont al. 2018, Leroy et al. 2019) found increased support for models of secondary contact or have demonstrated difficulty in distinguishing models of secondary contact from models of sympatric divergence with gene flow (i.e. IM models). Moreover, the role of linked selection is increasingly recognized as being implicated in observed patterns of genome-wide divergence Burri 2017, Burri et al. 2015, Wang et al. 2016, Stankowski et al. 2019). Here, our approach does not allow a fine quantitative estimation of linked selection and future studies could be undertaken to address this question.

Overall, our demographic reconstruction suggests a relatively long period of divergence, compatible with a level of divergence more typical of inter-specific rather than intra-specific variation (Hedges et al. 2015, Roux et al. 2016). This divergence has resulted in pronounced contemporary genome-wide differentiation randomly distributed across the genome. Accordingly, coalescent simulations suggested the existence of putative outliers (i.e., departing from the neutral envelop). These outliers were spread randomly among and within chromosomes, as expected under the genic view of speciation (Coyne & Orr 2004, Wu 2001). The divergence process has likely favored the establishment of a barrier to gene flow resulting in restricted gene flow along the genome. To our knowledge, such levels of restricted gene flow between parapatric lineages have rarely been documented in a strictly marine species (Le Moan et al. 2016, Gagnaire et al. 2018), thus raising the hypothesis that these three lineages may actually be cryptic species of capelin occurring in the North Atlantic and Arctic oceans. Further refinement would be required to improve our demographic reconstructions. In particular, three-populations models and the inclusion of ghost populations (Slatkin 2005) may provide a richer picture of the divergence history of capelin lineages. Moreover, further experimental studies should formally test the hypothesis that limited gene flow and the absence of current admixture (based our clustering analyses) among lineages result from pre- or post-zygotic reproductive isolation.

A potential chromosomal rearrangement in the NWA lineage

Our PCA revealed the existence of three haplogroups occurring at variable frequency among sampling sites. This observation suggests the presence of a chromosomal rearrangement characterised by a lower recombination rate compared to the rest of the genome, although PCA is not the most effective tool to detect structural variants (Li & Ralph 2019; but see Ma & Amos 2012). Indeed, the signal left by inversions cannot easily be distinguished from long haplotypes under balancing selection or simply regions of reduced recombination (Lotterhos 2019). Here however, our synteny analysis indicates that the loci included in the putative chromosomal rearrangement are located in Chr2 and Chr9 of the *Esox lucius* genome. The PCA executed for Chr2 and Chr9 separately also displayed the same haplogroup pattern. Assuming that Chr2 and Chr9 are completely fused in the capelin, this suggests the existence of a polymorphic chromosomal inversion in the NWA lineage. Alternatively, it could be two covarying chromosomal inversions or a polymorphic fusion.

Our analyses indicated that such genomic regions, regardless of their true nature, are absent in ARC and GRE lineages. Given the large N_e of these lineages, it is unlikely that polymorphism within this region was lost due to genetic drift during lineage divergence. Instead, we hypothesized that the putative rearrangement appeared in the NWA lineage ~ 2.8 MyA after the split from the ARC lineage but almost concomitantly with the GRE split from the NWA.

Future work based on whole genome resequencing and long-read data will allow clarifying the nature of the putative rearrangement (chromosomal fusion or inversion), and quantifying its exact size, and identify the genes that are located within this region. Furthermore, exciting questions regarding the age of the putative rearrangement, its ancestry, and the possible association with individual phenotype (e.g., morphological, physiological, and life history traits) could be addressed with such additional sequencing data.

High gene flow among spawning sites within lineages

Our study highlighted weak, yet significant genetic variation within NWA lineage. These results are also congruent with many studies on marine organisms (excluding reef species) that reported low genetic variation across large geographic areas (e.g., fishes: Lamichhaney et al. 2012; crustaceans: Benestan et al. 2015; molluscs: Van Wyngaarden et al. 2017; for a review, see Kelley et al. 2016). The weak intra-lineage genetic differentiation within NWA partly results from large historical effective population sizes in the three lineages. Furthermore, the low IBD detected using neutral markers both suggests high dispersal among spawning sites that led to pronounced gene flow.

Beyond weak genetic structure at the scale of the NWA geographic range of distribution, we detected loci under putative divergent selection associated with spawning habitats. As adult capelin were captured at spawning sites during spawning, this pattern could result from habitat match choice (Edelaar et al. 2008). This mechanism implies that individuals disperse towards reproductive sites that maximize their breeding performances, which allows local adaptation despite high gene flow (Edelaar et al. 2008, Jacob et al. 2017). As individual habitat choice does not always match optimal environment conditions (i.e., partial habitat matching) or because a population may contain both specialist and generalist individuals (Jacob et al. 2017), it may result in weak genetic structure due to large effective population size and/or gene flow between the two habitats.

Our results and those previous studies suggest that Capelin could be adapted to spawning site characteristics. The breeding sites of beach-spawning capelin located in the intertidal zone are exposed to weather variation (e.g., temperature, wind) that strongly affects embryonic and larval growth and survival, as well as swimming performance during larval dispersal (Frank & Leggett 1981a, 1981b, 1982; Leggett et al. 1984). Beach spawning sites also display more variable and higher water temperatures than demersal sites (Nakashima & Wheeler 2002), which may sometimes lead to dramatic mortality events before larval hatching (Leggett et al. 1984). Interestingly, a previous study has suggested that embryos from beach-spawning capelin have a faster growth rate than those of demersal capelins (Nakashima & Wheeler 2002), which might indicate local adaptation allowing an accelerated development in the intertidal zone. In parallel, our results indicate that candidate loci potentially under divergent polygenic selection (that is, co-varying markers of small effects detected both by pRDA and LFMM) and associated to spawning habitat are involved in a broad range of biological processes (**Table S9**) driving cell resistance to environmental stress. Taken together, these results raise the hypothesis of habitat matching choice and adaptation to local environmental conditions prevailing at beach and demersal spawning sites. Yet, as the number of demersal sites was limited (6 versus 18 beach sites) in our study due to sampling constraints, we encourage future work to perform additional genotypeenvironment associations to validate the putative candidate outliers detected in our study.

Our study also suggests a similar mechanism of adaptation to the conditions of temperature and trophic productivity experienced in beach spawning sites. We detected two markers under putative divergent selection associated with water temperature and localized in genomic regions (a narrow window of 1 Kbp) that contain genes involved in growth regulation (thermo-dependent in fishes, Pauly 1980) and response to UV radiation (**Table S4**). Overall, these results are in accordance with previous studies on capelin showing that water temperature and trophic productivity strongly affect embryonic and larval growth and survival, as well as the

swimming capacity of larvae during the drift phase following hatching (Frank & Leggett 1981a, 1981b, 1982; Leggett et al. 1984). However, although our findings are consistent with previous GBS studies suggesting environmentally-driven signature of molecular adaptation in marine organisms (e.g., Bradbury et al. 2010, Benestan et al. 2016, Stanley et al. 2018, Xuereb et al. 2018), they suffer from the limitations already reported in RAD-seq analyses, especially the non-exhaustive genome sampling that prevents a full assessment of candidate loci (Lowry et al. 2017) and are limited by the accuracy of genome scan approaches (Pavlidis et al. 2012). Further whole genome resequencing combined with functional validations and experimental work may provide a more exhaustive picture of candidate loci and could help validate our findings.

Contribution of genomic background to local adaptation

Our analyses suggested that shared ancestral polymorphism may play a central role in local adaptation to sea conditions, as predicted by theory (Barrett & Schluter 2008, Hedrick 2013). Indeed, 85% and 55% of the markers putatively under divergent selection and associated with the type of spawning site were polymorphic in the GRE and ARC lineages, respectively. In addition, 76% and 47% of the candidate outliers related to temperature, as well as 91% and 55% of the outliers associated with trophic productivity, were also polymorphic in the GRE and ARC lineages, respectively. Importantly, we consistently found an excess of shared polymorphisms with both lineages in this set of outlier loci. Therefore, these results, combined with the observed level of divergence (at least 2 Mya despite all uncertainties) and low migration rate among lineages, support the hypothesis that these shared variants are likely standing variants inherited from the last common ancestors of these lineages (Lai et al. 2019). Alternatively, they may have originated in a single lineage and then spread among lineages via introgression following the secondary contact (Welch & Jiggins 2014). Lastly, as it is the case for any other studies based reduced representation sequencing data, the SNP outliers are probably neutral hitchhiker loci linked to the direct target of selection (Bierne et al. 2011). To distinguish among the different hypotheses above, whole genome data, combined with methods to distinguish among soft versus hard selective sweep, would be needed (e.g. Kern & Schrider, 2018). Further use of coalescent methods would also allow to directly infer the age of the variants relative to the most recent common ancestor of the lineage (Albers & McVean 2020).

Our analyses also indicated that a non-negligible proportion of outlier markers associated with the type of spawning site (8%), temperature (7%), and trophic productivity (2%) is located in the putative chromosomal rearrangement. For both the type of spawning site and temperature, outlier loci were in excess in the genomic regions contained in the chromosomal rearrangement compared to the rest of the genome. These results thus

appear to be consistent with previous work suggesting that chromosomal polymorphism may facilitate thermal adaptation despite high gene flow in marine fishes (Barth et al. 2017, Wellband et al. 2019). However, future work based on whole genome resequencing data should be conducted to more accurately assess the contribution of the putative rearrangement to local adaptation, especially by identifying the type of rearrangement detected in our study and as well as its characteristics (i.e., size and content), and its effects on capelin phenotype and performance.

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Author contributions

H.C. and Q.R. made the statistical analyses and wrote the paper. M.L., C.M., E.N., and Y.D. contributed to the bioinformatics and statistical analyses. L.B., P.S., M.CA., and M.CL. initiated the project, and L.B. conceptualized and coordinated the work. T.J. collected the DNA samples in the GRE lineage. The generation of the draft capelin genome assembly was initiated and coordinated by S.J, and samples provided by K.P. The

whole genome sequencing DNA extraction was performed by S.N.K.H and construction of the draft genome assembly was done by O.K.T and S.N.K.H.

Data accessibility

Genome assembly and associated reads are available on NCBI under the accession number PRJEB38139. Raw sequencing data for GBS libraries are available under the accession number PRJNA631144. Filtered *vcf* file and environmental data are available on Dryad (https://doi.org/10.5061/dryad.hx3ffbgbp).

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Table 1. Parameter estimates under the best demographic models for each of the three lineages: Ne1 and Ne2, effective population size of the compared pair; m1 \leftarrow 2 and m2 \leftarrow 1, migration from population 2 to population 1 and migration from population 1 to population 2. me12 and me21, effective migration rate estimated in the most differentiated regions of the genome, Tsplit: Time of split of the ancestral population in the two daughter species; Tsc, duration of secondary contact; P, proportion of the genome freely exchanged (1-**P** provides the proportion of the genome non-neutrally exchanged); Q, proportion of the genome with a reduced Ne due to selection at linked sites (1-Q provides the proportion of the genome neutrally exchanged); hrf = Hill-Robertson factor representing the reduction of Ne in the region (1-Q) with reduced Ne. Values in brackets provide confidence intervals around the estimated parameters.

	NWA vs GRE	GRE vs ARC	NWA vs ARC		
best	SC2N2m	SC2N2m	IM2N2m		
Nref	131000 [11,000 – 250,000]	208, 000 [72,760 – 343,000]	155, 000 [102,000 – 208, 000]		
Ne1	828, 000 [397,000 – 1,259,000]	592,000 [521,000 – 662,000]	1,329,000 [1,166, 000 – 1,491,700		
Ne2	311,000 [162,000 - 460,000]	348, 000 [239,590 – 456, 000]	226,000 [211,000 - 241,000]		
m1	0,000001 [0-0,000003]	0,0000003 [0 - 0,000001]	0,0000006 [0 - 0,000002]		
m2	0,0000016 [0-0,000003]	0,0000002 [0 - 0,000003]	0,0000001 [0 - 0,0000003]		
me1	0,000009 [0–0,00004]	0,000002 [0 - 0,00003]	0,0000006 [0 - 0,000004]		
me2	0,00002 [0–0,00008]	0,000003 [0 - 0,00009]	0,000001 [0 - 0,000006]		
Tsplit	1,785,000 [811,000 – 2,758,000]	2,700,000 [2,119,900 – 3, 280,000]	3,760,000 [3,697,000 – 3,823,000		
Tsc	898,000 [0 - 1,807,000]	775,000 [732,000 – 818,000]	ΝΑ		
hrf	0,10 [0.03 - 0,17]	0.24 [0.23-0.25]	0.12 [0.00 – 0.24]		
Р	0.53	0.57	0.01		
Q	0.46	0.57	0.5		

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Table 2. Enrichment of SNP outliers in the potential chromosomal rearrangement in chromosome Chr2 and Chr9 and in shared polymorphisms with ARC and GRE lineages. Also presented are the results of Fisher tests (odds ratio and p-values).

				Type of spawning		
	Chlorophyll		Temperature		site	
Enrichment	Ratio	р	Ratio	р	Ratio	р
Chromosomal rearrangement	0.35	0.16	0.10	< 0.0001	0.09	< 0.0001
Shared polymorphism with ARC	0.31	< 0.0001	0.42	0.0001	0.31	< 0.0001
Shared polymorphism with GRE	0.10	< 0.0001	0.19	< 0.0001	0.18	< 0.0001
Shared polymorphism with GRE and ARC	0.12	< 0.0001	0.36	< 0.0001	0.22	< 0.0001

Figure caption

Fig.1. Map of the study area. A total of 31 sampling locations were considered in the distribution range of three glacial lineages (NWA, ARC and GRE) of capelin (*Mallotus villosus*). In the NWA lineage distribution range, we sampled individuals according to three types of spawning sites: beach-spawning sites, shallow-water demersal sites, and deep-water demersal sites. In the ARC lineage, the two sampling sites are very close (few kilometers) and cannot be distinguished on the map.

Fig.2. Genetic structure and differentiation among the three lineages (NWA, GRE, and ARC) of capelin (*Mallotus villosus*). (A) Principal Component Analysis (axes 1 and 2) showing the genetic variation among the lineages. (B) Heatmap of F_{ST} between and within three lineages. (C) Clustering analysis performed using LEA R package.

Fig.3. Spatial variation of the putative chromosomal rearrangement (involving chromosomes 2 and 9) within the NWA lineage of capelin (*Mallotus villosus*). (A) Principal Component Analysis (axes 1 and 2) showing the three clusters of individuals corresponding to the three haplogroups. (B) Relative proportion of the three haplogroups among the 25 sampled spawning sites throughout the distribution area of the NWA lineage. (C) Ancestry proportion ordered by haplogroups from clustering analyses. (D) Frequency of the rearrangement in the demersal (deep-water sites + shallow-water sites) and beach spawning sites; we provide the results of an ANOVA testing the effect of the type of spawning site on the rearrangement frequency. (E) Heterozygosity analysis focused on the contigs contained in the potential chromosomal rearrangement in chromosomes Chr2 and Chr9. We provide the results of the beta regression model: intercept for haplogroup1, and the slope coefficients for haplogroup2 and haplogroup3, and their associated standard error and statistical test (z statistic and p-value).

Fig.4. Manhattan plots of the F_{ST} values between the three haplogroups within the NWA lineage of capelin (*Mallotus villosus*). The external circle shows the F_{ST} between haplogroups 1 and 2. The medium circle shows the F_{ST} between haplogroups 2 and 3. The central circle shows the F_{ST} between haplogroups 2 and 3. Capelin contigs were placed into 24 ancestral chromosomes based on synteny with 4 related species. Contigs that were not assigned to a chromosome were aggregated in the group ChrU. As the order of the contigs within the chromosomes was not conserved and was not always consistent among genome comparisons, contigs were

ranked according to their size. SNPs having F_{ST} less than 0.25 are shown in grey, SNPs with F_{ST} ranging from 0.25 to 0.50 are shown in yellow, and SNPs with F_{ST} higher than 0.50 are shown in red.

Fig.5. Demographic divergence among lineages of capelin (*Mallotus villosus*): simplified representation of the best demographic model (left panel), observed and best fitted jSFS obtained from $\partial a \partial i$ to unravel the demographic histories of the three lineages (middle panels) and residuals of the fitted models (right panels). The best model was the SC2N2m (model of secondary contact including both linked selection and variable introgression rate among loci) in the comparison between ARC *vs* GRE and GRE *vs* NWA, whereas the IM2N2m model (model of isolation with migration including both linked selection and variable introgression rate among loci) best fits the data in the ARC *vs* NWA comparison. Left panel: N_1 , N_2 and N_A represent the respective effective population size of the two descending population and of the ancestral lineage. The black and red arrows (m_{ij}) represents the migration rate from population j into population i. Tsplit = splitting time. Tsc = Time of secondary contact.

Fig.6. Molecular adaptation to local environmental conditions in the capelin (*Mallotus villosus*). (A) SNP outliers associated with temperature: the manhattan plot show p-values from LFMM analyses for SNPs detected as outliers by both pRDA and LFMM (shown in red) and the others (shown in grey). Venn diagrams show the outlier loci detected with either LFMMs or pRDA, or both – the percentage corresponds to the proportion of outliers detected by each method and the two methods combined. The map shows spatial variation of temperature. (B) SNP outliers associated with chlorophyll concentration. (C) SNP outliers associated with spawning habitats. Note that contig were placed on chromosome but the order of contig themselves may not be accurate. The SNPs positions here should not be interpreted as physical position on the chromosomes.



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