

Seascape genomics reveals population isolation in the reef-building honeycomb worm, *Sabellaria alveolata* (L.)

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Abstract

Background Under the threat of climate change populations can disperse, acclimatise or evolve in order to avoid fitness loss. In light of this, it is important to understand neutral gene flow patterns as a measure of dispersal potential, but also adaptive genetic variation as a measure of evolutionary potential. In order to assess genetic variation and how this relates to environment in the honeycomb worm (*Sabellaria alveolata* (L.)), a reef-building polychaete that supports high biodiversity, we carried out RAD sequencing using individuals from along its complete latitudinal range. Patterns of neutral population genetic structure were compared to larval dispersal as predicted by ocean circulation modelling, and outlier analyses and genotype-environment associations tests were used to attempt to identify loci under selection in relation to local temperature data. **Results** We genotyped 482 filtered SNPs, from 68 individuals across nine sites, 27 of which were identified as outliers using BAYESCAN and ARLEQUIN. All outlier loci were potentially under balancing selection, despite previous evidence of local adaptation in the system. Limited gene flow was observed among reef-sites ($F_{ST} = 0.28 \pm 0.10$), in line with the low dispersal potential identified by the larval dispersal models. The North Atlantic reef emerged as an isolated population and this was linked to high local larval retention and the effect of the North Atlantic Current on dispersal. **Conclusions** As an isolated population, with limited potential for natural genetic or demographic augmentation from other reefs, the North Atlantic site warrants conservation attention in order to preserve not only this species, but above all the crucial functional ecological roles that are associated with their bioconstructions. Our study highlights the utility of using seascape genomics to identify populations of conservation concern.

Background

Conservation strategies in the context of climate change rest on the premise that populations can show three responses to avoid fitness loss: evade, evolve or acclimatise [1]. Characterisation of genetic diversity can shed light on these responses [2], by indicating dispersal and gene flow using neutral genetic markers (evolution), or adaptation using adaptive genetic variation, which inform us about the potential for an organism to evolve in altered conditions [3–5]. Separating neutral and adaptive causes of phenotypic differentiation, or lack thereof in a heterogeneous environment (*sensu* countergradient variation; [6]), can require logistically difficult experiments [7]. However, advances in genomic technologies and analyses have opened up new avenues for exploring genome-wide (neutral) versus locus-specific (adaptive) genetic variation between populations, allowing us to compare patterns of adaptive genetic diversity with environmental variation to infer agents of selection [8, 9]. These methods have already been successful in contrasting neutral and non-neutral genetic variation in a range of taxa including fish [5, 8, 10, 11], insects [4], and marine invertebrates [12, 13] and represent an important step in interpreting genetic diversity in relation to environment.

Despite genomic advances allowing investigations of genetic diversity in non-model species, which is particularly important for species of conservation concern, understanding of gene flow and local adaptation in marine environments had, until recently, lagged behind that for terrestrial species [14–16]. Recent advances in seascape genomics have facilitated consideration of the effect of landscape on neutral and adaptive genetic variation in marine environments, taking into account ocean circulation patterns and environmental variation to explain observed population structuring [8, 17]. For instance, Benestan *et al.* (2016)[17] found that ocean currents as well as geographic distance were key to explaining observed patterns of neutral genetic variation in the American lobster (*Homarus americanus*), whereas sea surface temperature acted as an agent for natural selection. Ocean circulation modelling for understanding dispersing larval stages has emerged as an important tool to explain neutral population structure where isolation-by-distance measures alone are uninformative [18–20].

Intertidal communities have been proposed as particularly suitable for studying the impacts of climate change, as many species already exist at the limit of their thermal tolerance ranges [21, 22]. Ecosystem engineers provide and modify habitat for other species and are thus disproportionately important to diversity, ecological functioning, and conservation

in a changing climate [1, 14, 23–25]. Therefore, understanding how littoral ecosystem engineers have adapted to the local environment they experience is key to understanding not only their future survival in a changing climate, but also the potential survival of the intertidal ecosystems as a whole.

The honeycomb worm (*Sabellaria alveolata* (L.)), is an ecosystem engineer that constructs biogenic reefs along temperate coastlines of North Africa and Europe [26]. Each individual constructs a tube from sand, which is glued together with biomineralised cement [27, 28]. Individuals are gregarious and tubes aggregate to form large bioconstructions or reefs, which can stretch over several kilometres [29–31]. These reefs provide a large diversity of microhabitats for other species and alter wave action, water circulation, erosion and sedimentation patterns [32], and phytoplankton concentration [33] and thus modify the availability of resources, leading to a high associated biodiversity compared to surrounding areas [34, 35]. Biogenic reefs, including those of *Sabellaria*, are listed under the EC Habitats Directive (92/43/EEC) Annex 1 and are thus of conservation interest [36]. Despite this, little is known about neutral genetic structure (but see [37, 38]) and to our knowledge nothing is known about adaptive genetic structure of *S. alveolata*. Neutral population structuring is expected because (i) adults are sessile, and (ii) during the dispersive planktotrophic larval stage of this species, individuals have been shown to demonstrate positive taxis towards the cement of sand grains and bioclasts of *S. alveolata* tubes [39] leading to site philopatry [40]. However, modelling of larval dispersal is needed to fully understanding how the environment influences gene flow within marine systems [17].

We have previously demonstrated that *S. alveolata* show local adaptation to temperature in terms of their membrane lipid composition [41] but the genomic basis of this adaptation is not known. Therefore, as their distributional range extends along a latitudinal thermal gradient from Scotland to Morocco, *S. alveolata* provide an excellent opportunity to investigate if/how adaptive population structure is related to temperature, and assess the role of neutral processes in gene flow.

In this study, we explore the extent of neutral and adaptive genetic variation among reef sites of the honeycomb worm, *S. alveolata*, and the relationship with the environment. We test: 1) if gene flow is restricted among *S. alveolata* reef sites and whether population structure can be predicted by ocean circulation modelling; and 2) whether *S. alveolata* show evidence of local adaptation and if this relates to local thermal conditions.

Results

SNP detection and summary statistics

Due to high variability in sequence coverage between samples (37,000 - 1,400,000 reads per sample), only 68 samples across the nine sites were retained for analysis (Table 1). Of these, 50 samples were from sites where temperature data was also available (see below; Figure 1). In total, 439,598 SNPs were identified across 273,397 RADtags with an average coverage depth of 14.85x per sample. Due to low read coverage overall, only 506 variable SNPs were identified, 482 of which were from independent RADtags and could be used for population genetic analyses. Marker independence across sites was confirmed with no loci flagged as showing linkage disequilibrium. Although some loci showed significant deviation from Hardy-Weinberg, the loci involved differed across sites and were thus included in further analyses [42].

All sites showed reasonable numbers of alleles per locus, nucleotide diversity and expected heterozygosity (N alleles = 2.44 ± 0.13 , $\theta = 0.26 \pm 0.05$, $H_e = 0.52 \pm 0.04$). Observed heterozygosity was low across all sites ($H_o = 0.11 \pm 0.02$) but was significantly lower than H_e only in the Balearic Sea site, which had a low sample size (Table 1). Most molecular variance was found within rather than between sites (within = 71.42%, between = 28.58%, $P < 0.01$) and within rather than between the genetic clusters identified by STRUCTURE (see below) (within = 84.08%, between = 15.92%, $P < 0.01$).

From the total number of SNPs, ARLEQUIN identified 191 as outliers (39.63%) and BAYESCAN identified 28 as outliers (Figure 2; 5.81%). Twenty seven of the 28 outlier SNPs identified by BAYESCAN as outliers were also identified by ARLEQUIN. Therefore, the 27 outlier SNPs (5.60%) were removed from the data set and 455 (94.40%) were used in the neutral genetic analyses.

Population structure

Pairwise F_{ST} values for neutral loci had an average of 0.28 ± 0.10 across sites, with 25 out of the 36 comparisons showing significance following Bonferroni correction, demonstrating a high degree of population substructure within the system as a whole (Table 2). In particular, the North Atlantic, Tyrrhenian Sea and Bay of Biscay sites were significantly differentiated from all other sites (Average $F_{ST} = 0.42 \pm 0.10$; 0.32 ± 0.12 and 0.25 ± 0.05 , respectively; Table 2), suggesting very low gene flow to and from these sites. Using the Bayesian cluster analysis in STRUCTURE, ΔK showed a clear peak at $K = 2$, suggesting two genetic clusters within the nine sites studied (Figure 3a). Likelihood probability profiles also showed that $K = 2$ had the highest likelihood with low variance. The barplot in STRUCTURE for $K = 2$ identified that individuals from the North Atlantic site formed a separate population from all the other sites combined (Figure 3b). This result did not vary regardless of whether sampling location was included as a prior and there was no evidence of hierarchical clustering ($K = 1$ when North Atlantic individuals were removed). F_{ST} based isolation by straight-line geographic distance and shortest ocean distance (Table 2) were not significant ($r = 0.14$, $P = 0.27$ and $r = 0.08$, $P = 0.27$, respectively).

Maps of larval densities from dispersal simulations show the range of potential dispersal pathways for a single generation released from each sample location: from the most conservative (one week of 6 hourly releases, 5 weeks maximum planktonic larval duration (PLD), Figure 4a), to the most optimistic (daily releases all year round, 12 weeks PLD, Figure 4b). The most optimistic dispersal scenarios are suggestive of isolation among sites (Additional file 1). Model simulations suggest isolation among the North Atlantic, northern Irish Sea, and southern Irish Sea sites (Additional file 1), which is reinforced by the F_{ST} distances (Table 2). Under the conservative scenario estimates, no larvae were exchanged among any sites in a single generation (Figure 4a). Under the most optimistic scenario, high densities of larvae were retained close to natal release sites (average within 133km, maximum 230km) and very few larvae exchanged among release locations (Additional file 1). Potential connectivity was only identified between the northern Irish Sea and southern Irish Sea sites (as per [43]) but $<0.1\%$ of the larval population made this transition. Dispersal distances away from source were shortest from the North Atlantic release site on the west coast of Ireland; this site exhibited the highest proportion of larval retention (79% self-recruited) and very few larvae reached nearby reefs, and none reached other release sites (Additional file 1). Dispersal resistance, as identified using ocean circulation modelling, did not show significant correlation with genetic distance, in terms of F_{ST} , for any of the PLD scenarios (5 weeks: $r = -0.02$, $P = 0.59$; 8 weeks: $r = -0.01$, $P = 0.59$; 10 weeks: $r = -0.006$, $P = 0.57$; 12 weeks: $r = 0.00$, $P = 0.57$).

Loci under selection

Both ARLEQUIN and BAYESCAN analyses suggested that the 27 loci identified as outliers were under balancing selection (Figure 2). Temperature data could not be collected from three of the study sites due to loss of equipment during the data collection period (Balearic Sea, Tyrrhenian Sea and North Africa). Temperature data was thus available for six of the study sites (Figure 1) and only these study sites were included in the environmental association analyses. Of the temperature parameters collected from six of the sites (Table 3), no environmental associations were found with any loci.

Discussion

Population structure

Population genetic analysis using genome-wide SNPs identified from nine *S. alveolata* reef sites spanning the latitudinal and longitudinal range of the species revealed low gene flow between sites, equating to low effective dispersal, with 25 out of 36 pairwise F_{ST} values showing significant differentiation and an average pairwise F_{ST} of 0.28 ± 0.10 across the system. The STRUCTURE analysis revealed the North Atlantic site to be a separate population from all the others ($F_{ST} = 0.42 \pm 0.10$). Genetic divergence in terms of F_{ST} was not significantly correlated with either straight-line distance or shortest ocean distance. Presence of low gene flow between reef sites is in line with a growing number of studies that have identified population structuring in marine species which, as with *S. alveolata* in our study, have a dispersing larval stage and no hard geographical barriers to dispersal [17, 18, 44–46]. For instance, Benestan *et al.* (2015) [46] identified strong neutral genetic structure (average pairwise $F_{ST} = 0.0019$) in the American lobster (*Homarus americanus*) along a sea surface temperature gradient using RAD genotyping despite a dispersing planktonic larval stage with a duration of 4–6 weeks. Similarly, the bat star (*Patiria miniata*) showed genetic population structure along a latitudinal gradient (average pairwise $F_{ST} = 0.061$) despite a longer larval dispersal period of 6–10 weeks [45], which is in line with the estimated larval duration of *S. alveolata* at 5–12 weeks, dependant on temperature and food availability [40, 47]. Indeed, a meta-analysis revealed that mean pelagic larval duration was not a good predictor of gene flow [48] and our findings support this interpretation.

Three sites were significantly differentiated from all other sites: North Atlantic (average pairwise $F_{ST} = 0.42 \pm 0.10$ and STRUCTURE), Tyrrhenian Sea (average pairwise $F_{ST} = 0.32 \pm 0.12$) and Bay of Biscay (average pairwise $F_{ST} = 0.25 \pm 0.05$). The northern Irish Sea, southern Irish Sea and English Channel sites were also all significantly differentiated from each other (Table 2). The ocean circulation modelling predicted high isolation (zero to low larval density; Additional file 1) among all release sites, even in the most optimistic (dispersive) scenario. Therefore, the F_{ST} results are supported, at least in part, by patterns of ocean circulation influencing passive larval dispersal along the Atlantic and Mediterranean coastlines of Europe and North Africa.

High population sub-structuring could be partially explained by two behavioural barriers to connectivity: first, spawning in polychaetes is triggered by the action of waves, especially during spring tides, which tends to push larvae toward the coast rather than offshore [49] and second, *S. alveolata* larvae act to reduce dispersal out of the bay where they were spawned by moving higher in the water column with an incoming- and lower with an outgoing-tide [40]. This ‘tidal stream transport’, has been suggested as a possible mechanism for facilitating/restricting advective transport in a number of taxa (e.g. [50, 51]). Yet despite this, relatively high levels of genetic diversity were observed across the sites ($H_e = 0.52 \pm 0.04$). Observed heterozygosity was very low at all sites and could be due to inbreeding [52], as a result of the site isolation predicted by the ocean circulation modelling and due to the reproductive strategy of gregarious colonial species, as juveniles settle together in patches and spawning of one individual triggers the spawning of those immediately adjacent [39]. Heterozygote deficiency has also been recorded in marine invertebrates as a consequence of the Wahlund effect, due to the coexistence of genetically distinct cohorts within a sampling site [18]. However, low heterozygosity estimations could also be a consequence of poor genome coverage and low sample size at some sites [53, 54]. Further research is needed to assess whether presence of null alleles is genome-wide, as evidence of inbreeding, or restricted to particular loci, as evidence of allelic dropout due to low coverage, in order to assess whether the species is at inbreeding risk.

Identifying the relative contribution of gene flow, genetic drift and natural selection to population structure is difficult in marine invertebrates due to their fluctuating population sizes [9]. This is particularly difficult in *S. alveolata*, as in-depth local ecological knowledge, such as population size and breeding behaviour, is lacking in the majority of sites (but see [31, 55–57]). That said, the identification by STRUCTURE of only the North Atlantic site as a separate population, with

all other sites forming a single population, supports the idea that low levels of gene flow are still maintaining genetic diversity within the system as a whole. Such wide-ranging genetic connectivity has also been observed in other polychaete species [58]. This is potentially a positive sign for survival of the species in a changing climate, as the maintenance of genetic diversity is key to facilitating rapid evolution when environmental conditions change [59]. However, *S. alveolata* reduce their larval duration in line with increasing temperatures [40] and our models show that shorter larval duration leads to reduced larval dispersal. Therefore, lower connectivity, and thus gene flow, is predicted for *S. alveolata* in a changing climate.

Despite some evidence of low levels of gene flow between the majority of the study sites within the system, both F_{ST} values and STRUCTURE analysis revealed isolation of the North Atlantic site and identified this reef as a separate population to all others. There are two possible processes that could, separately or in synergy, be causing the observed pattern of population structure: historical isolation and contemporary patterns of gene flow. During the last glacial maximum, geomorphology and fossil evidence suggests that southwest Ireland was partially unglaciated [60] and genetic data supports the presence of a glacial refugium in this area for both terrestrial [61, 62] and marine species [63–65]. In particular, Jolly *et al.* (2006) [63] found that two coastal polychaete worms (*Pectinaria koreni* and *Owenia fusiformis*) showed a private Irish Sea haplotype linking two ancestral haplotypes and they suggest this could have evolved in a small ice-free area along the southwest coast of Ireland. Therefore, one potential explanation is that this population was isolated in a different glacial refugium to the rest of the sites, but further study is needed to test this hypothesis.

The second hypothesis, that contemporary gene flow is very low between the North Atlantic and other *S. alveolata* sites, is supported by predictions of larval dispersal as seen in the ocean circulation modelling. In even the most optimistic scenario for larval dispersal, the North Atlantic site was not predicted to have interchange of individuals with any of the other sampled sites and had by far the highest predicted proportion of larval retention (79% of released individuals were retained compared to an average of $20 \pm 13\%$ for all other sites). This can be explained by the hydrodynamic modelling in terms of current patterns within Galway Bay limiting dispersal. The presence of the North Atlantic Current, which moves eastwardly towards Ireland and then continues northwards, is also an isolating factor for the North Atlantic reef. Any larvae that do move beyond their spawning reef at the North Atlantic site are drawn northwards, beyond the current northern range limit for the species (the Solway Firth, Scotland [41]), and there are no reef sites along the north coast of Northern Ireland to act as stepping stone populations (Firth, unpublished). This oceanic barrier is also a likely cause of the observed genetic divergence between the northern/southern Irish Sea sites and the English Channel, and the reason that the dispersing larvae from more southerly sites do not reach the North Atlantic site. Although larval dispersal of many other marine species has been found to be mediated by the North Atlantic Current (e.g. [66, 67]), isolation due to the North Atlantic Current has not previously been reported, likely because it depends upon species-specific occurrence range. Therefore, reduced contemporary gene flow is a likely cause of the observed population isolation of the North Atlantic *S. alveolata* reef. These findings highlight the North Atlantic reef site as at risk if population size reductions occur as recruitment and genetic augmentation is unlikely from elsewhere within the species' range. Therefore, conservation management is needed to ensure that population size does not decrease at this site.

Despite the importance of ocean circulation to larval dispersal in general [18, 19, 68] and to *S. alveolata* in particular, as identified by our data in regards to the isolated North Atlantic site, our ocean circulation models of larval dispersal did not show a significant correlation with observed genetic distance between sites. This is surprising as both F_{ST} values and larval dispersal models suggest low connectivity between sites. This lack of correlation between our larval dispersal modelling and genetic distance values is in contrast to studies on a number of species with a dispersing larval stage including the Mediterranean shore crab (*Carcinus aestuarii*) [19], the bat star (*P. miniata*) [45] and the American lobster (*H. americanus*) [17]; these studies found genetic structure was directly related to ocean currents or to estimates of

potential larval connectivity obtained with coupled physical-biological models. However, these studies were conducted over a smaller geographical area than our study. As per our study, Jorde *et al.* (2015) [69] found that when looking at large-scale differentiation patterns in the north Atlantic, geographic distance and larval drift alone explained only a minor portion (2.5–4.7%) of genetic isolation in the northern shrimp (*Pandalus borealis*). Galindo *et al.* (2010) [44] modified a biophysical model for Monterey Bay in California to simulate dispersal of the acorn barnacle (*Balanus glandula*) but it also did not match an observed genetic cline in the species. They discovered that their model fit was improved by including natural selection, larval retention, and input values from an additional source population [44].

There are several factors likely to reduce the similarity between modelled larval dispersal and observed gene flow. Gene flow is representative of multigenerational mixing, while the model used in this study is representative of the dispersal patterns of only one generation. It is possible to simulate multigenerational gene flow based on dispersal models [45, 70, 71] but this would be confounded by the lack of known sites that could be used as “stepping stone” sites, which may facilitate dispersive spread over multiple generations. Such sites are likely to exist but are currently undocumented. These undiscovered sites represent missing source populations within our model and may well be a cause of the observed mismatch between predicted larval dispersal and observed gene flow. The distribution and occurrence of *S. alveolata* reefs are not well documented, particularly in the southern range of the species (Firth, unpublished), and the prediction of our model that genetic differentiation between our sample sites would be high could be due to the absence of intermediate reef sites within the model that would create higher levels of admixture within the system as a whole [9, 72]. This is further complicated by the fact that reefs can vary temporally in their presence at a site [56]. Therefore, increased knowledge of the location of *S. alveolata* reefs and ongoing monitoring of reef sites is required to generate a more clear picture of the role of connectivity in meta-population maintenance in this species and to further inform the ocean circulation model. Our study highlights the importance of comparing bio-physical models with observed population structure of a species in order to create accurate dispersal predictions.

Loci under selection

Twenty-seven SNPs were identified as outlier loci, all of which were putatively under balancing selection. Identification of a higher number of loci potentially under balancing rather than divergent selection is common in genome scan studies, including those on marine invertebrates [12, 46, 73–75]. Our finding that 5.6% of SNPs were potentially under balancing selection is reflective of a growing body of evidence that suggests that balancing selection, which acts to preserve polymorphism, is more important in the genome than previously considered [12, 46, 76]. In their study of the sea anemone, *Nematostella vectensis*, covering a broad geographical range, Reitzel *et al.* (2013) [12] identified 37 polymorphic sites inferred to be under balancing selection, but none under divergent selection, as with our study. However, further evidence in the form of elevated polymorphism, reduced differentiation, and shifts towards intermediate allele frequencies are needed to confirm that these loci are indeed under balancing selection [52].

The lack of identification of loci under divergent selection using either outlier analyses or environmental association tests is surprising given that local adaptation to temperature is known to be present in this system [41]. We previously reported that *S. alveolata* individuals from sites along the latitudinal range of this species (all of which are included in this study) showed different responses to thermal regime changes in terms of their membrane lipid composition dependant on their site of origin [41]. This is likely a reflection of poor genome coverage leading to a low number of genotyped loci (as per [12]). Further studies are needed to search for adaptive loci and their functions in this system.

Conclusions

Modelling of larval dispersal predicted low gene flow between sites for *S. alveolata* across their range, which was in part supported by the genomic data. In particular, population genetic analyses identified the North Atlantic reef is an isolated

population, due to the effect on dispersal of the North Atlantic Current. These findings have implications for the management of *S. alveolata* engineered habitats in a changing climate and have highlighted the North Atlantic reef as at increased risk and in need of direct conservation action in order to preserve not only this species, but above all the crucial functional ecological roles (biodiversity hotspots, coastal protection, carbonate traps, *etc.*) that are associated with these bioconstructions. Our study highlights the utility of using seascape genomics to identify populations of conservation concern.

Methods

Sampling and molecular methods

In total, 180 mature individual worms were haphazardly collected from nine sites (20 per site) across the global geographic range between August 2013 and March 2014, from North Africa at the species' southern boundary, along the Atlantic Coast, to the Northern Irish Sea in the north, as well as from two sites on the Mediterranean coast (Figure 1). Individuals were placed immediately in RNA-later (Qiagen Inc., Crawley), maintained at room temperature for between 48 hours and one week and then stored at -20°C until DNA extraction.

Genomic DNA was extracted using chaotropic "Chaos" buffer (4M guanidine thiocyanate; 0.5% N-lauroyl sarcosine; 25 μM tris(hydroxymethyl)aminomethane (Tris), pH 8.0; 0.1M 2-mercaptoethanol, 0.5% N-lauroyl sarcosine) (modified from Fukami *et al.* 2004). Worms were homogenised within the buffer at room temperature for at least 48 hours before adding an equal volume of phenol extraction buffer (PEB) and double the volume of phenol:chloroform:isoamyl alcohol (25:24:1). From here the protocol followed a phenol-chloroform extraction with isopropanol precipitation [78]. An additional ammonium acetate (3M) precipitation step was added to further purify DNA. DNA was resuspended in sterile distilled H_2O , quality assessed using a NanoDrop 8000 spectrophotometer (Thermo Scientific, Wilmington, DE), visualised on 1% agarose gels, and quantified using the Quant-IT PicoGreen dsDNA Assay Kit (Life Technologies, Carlsbad, California). Single-end Restriction-site Associated DNA (RAD) libraries were prepared by digesting DNA from each individual using the EcoRI and MseI restriction enzymes. Samples were individually barcoded and sequenced (Ecogenics, Zurich-Schlieren, Switzerland) using an Illumina NextSeq 500 (Illumina, San Diego, California).

SNP detection and summary statistics

All analyses were carried out on the Analyses and Bioinformatics for Marine Science (ABiMS) Galaxy Platform at the Station Biologique de Roscoff (galaxy.sb-roscoff.fr; [79]). Samples with mean Phred scores of <25 or read coverage of $<150,000$ were removed from the dataset and sequences were trimmed to 60bp length using Trimmomatic version 0.36.4 [80]. STACKS version 1.46.0 [81] was used to *de novo* map the sequences and call SNPs by sequentially running "ustacks", "cstacks" and "sstacks" using the parameters suggested to be optimal for population genetic inference by [82] namely, allowing a minimum depth of coverage of three (ustacks: m), a maximum of two mismatches between reads for a single individual (ustacks: M), and four mismatches between primary and secondary reads (ustacks: N). These settings have been found to increase the number of loci and reduce the SNP and allele error rates within a dataset [82]. The number of mismatches allowed between loci was set at three (cstacks: n; [83]). The removal or break up of highly repetitive RAD tags was permitted within the program (ustacks: deleverage).

The programme *populations* from the STACKS suite, was used to process the SNP data [81]. RADtags with at least 10x depth of coverage were retained to ensure accuracy of heterozygous SNP calls (populations: p; [46]). A locus had to be present in 70% of the populations to be included in the analysis, which allows for mutations in restriction sites that may cause loci to dropout in certain lineages [46, 84] and the minor allele frequency within populations (populations: min_maf) was set at > 0.01 [83]. In order to maximise SNP discovery, the percentage of individuals required within a

population to process a locus was set at 50% (populations: r). Only a single SNP per RAD tag was retained for subsequent analyses to avoid problems of non-independence between markers [10]. The STRUCTURE file output by *populations* was transformed in PGD Spider version 2.1.0.3 [85] for use in all further analyses. Each population and locus was tested for deviation from Hardy-Weinberg equilibrium and linkage disequilibrium in ARLEQUIN version 3.5.2.2 [86] and significance was assessed following Bonferroni correction for multiple tests. Loci that showed null alleles in multiple populations, or with significant deviation from Hardy-Weinberg equilibrium in multiple populations, were removed from further analyses.

Genomic summary statistics were calculated in ARLEQUIN version 3.5.2.2 [86] as mean number of alleles per locus (N alleles), nucleotide diversity (θ), expected heterozygosity (H_e) and observed heterozygosity (H_o). Separate analyses of molecular variance (AMOVA) were carried out based on site of origin and genetic clusters identified in STRUCTURE (see below) using 16,000 permutations.

An outlier approach was used to identify loci that had a higher or lower F_{ST} than expected under a neutral model of selection (under positive or balancing selection, respectively). Outliers were estimated in two ways: firstly, using BAYESCAN version 2.1 [87], which uses a Bayesian approach to estimate the posterior probability that a locus is affected by selection, and was run using 20 pilot runs of 5,000 iterations each, a total of 1,050,000 iterations (sample size of 100,000 and a thinning interval of 10), and a burn-in of 50,000. Only loci with a posterior probability (P) ≥ 0.95 with a prior odd of 10 were considered as outliers. BAYESCAN has consistently outperformed other outlier detection methods in terms of lack of false positives [4, 17, 88]. Outliers were also calculated in ARLEQUIN using 100,000 simulations, 500 demes, and a maximum expected heterozygosity of 0.5 [89]. Outlier SNPs were identified using a P -value of ≤ 0.05 . Only SNPs identified as outliers by both BAYESCAN and ARLEQUIN were conservatively selected as candidate loci under balancing or directional selection. Subsequent analyses of population structure and seascape genomics were conducted separately for neutral and outlier loci.

Population structure

Pairwise F_{ST} were calculated in ARLEQUIN version 3.5.2.2 [86] and significance assessed following Bonferroni correction for multiple tests. Straight-line distances between sites were calculated using ArcGIS Version 10.1 (ESRI, 2011) and shortest ocean distances between sites were calculated using the R package *marmap* [90], where distance was calculated excluding positive elevation [89]. Isolation-by-distance (straight line and shortest ocean) was tested in ARLEQUIN (Version 3.5.2.2) using a Mantel test [91] with 10,000 permutations.

Genetic clusters (populations) were inferred using Bayesian analyses carried out in STRUCTURE version 2.3.4 [92]. The analyses assumed admixture, correlated allele frequencies, and were run with 100,000 burn-in cycles and 100,000 Markov Chain Monte Carlo runs. The number of populations (K) was considered between 1 and 10, with 10 replicates per K . This was repeated with and without location priors, and using a hierarchical approach to resolve fine-scale genetic structure by removing highly differentiated populations [42]. The most likely value of K was inferred by estimating ΔK using the Evanno method [93], as implemented in Structure Harvester version 0.6.94 [94].

Hydrodynamic models are increasingly used as predictors of larval dispersal (gene flow) in marine systems (e.g. [20, 95, 96] and many others). Probabilistic dispersal simulations of potential larval connectivity were run using the Connectivity Modeling System (CMS, Paris *et al.* 2013) paired with 3-hrly Hybrid Coordinate Ocean Model (HYCOM) and Navy coupled ocean data assimilation (NCODA) global $1/12^\circ$ daily outputs from 2004, 2010, and 2012 [98]. These years were selected to represent neutral, negative and positive phases of the North Atlantic Oscillation (NAO) respectively, and are thus likely to reflect average dispersal patterns over the last decade [99, 100]. Vertical velocities were calculated using the eulerian continuity equation, and a random diffusive kick was added at each hourly dispersal time-step to better

capture the potential effects of sub-gridscale turbulent diffusion (horizontal = $15 \text{ m}^2\text{s}^{-1}$, vertical = $0.05 \text{ m}^2\text{s}^{-1}$; after Kough *et al.* 2016). HYCOM is a global model appropriate for large landscape continental-scale dispersal simulations, but does not include tides. Larvae were considered as passive particles even though diel vertical migrations may occur [40]. Simulations can therefore be over-dispersive, as the inclusion of tides and behaviour is liable to be retentive [102].

In the simulations, 100 propagules, each representing an *S. alveolata* larva, were released in a Lagrangian framework (so that each theoretical larvae was tracked individually), from each of nine locations (Figure 1), and their dispersal under two different release scenarios predicted: (1) a *conservative* scenario with propagule releases every 6-h over one week in April for each year, centred on the spring tide, and (2) an *optimistic* scenario which modelled a daily release on every day of the year. The *optimistic* scenario was designed to simulate all the potential pathways of dispersal if reproduction occurs throughout the year, which has been suggested in *S. alveolata* [40], thus capturing the full variability of potential larval fates. Larval competency to settle was assumed after 4 weeks, and a maximum PLD of 12 weeks applied based on field and laboratory estimates [40, 47]. Larvae were permitted to settle when located within 5km of a known reef site (Additional file 1). Additional (non-release) sites were also included as reported in The Global Biodiversity Information Facility (GBIF.org) in which larvae could also settle if encountered. Larval exchange counts (i.e. the source-sink contribution) were generated after 5, 8, 10, and 12 weeks PLD. Model outputs were processed in Matlab (Mathworks, Version R2016a). Pairwise matrices of larval exchange counts, standardised using proportions, were generated from larval fates averaged across the three focus years to compare against genetic pairwise F_{ST} matrices. Daily positions of larvae throughout the simulation were used to generate cumulative 2-D maps of larval density to highlight the predicted pathways of dispersal and areas of potential settlement

The proportion of settlers reaching each study site from each release site under the four larval duration scenarios (5, 8, 10 and 12-weeks) in the ocean circulation modelling simulations were inverted and used to form a matrix of pairwise dispersal resistances (i.e. the proportion of larvae that did not reach each site). Mantel tests [91] were carried out in ARLEQUIN (Version 3.5.2.2) [86] using 16,000 permutations to assess the correlation between pairwise F_{ST} (using putatively neutral loci) and dispersal. A Bonferroni correction was applied to assess the significance of the Mantel tests [103].

Loci under selection

Locally collected temperature data provides insight into the conditions that *S. alveolata* experience at a spatial and temporal scale that cannot be achieved using satellite data [104] and can be used for comparison with genomic data, to test genotype-environment associations [105]. Temperature data were obtained from Seabra *et al.* (2015) [106] who used biomimetic temperature loggers placed in the intertidal zone of exposed shores [104, 107]. At five sites, temperature data were collected with a resolution of 0.5°C at mid-intertidal level every hour between July 2010 and July 2014. High- and low-tide temperatures were also identified to the nearest hour. At the remaining four sites (i.e. English Channel, Balearic Sea, Tyrrhenian Sea and North Africa), an iButton (Maxim, Munich) was placed in the mid-littoral zone, collecting temperature data with a resolution of 0.5°C every 3 hours between April 2014 and April 2015. Temperature readings closest to high- and low-tide were identified.

S. alveolata experience tidally-driven fluctuations in water and air temperatures. Sixteen metrics were calculated to describe variation in temperature at each site as follows: mean temperature, maximum temperature, minimum temperature, temperature range (maximum-minimum), standard deviation of the mean, 95th–5th percentile (to exclude outlier temperatures), average daily temperature range (maximum-minimum per day), and average daily standard distribution (standard deviation of the mean), per site. High-tide (water) and low-tide (air) temperature values were also used separately to calculate mean temperature, maximum temperature, minimum temperature, and temperature range (maximum-minimum) for water and air temperatures, respectively (Table 3). Larvae of *S. alveolata* can be found in the

water column all year round [40], suggesting that they are reproductively active throughout the year, thus temperature data from all months were included in the analyses.

We utilised a gene-environment association software, BAYENV2 [108], to test for covariance between SNP allele frequencies and the 16 thermal variables. BAYENV2 controls for neutral population structure by first estimating a covariance matrix among populations for all loci and then accounting for that covariance in the gene-environment association test [109, 110]. The programme was run with 100,000 iterations for both neutral parameterisation and association testing [111]. Outlier loci were classed as those with a Bayes Factor > 3 [8, 112, 113].

List Of Abbreviations

PLD: planktonic larval duration

RADseq: restriction-site associated DNA sequencing

Declarations

Ethics approval

Not applicable

Availability of data and materials

DNA sequences are available on NCBI SRA (Accession: xx). Sample site locations, temperature data and ocean circulation model outputs are available on Dryad Digital Repository at the DOI (DOI).

Competing interests

The authors declare that they have no competing interests

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Authors’ contributions

AM, FN and SD designed the experiment and carried out the fieldwork. AM carried out the laboratory work. RR and AK carried out the ocean circulation modelling. FL and RS collected the temperature data. LF provided occurrence data and carried out sample collection. AM and FN carried out the bioinformatics and statistical analyses. EC and GLC provided bioinformatics support. AM and RR wrote the manuscript. All authors read and approved the manuscript.

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Additional Files

Additional file 1: Additionalfile1_SeascapeGenomics_Muiretal.docx

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Tables

Table 1: Genomic summary statistics of populations sampled.

Site	n	N alleles	θ	π	He	Ho
Northern Irish Seat†	6	2.50 ± 0.50	0.28 ± 0.14	132.56	0.53 ± 0.34	0.14 ± 0.19
North Atlantict	8	2.45 ± 0.50	0.15 ± 0.08	73.08	0.5 ± 0.36	0.11 ± 0.16
Southern Irish Seat†	9	2.54 ± 0.50	0.26 ± 0.13	126.36	0.53 ± 0.28	0.01 ± 0.15
English Channel†	11	2.49 ± 0.50	0.29 ± 0.14	140.50	0.49 ± 0.33	0.09 ± 0.15
Bay of Biscay†	12	2.59 ± 0.49	0.28 ± 0.14	133.63	0.51 ± 0.32	0.11 ± 0.15
Tyrrhenian Sea	7	2.23 ± 0.42	0.22 ± 0.11	106.08	0.45 ± 0.36	0.08 ± 0.19
Balearic Sea	4	2.36 ± 0.48	0.29 ± 0.16	141.43	0.58 ± 0.31	0.11 ± 0.15*
Iberian Peninsula†	5	2.23 ± 0.42	0.26 ± 0.14	123.47	0.50 ± 0.34	0.09 ± 0.20
North Africa	6	2.53 ± 0.50	0.32 ± 0.16	152.42	0.55 ± 0.34	0.14 ± 0.19

Shown are site name, sample size for which usable DNA was available (n), mean number of alleles per locus (N alleles), nucleotide diversity (θ and π), expected heterozygosity (He) and observed heterozygosity (Ho). Standard deviations are indicated for mean values.

† Temperature data available for this site; * Ho significantly different to He

Table 2: Pairwise genetic distances between sites (FST; lower triangle) and coastline distances between sites (km; upper triangle).

	Northern Irish Sea	North Atlantic	Southern Irish Sea	English Channel	Bay of Biscay	Tyrrhenian Sea	Balearic Sea	Iberian Peninsula	North Africa
Northern Irish Sea	-	453	180	864	1286	4134	3075	1853	2447
North Atlantic	0.36	-	612	994	1332	3998	2939	1715	2311
Southern Irish Sea	0.30	0.42	-	768	1190	4041	2982	1760	2354
English Channel	0.23	0.43	0.25	-	834	3742	2683	1461	2056
Bay of Biscay	0.18	0.31	0.25	0.20	-	3390	2331	1109	1703
Tyrrhenian Sea	0.35	0.58	0.31	0.20	0.31	-	1177	2283	2150
Balearic Sea	0.22	0.31	0.25	0.22	0.22	0.34	-	1224	1091
Iberian Peninsula	0.32	0.53	0.28	0.19	0.29	0.24	0.27	-	597
North Africa	0.21	0.44	0.21	0.15	0.22	0.22	0.22	0.19	-

Numbers in bold are significant after Bonferroni correction ($p < 0.0056$)

Table 3: Temperature parameters per site.

Temperature data	Temperature parameter	Northern Irish Sea	North Atlantic	Southern Irish Sea	English Channel	Bay of Biscay	Iberian Peninsula
All temperatures	Mean temp	10.82	11.62	11.93	14.21	16.67	16.10
	Max temp	37.85	39.37	34.78	41.54	42.13	35.48
	Min temp	-3.31	-5.62	-0.46	0.70	-2.46	3.32
	Temp range	41.16	44.99	35.24	40.84	44.59	32.16
	SD of mean	5.61	5.94	6.10	5.87	8.36	7.79
	95th – 5th percentile	14.38	15.40	15.77	23.68	21.74	18.59
	Average daily range	7.51	7.11	6.77	8.15	8.28	8.28
	Average daily SD	5.50	5.50	4.51	5.76	5.30	5.42
	Water temperatures	Mean temp	10.55	11.53	11.76	14.00	16.76
Max temp		16.14	19.73	17.24	22.50	25.09	21.17
Min temp		4.52	3.67	5.88	5.91	7.90	11.18
Temp range		11.62	16.06	11.36	16.59	17.19	9.99
Air temperatures	Mean temp	11.06	11.84	12.04	14.59	16.58	16.59
	Max temp	37.76	36.93	34.53	41.54	40.74	34.49
	Min temp	-2.28	-4.73	0.00	0.70	-1.03	3.53
	Temp range	40.04	41.66	34.53	40.84	41.77	30.96

Shown are all temperature data combined, water (high tide) temperatures and air (low tide) temperatures (°C). The mean, maximum, minimum, temperature range and average daily temperature range are shown for all temperatures, water temperatures and air temperatures, separately. The standard deviation (SD) of the mean, 95th – 5th percentile, and average daily standard deviation (SD) are also shown for all temperatures only.

Figures



Figure 1

Sampling sites spanning the latitudinal and longitudinal range of *S. alveolata*, which were also used as the release sites for the Lagrangian model. Sites shown as triangles are those from which temperature was available.

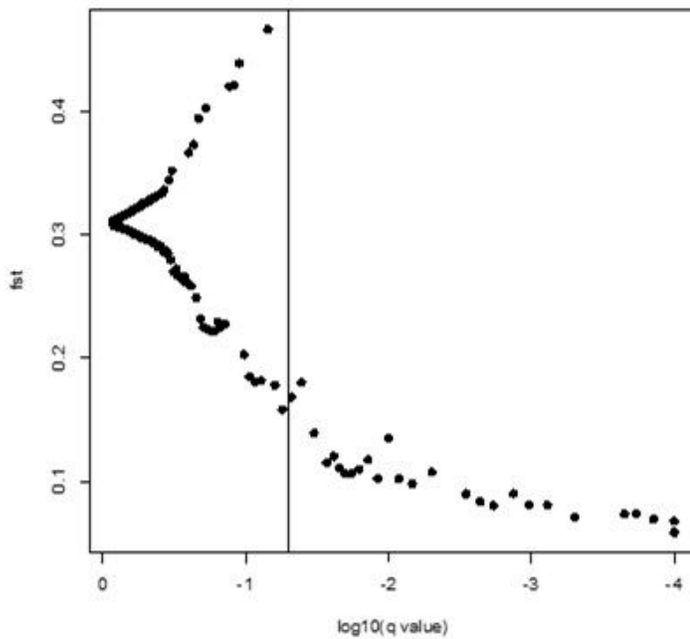


Figure 2

Posterior odds of the selection model and locus-specific F_{ST} for each SNP as calculated in BAYESCAN [87]. The vertical line represents the threshold for significance for a locus to be under selection of $P \geq 0.99$. Loci to the right of the threshold line are suggested to be outlier loci.

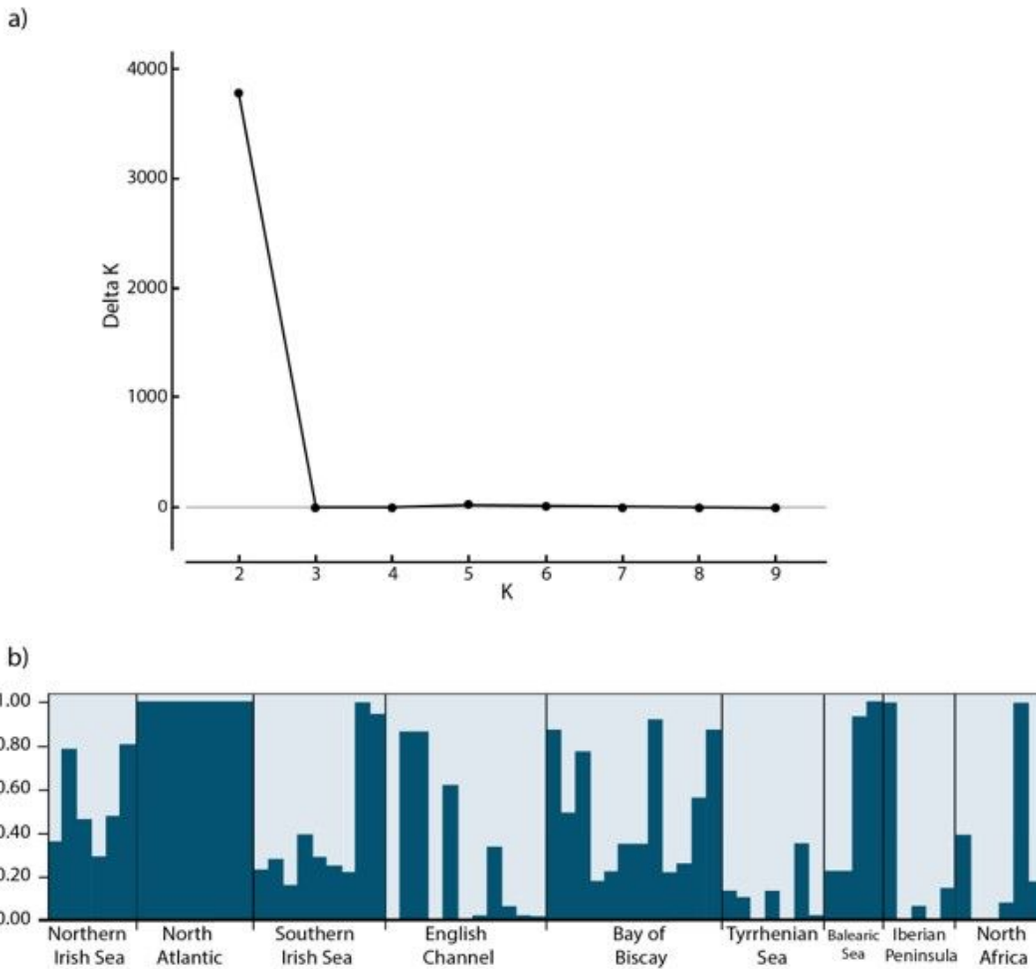


Figure 3

Population structure based on genetic clustering in STRUCTURE (Pritchard et al. 2000) using all loci. a) ΔK for each value of K (putative number of populations), averaged over ten replicates. The peak at $K = 2$ shows the most likely number of genetic clusters within the sample. b) STRUCTURE barplot for $K = 2$. Columns are individuals, with the proportion of an individual's genotype assigned to each cluster (K) denoted by dark blue or light blue.

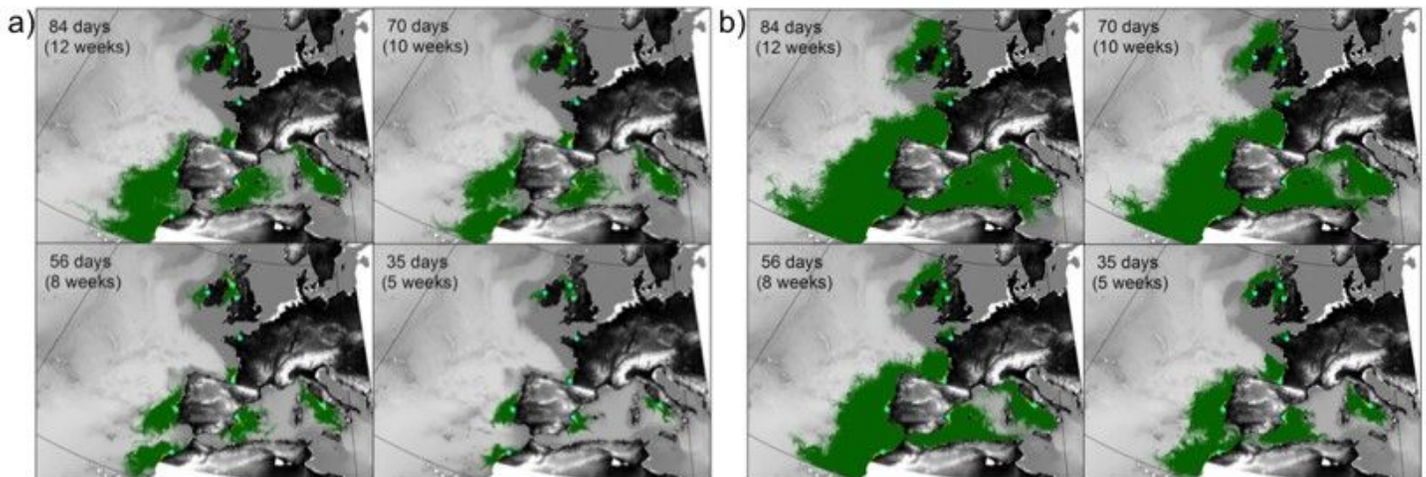


Figure 4

Maps of larval densities from dispersal simulations with a) releases every 6 hours over one week in April centred on the spring tide, representing the most conservative dispersal simulations; and b) daily releases throughout each year of simulation, represent the most optimistic dispersal simulations. They show larval densities ranging between red (high), yellow (medium), and green (low). Maps are shown for the four different planktonic larval durations ranging from 5 - 12weeks.

Supplementary Files

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