INTRODUCTION

Assessment of genetic connectivity among populations is widely used in conservation biology and species management to evaluate effective dispersal and the consequences of spatially-dependent evolutionary processes on species persistence, resilience and adaptability (Cayuela et al., 2018; Gagnaire, 2020). Using polymorphism data from individuals sampled across the landscape, the genetic approach can detect local barriers to gene flow separating distinct populations that exchange measurable amounts of migrants per generation. Demographic independence between such populations can be assumed when the fraction of exchanged migrants is too small to ensure population persistence in case of negative intrinsic growth rate (Lowe & Allendorf, 2010; Pulliam, 1988). Measuring
migratory exchanges using genetic data may, however, be a difficult task.

The power of the population genetics approach depends on the relative intensity of two opposed forces, migration (m) and genetic drift (1/N_e). When migration overwhelms the effect of genetic drift (i.e. when the number of migrants exchanged per generation [N_e*m] exceeds a few dozen), genetic differentiation between populations is close to zero. Null genetic differentiation can result from a variety of equilibrium scenarios ranging from high connectivity among small populations to nearly complete demographic independence among large populations. Consequently, assessing demographic connectivity in species with both large populations and high migration rates has been a long-standing challenge to population genetic approaches (Waples, 1998). Such combinations of biological parameters are frequently encountered in marine species with a dispersive larval stage (Hedgecock, Barber, & Edmands, 2007). Therefore, a persistent gap between evolutionary and ecological scales often prevents managers to use genetic connectivity information for fisheries management (Palumbi, 2003; Waples & Gaggiotti, 2006; Waples, Punt, & Cope, 2008).

Molecular markers influenced by selection can compensate the lack of signal at neutral loci when the migration-drift equilibrium is not informative. The use of adaptive differentiation signals has been proposed as a solution to improve the delineation of management units (Funk, McKay, Hohenlohe, & Allendorf, 2012). This approach is now greatly facilitated by the availability of large polymorphism data sets to scan genomes for outlier markers influenced by selection (Narum, Buerkle, Davey, Miller, & Hohenlohe, 2013; Savolainen, Lascoux, & Merilä, 2013; Stapley et al., 2010). Several studies in marine species that typically show weak to no genetic differentiation at neutral markers have found stronger signals of spatial structure at different scales using outlier loci (e.g., Bekkevold, Gross, Arula, Helyar, & Ojaveer, 2016; Benestan et al., 2016; Van Wyngaarden et al., 2017). Although these could help in delineating cryptic evolutionary management units, the variation patterns displayed by outlier loci often remain challenging to translate into quantitative assessments of demographic connectivity. One of the main reasons is that observed genetic variation patterns can be generally attributed to a variety of possible selective mechanisms (Bierne, Roze, & Welch, 2013), potentially involving complex relationships between local adaptation and gene flow (Tigano & Friesen, 2016). Contextualizing the ecoevolutionary history of the studied species thus helps to discern the mechanisms underlying the patterns of differentiation displayed by outlier loci, toward a better understanding of connectivity (Liggins, Treml, & Riginos, 2019).

Genetic admixture between differentiated lineages is a particularly informative evolutionary context to learn about connectivity. When two divergent taxa come into secondary contact and exchange genes, the spatial diffusion of foreign genetic material can be used to reveal the population genetic connectivity within each introgressed lineage (Bertl, Ringbauer, & Blum, 2018; Duranton, Bonhomme, & Gagnaire, 2019; Sedghifar, Brandvain, Ralph, & Coop, 2015). If the recipient lineage is not genetically structured, the spatial homogenization of foreign allele frequencies is a quick process. However, local barriers to gene flow may slow down the spread of foreign alleles and generate steps in the admixture and introgression gradients within the recipient lineage (Gagnaire et al., 2015). Although allele frequencies equilibrate quickly at neutral loci following secondary contact, barrier loci involved in partial reproductive isolation have a reduced effective migration rate between lineages and therefore retain their signal of differentiation for longer (Sedghifar, Brandvain, & Ralph, 2016). Semi-permeable species boundaries characterized by heterogeneous rates of introgression among loci thus provide favourable conditions to reveal cryptic barriers to gene flow within introgressed lineages, even after several thousands of generations of introgression (Gagnaire et al., 2015). Post-glacial lineages that are nowadays in contact and exchange genes through natural hybridization represent suitable cases for using spatial introgression gradients to study connectivity.

The European sea bass (Dicentrarchus labrax L.) is a marine fish species that ranges from northwestern Africa to southern Norway in the Atlantic, and throughout the Mediterranean and Black Seas (Vandeputte, Gagnaire, & Allal, 2019). The species is genetically subdivided into two distinct lineages, one Atlantic (ATL) and one Mediterranean (MED), which naturally hybridize at the Atlantic-Mediterranean transition zone (Lemaire, Versini, & Bonhomme, 2005; Naciri, Lemaire, Borsa, & Bonhomme, 1999; Souche et al., 2015). Population genomic studies based on RAD-sequencing (Tine et al., 2014) and whole-genome resequencing (Duranton et al., 2018) have revealed several important aspects of the of the European sea bass evolutionary history: (a) genetic divergence between ATL and MED lineages is the result of about 270 kys of allopatric divergence; (b) secondary contact has started after the last glacial retreat about 11.5 kys ago, and since then (c) gene flow occurs at variable rates across the genome due to partial reproductive isolation between the two lineages. (d) For a still unknown reason, postglacial gene flow has been more pronounced from the Atlantic into the Mediterranean than in the opposite direction. As a result, contemporary ATL genomes are made of ~5% of MED genetic material, while western and eastern MED genomes contain ~31% and 13% of ATL ancestry, respectively (Duranton et al., 2018). Such levels of introgression are sufficiently high to provide information on genetic connectivity within both sea bass lineages. In the Mediterranean, Duranton et al. (2019) analysed the neutral decay of introgressed haplotype length as a function of distance from the contact zone to estimate a mean per-generation dispersal distance of five to 50 km. This quantitative approach, based on a small number of whole-genome sequences, did not have the spatial resolution required to provide a detailed map of connectivity. Here, we used a different strategy based on moderate genome coverage but extensive spatial sampling to identify local barriers to gene flow within the Atlantic lineage. Our objective was to study both the neutral and non-neutral diffusion of MED alleles across Atlantic sea bass populations to evaluate their fine-scale spatial genetic structure.

Atlantic sea bass populations are of high economic importance to European fisheries, but overfishing probably combined with...
other factors have led to a decline in landings since 2009-2010, particularly north of the 48th parallel (ICES, 2019). Sea bass stock assessment by scientific authorities (ICES: International Council for the Exploration of the Sea) in the northeastern Atlantic currently relies on four presumably distinct stocks: northern Atlantic, southern Ireland/western Scotland, Biscay and Iberia. However, stock delineation remains poorly understood, making it difficult to properly assess population connectivity and implement effective management programs. Population genetic studies generally found non-significant genetic structure across the northeastern Atlantic (Coscia & Mariani, 2011; Fritsch, Morizur, Lambert, Bonhomme, & Guinand, 2007; Souche et al., 2015), possibly due to insufficient spatial and genomic coverage. In parallel, mark-recapture data indicated restricted individual movement between the Bay of Biscay and the English Channel-Celtic Sea region (Fritsch et al., 2007), a result partly explained by the fidelity to winter spawning areas recently evidenced with tagging data (de Pontual et al., 2019). A small number of effective migrants per generation may account for the absence of genetic differentiation at neutral markers despite high fidelity to spawning grounds. Therefore, it is possible that independent demographic units have remained undetected by previous population genetic studies. Here, we investigate fine-scale connectivity patterns among Atlantic sea bass populations using the spatial diffusion of Mediterranean alleles from southern Portugal toward the northern part of the species range. We document the existence of two significant steps in the Mediterranean ancestry gradient along the Atlantic coast, which most likely delineate demographically independent populations that slightly differ from current management units. We also find molecular signatures of a recent northward expansion, consistent with recently expanding sea bass fisheries in the northern part of the species range.

2 MATERIALS AND METHODS

2.1 Sampling

Fin clips from 846 individuals were sampled from southern Portugal to the Irish Sea and North Sea in the period October 2012 to May 2015 (Table S1). Samples were mostly collected in fish markets, only from fresh fish with a known geographical origin indicated by its ICES statistical rectangle (precision 0.5° in latitude and 1° in longitude) and a known fishing day. The quality of the spatial fishing information of each individual was checked with the name of the fishing vessel by tracking its position history on http://www.marintraffic.com.

Tissues were preserved in 85% ethanol at −20°C. Individual genomic DNA was extracted following cellular lysis and proteinase K digestion using the Qiagen DNeasy Blood & Tissue kit, and then conserved in TE. DNA extraction quality was controlled using a NanoDrop spectrophotometer keeping samples with an A260/A280 ratio between 1.8 and 2.0. Concentration of double-stranded DNA was then measured with the Qbit dsDNA Broad Range kit with a standard benchmark (0–100 ng/µl) on a qPCR BioRAD. Individual dsDNA were finally normalized at 25 ng/µl and randomly distributed across nine 96-well plates before genotyping.

2.2 Genotyping and data quality control

Genotyping was performed by the Labogena platform (Jouy-en-Josas, France) using an iSelect Custom Infinium Illumina array specifically developed in the European sea bass (Faggion, Vandeputte, Chatain, Gagnaire, & Allal, 2019). The chip contains 1,531 validated SNPs covering the whole genome while being homogeneously distributed along the recombination map, and presents no ascertainment bias between Atlantic and Mediterranean sea bass lineages. Individual genotypes and hybridization intensities generated by Illumina’s BeadStudio software were reanalysed using the R package Argyle (Morgan, 2016) to perform SNP quality control. Markers were filtered based on their rate of no-call genotypes (per locus missingness < 0.1), rate of heterozygosity (Het < 0.55), and minor-allele frequency within the whole Atlantic data set (MAF > 0.01), in order to include even variants that are frequent or fixed within the Mediterranean, but rare in the Atlantic. Individuals with an excess of missing genotypes were also excluded (per sample missingness < 0.1).

In order to remove a few miscalled markers which variation profile was correlated with samples arrangement in the 96-well plates, we performed a genotype-plate association analysis in Plink (Purcell et al., 2007, p-value exclusion threshold = 1e-4).

Reference Mediterranean genotypes of 10 MED individuals (four from eastern MED and six from western MED) were merged to our final data set to compare allele frequencies between ATL and MED populations. These reference genotypes were extracted from high-quality whole-genome resequencing data without any missing genotype at the retained loci (Duranton et al., 2018).

2.3 Spatial genetic variation on different data sets

The 827 quality-filtered specimens were stratified into three different data sets to reach variable levels of precision in spatial genetics analyses: (a) a “regional data set” of 827 specimens distributed into seven regions; (b) a “main data set” in which the 827 specimens were assigned to 21 localities corresponding to single ICES rectangles or groups of adjacent rectangles, with a minimum precision of 1.5° in latitude and 3° in longitude, and a number of individuals per locality ranging from 11 to 111 (mean = 40, Figures 1 and S1, Table S1 for more details); (c) a “refined data set”, more precise in space, consisting of 761 specimens distributed into 31 localities, with a minimum precision of 30’ in latitude and 1° in longitude, and a minimal number of eight individuals per locality (from eight to 43, mean = 24, Figure S2). The main data set was used for all analyses, unless stated otherwise.
The proportion of polymorphic markers among the 1,012 retained SNPs was calculated for each of the 21 localities. A regression model of the proportion of polymorphic markers as a function of sample size per location was fitted using a nonlinear least squares analysis with two parameters in order to account for the effect of sample size. We then tested for a linear relationship between the localities’ residuals to the fitted model and the latitude of localities. We also tested the existence of a linear correlation between the mean observed heterozygosity per sample locality and latitude. Sample groupings from the regional data set were then used to test for deviations from expectations under Hardy-Weinberg Equilibrium within each of the seven regions defined above with the Pegas R package (Paradis, 2010).

A redundancy analysis (RDA) was performed using the R package Vegan (Oksanen et al., 2019) in order to evaluate the extent to which SNP variation is influenced by geographic and temporal factors. We first fitted the model $Y$ (individual genotype) ~ (Latitude + Longitude + Month + Year of capture) and assessed the significance of each explanatory factor using 1,000 permutations of genotypic data before refitting the model.

Overall $F_{st}$ was calculated among all of the 21 Atlantic locations using all loci and pairwise $F_{st}$ were then calculated for each pair of sampling locations including the two reference MED populations, as well as for pairs of regions using the "regional data set". We used the Hierfstat R package (Goudet, 2005) to calculate $F_{st}$ values and assess their significance using 5,000 random permutations of genotypes among localities and among regions. A sequential Bonferroni correction (Hommel, 1989) was applied to empirical $p$-values with the function `p.adjust()` of the R-base package Stats. In order to assess the

**FIGURE 1** Sampling map of the 827 Atlantic sea bass individuals analysed in this study. In the ‘main data set’, the 827 specimens were assigned to 21 localities corresponding to single ICES rectangles or groups of adjacent rectangles represented with the same colour. In the ‘regional data set’, specimens were grouped into seven regions represented by coloured polygons. Total number of samples is indicated below each region name. Details about sample locations and the spatially ‘refined data set’ are provided in Figure S1 and Table S1.
presence of an isolation-by-distance pattern among ATL locations, we tested the linear correlation between \( F_{ST}/(1-F_{ST}) \) and geographic distance between pairs of localities. We also performed a Mantel test between matrices of pairwise genetic and geographic distances with 10,000 permutations using adegenet (Jombart, 2008). Pairwise geographic distances were calculated as the shortest path by the continental plateau between two localities (i.e., bathymetry < 200 m) using the R package Marmap (Pante & Simon-Bouhet, 2013).

We used the directionality index statistics \( \Psi \) (Peter & Slatkin, 2013) to test whether genetic differentiation among pairs of localities was due to isolation-by-distance at equilibrium or to a recent range expansion in the Atlantic distribution area of *D. labrax*. Briefly, the directionality index uses comparisons of two-dimensional site frequency spectra to capture deviations from symmetric migration, such as those generated during range expansions. We used the R package rangeExpansion (https://github.com/BenjaminPeter/rangeExpansion) to calculate \( \Psi \) for all pairs of Atlantic locations, and used pairwise geographic distances between locations to infer the most likely spatial origin of a possible range expansion.

### 2.4 MED introgression in ATL genomes

In order to estimate the fraction of MED alleles present in ATL individuals, we used the program Admixture (Alexander, Novembre, & Lange, 2009), which infers individual ancestry proportions from \( K \) ancestral populations (here \( K \) was set to 2). Our unsupervised ancestry inference included 10 MED reference individuals in addition to the 827 ATL samples, so that the two ancestral populations correspond to the ATL and MED sea bass lineages. We set the termination criterion for the optimization algorithm to 100 iterations to ensure convergence and used 1,000 bootstrap replicates to estimate the standard error of individual ancestry proportions.

The effects of sampling date and geographic distance to the southernmost Atlantic location (Sines in southern Portugal, code “SINE”) on the extent of Mediterranean introgression were tested using a global linear model (GLM, family = Gaussian), with the percentage of MED ancestry as the dependent variable. The date of capture (using the ‘as.date()’ class in R) and the distance to the SINE locality (calculated as the shortest path by the continental plateau) were used as explanatory factors.

The effect of geographic distance to the southernmost ATL location (SINE) on the extent of MED ancestry was then tested alone using a linear regression model. The distance to the SINE location was calculated as the shortest path by the continental plateau. In order to test for the existence of potential breaks in the admixture gradient, we then performed a piecewise regression of MED ancestry as a function of distance to SINE. Breaks separating different linear regression models were introduced one by one and, at each step, every possible break position was examined by calculating the residual standard error of the piecewise regression model. The significance of the reduction in the residual sum of squares of each piecewise regression model compared to the simple regression model was tested using an ANOVA, taking into account the number of additional parameters in the piecewise regression model.

### 2.5 Genome scans

We then evaluated the impact of variable rates of introgression between ATL and MED sea bass lineages on allele frequency gradients within the Atlantic. We more specifically tested the theoretical prediction that the genomic regions with reduced rates of introgression between lineages tend to exhibit the strongest spatial allele frequency patterns within the ATL range due to propitious ratios of between- to within-lineage gene flow intensities (Gagnaire et al., 2015). We thus expected that the loci showing the strongest genetic differentiation within the Atlantic would generally also be \( F_{ST} \) outliers between ATL and MED lineages. Given the relatively low density of markers in our data set (about 1 SNP per cM), the candidate loci detected by genomes scans for differentiation are most likely to be indirectly influenced by selection through linkage with a nearby selected locus. A previous quantification indicated that the genomic island regions under the influence of the strongest reproductive isolation barriers between Atlantic and Mediterranean lineages occupy about 4% of the sea bass genome (Duranton et al., 2018). Therefore, our SNP panel is expected to contain at least 40 SNPs in strong linkage with reproductive isolation barriers between the two sea bass lineages, plus variants showing intermediate degrees of linkage with them. To identify those loci, we used two different genome scan methods to detect \( F_{ST} \) outlier SNPs. Both methods were applied to the analysis of genetic differentiation at two different scales: (a) within the ATL range among the 21 localities; and (b) between the ATL and MED sea bass lineages.

The first method used was Lositan (Antao, Lopes, Lopes, Beja-Pereira, & Luikart, 2008; Beaumont & Nichols, 1996), which performs coalescent simulations under the symmetric island model to generate a neutral distribution of \( F_{ST} \) conditioned on the mean expected heterozygosity among populations. The mean \( F_{ST} \) calculated across all markers was used as a target value to simulate 1 million SNPs, from which we empirically determined the 99.5th percentile of the \( F_{ST} \) distribution to identify candidate outlier loci showing an excess of genetic differentiation among locations.

We then used BayeScan (Foll & Gaggiotti, 2008), a Bayesian outlier detection method that relies on the multinomial-Dirichlet model. The difference in allele frequency between every sampling location and a common theoretical gene pool was measured by a subpopulation-specific \( F_{ST} \) coefficient in order to account for differences in effective sizes and migration rates among subpopulations. Selection was introduced by decomposing for each locus the subpopulation-specific \( F_{ST} \) coefficient into a subpopulation-specific component shared by all loci and a locus-specific component shared by all subpopulations. Deviation from neutrality for a given locus was assumed when the locus-specific component was needed to explain the observed diversity pattern. The posterior probability that a locus is influenced by selection after accounting for multiple
testing was determined by the posterior odds (PO), which is the ratio of posterior probabilities between the selection and neutral models. The minimum false discovery rate (FDR) at which a given locus reached significance was determined using the q-value of each locus. A q-value threshold of 0.05 was used for outlier detection in most analyses, corresponding to an FDR threshold of 5%. BayeScan chain parameters were set to 5,000 outputted iterations with 5,000 steps for pilot runs and 50,000 burnin steps.

2.6 | Analysis of introgression gradients around barrier loci

Simulations of secondary contact between two genetically subdivided lineages exhibiting partial reproductive isolation show that neutral markers linked to barrier loci can display substantial allele frequency steps within the introgressed lineage (Gagnaire et al., 2015). These steps appear at the place where local barriers to gene flow slow down the diffusion of introgressed alleles. They are dynamically maintained by differences in flow intensity, and are amplified when gene flow between lineages is slightly higher than the rate of homogenization within the introgressed lineage. Depending on the age of the secondary contact, the steps in allele frequencies magnified by foreign introgression may appear at variable recombination distances from the barrier loci.

To explore that effect, we first focused on the subset of SNPs that were the most strongly associated to barrier loci in our data set, using only outlier loci that were detected in the BayeScan MED-ATL genome-scan with a stringent FDR threshold of 1%. Since genetic differentiation is strong between ATL and MED within genomic islands, the minor frequency allele found at each of these loci in the ATL most often corresponds to the major frequency allele in the MED. Therefore, gradients of introgression of Mediterranean alleles at those loci are expected to generate minor allele frequency (MAF) clines within the Atlantic distribution range. In order to test for a cryptic population structure within the Atlantic, we thus compared the mean MAF of MED-ATL outlier loci to that of neutral SNPs as a function of the distance to the southernmost ATL location (SINE).

Finally, we evaluated how different degrees of linkage to barrier loci affect MAF gradients within the Atlantic. We assigned SNPs to different categories defined by their level of allele frequency difference (Δp) between ATL and MED. Our rationale was that stronger Δp values between ATL and MED are associated to stronger linkage to barrier loci, which was supported from earlier works (Duranton et al., 2018).

3 | RESULTS

3.1 | SNP genotyping and quality filtering

After excluding nonvariable markers corresponding to polymorphisms private to the Mediterranean sea bass lineage as well as marker genotyping failures, 1,361 SNPs were successfully genotyped in 846 specimens over the 1531 markers present on the array, representing an overall call-rate of 90.7%. SNPs with a minor allele frequency (MAF) lower than 0.01 within the Atlantic were subsequently excluded, as were individuals with more than 10% of nonscored genotypes. A few additional SNPs showing variation profiles significantly correlated with the samples’ mapping within the 96-wells plates used for genotyping were also discarded to avoid experimental effects. After these steps of quality-filtering, 1,012 biallelic SNP markers that were genotyped in 827 individuals were retained in the final main data set (Table S1), resulting in an overall genotyping call rate of 96.87%.

3.2 | Genetic diversity gradient and spatial structure

The average fraction of polymorphic markers per sampling location calculated across the 1,012 SNPs was negatively correlated to latitude after standardizing for the number of specimens genotyped by locality (adjusted $R^2 = .77; p < .001$; Figure 2a), providing evidence of a latitudinal decrease in genetic diversity across the ATL sea bass range. Similarly, the mean observed heterozygosity per sample location was negatively correlated to latitude (adj. $r^2 = .66; p < .001$; Figure S3), decreasing by nearly 10% from Southern Portugal to northeastern UK (Table S2).

The proportion of the total genotypic variance explained by the redundancy analysis constrained by (latitude + longitude + month + year of capture) was low (0.64%) but highly significantly (p-value < .001). However, constrained ordination only revealed significant marginal effects on SNP variation for the two spatial variables (latitude: p-value < .001; longitude: p-value = .037). The RDA1 axis received a large contribution of latitude, as illustrated by the gradient in individual coordinates distributions among the seven regions from Portugal to the North Sea (Figure 2b).

The overall $F_{ST}$ calculated over all loci among the 21 Atlantic locations was not significant ($F_{ST} = 0.0002, p > .05$). Pairwise $F_{ST}$ values calculated between pairs of ATL locations or regions were consistently low and most often nonsignificant (Tables S3–S5), and only few pairwise $F_{ST}$ values reached significance among the seven ATL regions considered (Table S6). By contrast, genetic differentiation was much higher between ATL and MED sea bass lineages (average $F_{ST} = 0.1474$) using the same marker data set, consistently with previous studies (Duranton et al., 2018; Tine et al., 2014). A weak but significant positive correlation was found between geographic distance separating pairs of ATL localities and genetic distance estimated by $F_{ST}/(1-F_{ST})$ (adj. $r^2 = .078; p < .001$; Figure 2c). This isolation-by-distance (IBD) pattern was remarkably stronger when the analysis was restricted to the northern part of the Atlantic range, that is, excluding Portugal, Biscay and southwestern Channel locations (adj. $r^2 = .236; p$-value < .001; Figure S4). This was also confirmed by the Mantel test (Table S7).

The analysis of directionality index $\Psi$ based on the method by Peter and Slatkin (2013) detected a strongly significant deviation
Figure 2  Spatial genetic variation. (a) Negative correlation between standardized polymorphism corrected for sampling size and latitude of the 21 sampling localities. The blue line shows the linear regression (slope $p < .001$, $r^2 = .77$) and its 0.95 confidence interval (grey shade). (b) Distribution of individual coordinates on RDA1 axis as a function of latitude for each of the seven regions: Po, Portugal; SB, South Biscay; NB, North Biscay; CS, Celtic Sea; Ch, Channel; IS, Irish Sea; NS, North Sea. The RDA was constrained using only spatial coordinates. For each region, the boxplot horizontal line represents the mean, whiskers the standard deviation and vertical lines the 0.95 confidence interval. Dots indicate outlier individuals, and whisker width cover the latitudinal range of each region. (c) Positive correlation between genetic differentiation represented by $F_{ST}/(1-F_{ST})$ and distance by the plateau (in km) between pairs of localities. The blue line shows the linear regression (slope $p < .001$, $r^2 = .078$) and its 0.95 confidence interval (grey shade). (d) Results of the directionality index analysis showing the gradient in the relative fit as expansion origin (colour scale) and most likely origin of expansion in southern Ireland (blue cross). For each of the 21 sampling localities, the inner circle indicates the mean heterozygosity (grey scale) and the colour of the outer circle indicates the group identified with the admixture gradient analysis (blue, Portugal; pink, Biscay; green, Celtic Sea - Channel - North Sea, see Figure 3).
from an isolation-by-distance at equilibrium model. Instead, a spatial expansion scenario was supported with an inferred origin located in the northwestern part of the Celtic Sea ($p$-value < $10^{-10}$, Figure 2d). The gradient in the relative fit as expansion origin showed a steep decrease in likelihood location towards the southern Celtic Sea and northern Biscay, and a smoother gradient towards the northeastern part of the range (Irish Sea, English Channel and North Sea). Together with the decreasing latitudinal gradient in heterozygosity, these results support a recent northeastern range expansion from a region located in the northwestern Celtic Sea.

### 3.3 Spatial gradient of MED ancestry in the ATL

Our inferences of individual MED ancestry proportions showed a clear northward-decreasing gradient (Figure 3). The southernmost location in Portugal (SINE), which is the closest to the ATL-MED transition zone, showed a significantly higher average level of MED ancestry (11.2%) compared to all other locations and a stronger variance in MED ancestry among individuals.

The GLM analysis showed that the explanatory variable "distance to SINE" had a very significant negative effect ($p < 2e-16$) on the percentage of MED ancestry, whereas no significant effect could be detected for the date of sampling ($p = .135$). The spatial gradient in MED ancestry was even more obvious when individual ancestry proportions were averaged by sampling location (Figure 4a). The mean fraction of MED ancestry per location was negatively correlated with the distance to SINE in the simple linear regression model (adj. $r^2 = .6026$, $p$-value < .001), and the correlation was strengthened when the SINE location (which is highly admixed) was removed from the analysis (adj. $r^2 = .7405$; $p$-value < .001).

The piecewise regression approach detected two significant break points in the spatial admixture gradient. Taking into account the presence of two additional parameters, the model with three different linear regression displayed a significantly better fit to the data than the simple linear regression model (ANOVA for model comparison: $p$-value = $3.094e-9$; best model with three regression lines: adj. $r^2 = .93$, $p$-value = $1.3e-9$). The strongest break occurred in the southwestern English Channel near the GONB locality in the Gulf of Saint-Malo, more than 2000 km northward to SINE by the plateau (Figure 4b). The second break was detected at the tip of Galicia between CORU and ASTU sampling locations (Figure 4c). Consistent with the dilution of MED ancestry toward the north, the average fraction of MED alleles decreased from 5.7% [0.037, 0.076] in the Portugal region, to 2.6% [0.013, 0.041] in the Bay of Biscay and southwestern English Channel, and 1.2% [0.004, 0.022] in the north of Brittany from Celtic Sea to Irish and North Seas (Figure 3). Each of these two steps was characterized by a more than two-fold reduction in the mean MED ancestry from south to north (Figure 4d). Although we found negative regression slopes indicating a decreasing mean MED ancestry with increasing distance to SINE in both the Portugal and Biscay-southwestern Channel regions, the slope was significantly positive to the north of the Gulf of Saint-Malo breakpoint. This striking result, indicating an inverted latitudinal gradient in MED ancestry in the northern part of the range, remained visible separately in a western Channel-Irish Sea transect and an eastern Channel-North Sea transect using the more spatially "refined data set" containing 31 localities (Figure S5a).
Within-ATL outliers are enriched for between-lineages outliers

Genome scans for highly differentiated SNPs were performed both at the within-ATL scale to detect outlier loci within the northeastern Atlantic and between ATL and MED samples to detect outlier loci showing an excess of differentiation between the two European sea bass lineages.

Lositan detected 32 outliers exceeding the 99.5th percentile of the neutral distribution of $F_{ST}$ at the within-ATL scale (Figure S6a) and 183 outliers between ATL and MED lineages (Figure S6b). Within-ATL outliers were highly significantly enriched for between-lineages outliers (21 out of 32 within-ATL outliers belong to the 183 ATL-MED outliers, hypergeometric test $p$-value = 1.9e-9).

Comparatively, the BayeScan analysis was more stringent in detecting outlier loci showing $q$-values < 0.05, with only seven candidate outliers being found at the within-ATL scale (Figure S6c) and 74 outliers detected between ATL and MED lineages (Figure S6d). All of the outliers identified with BayeScan were also detected by Lositan. Here again, the outliers detected at the within-ATL scale...
were highly significantly enriched for between-lineages outliers (five out of seven within-ATL outliers belong to the 74 ATL-MED outliers, hypergeometric test $p$-value = 3.4e-5).

3.5 | Minor allele frequency steps at genomic islands associated loci

We used a stringent subset of 52 outlier loci that were detected in the BayeScan MED-ATL genome-scan with an FDR < 0.01 to characterize the gradient in mean MAF as a function of the distance to SINE. Major steps were observed directly northward to CORU and GONB in these mean MAF profiles, which were not detected using nonoutlier loci (Figure 5). In order to refine the spatial positions of these steps, the same analysis was made based on the spatially refined data set (31 localities). Mean MAF were represented separately along two transects: one from Portugal toward the Irish Sea and one toward the North Sea (Figure S5b). Both profiles revealed low MAF values in the western Channel area (after PBRE-CRNW and PBRE-GONB), with the lowest values being found in IRIS, CRNW2 and 3, ISWI2 and BSEI1. Northward to these regions, the mean MAF profiles steadily increased towards WALE-SGEO (Irish Sea) and DOVE-ANGL-NEUK (North Sea), on both sides of Britain.

Finally, the evolution of mean MAF profiles was evaluated as a function of the degree of marker linkage to MED-ATL islands of divergence assessed with difference in allele frequencies between ATL and MED ($\Delta p$). Progressively flattening MAF profiles were found for groups of loci showing from high to low $\Delta p$ values (Figure S7).

4 | DISCUSSION

Genetic connectivity assessment in high gene flow species generally provides little information on the degree of demographic connectivity among populations (Lowe & Allendorf, 2010; Waples, 1998). This shortcoming typically arises when genetic differentiation at neutral markers is null or close to zero, so that no real signal can be used to determine whether migration rates are sufficiently high to ensure demographic coupling among populations. Here, we found nonsignificant overall genetic differentiation across the vast majority of the European sea bass northeastern Atlantic range. The sea bass thus represents one of the many borderline cases where neutral markers may remain silent to conservation and management issues.

As for many other species, however, the spatial distribution of genetic diversity in sea bass has retained signals of historical and contemporary admixture. Two sea bass lineages, one Atlantic and one Mediterranean, have diverged in allopatry during ca. 300,000 years and have subsequently experienced a postglacial secondary contact which started <15,000 years BP (Duranton et al., 2018; Tine et al., 2014). Since that time, gene flow has been strongly asymmetrical from the Atlantic into the Mediterranean lineage. However, gene flow has been also occurring in the opposite direction, accounting for <5% of introgressed Mediterranean alleles within Atlantic sea bass genomes (Duranton et al., 2018). Here, we show that the northward diffusion of Mediterranean alleles within the Atlantic sea bass populations can be used to reveal two dispersal barriers separating demographically independent units, and an ongoing expansion in the northern part of the species range. These findings could provide useful new elements to improve the delineation of European sea bass populations in fisheries management programs.

**FIGURE 5** Loci linked to genetic barriers to gene flow between ATL and MED lineages display stronger genetic differentiation within the Atlantic. (a) Scaled distributions of per-locus FST values between ATL and MED lineages for nonoutlier ($N = 960$, red density curve) and outlier SNPs ($N = 52$, blue density curve). Candidate outliers for between-lineage divergence were identified with BayeScan using a q-value threshold of 0.01. (b) Spatial profiles of mean minor allele frequency (MAF) for nonoutlier ($N = 960$, red dots) and outlier SNPs ($N = 52$, blue triangles), as a function of the distance to SINE by the plateau (km).
4.1 | Cryptic dispersal barriers revealed by introgression clines

Introgression between differentiated lineages can be used as an alternate way to assess within-species genetic and demographic connectivity (Duranton et al., 2019; Gagnaire et al., 2011, 2015). Several marine species display large spatial scale allele frequency patterns that can be attributed to ongoing gene flow between divergent units (e.g., Dahle et al., 2018; Lehnert et al., 2018; Milano et al., 2014; Wennevik et al., 2019). These admixture gradients originate at the edge of contact zones and sometimes attenuate over large distances beyond the contact zones. They are dynamically maintained by continuing gene flow between lineages, combined with a delayed rate of homogenization within the introgressed lineages due to spatially limited dispersal. Selection against introgressed fragments may also contribute to maintaining these admixture gradients (Sedghifar et al., 2016). Spatial discontinuities in admixture gradients indicate local reductions in the spread of introgressed alleles (Gagnaire et al., 2015). Therefore, they can be used to identify cryptic dispersal barriers that are not strong enough to produce genetic differentiation at migration-drift equilibrium.

Here, the analysis of individual ancestry proportions within the Atlantic revealed the existence of a wide admixture gradient extending from southern Portugal to the British Isles. Our closest sampling location from the contact zone with the Mediterranean sea bass lineage (Sines, South Portugal) displayed the highest mean and variance in individual Mediterranean ancestry (mean = 0.11, SD = 0.09). This was mainly driven by the presence of admixed genotypes with significantly higher proportions of Mediterranean ancestry above the background population level (e.g., six individuals between 0.15 and 0.40 in Sines), which most likely represent late-generation backcrosses. Therefore, our results support that ongoing gene flow between the two sea bass lineages contributes to maintain the admixture gradient.

Northward to Sines (SINE), the level of Mediterranean ancestry mostly remained below 5%, displaying a general decreasing trend with increasing latitude. A detailed examination of this admixture gradient revealed the existence of two significant breakpoints located at the tip of Galicia (CORU) and near the Gulf of Saint-Malo (GONB). Two-fold reductions in mean Mediterranean ancestry were observed between sampling sites directly located in the southern versus northern sides of these two breaks. Such a magnitude of change cannot be easily explained by insufficient sampling coverage generating spurious breaks in an otherwise continuous gradient. Instead, it indicates the presence of cryptic barriers that significantly reduce gene flow over relatively short distances, on a scale of a hundred kilometres or less (i.e., the distance between two consecutive sampling sites).

The observed discontinuities in admixture proportions are predicted under spatial admixture models, and can be maintained as long as differences in allele frequencies exist between hybridizing lineages (Gagnaire et al., 2015). For marine species with large population sizes and a planktonic larval stage, it is often assumed that a rapid erosion of allele frequency differences occurs upon secondary contact. However, the existence of genetic barriers to gene flow between Atlantic and Mediterranean sea bass lineages have potentially delayed the dissipation of allele-frequency gradients for a substantial amount of time (Duranton et al., 2018; Tine et al., 2014). Our results support this interpretation since most outlier loci found within the Atlantic were probably explained by the introgression of Mediterranean alleles (Figure 5). Moreover, loci showing delayed rates of introgression between lineages also displayed the largest steps in allele frequencies within the Atlantic, as predicted by theory (Gagnaire et al., 2015; Figure S7).

Our results suggest that migration rates between populations on either side of the identified breaks are too low to ensure demographic coupling, otherwise the breaks would not have been observed. However, quantifying the rate of effective migration across these breaks would require additional data on the intensity of selection acting at the barrier loci and their recombination rate with the neutral markers we used here. This remains challenging, and therefore calls for the use of complementary information on individual movement, as inferred for instance from archival tags (de Pontual et al., 2019) or scale microchemistry (Cambié et al., 2016). Such a coupling between life-history migration and genetic data has the potential to refine our understanding of sea bass demographic connectivity in the northeastern Atlantic.

4.2 | Surfing of introgressed alleles owing to northward range expansion

Northward to the break identified near the Gulf of Saint-Malo, the mean Mediterranean ancestry surprisingly increased towards the northernmost localities of WALE-SGEO (Irish Sea) and DOVE-ANGL-NEUK (North Sea) on both sides of Britain (Figure S5). This result was explained by an increased variance in Mediterranean ancestry among individuals (Figure 3). The mean Mediterranean ancestry in the English Channel is close to 1%, and therefore the increased variance in ancestry in the Irish Sea and North Sea results in increased average ancestry due to border effect (i.e., individual ancestries can deviate more strongly when they lie above versus below the 1% average ancestry).

A possible mechanism underlying the increased variance in Mediterranean ancestry is a recent range expansion in the northern part of the species distribution area. This spatial expansion hypothesis was supported both by the decreasing latitudinal gradient in heterozygosity and the directionality index analysis. A number of direct observations also support the recent increase in sea bass abundance around Britain (Wright, Pinnegar, & Fox, 2020). This is exemplified by the rapid increase of Eastern Channel, North and Irish Seas populations attributed to increased seawater winter temperatures (Henderson, 2007; Pawson, Kupschus, & Pickett, 2007), and the new or expanding sea bass fisheries in UK and Irish seas (Cheung, Pinnegar, Merino, Jones, & Barange, 2012).
In a spatially expanding population, the intensity of genetic drift is amplified at the wave front, which leads to increased variance in allele frequencies compared to the core population (Edmonds, Lillie, & Cavalli-Sforza, 2004; Klopfstein, Curat, & Excoffier, 2006). Putting this in the sea bass context may help explain how range expansion affects Mediterranean ancestry in the northern part of the species range. Alleles of Mediterranean origin segregate at low frequencies at the place where spatial expansion occurs. Therefore, most of these low-frequency Mediterranean alleles are likely to be lost by random genetic drift or to remain at low frequency during range expansion. Some alleles of Mediterranean origin, however, can successfully surf on the wave of advance and reach significantly higher frequencies. These two opposite effects of genetic drift (i.e., small decrease in frequency for most Mediterranean alleles but successful surfing for part of them) is expected to increase the variance in ancestry among individuals living in the expansion zone. Thus, our results are consistent with the surfing of a fraction of the rare Mediterranean alleles, as a consequence of ongoing northward expansion of sea bass populations on both sides of Britain. A more thorough examination of this hypothesis would require denser genome-wide polymorphism data to test the surfing of local ancestry blocks of Mediterranean origin.

A possible alternative explanation for the observed ancestry patterns is selection against weakly deleterious introgressed fragments of Mediterranean origin. Since the rate of introgression is particularly reduced in low-recombining regions of the sea bass genome, selection against introgression is thought to rely on many loci of small individual effects (Duranton et al., 2018). In the large populations of Portugal, Biscay and western Channel, selection against introgressed tracts of Mediterranean origin could be sufficiently efficient to progressively decrease Mediterranean ancestry as the distance to the contact zone increases. By contrast, lower effective population sizes in the northern margins of the species range could reduce the efficacy of selection against introgression, resulting in slightly increased mean Mediterranean ancestry in these regions. The two proposed mechanisms (i.e., surfing of introgressed alleles and reduced efficacy of selection against introgression) are not mutually exclusive and may collectively explain the spatial ancestry patterns documented here.

### 4.3 Applications to conservation and management

Our results mirror previous findings that support the existence of dispersal barriers in the two identified regions of Galicia and Brittany for other marine taxa.

The northwestern coast of the Iberian Peninsula is characterized by complex seasonal oceanographic circulation patterns that may act as potential barriers to dispersal (Varela, Rosón, Herrera, Torres-López, & Fernández-Romero, 2005). Mixed evidence from genetic studies point out that, on the one hand, most marine species lack genetic structure across northwestern Iberia (reviewed in Gomez-Gesteira, Beiras, Presa, & Vilas, 2011). On the other hand, some exceptions have been found such as in *Littorina saxatilis* (Piñeira, Quesada, Rolán-Alvarez, & Caballero, 2008) or in *Fucus ceranoides* (Neiva, Pearson, Valero, & Serrão, 2012), which both lack planktonic dispersive stages. In these species, however, genetic breaks may also be attributed to past secondary contacts.

Brittany represents a less controversial biogeographic transition zone between marine biotas. In some species like *Fucus vesiculosus*, the genetic break occurs near the Ushant thermal front that lies off the coast of western Brittany (e.g., Almeida et al., 2017), and separates cold inshore surface waters from warm stratified offshore waters (Chevallier et al., 2014). Another important transition zone, visible in species assemblages of benthic macrofauna (Gaudin et al., 2018), and contact zones between divergence lineages (Bierne et al., 2003; Jolly, Jollivet, Gentil, Thiebaut, & Viard, 2005), corresponds to the Gulf of Saint-Malo that separates western from eastern English Channel.

Most interestingly, a genetic connectivity study in the marine trematode parasite *Bucephalus minimus*, whose final and most vagile host is *Dicentrarchus labrax*, revealed significant population structure along the northeastern Atlantic coastline (Feis et al., 2015). Population samples of parasites taken from both sides of the two dispersal barriers evidenced here exhibited genetic differentiation, which in turn suggested some level of isolation among their host sea bass populations. By contrast, population genetic studies conducted in sea bass generally failed to detect any significant genetic structure between populations of the Bay of Biscay and English Channel-Celtic Sea (Coscia & Mariani, 2011; Fritsch et al., 2007; Souche et al., 2015). On the other hand, mark-recapture data indicated restricted movement between these two populations, thus supporting their demographic independence (Fritsch et al., 2007). An important component in the maintenance of independent demographic units is the fidelity to winter spawning areas, recently evidenced with tagging data (de Pontual et al., 2019). Limited effective dispersal of larvae across barriers in Galicia and Brittany is also implicitly supported by our findings, and could be explained by the location of spawning areas in relation to the main oceanic current features.

The results presented here are thus broadly consistent with previous findings. They confirm that (a) Atlantic sea bass populations exhibit strong genetic connectivity at neutral markers, which implies that (b) gene flow is high enough to ensure the spread of adaptive variants among populations. (c) However, the number of migrants per generation among populations separated by barriers does not seem sufficiently large to homogenize the latitudinal gradient in Mediterranean ancestry across the whole genome. This supports the existence of already suspected demographically independent units, maintained by behavioral processes and spatially limited larval dispersal. Finally, (d) northern Atlantic sea bass populations seem to exhibit genetic signatures of ongoing northward expansion on both sides of Britain. The demographic independence of these expanding populations with respect to neighboring populations from Celtic Sea and English Channel could not be specifically addressed here and will need further examination. However, the finding of weak
but significant genetic differentiation between Irish-Celtic Seas and North Sea populations, indicates demographic independence over long distances.

These conclusions potentially have direct implications for the management of European sea bass populations in the northeastern Atlantic. The most recent report of the ICES advisory committee for sea bass populations (ICES, 2019) states that spawning-stock biomass has been declining since 2009–2010 and is now just above or below the maximum sustainability yield, depending on the region considered. The need for precise stock assessment is therefore more crucial than ever for a proper management of fisheries. To this end, the spatial delineation of at least three main populations would need to be revised to account for these new findings. Namely, the northern management unit comprising populations from the North Sea, Irish Sea, English Channel, Bristol Channel, and Celtic Sea includes areas that most likely belong to the northern part of the Bay of Biscay population. Moreover, the Bay of Biscay south, which is currently grouped with the Atlantic Iberian Waters, most likely belongs to the Bay of Biscay population.

To conclude, the approach presented in this study supports that introgression gradients consisting of <5% of introgressed ancestry are sufficient to detect cryptic contemporary barriers to gene flow, even using moderate genome coverage. Estimation of migration rates across these barriers and identification of ideal conditions for quantitative assessments will require further development. However, we believe that these will be encouraged by the frequent discovery of admixture in many species, thus expanding the possibilities of studying genetic connectivity in other taxa.

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AUTHOR CONTRIBUTIONS

T.R., and V.R. wrote the proposal, raised funds; T.R. collected specimens with many others; P.A.G. and T.R. conducted and supervised the analysis, with the help of V.R.; T.R., and K.C. managed collections and prepared DNA for genotyping; T.R., and P.A.G. analysed and interpreted the results; T.R., and P.A.G. wrote the paper, with the help of V.R.

DATA ACCESSIBILITY

Raw data, genotypes per site, variables associated with sampling sites have been deposited here: https://doi.org/10.5281/zenodo.3899247. All scripts used have been deposited here: https://github.com/tonyrobinet/introgession.

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