

## RESEARCH ARTICLE

# MHC class I- $\alpha$ population differentiation in a commercial fish, the European sea bass (*Dicentrarchus labrax*)

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**Abstract**

Identifying population structuring in highly fecund marine species with high dispersal rates is challenging, but critical for conservation and stock delimitation for fisheries management. European sea bass (*Dicentrarchus labrax*) is a commercial species of fisheries and aquaculture relevance whose stocks are declining in the North Atlantic, despite management measures to protect them and identifying their fine population structure is needed for managing their exploitation. As for other marine fishes, neutral genetic markers indicate that eastern Atlantic sea bass form a panmictic population and is currently managed as arbitrarily divided stocks. The genes of the major histocompatibility complex (MHC) are key components of the adaptive immune system and ideal candidates to assess fine structuring arising from local selective pressures. We used Illumina sequencing to characterise allelic composition and signatures of selection at the MHC class I- $\alpha$  region of six *D. labrax* populations across the Atlantic range. We found high allelic diversity driven by positive selection, corresponding to moderate supertype diversity, with 131 alleles clustering into four to eight superotypes, depending on the Bayesian information criterion threshold applied, and a mean number of 13 alleles per individual. Alleles could not be assigned to particular loci, but private alleles allowed us to detect regional genetic structuring not found previously using neutral markers. Our results suggest that MHC markers can be used to detect cryptic population structuring in marine species where neutral markers fail to identify differentiation. This is particularly critical for fisheries management, and of potential use for selective breeding or identifying escapes from sea farms.

**KEYWORDS**

aquaculture, fisheries, non-neutral markers

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## INTRODUCTION

With the decline of many traditional fisheries, accurate fish stock management has never been so critical (Beddington et al., 2007). However, identifying population structuring and genetic differentiation in marine populations using molecular markers can be challenging, particularly for highly fecund species with high dispersal rates and complex life cycles (Hedgecock et al., 2007). The use of large population genomic data sets and markers influenced by selection are increasingly revealing fine population structure and patterns of reproductive isolation even in highly dispersive marine species, with important implications for their conservation and management (Gagnaire et al., 2015). However, in many cases, lack of information about population structure results in a pragmatic approach to management (ICES, 2021) and a mismatch between biological and management units (Reiss et al., 2009), which can exacerbate the decline of stocks. Commercial fisheries are already contributing to the homogenisation of genetic diversity of marine fish species (Gandra et al., 2021) and ignoring fine scale local adaptation can further result in the loss of functionally important biodiversity (Limborg et al., 2012).

European seabass (*Dicentrarchus labrax*) is a traditionally important species in terms of commercial and recreational fisheries, and has more recently become a key species for aquaculture (Vandeputte et al., 2019). The extent of population structuring in European seabass is unclear, and in the Atlantic area has been arbitrarily divided by the International Council for the Exploration of the Sea (ICES) into four stocks: Stock West Ireland/West Scotland; Stock Irish Sea, Celtic Sea, English Channel, and North Sea; Stock Bay of Biscay; and Stock Iberian Peninsula (De Pontual et al., 2019; European Parliament, Directorate-General for Internal Policies of the Union et al., 2016). The seabass stock found in the English Channel, Celtic and Irish Seas, and North Sea is characterised by slow growth and late maturation and its productivity is greatly influenced by sea water temperature (Walker et al., 2020). This stock has declined markedly since 2010, despite stringent quotas and fishing bans (Walker et al., 2020), mainly due to overfishing and low recruitment since 2008, but also due to the influence of long-term climate change and local environmental factors that affect mostly the early developmental stages (Bento et al., 2016). Tagging studies indicate that there is strong site fidelity (Pawson et al., 2008) and frequent migrations between the English Channel and the Bay of Biscay areas, and between the North Sea and the English Channel, highlighting the need for more accurate information on fine population structuring, necessary for correct stock management (Pawson et al., 2007). For example, the Irish stock is exploited separately from the Atlantic stocks although its distribution spans more than one ICES division (ICES, 2021). Existing studies based on mitochondrial DNA suggest that there is population

structuring, not consistently supported by neutral markers. Mitochondrial DNA indicates that Atlantic and Mediterranean seabass populations are differentiated as a consequence of their isolation during the Pleistocene, that resulted in divergence in haplotype frequencies (Lemaire et al., 2005). However, a post-glacial secondary contact is thought to have eroded this divergence at the level of nuclear markers (Duranton et al., 2018), while differential introgression resulted in the creation of genomic islands of differentiation (Tine et al., 2014). Within the Atlantic seabass stock, there are three mitochondrial DNA lineages distributed predominantly in the Bay of Biscay (Atlantic 1; from where the Mediterranean lineage could have originated), European coast (Atlantic 2), and the British Islands and Norway (Atlantic 3) (Coscia & Mariani, 2011). In contrast, there is limited regional structuring based on nuclear markers (microsatellites and SNPs), with the exception of the populations in the south-eastern range of the Atlantic lineage (Portugal and Morocco) that display some Mediterranean influence in their SNP allelic composition (Souche et al., 2015). The allele distribution of markers under selection, such as allozymes or the somatolactin gene, allowed to identify some differentiation between the Bay of Biscay and the southern North Sea (Quére et al., 2010) and at the local scale (Castilho & McAndrew, 1998), suggesting that markers under selection may prove better candidates to identify finer scale structuring for this species than neutral markers.

The genes of the major histocompatibility complex (MHC) are some of the most studied and highly polymorphic genes in vertebrates (Edwards & Hedrick, 1998). MHC genes encode for proteins that present antigens to T-cells, triggering the adaptive immune response (Janeway et al., 2004) and most of their diversity is concentrated in the region that binds antigens from pathogens, the peptide binding region (PBR) (Hedrick & Kim, 2000). Polymorphism within the PBR determines to which pathogens an individual can respond (Radwan et al., 2020). PBR diversity is maintained by pathogen selection (Eizaguirre et al., 2012; Slade & McCallum, 1992) and in some species by mate choice (Consuegra & Garcia de Leaniz, 2008; Milinski, 2006), and can be influenced (directly or indirectly) by local environmental factors such as temperature (Dionne et al., 2007). We developed an Illumina sequencing-based protocol to genotype the PBR of the European seabass MHC class I- $\alpha$  gene, and investigated its potential for detecting fine scale population structuring and signatures of local selection pressures across the species' Atlantic range.

## METHODS

Genomic DNA was extracted using the Qiagen DNeasy Blood & Tissue Kit from 62 wild seabass individuals (caught between 2013 and 2015) from six areas of the

North Eastern Atlantic (Biscay: ICES rectangle 20 E8,  $n = 6$ , Dover: ICES 30 F1,  $n = 7$ , Irish Sea ICES 34E4,  $n = 5$ , Celtic Shelf: ICES 28 E1,  $n = 22$ , North Sea: ICES 38 E9,  $n = 3$ , and Portugal: ICES 4 E1,  $n = 19$ ). Fish were genotyped using a custom designed primer pair: sbMHC\_F2 5' CTGGAGTCCCAAACCTCC 3', sbMHC\_R2 5' AGGTGGACACCTCCAGTTTG 3', amplifying a 241-bp fragment of the protein coding sequence region of the MHC class I- $\alpha$  gene (Methods S1).

Libraries were prepared in a two-step PCR protocol, using Platinum™ Hot Start PCR Master Mix (2 $\times$ ; Thermo Fisher Scientific). Initially, 2.5  $\mu$ l of DNA per individual was amplified in a 25- $\mu$ l reaction (12.5  $\mu$ l taq, 0.5  $\mu$ l each of forward and reverse primer, 9  $\mu$ l water), using the above primers with Illumina overhang adapter sequences added (Illumina, Inc., 16S Metagenomic protocol). PCR amplification consisted of the following: 2 min at 94°C, 28 cycles of 30 s at 94°C, 30 s at 55°C (locus specific annealing temperature), 1 min 72°C (extension). Subsequently, 2  $\mu$ l of product from the initial reaction was used as template in a second PCR (12.5  $\mu$ l taq, 1.25  $\mu$ l each of a sample-specific combination of Nextera XT adapters (Index Kit v2; Illumina, Inc) and 10  $\mu$ l water) and eight cycles of this second PCR was run with identical conditions to the first. A 5- $\mu$ l aliquot of each indexed PCR product was pooled and the final pool was cleaned using Agencourt AMPure XP beads (Beckman Coulter), using a 1:1 ratio of beads to pooled template. The pool was then quantified using quantitative PCR (NEBNext® Library Quant Kit for Illumina®, protocol according to manufacturer's instructions). Blanks (where molecular grade water was added in the place of template) were used to check for any cross-contamination between samples. Pair-end sequencing was carried out on the MiSeq platform, using a V3 reagent kit (Illumina, Inc.). Samples were sequenced on two different MiSeq runs (run 1 = 36 individuals, run 2 = 26 individuals, plus six re-sequenced individuals from run 1 to test repeatability).

## Bioinformatics and data processing

De-multiplexed raw pair end sequences were processed using the ampliSAT suite of tools (Sebastian et al., 2016) (available at: <http://evobiolab.biol.amu.edu.pl/amplisat/>). For full methodology see Methods S1. Any samples with less than 1000 reads coverage were discarded from downstream analysis. Parameters were optimised to ensure maximum reliability of genotyping for the entire dataset. Filtering per amplicon frequency was set at 1% (the minimum per amplicon frequency for variants that appeared in both replicates). The minimum dominant frequency was set at 10%, in order to keep true similar variants, whilst removing high frequency motive specific errors, meaning that only sequences with a frequency below that threshold were clustered with a parental sequence (Biedrzycka et al., 2017; Sebastian et al., 2016).

A maximum of 16 alleles per individual was considered, based on the *D. labrax* MHC class I alleles identified by (Pinto et al., 2013) from cDNA cloning, which indicated a minimum of six and maximum of eight loci per individual. The following manual filtering steps were then applied to the AmpliSAS output: all singletons (variants that appeared in one fish only) and any sequences with <10 reads overall were removed from downstream analysis to control for the presence of sequencing errors (Migalska et al., 2019). Repeatability (number of identical variants identified in both PCRs) in the six re-sequenced samples was then calculated following (Biedrzycka et al., 2017), where the number of identical variants are divided by the total number of alleles called in both replicates in a given individual to ascertain a repeatability proportion.

## Polymorphism analysis

Initial screening of aligned variants was carried out in DnaSP (Rozas et al., 2017), using the 'polymorphism data' tool and the Codon-based Z-Test of Selection (MEGAX, Nei-Gojobori method). The seabass amino acid sequences were then aligned with other teleost MHC class I sequences of the U lineage described in Grimholt et al. (2015), and a neighbour-joining tree based on  $p$ -distance and pairwise deletion was constructed using 1000 bootstrapping iterations to assess their relationship.

## Recombination and selection analysis

Recombination was assessed using single break point analysis using the '010021' model of nucleotide substitution. Positive and negative selection were assessed using the HyPhy package (datamonkey.org) using three selection models. To detect pervasive positive/diversifying selection, FEL (fixed effects likelihood; Kosakovsky Pond & Frost, 2005) and FUBAR (Fast, Unconstrained Bayesian AppRoximation; Murrell et al., 2013) were used. To detect episodic positive/diversifying selection, MEME (mixed effects model of evolution; Pond et al., 2006) was used (for full methodology see Methods S1). The results of the three models were combined to identify positively selected sites for downstream analysis. A site was only considered to be under positive selection if it was selected by all three models. Recombination sites were compared with positively selected sites to avoid confounding effects, and, as they did not overlap, positively selected sites were analysed without removal of recombination sites.

In addition, alleles were analysed for conservative or radical amino-acid changes using TreeSAAP (Woolley et al., 2003), using a neighbour joining tree with 1000 bootstrap replicates (amino acid sequences). A sliding window value of 1 was used to obtain codon level magnitude of change scores. Significant differences in the top

two magnitude categories (7 and 8) were considered as 'radical' changes.

## Supertype identification

The amino acids of positively selected sites as identified by FEL, FUBAR and MEME were assigned five z-descriptors (Doytchinova & Flower, 2005): z1 (hydrophobicity), z2 (steric bulk), z3 (polarity), z4 and z5 (electronic effects) (Sandberg et al., 1998) and translated into a mathematical matrix. Gaps in the alignment of positively selected sites were assigned zeros in the matrix.

Supertype clustering was performed in discriminant analysis of principle components (DAPC) with the 'adegenet' package (Jombart, 2008) in R (Version 4.0.4; R Core Team, 2019). Cluster values between the minimum number of clusters after which Bayesian information criterion (BIC) decreased by a negligible amount, and the lowest BIC value were then evaluated using stepwise DAPC runs, plotted, and inspected visually. The number of PCs was chosen using the function 'optim.a.score' and kept consistent for each DAPC analysis.

## Loci identification via allele clustering

Phylogenetic trees were generated using the following methods: (1) neighbour joining (codons), using *p*-distance with 1000 bootstrap support (MEGAX, Kumar et al., 2018); (2) Ward's (Euclidian distance) using the z-matrix (Doytchinova & Flower, 2005); (3) UPGMA using the z-matrix, with three distances: Euclidian, Cosine and Pearson Correlation algorithms (PAST 4; Hammer et al., 2001). In addition, a Neighbor-Net (1000 bootstrap support) was generated in SplitsTree4 (Huson & Bryant, 2006), because networks may be more appropriate than trees for identifying relationships between alleles where duplication, recombination and gene conversion are common (Biedrzycka et al., 2017).

The performance of all methods in identifying loci was evaluated by testing whether any given individual exhibited more than two variants/allele per cluster. This was based on the assumption that, as a diploid organism (Felip et al., 2001), no individual should exhibit more than two alleles at a given locus.

## Population analysis

Fish from the two regions with the largest sample size, Portugal ( $n = 19$ , ICES rectangle 4 E1) and Celtic Shelf ( $n = 22$ , ICES rectangle 28 E1), were examined to identify private alleles to each region and potential population structuring. To further explore population structure between these two populations, we used DAPC (Jombart, 2008; Jombart & Collins, 2015), using a presence/absence

matrix where each column represented an allele, and each row an individual fish. Presence/absence of alleles was then recoded as frequencies, where alleles were represented as a proportion of the total alleles for each individual. This matrix was then used to calculate pairwise Jost's  $D$  (pairwise  $D_{\text{est}}$ , [Jost, 2008], 'mmod' package in R [Winter, 2012]). Jost's  $D$  is appropriate for genes that exist in multiple copies with unknown locus affiliation of alleles (Jost, 2008; Lighten et al., 2017). We used the function 'confusionMatrix' in the 'caret' package (Kuhn, 2009) in R, to assess the accuracy of the assignment of individuals to their stock based on DAPC.

## RESULTS

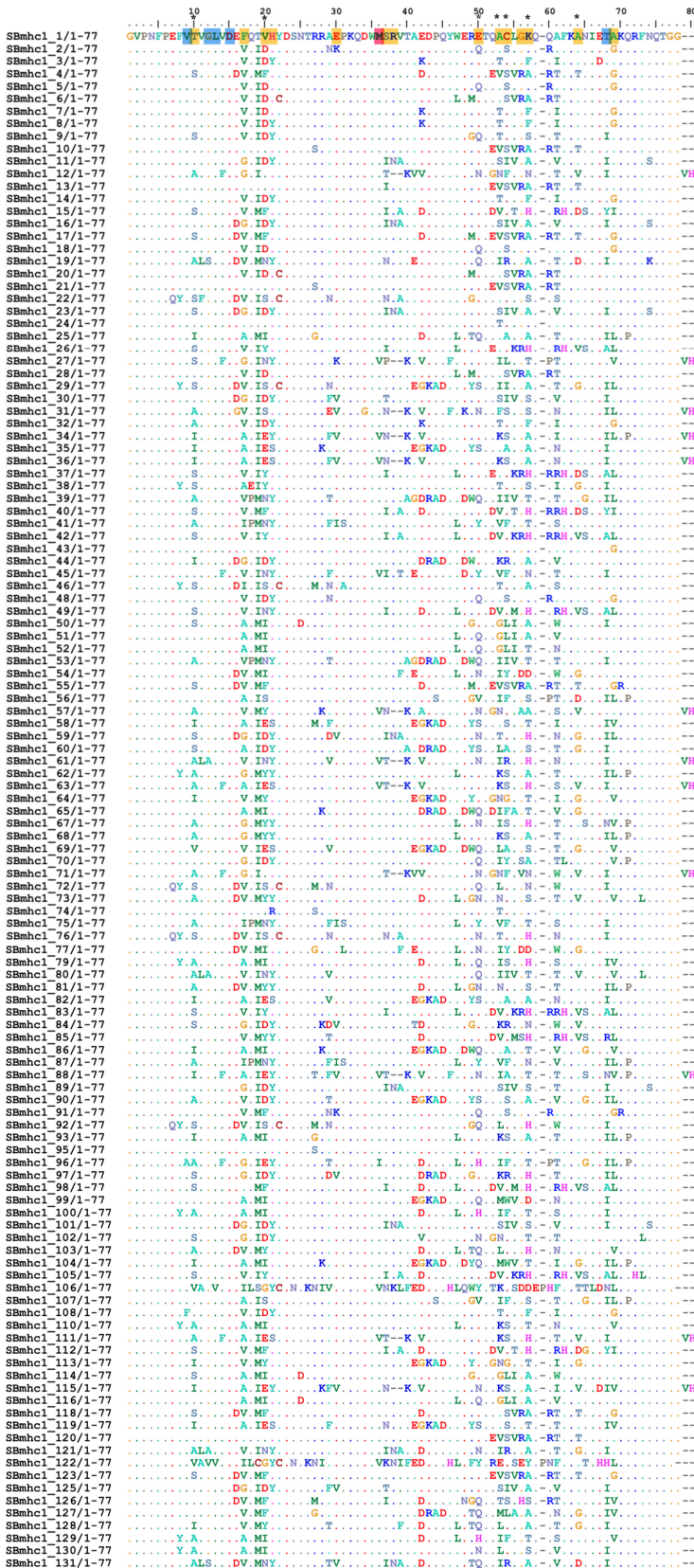
### MHC class I $\alpha$ polymorphism in seabass

A total of 62 individual fish were sequenced with sufficient depth for downstream analysis (>1000 reads). Samples were capped at a maximum of 5000 reads for AmpliSAT processing reasons and, after processing (prior to removal of singletons), had an average coverage of 3140 reads (STDEV = 1346 reads). Singleton removal reduced the number of variants/alleles from 332 to 133, and removal of two variants <200 bp in length resulted in 131 variants retained in the dataset for selection analysis. None of the variants was an exact match to already published alleles (Pinto et al., 2013); however, SBmhc1\_111 has only one nucleotide difference to HQ290120.1/1-825 clone\_15 and HQ290116.1/1-825 clone\_1 (which are identical in the target region). Repeatability (number of identical variants identified in both PCRs) in the six re-sequenced samples was estimated as 0.89 (based on those samples as processed with the whole dataset). Individual copy number varied between seven and 16 alleles (mean = 12.6, mode = 13).

The 131 different nucleotide sequences identified corresponded to 123 unique amino acid sequences (Figure 1). An indel of two amino acids resulted in sequences of two lengths, 240 bp and 234 bp. From the 240 sites, 137 were variable and there were a total 228 mutations (Eta). Nucleotide diversity per site ( $P_i$ ) was: 0.13216. The codon-based Z-test of selection indicated the presence of positive selection ( $p = 0.03$ ,  $Z = 1.85$ ).

### Recombination and selection analysis

Strong support for recombination was found at position 109, inferred from both Akaike information criterion and BIC with 100% support. MEME analysis indicated that there was possible episodic positive/diversifying selection at 23 codon sites (Table S1, Figure 1). FEL analysis indicated that there was pervasive positive/diversifying selection at 19 sites and pervasive negative/purifying selection at six sites (Table S1). FUBAR found evidence



**FIGURE 1** Codon alignment (Clustal W) of all alleles sequenced in this study. Yellow highlighting indicates pervasive/episodic positive/diversifying selection detected by mixed effects model of evolution (MEME), fixed effects likelihood (FEL), and Fast, Unconstrained Bayesian AppRoximation (FUBAR). Blue highlighting indicates negative episodic negative/purifying selection detected by both FEL and FUBAR. Pink highlighting indicates breakpoint (nucleotide 109) detected by single break point (all analyses performed using datamonkey.org tools). Asterisks represent codons which align to residues under selection at the peptide binding region of the human HLA-A2 gene (Grimholt et al., 2015)

of episodic positive/diversifying selection at 20 sites episodic negative/purifying selection at eight sites. In total, 14 sites were identified as being under positive selection by all three models (Table S1; Figure 1) and a further five

sites that were identified as under negative selection by both FUBAR and FEL (Table S1). Of the positively selected sites, seven coincided with residues contributing to the peptide binding pockets of the human HLA-A2 gene,

based on the alignment of the sequences with those from (Grimholt et al., 2015) (Figure 1).

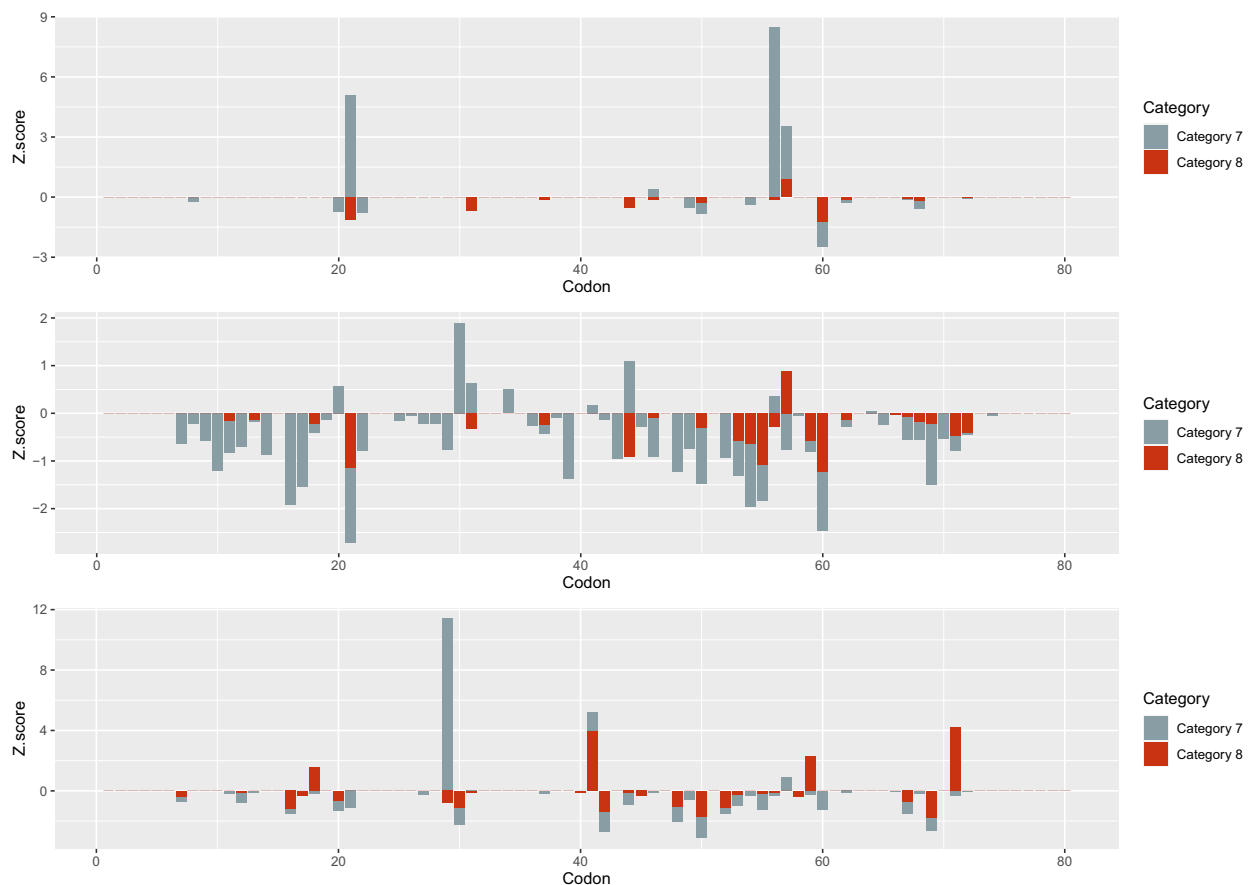
### Amino acid properties under selection (TreeSAAP)

TreeSAAP analysis identified significant deviations from neutral expectations (categories 7 and 8, Table S2, Figure 2), within the whole sequenced fragment, for the following physicochemical amino acid properties: ‘power to be at the c-terminus’ (energy potential of the c-terminus of an  $\alpha$  helix to interact with other residues); ‘power to be at the middle of the  $\alpha$ -helix’ (energy potential of the middle of an  $\alpha$  helix to interact with other residues); ‘mean r.m.s. fluctuation displacement’ (the ability of a residue to change position in three-dimensional space); ‘compressibility’ (the contribution of a residue to the local density of protein secondary structures); ‘solvent accessible reduction ratio’ (the reduction ratio of cross peak intensity when residues are or are not irradiated with aliphatic protons); and ‘thermodynamic transfer hydrophobicity’ (the difference in the solubility of amino acids in water and ethanol;

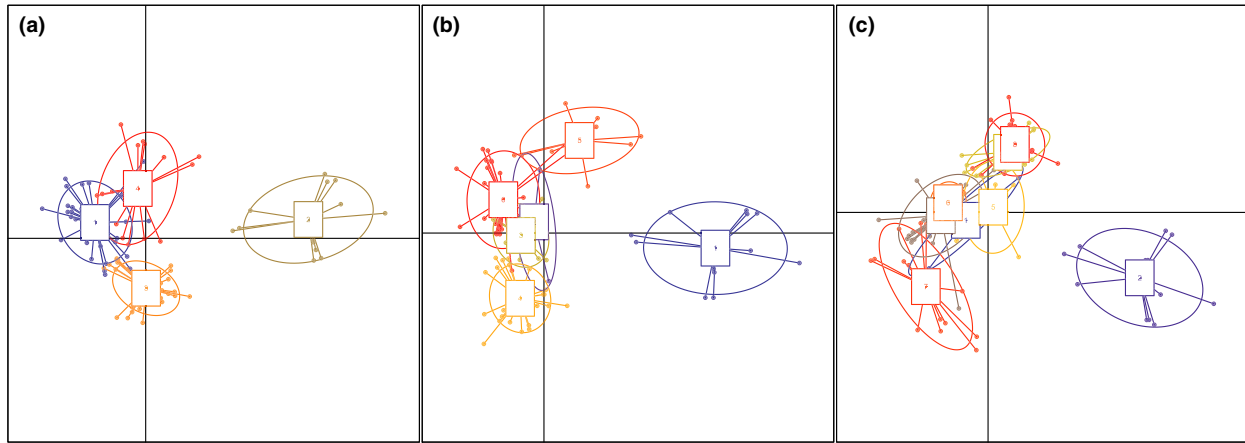
Figure 2, Table S2). This indicates that selection pressure on these properties have driven changes at this region (Woolley et al., 2003).

### Supertype and loci clustering

BIC values indicated that there were four main supertype clusters (Figure 3) before BIC values decreased by a negligible amount (Figure S1). We also explored a finer scale clustering, up to eight clusters, which represented the lowest BIC value (Figure 3). Assignment of potential alleles to individuals indicated that there could be between nine and 18 loci, depending on the method used (Figures S2, S4–S8; Table S4) but none of the clustering methods resulted in all individuals in the dataset having only two alleles in a cluster (Table S4), even when fine clustering (18 clusters) was applied. Seabass sequences clustered closer to class I- $\alpha$  lineage I than to the other lineages (Figure S3). Two sequences, SBmhcl\_106 and SBmhcl\_122, clustered with lineage I sequences from salmonids (Atlantic salmon, brown trout, and rainbow trout), and close to Medaka sequences, also from lineage I (Figure S3).



**FIGURE 2** Z-scores relating to phenotypic amino acid properties/traits (Power to be at the middle of the  $\alpha$ -helix, power to be at the c-terminus and compressibility). Categories represent magnitude of amino acid property change at nonsynonymous residues on a site-by-site basis. Class 8 represents changes of the highest magnitude, followed by class 7. Low and 0 z score values represent areas of conservation in terms of amino acid properties, in contrast to high values, which represent areas of selection



**FIGURE 3** (a–c) Clustering of major histocompatibility complex alleles in *Dicentrarchus labrax*, based on the physicochemical properties of amino acids (Z scores) of positively selected sites, using discriminant analysis of principal components. Each point represents the positioning of each major histocompatibility complex allele within corresponding number of discriminant functions (Eigenvalues insets) and the circles represent the supertypes

## Population analysis

We found 30 unique alleles (representing 29 unique amino acid sequences) in the Celtic Shelf population that were not represented in the Portuguese population and 22 alleles (21 unique amino acid sequences) unique to the Portuguese population (Figure 4a), including the two alleles with an insertion at codon 59 in the alignment (SBmhc1\_106 and SBmhc1\_122). Private alleles were present across the phylogenetic tree, but Celtic shelf alleles were more frequent in recently diverged clusters than those from Portugal (Figure 4a). Most unique alleles appeared in only one or two individuals in the respective populations (Table S4), with the exception of allele SBmhc1\_25 and SBmhc1\_38, which appeared in six and three Portuguese individuals respectively, and SBmhc1\_54, present in three Celtic Shelf individuals. There was no difference in the number of alleles per fish between the two populations (mean of Celtic population = 12.64 alleles per individual, mean of Portuguese population = 12.79 alleles per individual; Welch two-sample  $t$ -test  $t = -0.26442$ ,  $df = 38.999$ ,  $p = 0.7928$ ). DAPC analysis indicated low differentiation in allele frequencies as a whole (Figure 4b) between the two regions ( $D_{est} = 0.05$ ), with most of the variance described by the first principal component. Based on all the alleles, we were able to assign 75% of the fish (31/41) to their region of origin (Celtic Shelf or Portugal). The accuracy of the assignment was significantly greater than chance (aAccuracy = 0.756; 95% CI: 0.597, 0.8764;  $p = 0.003$ ).

## DISCUSSION

The high variability of MHC genes is known to be driven by natural and sexual selection and, for this reason, they represent good candidates to assess fine-scale structuring potentially derived from local selective pressures

(Larson et al., 2019). We found that the MHC class I- $\alpha$  gene in the Atlantic seabass displays high allelic diversity, evidence for selection and, importantly, potential for detecting fine scale structuring within stocks largely considered panmictic (ICES, 2021). Our study represents the first population study of MHC class I in seabass and indicates that, as for other fishes, the *D. labrax* MHC class I- $\alpha$  locus has undergone strong positive selection and recombination, while maintaining intermediate copy numbers and relatively few supertypes. This is similar to the diversity of MHC class II in guppies, where supertypes are maintained by balancing selection and consist of groups of alleles with similar functionality that are subject to positive selection (Lighten et al., 2017).

Most seabass MHC-I sequences clustered close to the lineage I of the U-type classical alleles from different fish species, two of them grouping with salmonid alleles (Grimholt et al., 2015). Close clustering between seabass and salmonid alleles might indicate trans-species polymorphism (the occurrence of similar alleles in related species) typical of MHC genes (Klein et al., 1998), but this must be interpreted with caution as it is based on the  $\alpha$ -1 domain only and the bootstrapping support was relatively low. We found an average of 13 copies per individual in *D. labrax*, which is similar to other teleost species with U lineage genes, such as the African moonfish, *Selene dorsalis* and Sand roller *Percopsis transmontana* (Malmström et al., 2016). In general, the teleost MHC class I genes are highly polymorphic, with copy number varying between families and species (Malmström et al., 2016). For instance, Atlantic salmon (*Salmo salar*) has a single MHC I locus (Grimholt et al., 2003), while three-spined stickleback, *Gasterosteus aculeatus* possess between three and nine alleles per individual, which are difficult to assign to loci probably due to high allele similarity (Aeschlimann et al., 2003). In contrast, most gadoids have lost classical MHC II genes and possess a largely expanded number of MHC I copies; for example,





Atlantic cod (*Gadus morhua*) which has approximately 100 (Star et al., 2011). The observed diversity in *D. labrax* was, therefore, within the range expected for a species with a classical MHC II and fitted well with the previously predicted six to more than eight MHC-I loci for the species (Pinto et al., 2013), although none of them were an exact match to those previously published, likely due to the different origin of the populations and the high variability of the MHC genes. However, as our analysis was based on DNA sequences and not on expressed alleles (mRNA), we could not identify the exact number of loci and it is possible that some of the sequences could represent pseudogenes.

Positive selection was identified at 14 polymorphic sites across the 80 codons in this study, some of them coinciding with PBR sites previously defined in fish based on the alignment with human sequences (Grimholt et al., 2015) (Figure S3). This level of selection is comparable to other fishes (Wegner, 2008) for example sockeye salmon (*Oncorhynchus nerka*) where four codons are under positive selection in a 32-codon fragment (McClelland et al., 2011). High rates of positive selection are almost always observed in MHC alleles, driven by parasite-mediated balancing selection or sexual selection (Bernatchez & Landry, 2003). In addition, recombination can also act as a significant force to rapidly increase diversity in MHC alleles (Consuegra et al., 2005), and we found strong support for a breakpoint at position 109 bp, possibly contributing to the observed allelic diversity.

While codon-based selection models give an indication of the areas of the gene under selection, the ability to bind antigens from specific pathogens is determined by changes in the shape of the cleft of the PBR (Dionne et al., 2007). Nucleotide differences, or even amino acid differences, may not directly translate to protein binding differences, and additional alleles may not necessarily confer the ability to respond to a greater range of pathogens, unless the changes in the protein also change the binding properties of that allele (Ellison et al., 2012; Lighten et al., 2017). The analysis of functional differences based on binding properties analysis indicated that the energy potential of the C-terminus, the position of amino acids at the middle of an  $\alpha$  helix (where they can interact with other residues), and the ability for a residue to change position in three-dimensional space were under selection, supporting the role of selective pressures (Doytchinova & Flower, 2005; Lighten et al., 2017). Alleles within a supertype are predicted to bind similar antigenic 'supermotifs' (Phillips et al., 2018). We identified between four and eight clusters, although the precise number of clusters is not clear. Clustering can be challenging when loci share identical alleles, there are null alleles or, as found here, copy number varies between haplotypes (Huang et al., 2019). Even with an upper estimate of eight clusters, we found relatively low supertype diversity in comparison to other fishes, despite a relatively diverse allele pool of 131 alleles. For instance,

in guppies, 66 alleles were clustered to 13 supertypes (Smallbone et al., 2021). This could be the result of convergent evolution resulting in a relatively small number of overlapping peptide-binding motifs, as for the highly diverse human class I alleles that may be clustered into as few as nine supertypes (Sidney et al., 2008).

Despite loci remaining undefined, we observed allelic differentiation in the two seabass populations we compared, confirmed by DAPC analysis, although private alleles were detected in relatively few individuals within each region. Overall, the differentiation between regions was small but, based on the MHC sequences identified here, we were able to assign 75% of the fish to their region of origin (Celtic Shelf or Portugal) with high confidence. Previous studies of Eastern Atlantic seabass using microsatellite and SNP analysis failed to identify consistent population structuring (Souche et al., 2015), and tagging studies have also shown that *D. labrax* are capable of swimming large distances, e.g. 1200 km within 2 months of tags being deployed, and migrate considerable distances offshore to spawn (Pawson et al., 2007). It had therefore been assumed that Eastern Atlantic seabass formed one single panmictic population. Seabass seems to display a shallow population structuring based on neutral markers but, as for other marine species (Milano et al., 2014), a finer level of structuring resulting from local selective pressures can be detected when markers under selection are used. The difference in private alleles between the Celtic shelf and Portuguese uncovered population structuring not observed using neutral markers (Souche et al., 2015), and indicates the potential for MHC-I to detect fine-scale population variation. Markers in the MHC region can be particularly good for showing fine scale differentiation, as changes can accumulate relatively quickly and are affected by local selection (Consuegra et al., 2005), and genome-wide high-throughput methods (Griot et al., 2021; Penaloza et al., 2021) can help identify additional markers under selection (Carreras et al., 2017). Our study provides the first detailed analysis of MHC class I in seabass, a species that supports important commercial and sport fisheries, as well as aquaculture. Our approach provides a quick tool to screen the highly variable MHC region of Atlantic seabass for fisheries management purposes, that can be adapted to other marine species. Moreover, given the increasing importance of seabass aquaculture (Vandeputte et al., 2019), and the potential risk that escapes pose to wild populations (Arechavala-Lopez et al., 2018), the same approach employed here could be used for genotyping farmed species, potentially for selective breeding for disease resistance (Pawluk et al., 2019) and/or to monitor the incidence of escapees from fish farms (Monzón-Argüello et al., 2013).

Environmental complexity has the potential to create refugia for marine species that may result in local adaptation and create cryptic population structuring, essential for the long-term persistence of exploited stocks

(Midway et al., 2018). In many marine species with high fecundity and dispersal rates, this fine population structure can only be identified by markers under natural selection (André et al., 2011; Jorde et al., 2018; Lamichhaney et al., 2012). Based on our initial results from sea bass, we propose that MHC markers can be used for the management of marine species with cryptic population structure, for which preserving their fine population structuring is needed to maintain their functional biodiversity (Limborg et al., 2012).

## CONFLICT OF INTEREST

The authors declare no conflict of interest.

## ACKNOWLEDGEMENTS

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## ETHICAL APPROVAL

All work was carried out with approval from Swansea University Ethics Committee (Reference number SU-Ethics-Student-280621/4326).

## DATA AVAILABILITY STATEMENT

Sequences have been submitted to GenBank and are accessible under Accession numbers [MZ466411-MZ466541](#).

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