Genetic structuring in farmed and wild Gilthead seabream and European seabass in the Mediterranean Sea: implementations for detection of escapees

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Abstract – Microsatellite markers were used to investigate the genetic structure of the two most important cultured fish in the Mediterranean Sea, the gilthead seabream (Sparus aurata) and the European seabass (Dicentrarchus labrax), from two (one wild and one farmed) populations in Western Mediterranean (Spain) and from two (one wild and one farmed) populations Eastern Mediterranean (Greece). All populations were in Hardy-Weinberg disequilibrium. Interestingly, wild and farmed populations for both species from Greece were genetically differentiated and could be distinguished from each other. We used Bayesian methods for cluster analysis of farmed and wild populations. Our analysis has implications for the identification of escapees from fish farms to the wild.

Keywords: Seabream / seabass / genetic structure / escapees

1 Introduction

Gilthead seabream (Sparus aurata) and European sea bass (Dicentrarchus labrax) are the two most important species for finfish aquaculture in the Mediterranean Sea. Both species are found in the Mediterranean Sea and in the eastern North Atlantic, with the European sea bass having a more northerly distribution up to Norway. They support commercial fisheries in some areas, and are well-established as aquaculture species, particularly in the Mediterranean Sea. The two species are usually reared in the same structures and the same farms. The majority of the fish reared in Europe are maintained in floating sea cages, into near shore locations, although few farms in Spain and Italy are using salt marshes to produce fish in ponds (ICES, 2006). With the global number of cultured fish being in a very high proportion compared to the wild populations (mainly in areas where fish farms are located, a high number of escapes might have the potential to lead to displacement of wild population in some specific sites. This is the reason why mitigation strategies based on sound scientific criteria in relation to both species need to be prepared and rapidly taken into consideration. ICES (2015) suggests that for these two non-highly migratory fish species with small localized populations, studies seem to be of particular importance to consider the status of the natural stocks in the area, the potential genetic, trophic and behavioral interactions, and, foremost and specifically, the development of methods for recovery of escaped fish in the event of large-scale escapements.

Depending on the strategy of the fish farms to manage their broodstocks, farmed populations are expected to differ from the local natural populations for three main reasons. First, breeders may be wild-caught fish, according to a common practice for re-stocking the farms. These breeders represent a small sub-sample of the wild population, thus farmed stocks are expected to contain less diversity than the wild ones. This has been observed in several species, such as salmon (Norris et al., 1999; Skaala et al., 2004), seabream (Karaiskou et al., 2009; Loukovitis et al., 2011) and sea trout (Bernas et al., 2020) (but see de Oliveira et al., 2018). Second, in some cases, farms located in different geographic regions exchange fry and the farmed fish have different genetic backgrounds from the wild fish outside cages. For instance, due to increased demand for juveniles (fingerlings) in the Eastern Mediterranean countries (Greece, Turkey and Croatia), there are often imports of fry from the Western Mediterranean countries such as France and Spain (Barazi-Yeroulanos, 2010; Segovic-Bubic...
et al., 2011). Finally, various programs of selective breeding already in place for multiple consecutive generations, might reduce in the long term the heterozygosity of the farmed compared to the wild populations (for example see D’Ambrosio et al., 2019).

Due to technical or operational failures, farmed fish can escape in natural environment (Dempster et al., 2007). The impact of escaping fish to local populations has not been assessed in detail and there are different lines of evidence, which suggest that escapes might have important effects on native populations (Arechavala-Lopez et al., 2018). A potential escaping event entails the risk of genetic admixture of local populations with fish from the aquaculture facilities. The foreign DNA can genetically blight local wild gene pool and damage local adaptations which have been reported in many fish species, even under high gene flow (see Barth et al., 2017). For example, farmed fish strains can be genetically different from the wild populations in traits such as growth rate, physiology, and behaviour (Glover, 2010). Then, the hybrids from crosses between farmed and wild fish can have lower fitness than the wild fish in natural environments, as it has been reported in several cases (McGinnity et al., 1997; Fleming et al., 2000; McGinnity et al., 2003; Reed et al., 2015; Skaala et al., 2019). It has been reported that gilthead seabream escapees can survive successfully to the natural environment posing a risk of hybridization with individuals from the wild (Segvic-Bubic et al., 2018). The risk from this genetic contamination has not yet been fully assessed (Yang et al., 2019).

Identifying the escapees could be an efficient approach for mitigating the effects of genetic admixture of farmed with wild populations. Such identification has been applied by using physical marking of the farmed fish with tags (Glover, 2010). This method is prohibitively expensive and has animal welfare concerns. An alternative method which has been used for salmon, Atlantic cod and rainbow trout is the identification of escapees using the DNA-stand-by method (Glover, 2010). This method assigns an individual to a certain population — in this case farmed or wild — according to its genetic profile. The genetic profile of the individuals is usually retrieved using genetic markers such as microsatellites, AFLPs and single nucleotide polymorphisms (SNPs). The assignment of an individual to a certain population requires prior good knowledge of the genetic structure of the species, and its reliability depends on the level of divergence between populations/stocks and the number of markers used. If the genetic distance between populations is low or/and only few genetic markers have been used, then it is difficult to assign an individual to a specific population. The advantage of a DNA-based method compared to other methods is that it can identify individuals even if they have escaped very early in their life or even if the eggs and/or gametes have escaped from net-cages into the wild and probably drifted for long distances. Differences in morphological characteristics, skeletal shape and chemical composition of the body between farmed and wild fish, can also be used to identify escapees (Arechavala-Lopez et al., 2012). However, these methods are based on characteristics that are affected by the environment. If farmed individuals have escaped early in their development, they would be hardly distinguishable from the wild fish in terms of morphology but they would remain genetically distinct if farmed stocks were genetically different from the wild. The DNA stand-by method has given satisfactory results for many species such as salmon (McGinnity et al., 1997), cod (Jensen et al., 2010), sea trout (Bernas et al., 2020), rainbow trout (Glover, 2008) and red sea bream (Sawayama et al., 2019) so far but has been used to a lesser extend for gilthead seabream and not at all for European sea bass.

The aim of this study was to investigate the genetic differences between farmed populations of seabream and seabass and their neighbouring wild populations in two countries (a fish farm from each country) and to assess the probability of identifying escapees from fish farms into the wild using DNA stand-by methods. These samples have been previously used by Arechavalla-Lopez et al. (2012), who they investigated the morphological differences between the specific populations.

2 Materials and methods

Farmed individuals of gilthead seabream and common seabass were sampled from a fish farm in Spain (Alicante bay) and from a fish farm in Greece (Mesologgi bay) (Fig. 1). Additionally, individuals from the wild from places that were adjacent to fish farms were sampled (Fig. 1). Each sample consisted of 100 individuals. For details of the location and the characteristics of the samples see Arechavala-Lopez et al. (2012b). From each individual, we cut a small piece of fin tissue which was stored in 90% ethanol. Total DNA was extracted from fin-clips according to the salt extraction method (Miller et al., 1988).

We used two multiplex PCR reactions to amplify 16 microsatellite loci for each species. The loci and the primers are described in Table 1. The final volume of all the reactions was 10 μl. Each reaction contained 1 μl template DNA (10–20 ng/μl), 1 μl Taq DNA polymerase buffer 10×, 0.3 μl from each primer (forward and reverse from a stock of 10 pmol/μl), 3 mM MgCl2, 0.3 μl dNTPs from a stock of 10 mM each) and 0.1 U Taq DNA polymerase. PCR conditions were 15 min at 94 °C followed by 35 cycles each of 30 s in 94 °C, 90 s in 57 °C (58 °C for D. labrax) and 60 s in 72 °C. There was a final extension step for 10 min at 72 °C.

The genetic loci that we examined for S. aurata were 172EP, Ad-10, B13b, Bld-10, Bld-04, C67b, C77b, Cld-29, Cld-35, Ctd-27, Dc23, F6, F7b, Fd-92, Hc-33, P20, Saimbb25,
Table 1. Observed and expected heterozygosity for *S. aurata* (A) and *D. labrax* (B).

A.

<table>
<thead>
<tr>
<th>Locus</th>
<th>FarmedSP (N = 86)</th>
<th>FarmedGR (N = 100)</th>
<th>WildSP (N = 88)</th>
<th>WildGR (N = 80)</th>
<th>No. of Alleles</th>
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<tbody>
<tr>
<td>Ad-10</td>
<td>0.314</td>
<td>0.266</td>
<td>0.480</td>
<td>0.450</td>
<td>0.250</td>
</tr>
<tr>
<td>B13b</td>
<td>0.884</td>
<td>0.801</td>
<td>0.750</td>
<td>0.739</td>
<td>0.807</td>
</tr>
<tr>
<td>Bld-04</td>
<td>0.640</td>
<td>0.649</td>
<td>0.740</td>
<td>0.675</td>
<td>0.625</td>
</tr>
<tr>
<td>Bld-10</td>
<td>0.372</td>
<td>0.376</td>
<td>0.090</td>
<td>0.086</td>
<td>0.216</td>
</tr>
<tr>
<td>C77b</td>
<td>0.547</td>
<td>0.486</td>
<td>0.50</td>
<td>0.496</td>
<td>0.598</td>
</tr>
<tr>
<td>Cld-29</td>
<td>0.733</td>
<td>0.759</td>
<td>0.720</td>
<td>0.710</td>
<td>0.667</td>
</tr>
<tr>
<td>Cld-35</td>
<td>0.651</td>
<td>0.639</td>
<td>0.430</td>
<td>0.430</td>
<td>0.727</td>
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<tr>
<td>Ct27</td>
<td>0.907</td>
<td>0.866</td>
<td>0.950</td>
<td>0.869</td>
<td>0.965</td>
</tr>
<tr>
<td>D23</td>
<td>0.907</td>
<td>0.895</td>
<td>0.900</td>
<td>0.861</td>
<td>0.909</td>
</tr>
<tr>
<td>F6</td>
<td>0.942</td>
<td>0.897</td>
<td>0.870</td>
<td>0.777</td>
<td>0.909</td>
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<tr>
<td>F7b</td>
<td>0.767</td>
<td>0.801</td>
<td>0.850</td>
<td>0.76</td>
<td>0.793</td>
</tr>
<tr>
<td>Fd-92</td>
<td>0.733</td>
<td>0.826</td>
<td>0.890</td>
<td>0.793</td>
<td>0.727</td>
</tr>
<tr>
<td>Hd-33</td>
<td>0.659</td>
<td>0.634</td>
<td>0.760</td>
<td>0.626</td>
<td>0.614</td>
</tr>
<tr>
<td>P20</td>
<td>0.779</td>
<td>0.777</td>
<td>0.650</td>
<td>0.645</td>
<td>0.807</td>
</tr>
<tr>
<td>Sambil25</td>
<td>0.930</td>
<td>0.859</td>
<td>0.830</td>
<td>0.809</td>
<td>0.852</td>
</tr>
<tr>
<td>SauE82</td>
<td>0.721</td>
<td>0.675</td>
<td>0.870</td>
<td>0.697</td>
<td>0.796</td>
</tr>
<tr>
<td>SauI47</td>
<td>0.643</td>
<td>0.650</td>
<td>0.697</td>
<td>0.705</td>
<td>0.580</td>
</tr>
<tr>
<td>172EP</td>
<td>0.535</td>
<td>0.483</td>
<td>0.520</td>
<td>0.461</td>
<td>0.534</td>
</tr>
</tbody>
</table>

B.

<table>
<thead>
<tr>
<th>Locus</th>
<th>FarmedSP (N = 52)</th>
<th>FarmedGR (N = 93)</th>
<th>WildSP (N = 86)</th>
<th>WildGR (N = 87)</th>
<th>No. of Alleles</th>
</tr>
</thead>
<tbody>
<tr>
<td>DLA0041</td>
<td>0.950</td>
<td>0.869</td>
<td>0.756</td>
<td>0.741</td>
<td>0.824</td>
</tr>
<tr>
<td>DLA0044</td>
<td>0.684</td>
<td>0.576</td>
<td>0.943</td>
<td>0.722</td>
<td>0.904</td>
</tr>
<tr>
<td>DLA0051</td>
<td>0.905</td>
<td>0.734</td>
<td>0.895</td>
<td>0.614</td>
<td>0.648</td>
</tr>
<tr>
<td>DLA0060</td>
<td>0.654</td>
<td>0.538</td>
<td>0.511</td>
<td>0.487</td>
<td>0.435</td>
</tr>
<tr>
<td>DLA0061</td>
<td>0.529</td>
<td>0.437</td>
<td>0.604</td>
<td>0.443</td>
<td>0.709</td>
</tr>
<tr>
<td>DLA0064</td>
<td>0.920</td>
<td>0.835</td>
<td>0.730</td>
<td>0.814</td>
<td>0.813</td>
</tr>
<tr>
<td>DLA0066</td>
<td>0.837</td>
<td>0.780</td>
<td>0.554</td>
<td>0.579</td>
<td>0.913</td>
</tr>
<tr>
<td>DLA0068</td>
<td>0.608</td>
<td>0.551</td>
<td>0.771</td>
<td>0.616</td>
<td>0.536</td>
</tr>
<tr>
<td>DLA0073</td>
<td>0.673</td>
<td>0.717</td>
<td>0.796</td>
<td>0.681</td>
<td>0.895</td>
</tr>
<tr>
<td>DLA0075</td>
<td>0.808</td>
<td>0.694</td>
<td>0.677</td>
<td>0.601</td>
<td>0.759</td>
</tr>
<tr>
<td>DLA0078</td>
<td>0.955</td>
<td>0.869</td>
<td>1.000</td>
<td>0.758</td>
<td>0.883</td>
</tr>
<tr>
<td>DLA0081</td>
<td>0.820</td>
<td>0.783</td>
<td>0.857</td>
<td>0.706</td>
<td>0.895</td>
</tr>
<tr>
<td>DLA0086</td>
<td>0.955</td>
<td>0.843</td>
<td>0.351</td>
<td>0.686</td>
<td>0.922</td>
</tr>
<tr>
<td>DLA0089</td>
<td>0.275</td>
<td>0.239</td>
<td>0.394</td>
<td>0.371</td>
<td>0.454</td>
</tr>
<tr>
<td>DLA0096</td>
<td>0.659</td>
<td>0.624</td>
<td>0.845</td>
<td>0.693</td>
<td>0.653</td>
</tr>
<tr>
<td>DLA0097</td>
<td>0.800</td>
<td>0.794</td>
<td>0.765</td>
<td>0.660</td>
<td>0.861</td>
</tr>
</tbody>
</table>

SauE82, SauI47 (Franch et al., 2006; Guinand et al., 2015), whereas for *D. labrax* the genetic loci were DLA0041, DLA0051, DLA0044, DLA0060, DLA0061, DLA0064, DLA0066, DLA0068, DLA0073, DLA0075, DLA0078, DLA0081, DLA0086, DLA0089, DLA0096, DLA0097, (Chistiakov et al., 2008; Guinand et al., 2015). These are commonly used microsatellite polymorphic loci in both species.

PCR products were run on an ABI PRISMs 3700 DNA Analyzer (Applied Biosystems, Carlsbad, CA, USA), using 5’-labelled reverse primers and the GeneScanTM- 500 LIZs Size Standard (Applied Biosystems) as an internal size standard for allele sizing. Alleles were sized using the software STRAND 2.3.79 (http://www.vgl.ucdavis.edu/informatics/STRand/).

### 3 Statistical analysis

Adegenet R package (Jombart and Ahmed, 2011) was employed to calculate observed and expected heterozygosities, and GENETIX 4.05 (Belkhir et al., 2000) for gene diversity and number of alleles. Detection of loci with excess in null alleles, F statistics, departure from Hardy-Weinberg Equilibrium (HWE) as well as linkage analysis between loci were performed with GENEPOP 3.4 (Raymond and Rousset, 1995). Adegenet (Jombart and Ahmed, 2011) was used to perform a discriminate analysis of principal components (DAPC). The analysis produced the same results independently of the number of axes that we retain in the principal component analysis step or in the discriminant analysis. Finally, we performed a Bayesian clustering analysis using STRUCTURE.
2.3 (Pritchard et al., 2000; Falush et al., 2003), assuming a number of K populations (maximum value set K = 7), using the default settings with twenty independent runs for each K, 1,000,000 iterations and a burn-in period of 100,000. Then we used the Harvest online software (Earl and Vonholdt, 2012) which uses the Evanno’s algorithm (Evanno et al., 2005), to find the best K value. Given that we would like to identify escapees from the fish farms, we also run Structure for Greek farmed and wild samples as well as for Spanish farmed and wild samples, using K = 2 as a default value. We visualize the STRUCTURE results with the program STRUCTURE PLOT v.2 (Ramasamy et al., 2014). We have run the software multiple times with membership coefficient ranging from 70% to 80%; results did not change qualitatively (data not shown) and the 75% was finally used as a threshold for the analysis of current data. Therefore, an individual belongs to a certain group if the membership coefficient is higher than 75%; otherwise, that individual is assigned as belonging to the alternative group.

4 Results

4.1 Genetic population structure

4.1.1 Sparus aurata

We genotyped 19 loci for S. aurata but we excluded locus C67b from the analysis as having significant excess of null alleles in all four samples; thus, 18 loci were finally retained. We detected 262 different alleles in all samples of S. aurata (Tab. 1A). The number of alleles per locus varied from 2 (locus 172EP) to 30 (locus C127). We also detected 45 private alleles (alleles appeared only in a single sample), which represent 17% of the total number of alleles. Seven and 15 of the private alleles were found in wild and farmed Greek samples, respectively, and two and 21 were found in Spanish wild and farmed samples, respectively. Private alleles were equally distributed across populations (Kruskal-Wallis, p = 0.1322). Gene diversity was highly variable across loci (Tab. 1A), and ranged from 0.086 in locus Blt-10 in Greek farmed sample to 0.932 in locus Dt23 in the wild Spanish sample. Nevertheless, different samples had not significantly different gene diversities (Kruscal-Wallis, p = 0.4451).

All four samples appear to be in Hardy Weinberg disequilibrium (p = 0.000). Three loci in Spanish farmed sample, eight loci in Greek farmed sample, two loci in Spanish wild sample and two loci in Greek wild samples were significantly departed from HWE. In eight of 18 loci that showed significant departure from HWE, FIS was greater than zero indicating inbreeding. In some cases, FIS was high (e.g. 0.101 in locus SauE82 for the Greek wild samples (Tab. 2A). In cases that FIS was significantly lower than zero, their values were close to zero except from the Greek farmed population that FIS was −0.244, −0.209, −0.115 and 0.114 in four loci.

The four samples were distinct from each other, as the FST index is significantly different from zero between all pairs of samples except for the wild Spanish and Greek samples (Tab. 3A). This is confirmed by Nei distance, which shows that the closest samples were the wild Spanish and Greek ones. The more distantly related samples were the farmed Spanish and the farmed Greek (FST = 0.0778, Nei distance = 0.2020).

The DAPC analysis showed a similar picture; wild Spanish and wild Greek samples could not be distinguished from each other, whereas both the farmed Greek and Spanish samples formed distinct groups (Fig 2A).

4.1.2 Dicentrarchus labrax

All the 16 loci that we used for D. labrax were polymorphic with a total number of 243 alleles in all samples, and mean number of alleles greater than 15 (Tab. 1B). The total number of alleles per locus varied from 6 alleles in the locus DLZ-377 to 24 alleles in locus DLA0051. We identified 48 private alleles, which correspond to 28% of the total number of alleles. Private alleles are not equally distributed across samples (Kruskal-Wallis test, p = 0.000122).

The expected heterozygosity ranged from 0.239 (locus DLA0089, in farmed Spanish sample) to 0.915 (in locus DLA0073, in wild Spanish sample) (Tab. 1B). The differences in gene diversity among samples were not significant (Kruskal-Wallis test, p = 0.51).

All four samples were not in Hardy-Weinberg equilibrium (p = 0.000 for each of the populations). However, not all individual loxi were significantly departed from Hardy-Weinberg equilibrium as shown in FIS, the inbreeding coefficient, which was significantly different from zero only in few loci for each population (in 6 and 5 loci in farmed and wild samples from Spain and in 10 and one loci in farmed and wild Greek samples, respectively) (Tab. 2B). When FIS was significantly different from zero, it was negative in most of the cases (in 13 out of 22). Interestingly, the farmed Greek sample had 10 loci departed from HWE, and in eight of them the FIS was negative. On the other hand, in this population FIS had the highest positive FIS value (0.493).

All samples were significantly differentiated from each other as indicated by the FST index (Tab. 3B). The higher differentiation was between the farmed Greek and the farmed Spanish samples according to FST (FST = 0.199), whereas according to Nei distance the higher differentiation was between the wild Spanish sample and the farmed Greek sample (Nei distance = 0.697). However, the distance between this pair of samples and farmed Greek and Spanish samples was very similar (Nei distance = 0.679). The lowest differentiation was between the wild and farmed samples from Spain (FST = 0.038; Nei distance = 0.123). DAPC analysis produced similar results (Fig 2B); farmed and wild Spanish samples formed an indiscriminate group, whereas farmed and wild Greek samples formed two highly distinct groups.

4.2 Genetic assignment

4.2.1 S. aurata

We run STRUCTURE until convergence and the Bayesian analysis revealed that all four populations of S. aurata could be assigned into two clusters as indicated by Evanno’s test (ΔK = 172.7955). In the next alternative (K = 3), the ΔK was 74.3202. This clustering is different from the DAPC analysis in which both Spanish samples and the Greek wild sample formed a single group, whereas the Greek farmed samples formed a different cluster (Fig. 3A).
We then run separately the Spanish (Fig. 3B) and the Greek samples (Fig. 3C), inferring two clusters in each location, which corresponded to farmed and wild samples. This correspondence was poor for Spanish samples but stronger for Greek samples. The proportion of membership for Spanish farmed samples to belong to one cluster was 0.618 and for Spanish wild samples to belong to the second cluster was 0.892. For the Greek samples, the proportion of membership to one cluster for the farmed samples was 0.926 and for the wild samples the proportion of membership to the second cluster was 0.146.
was 0.965 indicating strong grouping. When we considered the probability of participating to one or the other cluster for each individual separately, 27 individuals from Spanish farmed population failed to participate in any of the two clusters (probability of participation less than 0.75), 42 individuals belonged to one cluster and 17 to the other. Thus, the origin of the farmed Spanish sea-bream sample seems to be obscure. On the other hand, the wild population seems to be genetically more concrete. Only ten out of 88 individuals failed to be assigned to a cluster, with probability higher than 0.75 and the rest 77 formed a distinct cluster. From these results, we do not have clear evidence for escapees from the farmed samples to the wild. However, given that the Spanish farmed samples do not form a concrete group, the genetic characterization of escapees is not possible.

For the Greek farmed sample, 11 individuals failed to be assigned to a single cluster. Eighty-eight individuals formed a well-defined cluster and one individual belonged to the alternative cluster. For the Greek wild sample, 79 individuals belonged to a single cluster (different from that of the farmed samples) and one individual failed to be assigned to a cluster. Thus we have no evidence for escapees from the farm to the wild in this sample.

4.2.2 *D. labrax*

We run together all four samples of *D. labrax* for Bayesian clustering. The Evanno’s algorithm suggested three groups (Evanno et al., 2005) ($\Delta K = 219.7664$) (Fig. 4A). The farmed and wild Greek samples formed two distinct groups, whereas the Spanish samples formed a single group. This is consistent with the DPAC Analysis (Fig. 3B).

Given that we would like to test whether there were escapees from the farmed to wild populations we run STRUCTURE separately for the Spanish (Fig. 4B) and the Greek samples (Fig. 4C) defining $K = 2$ for farmed and wild samples in each location. The proportion of membership to the farmed and wild groups for Greek individuals was on average 0.937 and 0.985, respectively. Similarly, for the Spanish samples the proportion of participation to farmed and wild samples was on average 0.875 and 0.855, respectively. There were two individuals from the Spanish farmed samples that showed probability of participation to the wild group higher than 0.75 (probabilities 0.810 and 0.806). There were also two individuals from the wild sample that appeared to come from the farmed population. One of these individuals had probability of participation to the farmed sample 0.821 and the other 0.977. These two individuals are potentially escapees.

There were five (out of 52) individuals from the Spanish farmed sample and 12 (out of 86) individuals from wild Spanish sample, as well as five (out of 93) individuals from Greek farmed sample and none (out of 87) individuals from Greek wild sample that could not be assigned to one or the other group, i.e. the probability of participation to any group was lower than 0.75.

5 Discussion

5.1 Wild populations

Large-scale genetic studies suggest that wild populations of *D. labrax* are clustered into three large genetic pools: the Atlantic, which is separated from western Mediterranean in the Almeria-Oran front, the western Mediterranean and the eastern Mediterranean. Some studies show low but significant $F_{ST}$ values between local populations (Naciri et al., 1999; Lemaire et al., 2000; Tine et al., 2014; Souche et al., 2015; Vandeputte et al., 2019). Even though our samples are not representative of the variation in all populations from the two countries, our
results confirm that the wild Greek and Spanish seabass samples are distinct with significant $F_{ST}$ value.

Most of the previous studies suggest that seabass wild populations in the three genetic pools are in Hardy-Weinberg equilibrium if some loci were excluded (Garcia De Leon et al., 1997; Naciri et al., 1999; Lemaire et al., 2000; Lemaire et al., 2005). Other studies using a different set of markers, show that local wild populations are in Hardy-Weinberg disequilibrium (Castilho and Ciftci, 2005). This difference between studies could be probably due to genetic markers used in each study.

Population genetics of *S. aurata* show a different pattern than *D. labrax*. Low but significant $F_{ST}$ values indicate that seabream has genetically distinct local populations rather than large genetic (Ben Slimen et al., 2004; Rossi et al., 2006; Gkagkavouzis et al., 2019). Through the Aquatrace project (https://cordis.europa.eu/project/rcn/105481/reporting/en), clustering analysis suggested a relatively strong subdivision between Atlantic and Mediterranean basins ($F_{ST}$ values 2%-3%) and a less strong, though significant subdivision within the Mediterranean in three “sub-basins” (West Mediterranean, Ionian and Aegean) ($F_{ST}$ values from 0 to 1.8%) in sea bream wild populations. In this context, Gkagkavouzis et al. (2019) analyzed wild sea bream populations around Greece and they report that the populations appeared to be highly admixed and no geographic clusters are evident in structure plots. Our results confirm the differentiation between wild Greek and Spanish Mediterranean populations as well as the departure from HWE.

Natural populations could depart from HWE for several reasons such as inbreeding, admixture of previously separated populations, thus increase of heterozygosity (Wahlund effect), selection or presence of null alleles that artificially increases homozygosity. In our populations, selection could be excluded because it would seem highly unlikely a large number of random microsatellite loci to be under selection. The effect of null alleles can also be excluded because in our analysis we have tested for excess of null alleles. The negative $F_{IS}$ values indicate that departure from HWE in our study us most possibly is explained by the admixture of previously separated populations.

![Fig. 3. Bayesian clustering of *S. aurata* samples. A. All four samples form two clusters. B. Spanish samples. C. Greek samples. The different colors represent the groups that the program STRUCTURE has clustered the individuals. The length of the colored bar shows the probability that a specific individual belongs to the respective group. The length of the colored bars per individual adds up to 1.](image-url)
5.2 Farmed populations

The Greek farmed samples that we examined were significantly different from the wild samples both for gilthead seabream and for European seabass. This was obvious both from the $F_{ST}$ and from the Bayesian analyses. For the Spanish farmed samples, even though $F_{ST}$ showed low but significant differentiation, Bayesian analysis grouped the two samples into a single cluster both for seabream and for seabass. Previous studies have compared wild and farmed Greek populations of $S. aurata$ and revealed that the farmed populations were distinct from the wild ones (Alarcon et al., 2004; Karaiskou et al., 2009; Loukovitis et al., 2011). Interestingly, Karaiskou et al. (2009) found that among distantly related wild Greek seabream populations there was no genetic differentiation, while farmed fish from different cages of the same fish farm formed genetically distinct groups. Likewise, Loukovitis et al. (2011) using microsatellites and Alarcon et al. (2004) using both microsatellites and allozymes showed significant genetic differentiation of farmed sea bream from their proximal wild populations in different sites across north Mediterranean. Last, in the Aquatrace project, the report cites that the farm broodstocks are generally less genetically diverse than wild populations, and additionally broodstocks are more differentiated among each other ($F_{ST}$ ranging between 1.2% and 5.7%), which might be the results of strong genetic drift for many generations of selection or founder effects. However, when these broodstocks are compared to the wild counterpart, most of them are markedly different from them with the exception of broodstocks with a more recent history of hatchery selection which display similarities with some wild populations.

To our knowledge, there are limited previous studies for seabass to compare the genetic structure of farmed and wild populations. In such a study they found that the two of three aquaculture stocks were outbred (Bahri-Sfar et al., 2005). The results for gilthead seabream corroborate our findings, which suggest that at least the Greek farmed sample was significantly different from the wild samples.
distinct from the wild populations. Greek farmed sample was highly differentiated from all the other populations particularly for seabass. $F_{CT}$ values are high and $F_{IG}$ values are small. There are three potential scenarios to explain the genetic differentiation between farmed and wild samples in Greece but not in Spain. First, farmed samples might have different geographic origin from the proximal wild samples. Second, farmed sample might have been originated from local populations but they carry a small proportion of the genetic diversity of wild populations. Finally, farmed sample might have been originated from selective breeding programs, where artificial selection might have changed the genetic structure of the selected population.

From these potential explanations we can exclude the sampling effect because if farmed sample was a subsample of the adjacent wild populations, it should not contain private alleles, given that sample size for both farmed and wild samples is high enough. Selection alone would also have fewer odds to explain the results, because selective breeding has been very recently applied in both species. Nevertheless, it would seem unlikely that an artificial selection program could change the genetic structure of a species for neutral loci in few generations. The most plausible explanation would be that the fingerlings come from a different geographic region. A combination of different scenarios also cannot be excluded. For example, fingerlings might have been imported from a different country and be part of a selective breeding program.

The Messolonghi-Etoliko lagoon complex in Western Greece coat (Ionian Sea) is composed of six lagoons and represents one of the most important coastal lagoon systems in the Mediterranean Sea (Dimitriou et al., 2007). More than 25% of Greek fish farms and 20% of Greek hatcheries of gilthead sea bream and sea bass are located in this lagoon system (source: Mariculture Federation of Greece). It is known that for both species, strains of foreign origin are bought and raised in fish farms because they are considered to grow faster. Furthermore, during the last decade, a spectacular increase in the number of farms and their production, accompanied by a substantial price decrease resulted in the maintenance in cages of large individuals (more than 500 g) of gilthead seabream, which are potential spawners. Extensive histological investigation of gonads sampled from different farms in the two major regions for aquaculture activities in Greece revealed that female sea bream cultivated beyond the size of sex reversal matured, ovulated and released eggs during the normal spawning period of the species; moreover, the genetic identification of sparsid eggs demonstrated that sea bream eggs were spawned in farms and released to the environment (Somarakis et al., 2013). Escapes from cages have a rather accidental character and fluctuate from year to year due to invasion of large pelagic species or adverse weather conditions. Considering that only farms covered by a specific insurance provide these data, it is clear that the number of escaped individuals was even greater (Dimitriou et al., 2007).

Are the populations different enough to allow us correctly allocating each individual to a certain population? Bayesian analysis suggests that Greek but not Spanish farmed and wild samples are distinct well enough, and such allocation is possible with high probability. This is particularly interesting to find the escaped fish. The genetic method for allocation of escapees has been successfully used for tracing escapee salmon, rainbow trout and Atlantic cod back to their farm of origin (Glover, 2010). It seems a promising method, which has not been validated either for sea bream or for sea bass (but see Segvic-Bubic et al., 2011). Our results are encouraging, at least for the Greek populations. The common practice to move seed from a geographic location to other for farming as well as the recent programs for selective breeding of these species increase the probability of identifying genetically the escaped fish but increases the danger for the escapees to “genetically contaminate” the local populations. In certain cases, like in the Messolonghi-Etoliko lagoon which is a typical nursery for the species, the unintentional escapes in the natural environment may have devastating consequences, since alleles or genes of “alien” origin could be imported and propagate faster in the wild gene pool.

Arechavala-Lopez et al. (2012a) recommended the use of molecular genetic markers as a suitable tool for genetic discrimination of wild and farmed fish, enabling monitoring of the genetic impact of fish farm escapes and/or restocking releases. Broodstocks and their offspring could be genotyped in hatcheries before going to open-sea cages, and such information should be available to the scientific community and managers, in order to improve the accuracy and suitability of genetic tools.

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