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Splendiani A., Righi T., Fioravanti T., Sabatini A., Palmas F., Tougard C., Berrebi P., Talarico L., Caputo Barucchi V.; POPULATION GENETICS, DEMOGRAPHY AND CONSERVATION OF MEDITERRANEAN BROWN TROUT 2 FROM SARDINIA; Aquatic conservation – Marine and freshwater conservation; 34 (2) Art. n. e4099; John Wiley & Sons; 25 pp

**The publisher's version is available at:**

<http://dx.doi.org/10.1002/aqc.4099>

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30 POPULATION GENETICS, DEMOGRAPHY AND CONSERVATION OF MEDITERRANEAN BROWN TROUT  
31 FROM SARDINIA

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51  
52 Keywords: *Salmo trutta*, invasive species, conservation genetics, biogeography, conservation policy, extinction risk

53  
54 Abstract

55 1. Brown trout is a species complex (*Salmo trutta* complex, L., 1758) including both widespread invasive (non-native  
56 hatchery strains) lineages and endangered local-endemic lineages, among which is the Sardinian trout, the only native  
57 salmonid present in Sardinia. Multiple stressors (e.g., the spread of stocked brown trout of Atlantic origin, habitat  
58 alteration, and climate change) combine to seriously threaten the persistence of wild native populations.

- 59 2. In this study, the origin, population genetics, and demography of wild Sardinian brown trout populations were  
60 extensively investigated. A total of 274 trout individuals collected from 12 hydro-geographical basins were analysed  
61 using both mitochondrial (Control Region) and nuclear (*LDH-C1\** locus and 10 microsatellites) markers.
- 62 3. Although stocking activities have altered the native genetic makeup of some populations in the study area, several  
63 (almost) uncontaminated populations showing strong genetic structure were detected. Eroded intra-population  
64 diversity, and small effective population size, sometimes associated with a bottleneck signal were also found.
- 65 4. The genetic characteristics of Sardinian trout populations described in this study are probably due, at least partly, to the  
66 peculiarity of local environmental conditions at the margin of the ecological niche for salmonids. Based on the results  
67 of this study, the need for urgent measures of conservation aimed to ensure the near future viability of the last wild  
68 Sardinian trout populations was discussed.

## 69 **1 INTRODUCTION**

70 The delineation of spatial population structure represents a crucial step in understanding the demography and evolution  
71 of species (Waples & Gaggiotti, 2006). This implies understanding the spatial scales over which populations are connected  
72 through dispersal and gene flow and the role of environmental characteristics underlying the pattern of connectivity between  
73 populations. Obtaining this kind of information helps to plan biodiversity management in a rational manner. For example  
74 through the delineation of conservation categories (i.e. Conservation units CUs, Evolutionary Significant Units, ESUs and  
75 Management Units, MUs), assessment of population and meta-population viability, and strategic enhancement of landscape  
76 connectivity (e.g. Palsbøll, Bérubé & Allendorf, 2007; Robertson et al., 2013). Since pioneering reflections on protecting  
77 species' evolutionary potential (Mayr, 1960), the debate on the delineation of intra-specific entities of conservation and  
78 management has become of crucial interest mainly for heavily managed species attracting socio-economic interests, as in  
79 the case of the fisheries and/or game-fisheries-species (e.g., Fraser & Bernatchez, 2001). Thanks to a plethora of  
80 conservation genetics studies, protection of local populations is nowadays considered pivotal for local managers intending  
81 to restore and/or conserve species diversity (e.g. Bruce et al., 2019).

82 Brown trout (*Salmo trutta* complex L., 1758) is a fish of great economic (mainly in aquaculture) and recreational value  
83 both in its original range and worldwide. Habitat degradation coupled with massive and uncontrolled stocking activities  
84 with non-native lineages (mainly from northern Europe), have compromised the conservation status of native populations in  
85 several European countries (Weiss et al., 2001; Caputo et al. 2004; Araguas et al., 2017; Vera, Martinez & Bouza, 2018;  
86 Splendiani et al., 2019a; Prunier et al., 2021). Brown trout is an appealing and iconic species for scientists because of  
87 taxonomic controversies that are still unresolved, the complex evolutionary history, and the intricate patterns of life-history  
88 traits (Lobón-Cerviá & Sanz, 2018), as well as for its biological conservation needs (Piccolo et al., 2018).

89 Early phylogenetic studies identified five main mitochondrial (mtDNA) evolutionary lineages: the Atlantic (AT),  
90 Mediterranean (ME), marmoratus (MA), Adriatic (AD), and Danubian (DA) lineages (Bernatchez, Guyomard &

91 Bonhomme, 1992). Subsequently, other lineages were proposed, such as Duero (DU, Cortey et al., 2009; Vera et al., 2010),  
92 Tigris (TI, Bardakci et al., 2006), North African (NA, Tougard et al. 2018) and Dades (Snoj et al. 2011). However,  
93 mitochondrial lineages often show an overlapping natural distribution, with even more mitochondrial lineages observed in a  
94 single population (Hashemzadeh Segherloo et al., 2021). Therefore, if on the one hand, the phylogenetic and  
95 phylogeographic approach has failed to resolve taxonomic controversies to date, on the other side, molecular  
96 phylogeography has allowed the identification of the paleo-climatic and environmental events that played the most crucial  
97 roles in shaping brown trout biogeography (Splendiani et al., 2013; 2016a; 2020). For this reason and because the  
98 identification of brown trout taxonomic status is not the purpose of the present study, only mtDNA lineages and sub-  
99 lineages of *Salmo trutta* will be considered here.

100 In the Mediterranean area, the Italian Peninsula and its major islands represent a biodiversity hotspot for the genus  
101 *Salmo*. Here, at least five valid nominal species have been recognized (*S. ghigii* Pomini, 1941; *S. cettii* Rafinesque-  
102 Schmaltz 1810; *S. marmoratus*, Cuvier, 1829; *S. carpio*, Linnaeus 1758; and *S. fibreni*, Zerunian & Gandolfi, 1990; e.g  
103 Polgar et al., 2022), whose biogeographic history has been moulded by complex colonization routes and ecological  
104 adaptation driven by paleo-climatic changes and paleo-hydrological re-arrangements of river networks (Lerceteau-Köhler et  
105 al., 2013; Sanz 2018; Splendiani et al., 2020). A very high genetic differentiation was detected among insular populations  
106 (Sardinia and Corsica), especially in Corsican populations (Berrebi et al., 2019). The Corsican trout populations showed a  
107 certain degree of similarity with Sardinian brown trout populations when compared with other Italian peninsular trout  
108 populations, although Sardinian trout sampling sites were from two river basins only (Flumendosa and Cixerri). More  
109 recently, in a genome-wide based phylogenetic revision, Hashemzadeh Segherloo et al. (2021) highlighted the high  
110 distinctiveness of native trout populations from Sardinia with respect to other Mediterranean trout taxa, suggesting to  
111 recognize Sardinian trout populations as a distinct species.

112 Mediterranean brown trout is the only native salmonid in Sardinia. However, since the beginning of the 20<sup>th</sup> century,  
113 notably, from the 1960s onward, stocking activities became a common management practice and introduced into the rivers  
114 of this Mediterranean island two exotic species: *S. trutta* from Central Europe (*i.e.*, the Atlantic trout of hatchery origin) and  
115 *Oncorhynchus mykiss* from North America (Sabatini et al., 2006; Orrù et al., 2010). The introduction of non-native species  
116 were banned in Sardinia since the early 2000s, in compliance with Presidential Decree 357/97.

117 Habitat/trophic competition and the rapid adaptive plasticity of salmonids coupled with hybridization between native  
118 and Atlantic brown trout lineages had progressively reduced local wild populations and altered the original Sardinian gene

119 pool (Sabatini et al., 2006; 2011). As a consequence of genetic introgression, habitat alteration, and fishing, the  
120 Mediterranean trout is listed as critically endangered in the Italian IUCN Red List (e.g. *Salmo ghigii*, Rondinini, Battistoni  
121 & Teofili, 2022).

122 Although earlier data from the 20<sup>th</sup> century (Cottiglia, 1968) reported an almost homogeneous brown trout distribution  
123 throughout the island rivers, they were unfortunately not able to distinguish between Mediterranean-native and Atlantic-  
124 exotic trout of stocking origin. In subsequent studies (Massidda et al., 1996; Cau, 1997; Zanetti et al. 2007), the presence of  
125 native trout populations was proposed for a very small fraction of the investigated sites (11 out of 160). Genetic studies in  
126 the last two decades revealed that populations of pure Sardinian trout could be found in the Cixerri, Pula and Flumendosa  
127 basins (Sabatini et al. 2006; 2011; 2018; Zaccara et al. 2015; Berrebi et al. 2019; Palmas et al., 2020; Hashemzadeh  
128 Segherloo et al., 2021). Despite a number of studies focusing on Sardinian trout populations, to date, none has provided a  
129 comprehensive characterization of the genetic population structure and diversity, demography and conservation status of  
130 wild populations. This is especially relevant as wild Sardinian trout populations are known to inhabit peculiar, sometimes  
131 even extreme, environments as, for instance, creeks subject to extreme water flow fluctuations and small ponds  
132 characterized by relatively high seasonal temperatures (Mulas et al., 2009; Zaccara et al., 2015). In this Mediterranean  
133 island, up to 90% of all streams present a non-perennial hydrological regime (Mulas *et al.*, 2009). In most cases, the  
134 hydrology of the streams involved in this study was unstable or even intermittent with frequent severe summer droughts.  
135 (Table 1). Yearly, during the warmest and driest months, the water discharge is absent and the trout survive in small and  
136 isolated pools where the water temperature can exceed 25° C for several days or weeks (Table 1).

137 Here samples from various Sardinian rivers generally thought to be representative of the local Mediterranean brown  
138 trout variability (plus additional samples from Corsica and from hatcheries of the Italian Peninsula rearing trout of Atlantic  
139 origin) were collected and genotyped at multiple molecular markers (mtDNA, *LDH-CI*, and microsatellites) with respect to  
140 native/exotic lineages and/or fine-scale population distinctiveness. The aims of this study were to: i) infer population genetic  
141 structure while controlling for admixture from hatchery-reared Atlantic strains; ii) provide insight into demography  
142 (effective population size, occurrence of bottlenecks) of wild populations; iii) identify units for management and evaluate  
143 their conservation status to provide an appropriate baseline for restoring strategies.

144

## 145 2 MATERIAL AND METHODS

### 146 2.1 Sampling and DNA extraction

147 A total of 274 wild brown trout individuals were collected in 20 sampling sites between May and October from 2016 to  
148 2019, representing 12 Sardinian river basins (Table 1 and Figure 1). To introduce comparative (reference) populations, a  
149 total of 39 specimens from two pure wild Corsican sites (collected in 2015) and 46 specimens from two hatcheries-rearing  
150 Atlantic trout strains (collected in 2006) were also included. Overall, 359 individuals were analyzed in this study (Table 1).  
151 Unfortunately, the Atlantic strains from local Sardinian hatcheries, used for stocking in recent years were not available, as  
152 the only working Sardinian hatchery currently breeds only rainbow trout (*Oncorhynchus mykiss*). However, the Atlantic  
153 strains were obtained from two hatcheries in Central Italy which is an important trout aquaculture region along the Italian  
154 Peninsula (ISPRA, 2022). The wild fish were captured by electrofishing and subsequently housed in appropriate tanks  
155 during the field job. A small piece from the adipose fin was clipped from every individual and stored in absolute ethanol,  
156 before releasing the specimens into nature. Total genomic DNA was extracted using specific cartridge 401 in the  
157 *MagCore*® automated Nucleic Acid extractor (*MagCore*®, *Genomic DNA Tissue Kit, n° 401*).

## 158 2.2 Mitochondrial DNA

159 The CR sequence was used to detect the diagnostic sites of the major mitochondrial lineages of *Salmo trutta* complex,  
160 and therefore to assess the frequency of allochthonous (e.g. Atlantic and Danubian lineages, respectively AT and DA) and  
161 native (Adriatic, Mediterranean, and marmoratus lineages, respectively AD, ME and MA) Mediterranean haplotypes. A  
162 Polymerase chain reaction-restriction fragment length polymorphism-single-strand conformational polymorphism (PCR-  
163 RFLP-SSCP) analysis was performed to screen mitochondrial DNA (mtDNA) genetic variability. The mitochondrial control  
164 region (CR) was PCR amplified using the primers 28RIBa (Su-nik, Snoj & Dov , 2001) and HN20 (Bernatchez &  
165 Danzmann 1993), following procedures described in Bernatchez & Danzman (1993). Single strand conformation  
166 Polimorphisms (SSCP) (Orita et al., 1989) was analyzed following the method reported in Righi & Fasola (2023). Sanger  
167 sequencing of the CR (~1 Kbs) was performed, using the same primers of amplification, on a subsample for each different  
168 SSCP detected profile on an Applied Biosystems ABI 3730XL DNA by a service facility (BMR-Genomic, Padua).  
169 Sequences were aligned using ClustalW (Thompson, Higgins & Gibbons, 1994), checked by eye in BioEdit (Hall 1999) and  
170 assigned to sequences of *S. trutta* available in GenBank using Blast (Altschul et al., 1990). Levels of population genetic  
171 introgression were estimated by calculating the cumulative percentage of allochthonous haplotypes in each population.  
172 Phylogenetic relationships among 68 CR haplotypes (Table S1) were inferred using two approaches: i) a 95% parsimony  
173 network estimated by the software TCS version 1.18 (Clement et al., 2000) and ii) a phylogenetic tree using a Bayesian  
174 inference (BI) as provided in MrBayes 3.1.2 (Ronquist & Huelsenbeck, 2003). For the BI approach the HKY85 substitution  
175 model (i.e., the optimal model for our data, as identified by the selection procedure implemented in MEGAX; Kumar et al.,

176 2018), the invgamma rate variation and 5-gamma categories were used. A sequence of *S. salar* (GenBank accession number  
177 LC012541) was used as an outgroup. Divergence time estimation was carried out in Beast2 v.2.7.3 (Bouckaert et al., 2014).  
178 As calibration points, the more recent common ancestor (MRCA) of *Salmo* (*S. immigratus*) and of brown trout (*S.*  
179 *derzhavini*) was used by applying lognormal constraints following Veli kovi et al. (2023). Moreover, *S. orhidanus*, each  
180 brown trout lineage (AD, AT, MA, ME, DA) and groups supported by BI posterior probabilities = 1 were treated as *a priori*  
181 monophyletic. Divergence time estimations were done with an optimized lognormal relaxed clock (Douglas, Zhang &  
182 Bouckaert, 2021) and by applying a birth-death (Gernhard, 2008). Computations were performed for three independent runs  
183 for 100 million generations sampling every 10,000th generation using the Beagle library (Ayres et al., 2012). Adequate  
184 sampling and run convergence were verified in Tracer v.1.7.1 (Rambaut et al., 2018), and then the tree files were combined  
185 with LogCombiner. Finally, the maximum clade credibility tree was calculated in TreeAnnotator discharging 1,000,000  
186 states as burn-in. Posterior summaries were only calculated for the nodes having a posterior probability greater than 0.9. The  
187 final tree was drawn using FigTree v.1.4.4 (<http://tree.bio.ed.ac.uk/software/figtree/>).

188 Finally, isolation and contacts among trout populations, driven by past climate phases enhancing resident or  
189 anadromous lifestyle, were investigated using the analysis of molecular variance (AMOVA). Genetic variance was  
190 estimated by grouping populations according to i) 12 river basins and ii) four sea drainages: Gulf of Asinara, Tyrrhenian  
191 Sea, Gulf of Cagliari and the Mediterranean Sea. Tests were carried out with ARLEQUIN version 3.5.1.3 (Excoffier &  
192 Lischer, 2010), using conventional  $F$ -statistics and testing the statistical significance with 5,000 permutations.

193 A significant and substantial amount of variance explained by differences among river basins would suggest inter-  
194 watershed population isolation which likely occurred during the last glacial maximum, i.e. when the warmer conditions of  
195 the Mediterranean basin resulted in non-optimal environmental characteristics for anadromous Mediterranean trout.  
196 Conversely, a large amount of variance explained by differences among sea drainages would imply ancient gene flow  
197 among river basins flowing into the same sea drainage. In fact, lower water temperatures during colder climatic phases of  
198 the Pleistocene coupled with an anadromous brown trout lifestyle may have favored migrations along the coast through sea  
199 outlets of close river basins (e.g. Splendiani et al., 2016b and references therein). Note that for the above-mentioned  
200 mtDNA-based analyses, the dataset was enhanced including CR information of additional 15 trout individuals from three  
201 Corsican sites (i.e., LTT, CTT and HBT; see Figure 1, Table 1 and Table 2) from grey literature (Reynaud, Tougard &  
202 Berrebi, 2011).

203 2.3 Nuclear DNA



204 A PCR-RFLP analysis of the eye-specific lactate dehydrogenase protein-coding locus (*LDH-CI\**) was performed  
205 following the procedure described in McMeel, Hoey & Ferguson (2001). This analysis allows discrimination between  
206 diagnostic alleles for the north Atlantic (allele \*90) and Mediterranean populations (allele \*100) of the *Salmo trutta*  
207 complex. Conformity with HardyóWeinberg equilibrium was tested as described for microsatellite DNA (see below) and  
208 levels of genetic introgression were estimated by calculating the percentage of the allochthonous allele \*90 in each  
209 population.

210 Ten non-coding microsatellite loci (di- and tetra-nucleotide repeats) were labelled with fluorescent dyes and amplified  
211 following Splendiani et al (2019) in two separate multiplex reactions as reported in Table S2. Genotyping was performed  
212 using an ABI-PRISM 3130xl Genetic Analyzer (Applied Biosystems), with the LIZ 500 size standard, and allele sizes were  
213 manually scored using Peak Scanner<sup>®</sup> Software v1.0 (Applied Biosystems).

214 The microsatellite dataset was screened for false positives, null alleles or other genotyping errors with CERVUS v3.03  
215 (Kalinowski, Taper & Marshall, 2007), ML-NUllFreq (Kalinowski & Taper 2006) and MICRO-CHECKER 2.2.3 (Van  
216 Oosterhout et al. 2004). FreeNA (Chapuis & Estoup 2007) was used to control the effect of null alleles on  $F_{ST}$  estimate. The  
217 bootstrap 95% confidence intervals (CI) for the global  $F_{ST}$  value were estimated using 1,000 replicates over all loci. The  
218 allelic richness ( $A_r$ ) and inbreeding coefficient ( $F_{IS}$ ) were estimated using FSTAT 2.9.3 (Goudet 2001). The estimates of  
219  $A_r$ , were adjusted for the smallest sample size, i.e. COG at locus *Str60* ( $n = 3$ ). The observed ( $H_o$ ) and expected ( $H_e$ )  
220 heterozygosities for each sampling site were calculated in ARLEQUIN. The genotypic linkage disequilibrium between loci  
221 and population pairs, and the exact test for HardyóWeinberg equilibrium deviation per population were evaluated using the  
222 online software GENEPOP ON THE WEB (Raymond & Rousset, 1995; Rousset, 2008) with 10,000 de-memorizations and  
223 400 batches with 10,000 iterations each. The nominal level of significance (5%) was adjusted following a Bonferroni  
224 procedure (Rice, 1989).

225 The pairwise genetic differentiation among trout populations (i.e.,  $F_{ST}$  *sensu* Wright) was computed in FSTAT. As  
226 described for mtDNA (see section 2.2), the analyses of genetic variation (AMOVA) were performed in ARLEQUIN to  
227 investigate the partitioning of genetic variance under the two hypothesized hierarchical grouping tested above using CR  
228 haplotypes: populations groups were based on i) the 12 river basins of origin and ii) four sea drainages (Table 1).

229 The population genetic structure was investigated using the Bayesian clustering method implemented in STRUCTURE  
230 2.3.4 (Pritchard, Stephens & Donnelly, 2000) using a  $\Delta$ -hierarchical STRUCTURE approach (e.g. Vähä et al. 2007;  
231 Warnock, Rasmussen & Taylor, 2010; Mari et al., 2017; Berrebi et al. 2019; García-De León et al., 2020) performing

232 subsequent rounds on each subgroup identified by Evanno method. The STRUCTURE parameters were setup as follows: 10  
233 serial runs for each number of clusters (K) between 1 and sampling sites number +1; admixture model with correlated allele  
234 frequencies; burn-in period of 50,000 steps followed by 200,000 Monte Carlo replicates. The optimal K was chosen  
235 according to the  $\Delta K$  method (Evanno, Regnaut & Goudet, 2005) as estimated in STRUCTURE SELECTOR  
236 (<https://lmmme.ac.cn/StructureSelector/>) (Li & Liu, 2018). Finally, genetic differentiation among individuals and populations  
237 was also explored through a discriminant analysis of principal components of genetic variability (DAPC; Jombart, Devillard  
238 & Balloux, 2010), implemented in the package adegenet 2.0 (Jombart, 2008) for the R software (R core team 2021), by  
239 setting sampling locations as pre-defined groups.

240 Maximum likelihood method implemented in COLONY 2.0.6.1 (Jones & Wang, 2010) was used to evaluate family  
241 structure within sites, as it may affect the results of population structure analyses (Anderson & Dunham, 2008). Sib-ship  
242 probabilities were estimated by setting: random mating, polygamy for both sexes (e.g. Serbezov et al., 2010; Rossi et al.,  
243 2022), no prior for sib-ship assignments, long-length runs, and high likelihood precision (other settings were as default). To  
244 check for consistency among results, each run was replicated three times.

245 The effective population size ( $N_e$ ) for each site/drainage was estimated using both the programs NeESTIMATOR 2.01  
246 (Do et al., 2014) and COLONY. The first approach ( $N_{e1}$ ) is based on linkage disequilibrium and adjusts for missing data  
247 (LDNe method implemented in NeESTIMATOR). The  $N_{e1}$  estimation with the lowest allele frequency of 0.02 was  
248 reported as recommended for microsatellite markers (Do et al., 2014). The second approach ( $N_{e2}$ ) uses the sib-ship  
249 assignment methods (Wang, 2009) based on the frequencies of sib-ship estimated from a sib-ship assignment analysis, using  
250 the multi-locus genotypes of a sample of offspring taken at random from a single cohort in a population.

251 Recent and substantial demographic reductions were evaluated for each population using BOTTLENECK (Piry,  
252 Luikart & Cornuet, 1999) whose method relies on the assumption that the mutation-drift equilibrium is transiently disrupted  
253 and the heterozygosity measured at a locus ( $H_e$ ) will exceed the heterozygosity ( $H_{eq}$ ) computed from the number of alleles  
254 sampled (Cornuet & Luikart 1996). Both the infinite allele mutation model (IAM, Kimura and Crow, 1964) and the Two-  
255 Phased model (TPM: 90% of single-step mutations with variance set to 30%, Di Rienzo et al., 1994) were applied, as  
256 recommended for microsatellite data (Luikart et al. 1998), setting 5,000 replicates. The heterozygosity excess was evaluated  
257 according to the 1-way Wilcoxon signed-rank test (which is recommended in the event of limited sample sizes and/or loci;  
258 (Piry, Luikart & Cornuet, 1999) and the allele frequency distribution mode-shift method (Luikart et al. 1998).

259 Finally, the association between the amounts of introgression from Atlantic lineages within sampling sites/hatcheries,  
260 as revealed by employed diagnostic or semi-diagnostic molecular markers (microsatellites, *LDH-CI\** and mitochondrial  
261 CR) was investigated using the Pearson's linear correlation (*cor.test* function in R;). The relationship between measures of  
262 genetic diversity (*Ar* and *He*) and introgression of hatchery-Atlantic lineages (as estimated by the frequency of the *LDH-*  
263 *CI\*90* allele) across sites/hatcheries was also tested using the *lm* function in R: in this case, a quadratic model was used  
264 (second-degree polynomial) as diversity is expected to be higher at intermediate levels of introgression (Rossi et al., 2022).

### 265 3 RESULTS

#### 266 3.1 Mitochondrial DNA

267 A total of 18 CR haplotypes in 359 individuals were detected, belonging to both native and exotic mitochondrial  
268 lineages (Table 2). The latter included six AT haplotypes and a single DA haplotype. The AT haplotypes were already  
269 observed in European hatcheries ó i.e., *haplotype-1*, 2, 3 and 4 (Cortey & García-Marín, 2002), *AT-Tyrrh1* (Berrebi et al.,  
270 2019) and *At1e* (Meraner et al., 2007). The *haplotype-1* was observed in both reference Atlantic hatcheries (HATa and  
271 HATb), and in the wild sites GOG and FMCb, the *haplotype-2* was observed in HATb and in the wild site FMCb, the  
272 *haplotype-3* was observed in HATb, the *haplotype-4* was observed in the wild sites CDL and RMN, *AT-Tyrrh1* was  
273 observed in HATa, and *At1e* was observed in the wild site POSb. The single DA haplotype resulted identical to the  
274 haplotype *Da1a* (Duftner et al., 2003) and detected as dominant (90%) in FLUa. As indicated above, this Danubian  
275 haplotype was considered to be of stocking origin (see section 4 below).

276 The other 11 haplotypes belonged to the native AD phylogenetic lineage: four were previously described ó *A\_2*  
277 (Zaccara et al 2015), *AD-Tyrrh1* (Berrebi et al., 2019), *AD-Tyrrh4* (Berrebi et al., 2019, Zaccara et al. 2015 [C69]), *AD-*  
278 *Tyrrh7* (Palmas et al., 2020), while seven haplotypes were detected for the first time in this study (*AD-Tyrrh8* ó *AD-*  
279 *Tyrrh14*, Genbank accession numbers OR972382-OR972391, Table 2). Among AD haplotypes, sequence lengths ranged  
280 from 996 to 1324 bp. This polymorphism, observed in 5 (*AD-Tyrrh9* - *AD-Tyrrh13*) out of 11 haplotypes, was caused by  
281 one to five tandem duplications of an 82 bp motif located in the 3'-end of the CR. As the elongation model of this repetition  
282 is generally thought to be the result of intra-molecular processes (Buroker et al., 1990; Sell & Spirkovski, 2004), and the use  
283 of the number of repetitions may not be appropriate for phylogenetic reconstruction, only the first copy was kept in the  
284 analysis ó but note that after excluding the tandem repeat structures, haplotypes *AD-Tyrrh9* and *AD-Tyrrh13* collapsed into  
285 the haplotype *AD-Tyrrh4*. The phylogenetic tree (Figure 2) and the TCS network (Figure 3) roughly provided consistent  
286 results. In particular, 1) haplotypes *AD-Tyrrh10*, *AD-Tyrrh4* and *AD-Tyrrh12* formed a strongly supported clade (posterior  
287 probability = 1, Figure 2) along with the *ADcs-23/24/25* Corsican haplotypes detected in the west-flowing river basins

288 Seccu and Liamone (e.g. Reynaud, Tougard & Berrebi, 2011, Table 1 and Table 2) ó given their geographic distribution and  
289 remarkable differentiation within the AD lineage, they will hereafter be referred to as belonging to the òCorso-Sardinian  
290 sub-lineageö; 2) other AD haplotypes detected in this study were similar to each other (i.e. showing 1-4 mutations; Figure  
291 3), although mutual relationships were poorly resolved, except for the clade including *AD-Tyrrh8* and *AD-Tyrrh11*  
292 haplotypes (BI posterior probability value = 0.77, Figure 2). Time to the most recent common ancestor ( $T_{MRCA}$ ) of brown  
293 trout was dated to 3.82 Ma [95% HPD 1.83-8.54] and  $T_{MRCA}$  of AD lineage can be dated to 2.52 Ma [95% HPD 0.85-5.84]  
294 (Figure 2, Table S3). The AD lineage appeared ramified into three groups, in which only the Corso-Sardinian sub-lineage  
295 was highly statistically supported and its origin was dated around 1.05 Ma [95% HPD 0.24-2.72].

296 A total of 1-3 haplotypes per site were found in Sardinian locations. In a total of 20 sites, 13 and 3 sites were,  
297 respectively, entirely, or mainly (>70% frequency) composed of native AD haplotypes, whereas the remaining three sites  
298 (i.e. FLUa, FMCb and RMN) showed the prevalence of allochthonous haplotypes. A clear geographic pattern of  
299 differentiation was suggested by the distribution of AD haplotypes. The most widespread haplotype was *AD-Tyrrh1*, being  
300 detected with high frequencies (from 54 to 100%) in one-third of Sardinian rivers and two Corsican sites (VES and VIV).  
301 This haplotype was shared among all of the north-eastern basins investigated apart from the Padrogiano basin (PAD - Table  
302 2). On the other hand, the haplotypes of the Corso-Sardinian sub-lineage (both from this study and from literature) showed a  
303 western distribution (Table 2 Table S1 and Figure 1). The other AD haplotypes were found in very restricted areas (1-2 sites  
304 each) where they were generally present at high frequencies. In detail, the haplotype *AD-Tyrrh7* was observed only in the  
305 Flumendosa basin (FLUa and FLUc). Haplotypes *AD-Tyrrh8* and *AD-Tyrrh11* presented a northern distribution with the  
306 haplotype *AD-Tyrrh8* private and fixed in PAD and the haplotype *AD-Tyrrh11* detected in POSa and in COG. Finally, *AD-*  
307 *Tyrrh14* was private in RMF and the haplotype *A\_2* was fixed in all Pula Basin sampling sites (PULa, PULb1 and PULb2)  
308 and the most abundant in CIX (Table 2).

309 The AMOVAs (Table 3) revealed that grouping samples according to the river basin of origin explained most of the  
310 among-group genetic variance (i.e. 83.37%). When sites were grouped according to the location of the catchment outlet, the  
311 among-group component decreased to approximately 56%.

### 312 3.2 Nuclear DNA

313 Besides hatcheries, the exotic Atlantic *LDH-CI\*90* allele was found at high frequencies in FLUa (85%), FMCb (83%)  
314 and RMN (77%). On the other hand, the *LDH-CI\*90* allele was absent in several Sardinian sampling sites Canale  
315 dell'Iserno (POSa), Riu Flumineddu (CED - except for one hybrid specimen), Riu Bau Mandara (FLUb), Riu Furittu  
316 (FLUc), Pula basin (PULa, PULb1 e PULb2), Riu Piras (FMPa) and Riu Is Abius (CIX). Also, in the Corsican sites (VES

317 and VIV), the *LDH-CI\*90* allele was absent. In remaining Sardinian populations (COG, PAD, POSb, CDL, FMCa, FMPb,  
318 TEM), the *LDH-C\*90* allele showed moderate frequency (values between 12 and 36%)  
319 Regarding microsatellites data, the presence of null alleles was suggested by all three software used in this study (CERVUS,  
320 ML-NullFreq and MICRO-CHECKER) in 14 tests over 220. The loci *Ssa85* and *OMM1064* were affected by null alleles in  
321 respectively, three (FMCa, PULa and FMPb) and two sampling sites (FMCb and HATb). All other loci showed evidence of  
322 null alleles in just one population. However, global  $F_{ST}$  values, obtained including or excluding null alleles (i.e., the ENA  
323 correction method; Chapuis & Estoup, 2007), returned comparable results by using all loci screened, respectively, 0.422 (CI  
324 0.388-0.465) and 0.428 (CI 0.395-0.470). As null alleles negligibly affected estimates of the population genetic  
325 differentiation, all loci for downstream analyses were retained.

326 Results of genetic variability within populations were reported in Table 2. In total, 198 alleles were detected using 10  
327 microsatellite loci. The number of alleles per locus ranged from 5 (*Str60*) to 38 (*Ssa410UOS*). Measures of genetic diversity  
328 substantially differed among Sardinian sites: allelic richness ( $A_r$ ) and expected heterozygosity ( $H_e$ ) ranged from 1.28  
329 (PULb2) to 3.43 (FLUa) and 0.29 (CIX) to 0.74 (FLUa), respectively. Models revealed that *LDH*-based introgression  
330 explained a substantial fraction of both  $A_r$  ( $R^2 = 0.715$ ,  $F_{2,21} = 26.33$ ,  $P < 0.001$ ) and  $H_e$  ( $R^2 = 0.675$ ,  $F_{2,21} = 21.82$ ,  $P <$   
331  $0.001$ ), although suggesting roughly linear rather than quadratic relationships in our dataset (Figure S1). In other words,  
332 intra-population genetic diversity was higher in sites affected by deep introgression from Atlantic strains rather than in  
333 purely native sites.

334 Significant ( $P < 0.05$ ) deviations from Hardy Weinberg expectations were observed in three Sardinian (PULa, FMCa,  
335 and RMF) sampling sites, HATb and one Corsican location (VIV), although only the latter remained significant after  
336 Bonferroni correction. Tests for linkage disequilibrium (LD) at the population level revealed 3 significant associations  
337 ( $P < 0.001$ ) out of 1035 comparisons, namely between *Ssa410UOS* and *Ssa408UOS* loci in CIX and HATa, and between  
338 *SSsp2213* and *Ssa408UOS* in HATa.

339 The Wilcoxon one-tailed test revealed the signal of a recent bottleneck in four sampling sites (FLUa, FMCa, FMCb,  
340 and PULa) when using the TPM model, and in seven sites (FLUa, FMCa, FMCb, PULa, FMPa, RMN and VES) in the case  
341 of IAM. However, the shifted mode method confirmed the possibility of a bottleneck only in FLUa and PULa, while  
342 suggesting a possible bottleneck also for PULb (Table 4).

343 Both methods of effective population size estimation (Table 4) failed (confidence intervals including infinity) to  
344 determine  $N_e$  in several sampling sites caused by the small sample size. For the rest of the cases, the comparisons of the

345 output from both methods suggest that the Sardinian populations are particularly small ( $1.6 \leq N_e \leq 25.8$ ;  $10 \leq N_e \leq 29$ ).  
346 In general,  $N_e$  estimations based on the linkage disequilibrium method were lower compared to those based on the sib-ship  
347 assignment method. Estimates were partly related among methods (Spearman correlation:  $r_s = 0.52$ ,  $P = 0.039$ ), in any  
348 event both tests reported the lowest effective population size for CIX and the highest for POSb.

349 The global  $F_{ST}$  was 0.431 ( $P < 0.001$ ) implying remarkable genetic differentiation among populations. Pair-wise  $F_{ST}$   
350 values and their significance are reported in Table 5. The differentiation among sampling sites was substantial ( $P < 0.05$   
351 after adjustment for multiple comparisons) in 160 out of 253 comparisons. Lower pair-wise values ( $F_{ST} > 0.1$ ) were detected  
352 between the two hatcheries, between hatcheries and three wild sites (RMN, FLUa, FMCb), and between Posada Basin sites  
353 (POSa and POSb). Notably, three sites (i.e., COG, FLUc and PULa) were not statistically differentiated ( $P > 0.05$ ) from all  
354 other sampling sites.

355 AMOVAs provided similar outcomes, irrespective of the two tested partitioning of sites (Table 3): differentiation  
356 among sea drainages and river basins explained approximately 16 and 13% of the overall variance, both significantly ( $P <$   
357  $0.001$ ); the intra-population differentiation accounted for most of the variation ( $> 52\%$ ), as expected when dealing with  
358 hypervariable markers.

359 The sequential analysis of genetic structure investigated with STRUCTURE identified a total of 21 genetic cluster (K)  
360 populations (Figure 4). In the first round of analysis, involving the entire data set, multiple K values were supported,  
361 therefore, the uppermost structure was chosen corresponding to  $K = 13$  (Figure 4). As 7 out of 13 genetic clusters included  
362 more than a single sampling location, a second round of STRUCTURE analysis for each "multi-sample" genetic cluster was  
363 conducted: most of the sampling sites grouped together in the first step were split as single clusters. Finally, a third analysis  
364 round allowed distinguishing between POSa and POSb within the "Posada cluster" identified in the second round of  
365 analyses (Figure 4).

366 To specifically explore the presence of hybrid/Atlantic trout across 20 Sardinian and two Corsican wild sampling sites,  
367 while quantifying their admixture degree, a  $K = 2$  was forced in the Bayesian STRUCTURE analysis: because  
368 Atlantic/Mediterranean opposition is the first structure in these populations, the individual membership coefficients  
369 obtained (i.e.  $q$  values) were ranked from the highest ( $q = 1$ , indicating a pure native trout individual in this study) to the  
370 lowest ( $q = 0$ , namely a pure hatchery-Atlantic trout) and their 90% credible intervals (CIs) were plotted against rank  
371 (Figure S2). Based on admixture ( $q$ ) values and their CIs, frequency of *LDH-C1\*90* allele and AT-DA haplotypes, four  
372 groups of individuals were arbitrarily identified. In the first group (*pure native trout*, 25.00% of sites), the mean  $q$  values

373 were  $\neq 1$  with very narrow CIs (the mean lower CI was 0.982); here (FLUc, PULb1, PULb2, FMPa, and CIX), neither  
374 allochthonous haplotypes nor the *LDH-CI\*90* allele were detected. In the second group (*low introgressed trout*, 40.00%),  
375 mean  $q$  values were still high ( $\neq 1$ ), while contextually associated with lower mean CIs (mean lower CI = 0.912, range  
376 0.912  $\pm$  0.964); here (CED, PAD, FMCa, FMPb, COG, RMF, TEM and PULa), the frequency of allochthonous haplotypes  
377 ranged from 0.00 to 0.14 and the frequency of the *LDH-CI\*90* allele ranged from 0.00 to 0.33. In the third group  
378 (*moderately introgressed trout*, 25.00%), mean  $q$  values were even lower (mean  $q$  = 0.94), while the mean lower CI was  
379 0.850 (range = 0.761  $\pm$  0.891); in this group (CDL, POSb, RMN, POSa, and FLUb), the frequency of allochthonous  
380 haplotypes ranged from 0.00 to 1.00 and the frequency of the *LDH-CI\*90* allele ranged from 0.00 to 0.77. The fourth group  
381 (*non-native trout*, 10.00%) included pure or almost pure Atlantic trout (FMCb and FLUa), showing mean  $q$  values  $\neq 0$ ; in  
382 this latter group the frequency of allochthonous haplotypes ranged from 0.89 to 1 and the frequency of the *LDH-CI\*90*  
383 allele ranged from 0.83 to 0.85 (Table 2 and Figure S2).

384 Estimates of Atlantic brown trout introgression across sites/hatcheries strongly correlated between molecular markers:  
385  $r = 0.96$  and  $P < 0.001$  for *LDH-CI\*90* allele vs. Atlantic haplotypes;  $r = -0.93$  and  $P < 0.001$  for Atlantic haplotypes vs.  
386 coefficient of hatchery ancestry ( $q$  of STRUCTURE);  $r = -0.88$  and  $P < 0.001$  for *LDH-CI\*90* allele vs. hatchery ancestry.

387 The DAPC analyses showed a pattern of genetic differentiation quite similar to the scenario depicted by  
388 STRUCTURE. The first plot (Figure 5a), which included all sampling sites, pointed to the distinctiveness of Pula River  
389 (PULa, PULb1-2), CIX, FMPa and VIV while the rest of the other sites were grouped together. After removing such  
390 distinctive locations (Figure 5b), CED, FMPb and VES diverged from other sites, which were roughly arranged along a  
391 gradient: from Atlantic strains in the left (HATa, HATb, FMCb, FLUa), to Mediterranean-native ones at the center of the  
392 plot (e.g. CDL, FLUc, FLUb, FMCa, and RMF). The third plot (Figure 5c), which was obtained after removing the most  
393 divergent sites of the previous step (i.e. CED, FMPb, and VES), highlighted the presence of three groups of populations.  
394 Northern populations (TEM, COG, PAD, POSa, and POSb), located at the top left part of the scatterplot, form a group well  
395 separated from the remaining highly pure populations from the South-eastern side (FLUa, FLUb, FMCb) located at the bottom  
396 right portion. At the top center of the graph the hatchery-reared Atlantic strains and highly introgressed wild sampling sites  
397 FLUa and FMCb are overlapped identifying an homogeneous cluster, quite close to the wild sites RMN, CDL, and RMF.  
398 Generally, except for FLUa and FMCb, each sampling site was identified as a separated cluster.

399 The number of families per population identified by the parentage analyses performed with COLONY software  
400 identified very few siblings ( $>0.80$  inclusion and exclusion probability in most cases, see Table S3).

#### 401 4 DISCUSSION

402 In this study, the origin, population genetics, and demography of wild brown trout populations from Sardinia were  
403 investigated, and the role of Sardinia as a hotspot of *Salmo* (genetic) diversity within the Mediterranean basin was  
404 eventually demonstrated. In addition, the presence of a new distinctive Corso-Sardinian mtDNA sub-lineage characterized  
405 by haplotypes endemic to the Sardinian and Corsican rivers was described (Figures 2 and 3). Nuclear markers  
406 (microsatellites) also pointed out strong differentiation between wild native populations. At the same time, the reduced  
407 intra-population genetic variability coupled with small effective population sizes suggested the potentially severe  
408 vulnerability of such Sardinian-native populations inhabiting extreme habitats for salmonids. A similar pattern has been  
409 observed in Corsica, leading to the same interpretation (Berrebi et al., 2019). The need for the definition of appropriate  
410 categories of conservation applicable in the implementation of correct and concrete conservation actions appears crucial for  
411 the near future conservation of the last population of Sardinian trout.

##### 412 4.1 Population genetic variability and demography

413 The levels of genetic variability detected within most Sardinian sampling sites appeared generally low. If one takes into  
414 account only pure wild locations (i.e., absence of the *LDH-C1\*90* allele and AT mtDNA haplotypes, coupled with mean  
415  $q$ -values  $\leq 1$ ; Table 2), a mean value of observed heterozygosity of 0.41 (SD = 0.11) and a mean value of allelic richness of  
416 1.86 (SD = 0.55) were estimated. Generally, higher values of observed heterozygosity ( $H_o > 0.60$ ) and allelic richness ( $A_r >$   
417 4.0) are typically observed in the hatchery-reared Atlantic strains (Bohling, Haffray & Berrebi, 2016), or in native  
418 Mediterranean brown trout populations highly impacted by the latter (Vera et al., 2023). In fact, similar values of low intra-  
419 population genetic diversity have been observed in almost purely native, small and naturally isolated populations from  
420 central Italy – such as those inhabiting the Tenna River (Adriatic drainage; Splendiani et al., 2019a) or the Rio Santa Croce  
421 (Tyrrhenian drainage, Rossi et al., 2022) – or elsewhere, in the Mediterranean basin: Corsica (Berrebi et al., 2019); the  
422 upper part of the Do-nica, and Konjarska rivers in Macedonia (Aegean drainage; e.g. Mari et al., 2016), two localities from  
423 the Mijares and Turia basins (e.g. Vera et al., 2013), and the Ter River (e.g. Araguas et al., 2017) of the Iberian Peninsula.  
424 The above cases mostly represent typical freshwater environments where the last native trout populations still survive in the  
425 Mediterranean area, such as in small creeks or streams naturally and/or artificially isolated from the other river basins,  
426 showing stable hydrological conditions and suitable spawning habitats. Generally, the native trout populations inhabiting  
427 these sites benefit from high conservation priority and these habitats are managed, or present themselves to be managed, as  
428 genetic refuges. These kinds of river ecosystems are likely to become thermally crucial for the future viability of salmonids  
429 in the Mediterranean rivers where, in the next two decades, half of the suitable habitat is expected to be lost (e.g. Almodóvar



430 et al., 2012). However, regarding the present case of study, the water courses where the last pure Sardinian trout populations  
431 still survive are very far from the concept of ideal thermal refuge for brown trout. As described above (section 1), most  
432 water courses investigated presented a non-perennial hydrological regime, with trout populations surviving in small and  
433 isolated pools where the water temperature can exceed 25° C for several days or even weeks during the driest months. For  
434 brown trout, an upper critical temperature range of 25 ó 30° C with an incipient lethal temperature of approximately 25° C  
435 was reported (e.g. Jonsson & Jonsson 2009). Thermal stress together with low discharge can also affect size, fecundity and  
436 population density due to the increased metabolic costs of growth at elevated temperatures in south salmonid habitats (e.g.  
437 Jonsson & Jonsson, 2009). Furthermore, intermittent discharge is likely to contribute to the fragmentation of Sardinian trout  
438 populations within basins, leading to multiple isolated patches of small effective population sizes.

439 Estimates of  $N_e$  (Table 4) resulted dramatically low, irrespective of the adopted method (considering only  $N_e$  estimates  
440 with finite CIs: 1.6  $N_{e1}$   $N_{e2}$  25.8; 10  $N_{e1}$   $N_{e2}$  29). Furthermore,  $N_e$  could be even lower if only native individuals are taken  
441 into account, as revealed by previous studies on introgressed populations (Splendiani et al., 2019a). Assuming  $N_e$  estimates  
442 to correspond approximately to ½ of the census population size (according to models based on Norwegian river-resident  
443 brown trout populations; Serbezov et al., 2012), actual spawners would range between 3.2 and 20 in the smallest population  
444 (CIX), and between 51.6 and 58 in the largest population (POSb) according to  $N_{e1}$  and  $N_{e2}$  estimates, respectively. Such a  
445 low estimation of the number of spawning adults appears quite realistic and consistent with low densities of trout  
446 individuals recorded in the most recent regional freshwater fish census (e.g. AA. VV., 2022, Table 1). Furthermore, also the  
447 difficulty encountered during the sampling activities of this study in obtaining a sufficient number of adult specimens in  
448 most localities corresponds to the detection in wild Sardinian trout sites of a very low census size.

449 In addition to generally low levels of genetic diversity and effective population size, some Sardinian trout populations  
450 analyzed in this study showed signals of a recent bottleneck. In particular, in the Riu Litteras from the Pula River (PULa), a  
451 significant excess of heterozygosity and an L-shifted mode of the allele frequency distribution were observed. Here, very  
452 low values of effective population size ( $N_{e1} = 2.6$  and  $N_{e2} = 12$ , Table 4) were observed and the concomitant detection of a  
453 recent bottleneck could be related to an extreme flash flooding event that occurred in November 2015 in the area of the Pula  
454 River basin(see below, section 4.3.2). Elsewhere in Sardinia, FLUa also showed both a significant excess of heterozygosity  
455 and an L-shifted mode of the allele frequency distribution. This sampling site, however, is largely represented by non-native  
456 individuals (DA lineage and individual  $q$  values close to zero), then bottleneck signals might be related to a founder effect  
457 occurred by introducing a restricted number of hatchery origin individuals. Moreover, hybridization can severely influence

458 the outcome of the bottleneck tests (Zhang et al., 2017), so the significant heterozygosity excess of the FLUa is possibly due  
459 to hybridization between native and allochthonous stocks as suggest by co-presence of AD and DA haplotypes.

#### 460 4.2 Genetic structure and phylogeographic inferences

461 Genetic analyses carried out in the present study revealed strong differentiation among the wild Sardinian brown trout  
462 populations (global  $F_{ST} = 0.43$ ), which is remarkable even compared to the values observed in similar extreme  
463 environments for salmonids as, for example, in trout populations (*Oncorhynchus* sp.) from Northern Sierra Madre  
464 Occidental in Mexico ( $F_{ST} = 0.33$ ; Abadía-Cardoso et al., 2021). Considering that several investigated Sardinian sampling  
465 sites were collected above artificial barriers and were characterized by an elevated degree of isolation created by an  
466 intermittent water flow (Table 1), it could be argued that such a high degree of genetic differentiation can be due to the  
467 stochastic effects of strong genetic drift acting on very small populations. Similarly, Pujolar et al. (2011) argued that  
468 reduced genetic diversity, low  $N_e$  sizes and serial bottleneck events revealed in marble trout populations from Slovenia  
469 imply a strong impact of genetic drift, limited gene flow, and high genetic differentiation which could have been  
470 exacerbated by recurrent mortalities due to flash floods and debris flows. Genetic drift has been proposed also to explain the  
471 high level of genetic differentiation observed both between and within the basin level in Mexican trout species of the genus  
472 *Oncorhynchus* living at the extreme southern margin of the genus's range (Abadía-Cardoso et al., 2021).

473 Besides genetic drift, ancient climatic fluctuations (with implications in connectivity among drainage basins) coupled  
474 with the anadromous behavior of ancestral Mediterranean brown trout (Splendiani et al. 2016b; Splendiani et al., 2019b) can  
475 partly explain the current geographical pattern of genetic structure. Based on the time-calibrated molecular phylogeny of the  
476 Sardinian trout,  $T_{MRCA}$  suggests that the haplotypes belonging to the Corso-Sardinian sub-lineage (Figure 2, Table S3)  
477 originated during the Menapian-Bavelian periods (c. 1.1 Ma; Middle Pleistocene). The alternation of glacial and interglacial  
478 phases that characterized the Pleistocene has had an important role in shaping the biogeographic characteristic of  
479 Mediterranean trout populations through the alternating promotion of different lifestyle tactics, promoting migratory  
480 propensity during the cold phases or a more sedentary lifestyle during the warmest phases. Thus, isolation in thermal  
481 refuges during warmest periods may have promoted the observed haplotype diversification and, colder phases may have  
482 played a role in shaping the geographic distribution of the mtDNA diversity. During the colder phases of the Pleistocene  
483 Corsica and Sardinia were connected (Grill et al., 2007) and therefore the presence of the two routes (west and east) of  
484 colonization along the paleo-Corso-Sardinian coasts is conceivable.

485 The effect of historical colonization patterns and isolation driven by past climatic phases on Sardinian trout genetic  
486 diversity is corroborated by AMOVA analysis based on both mtDNA and microsatellites. Significant genetic differentiation

487 among river basins support the hypothesis of long periods of isolation between trout populations (Table 3). Strong  
488 population differentiation was also detected by hierarchical analyses carried out by using both STRUCTURE (Figure 4) and  
489 DAPC (Figure 5a,5b,5c).

490 Moreover, AMOVA detected significant genetic variance even when sites were grouped based on the coastal river  
491 mouth orientation suggesting also the presence of a geographic genetic structure related to periods of contact between  
492 neighboring rivers that occurred thanks to the anadromous behavior of trouts in defined periods of time. Anadromy, in the  
493 Mediterranean basin, appeared periodically during the cold phases of the Pleistocene when the lower part of the river was a  
494 more suitable habitat for salmonids (Muñoz & Casadevall, 1997) and seaward migration propensity more likely (e.g.  
495 Splendiani et al., 2019b). Contacts was emphasized by the geographic distribution of the mtDNA haplotypes. In particular,  
496 Corso-Sardinian sub-lineage showed a western distribution in Sardinia that points to the role played by the last glacial  
497 marine regression. During the last glacial maximum, Corsica and Sardinia were connected due to the closure of the  
498 Bonifacio strait (Figure 1) and, as a consequence, the populations inhabiting rivers flowing towards the Western  
499 Mediterranean Sea were more likely to be interconnected along the western Corso-Sardinian paleo-shoreline. Here, the  
500 spread of the Corso-Sardinian sub-lineage probably occurred through migratory trout (i.e. sea trout). In addition, as  
501 mentioned above (section 2.2), sea trout generally feed chiefly in estuaries and along coasts (Jonsson & Jonsson 2006) and,  
502 as a consequence, it is possible to hypothesize that gene flow between Sardinian populations was more likely between  
503 populations with a close sea outlet. According to this hypothesis, gene flow between sea trout populations from northern  
504 Spain was negatively related to the distance between river mouths (Moran et al., 2005). Furthermore, as regards rivers  
505 flowing in a close bay, as in the cases in this study of the Gulf of Asinara and the Gulf of Cagliari, it is reasonable to expect  
506 that from an initial population of pioneers a successive source population arises later. This will first colonize the closest  
507 rivers in the bay as suggested by shared A\_2 haplotype between closer basins Cixerri (CIX) and Pula (PULa, PULb1 and  
508 PULb2) and, as was recently observed in brown trout populations from the Kerguelen archipelago in the District of the  
509 French Southern and Antarctic Lands, introduced here during the second half of the twentieth century (Launey et al., 2010).  
510 Moreover, the occurrence of the Corso-Sardinian sub-lineage at mid to high-elevation Corse sites and above impassible  
511 waterfalls (e.g. Berrebi, 2015), suggests a role as refuge played by the Corsican rivers for this sub-lineage during the severe  
512 interglacial warming periods of the Pleistocene. Subsequently, during the colder phases of the Pleistocene (the last glacial  
513 phase during the late Pleistocene, c. 100,000 - 15,000 years ago), the Corso-Sardinian sub-lineage could have reached the  
514 Sardinian rivers thanks to migratory tactics along the western Corso-Sardinian paleo-shoreline.

515 Similarly, on the Tyrrhenian side, the distribution of the haplotype *AD-tyrrh1* (and related ones) appears in accordance  
516 with a peri-Tyrrhenian past route of colonization connecting Corsica and Sardinia along the eastern Sardinian-Corsican  
517 paleo-shoreline during the last glacial maximum (Figure 1). This haplotype spread mainly along the eastern side of Corsica  
518 and Sardinia (e.g. Berrebi et al., 2019 and Figure 1). Exception is the Corsican Eze River (VES), a tributary of the Prunelli  
519 River flowing into the western side, where haplotype *AD-tyrrh1* resulted rare both in Sardinian and Corsica (e.g. Berrebi et  
520 al. 2019). Here, the presence of this haplotype could either represent the consequence of the wider past distribution of this  
521 Tyrrhenian AD haplotype or, alternatively, the consequence of ancient river captures that occurred between the two sides of  
522 the west-Mediterranean and Tyrrhenian catchments, similarly to what was suggested elsewhere in the Mediterranean area  
523 (e.g. Splendiani et al., 2006; Berrebi, Jesens k & Crivelli, 2017).

524 Finally, the AD sub-cluster formed by the haplotypes *AD-Tyrrh8* and *AD-Tyrrh11* (Figures 2 and 3) showed a  
525 north-eastern distribution partially overlapping the distribution of the common haplotype *AD-Tyrrh1*, thus suggesting the  
526 occurrence of an eastern biogeographic route adopted by multiple waves of colonization of the AD lineage (Figure 1 and  
527 Table 2). Interestingly, the co-occurrence of the above haplotypes in the Coghinas basin (North-Western Sardinia; e.g. COG  
528 in Figure 1) suggests that waves of colonization involving these AD Tyrrhenian haplotypes is likely to have occurred when,  
529 thanks to the sea level rising at the end of the last glacial maximum, the reopening of the Bonifacio strait allowed the  
530 formation of a biological corridor for these eastern AD haplotypes. In the southern part of the island, *A\_2* represents the sole  
531 haplotype observed in the Pula basin and the most common in the Cixerri basin; this haplotype probably reached the Gulf of  
532 Cagliari through a further wave of colonization.

#### 533 4.3 Major threats acting on native trout populations in Sardinia

##### 534 4.3.1 Stocking and fishing activities

535 This study has revealed the presence of several severe threats to the survival, in the near future, of native trout  
536 populations in the Sardinian rivers. A first menace has been highlighted by the detection of clear signals of hybridization  
537 between native trout and Atlantic brown trout of hatchery origin. Admixture from Atlantic strains in Sardinian trout has  
538 been already observed (Sabatini et al., 2011; Zaccara et al., 2015; Berrebi et al., 2019), although based on a limited number  
539 of examined individuals and/or populations, as compared to the present study. Here, two sites comprised almost exclusively  
540 allochthonous alleles and/or haplotypes (FLUa and FMCb). Conversely, the rest of the locations revealed genetic  
541 introgression from Atlantic gene pools ranging from 0%, in about a third of sampling sites, to low-medium amounts in the  
542 rest of the locations (Table 2). In Italy, stocking activities by using non-native species and/or populations have been strictly  
543 banned since 2003 (DPR n. 197/2003), although this law has been systematically neglected by local administrations as well

544 as by fishing clubs. (Splendiani et al., 2016a, 2019a, 2020). More recently (since 2020), as indicated below (section 4.4),  
545 stocking activities using non-native trout are admissible upon an official request to the Italian Ministry of the Environment.  
546 However, as far as it is known, only a few regional administrations have obtained this permission and illegal stocking  
547 activities using non-native trout are still popular in some regions (personal communications from local anglers).

548 Nevertheless, limited evidence of very recent stocking in Sardinia was found, as only a single specimen characterized  
549 by a  $q$  value of 0.03 (corresponding to a pure Atlantic trout) was observed in RMN (Figure S2). However, because of the  
550 low effective sizes of wild populations, the deleterious effects of stocking activities should be taken into account more  
551 seriously than elsewhere: even though negative selection is expected to purge exotic maladaptive alleles from wild  
552 populations, mildly deleterious alleles may reach fixation in small populations where the action of the purifying selection is  
553 weaker as compared to the larger ones (Moran et al., 2021). This implies that particular attention should also be paid in any  
554 planning of supportive breeding programs based on native trout populations with very low  $N_e$  sizes, as in the case of  
555 Sardinian trout, because of the concrete risk of promoting (albeit unintentionally) the fixation of deleterious alleles.

556 Conversely to almost everywhere else in Italy, a relevant proportion of genetically pure native populations in Sardinian  
557 rivers were found. It could be argued that the absence of traditional (or intensive) brown trout farming on the island ó  
558 officially, only few small family-owned companies exist where the farming of rainbow trout is allowed by law,  
559 (Autonomous Region of Sardinia ó RAS Det. N.3/22.01.2020) would have facilitated preserving the genetic integrity of  
560 wild native populations. In addition, the occurrence of major trout fishing tournaments has been (and still is) rare in  
561 Sardinia, when compared with the rest of the Italian Peninsula, probably because the severe environmental characteristics of  
562 most Sardinian salmonid waters are inappropriate or unattractive to carry out fishing competitions. As reported in Table 1,  
563 most sampling sites of the present study come from streams experiencing long periods of severe droughts during the driest  
564 months. If, on the one hand, the risk of stocking activities with allochthonous trout is averted, at least temporarily, other  
565 threats related to fishing activities are still present. For example, fishing activities are allowed in most of the sampling sites  
566 investigated (Table 1). In Sardinia, a five-fish daily limit is set; however, based on a Regional law (óDecree of the Assessor  
567 of the Defense of the Environmentö 10.05.1995 n. 412) the fishing of pure native trout individuals is forbidden everywhere.

568 In addition, in Sardinia, the Autonomous Region designated several river segments as "genetic sanctuaries" (GS), such  
569 as Riu Furittu, Riu Piras, and Riu Flumineddu, and here, fishing activities are totally banned (DR n.314/Dec.A9 -  
570 07.02.2019). Therefore, based on the outcomes of this study, fishing activities should be totally banned also in those basins

571 hosting exceptionally pure or nearly pure native trout populations that have not yet been ad hoc normative. Therefore, the  
572 updating of regional norms regulating fishing activities in freshwaters appears desirable.

#### 573 4.3.2 Environmental and climate characteristics

574 The very low values of effective population size observed in most populations are in accordance with the hydrographic  
575 fragmentation of the Sardinian rivers and with the very high summer water temperatures characterizing these south  
576 salmonid waters (e.g. Jonsson & Jonsson, 2009; Shirmpton & Heath, 2003). Moreover, extreme and repeated flood episodes  
577 can create demographic and genetic bottleneck in salmonids (e.g. Pujolar et al., 2011) or even extinction of local  
578 populations as in the case of the *Salmo marmoratus* population from Predelica (So a River) that was extirpated by a  
579 landslide triggered by intense rainfall in 2000 (Vincenzi et al., 2016; 2017). In the last two decades (2000-2020), Sardinia  
580 has been affected by 13 extreme flooding events, 62% of which involved the Sardinian rivers flowing toward the Gulf of  
581 Cagliari (e.g. Faccini et al., 2021), while the others involved the northeastern part of Sardinia (De Waele et al., 2010): the  
582 detection of a bottleneck signal in both Riu Bizzolu (COG) and Flumendosa River (FLUa) appears consistent with such a  
583 scenario, although speculative. Similarly, the very low  $N_e$  values coupled with bottleneck signals in the Pula Basin (see  
584 above, section 4.1) could be related to an extreme flash flooding event that recently occurred in south Sardinia. Forecasts for  
585 the near future are even worse, as a 30% increase in extreme precipitation is foreseen. (e.g. Faccini et al., 2021; Marras et  
586 al., 2021), Therefore, the need for a comprehensive  $N_e$  size monitoring of the last Sardinian brown trout populations  
587 appears as a crucial and concrete conservation action also in light of the  $N_e$  values observed in this study ( $1.6 < N_{e1} < 42.6$ ,  
588  $\text{mean} = 13.2$ ;  $10 < N_{e2} < 56$ ,  $\text{mean} = 23.28$ ) being well below the safe threshold from the 50/500 rule proposed by  
589 Frankham et al. (2014). This rule suggests that an effective population size of 50 is desirable to contrast the short-term  
590 likelihood of extinction due to the harmful effects of inbreeding depression on population demography, while a  $N_e$  of 500 is  
591 required for mutation to provide genetic diversity back into a population at a similar rate to loss caused by genetic drift,  
592 thereby maintaining a population's long-term evolutionary potential.

#### 593 4.4 IMPLICATION FOR CONSERVATION

594 High isolation of Sardinia rivers, due to both natural and anthropogenic factors, is likely to have played a  $\delta$ Dr. Jekyll  
595 and Mr. Hyde role towards the current status of conservation of wild trout population. The severe degree of isolation of the  
596 wild populations likely played a role in hindering the spread of phenomena of introgressive hybridization between native  
597 trout and Atlantic trout of hatchery origin, however, at the same time, isolation determined the very low level of genetic  
598 variability observed in Sardinian trout populations. Improving river connectivity, through the mapping and removal of those  
599 artificial barriers hindering within-basin natural gene flow, is necessary to counteract the low levels of effective population

600 size observed in wild Sardinian trout populations. However, such a process should be carried out carefully since these  
601 barriers are also crucial to prevent the spread of alien Atlantic trout (e.g. Splendiani et al., 2019a).

602 The first step to design appropriate and effective conservation action should be the identification of correct  
603 management units. Based on high genetic differentiation observed in this study, preservation of Sardinian trout diversity  
604 should be start from the protection of local populations and the management of wild local populations should be focused on  
605 the conservation of genetic diversity at an intraspecific level (e.g. Ferguson 2004; Bruce et al., 2019; Vera et al., 2023).  
606 However, in light of the results obtained, more detailed genetic and/or genomic studies would contribute to the acquisition  
607 of sound data in order to support the need for a taxonomic revision of Sardinian trout (e.g. Hashemzadeh Segherloo et al.,  
608 2021), the individuation of evolutionarily significant units and the delineation of management units. Within the near future,  
609 an advisable long-term conservation strategy of Sardinian brown trout populations should foresee the acquisition of  
610 knowledge about the genetic diversity of several wild Sardinian trout populations not yet studied, with as large as possible  
611 coverage, as already accomplished for instance in Corsica (> 200 sites analyzed; e.g. Berrebi, 2015). Moreover, in-depth  
612 studies are needed to better understand the pattern of intra-basin genetic diversity, as well as the association between genetic  
613 diversity and environmental features of Sardinian salmonid freshwaters.

614 Together with the delineation of units of conservation and management hopefully by an authoritative scientific  
615 committee, it is of paramount importance that these management units receive a legal value in a similar way to what has  
616 been achieved elsewhere, as in Canada where the delineation of conservation units is performed by the Committee on the  
617 Status of Endangered Wildlife (e.g. Bernard et al., 2009). On the contrary, in Italy, wildlife species management is still  
618 merely based on the definition of Linnean species (e.g. Splendiani et al., 2019c) and furthermore, freshwater fish fauna (as  
619 the rest of the ectotherms) is not considered the property of the State, and the management of local fish fauna is mainly  
620 delegated to fishing clubs. In this context, the risks of underestimating native trout genetic diversity are significantly high.

621 Finally, the recent modifications to the Italian national legislation if, on the one hand, are open to the introduction of  
622 allochthonous fish in nature (decree of 2 April 2020), on the other hand, completely ignore the regulation of the  
623 management of native species. Therefore, in the present normative context, the legal designation of management units  
624 appears of crucial importance.

625 In conclusion, the need to proceed toward the realization of an international strategy of conservation for Mediterranean  
626 salmonids appears therefore clear. A fundamental first step should be the recognition of freshwater fish species as national  
627 property of the sovereign states and, consequently, the provision of a legal value to other categories of conservation (*i.e.*,

628 ESUs, MUs, etc). This will significantly help the planning of conservation strategies toward the populations that are most  
629 vulnerable to climate change, and therefore, for which conservation measures should be prioritized.

### 630 **Acknowledgements**

631 Many thanks to the "Fédération de Pêche de Corse" and especially to Stéphane Muracciole for providing the Corsican  
632 samples. A special thanks to Dr. Stephanie Susan Ames (CSAL Centro di Supporto per l'Apprendimento delle Lingue  
633 dell'Università Politecnica delle Marche) for the linguistic revision of the manuscript.

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**TABLE 1.** Sites of the 20 wild Sardinian brown trout sampling sites analyzed in this study. N, represents the whole sample size. VES and VIV represent two wild brown trout samples from Corsica analysed in the present study, while LTT, CTT and HBT are Corsican samples from Reynaud et al. (2011) (see material and methods section for more details). HATa and HATb represent two traditional hatchery strains used here as reference samples of the Atlantic genome. Environmental parameters: Elevation; mean monthly highest water temperature (JN = June, JL = July, AG = August, SP = September); number (between bracket) of impassible natural and or artificial barriers between the sampling site and the stream/river outflow (W = weir, D = Dam, F = ford, WF = waterfall; see also Table S4 for more details); mean summer discharge; duration of drought in days; length in meters of the dry river portion, rivers total length. Demographic parameters: trout density, estimated by applying the two-pass sampling removal method (Zippin 1956). Protected areas (RP = Regional Park, SCI = Site of Community Importance based on the Habitat Directive, \*\* denoted protected areas where the fishing activities are prohibited (DR n.314/Dec.A9 07.02.2019).

Location code	N	Region	Stream/River	Basin	Sea drainage	Elevation (m.a.s.l.)	Highest mean summer water temperature (°C) *	Barriers	Mean summer discharge (m <sup>3</sup> s <sup>-1</sup> )	Drought duration (days)	Drought length (m) <sup>§</sup>	River length (m)	Trout density (ind m <sup>-2</sup> )	Protected areas	
Sardinia	COG	7	Sardinia	Riu Bizzolu	Coghinas	Gulf of Asinara	276	23.43 (JL)	W (3)	0.0463		16284		RP	
	PAD	13	Sardinia	Riu de su Piricone	Padrogiano	Tyrrhenian Sea	140	23.86 (SP)	D (1)	0.1105		32190	0.0163		
	POSa	7	Sardinia	Canale dell'Iserno	Posada	Tyrrhenian Sea	569	23.40 (JL)	WF(1)	0.0213		11443	0.0047		
	POsb	18	Sardinia	Riu s'Abba e Salinu	Posada	Tyrrhenian Sea	507					6194	0.0210		
	CED	30	Sardinia	Riu Flumineddu	Cedrino	Tyrrhenian Sea	189	23.54 (JN)		0.4870	330	10000	35097	0.1369	SCI (**)
	CDL	8	Sardinia	Riu Codula de Luna	Riu Codula de Luna	Tyrrhenian Sea	254	19.00 (JN)		0.2025			21855	0.0257	SCI
	FLUa	10	Sardinia	Flumendosa	Flumendosa	Tyrrhenian Sea	802	19.80 (JN)	D (1)	0.0308			147878	0.0619	
	FLUb	9	Sardinia	Riu Bau Mandara	Flumendosa	Tyrrhenian Sea	977	20.32 (JL)	WF (1)	0.0375			13689	0.0090	
	FLUc	11	Sardinia	Riu Furittu	Flumendosa	Tyrrhenian Sea	390			0.0290	120	8848	14043	0.0504	(**)
	FMCa	8	Sardinia	Riu Cannisoni	Flumini Mannu di Cagliari	Gulf of Cagliari	380	23.90 (JL)	W (4)	0.0215			9346	0.0179	SCI
	FMCb	12	Sardinia	Riu su Salixi	Flumini Mannu di Cagliari	Gulf of Cagliari	425	20.65 (JL)	D (1)	0.0300			4536	0.0750	
	PULa	12	Sardinia	Riu Litteras	Pula	Gulf of Cagliari	296	21.90 (JL)		0.0328	120	2641	2848	0.1280	SCI
	PULb1	8	Sardinia	Rio Pula	Pula	Gulf of Cagliari	170			0.1950	120	13282	30832	0.0083	SCI
	PULb2	23	Sardinia	Rio Pula	Pula	Gulf of Cagliari	144		W (1)	0.1950	120	13282	30832	0.0792	RP
	FMPa	30	Sardinia	Riu Piras	Flumini Mannu di Pabillonis	Mediterranean Sea	324	26.27 (JL)	W (19)		120	6208	12293	0.2057	SCI (**)
	FMPb	17	Sardinia	Riu Sitzedda	Flumini Mannu di Pabillonis	Mediterranean Sea	323					4600	7001	0.0653	SCI
	TEM	6	Sardinia	Riu Matta Giuanna	Temo	Mediterranean Sea	722	27.00 (JL)	WF (1)	0.0475			12129	0.0200	
	RMN	10	Sardinia	Riu Mannu	Mare Foghe	Mediterranean Sea	465	22.15 (JL)	WF (1)	0.2283			25160	0.3200	
	RMF	5	Sardinia	Riu di Mare Foghe	Mare Foghe	Mediterranean Sea	192						33000	0.0420	
CIX	30	Sardinia	Riu Is Abius	Cixerri	Gulf of Cagliari	308	21.20 (AG)	F(3), D (1)	0.0078	120	2500	3421	0.2816		
Corse	LTT	5	Corsica	Lette	Seccu	Mediterranean Sea									
	CTT	5	Corsica	Ciuttare	Liamone	Mediterranean Sea									
	HBT	5	Corsica	Haut Botaro	Liamone	Mediterranean Sea									
	VES	19	Corsica	Ese	Prunelli	Mediterranean Sea									
	VIV	20	Corsica	Speluncello	Vecchio	Tyrrhenian Sea									
Hatch.	HATa	26	Central Italy	Hatchery a	Cantiano	Adriatic Sea									
	HATb	20	Central Italy	Hatchery b	Visso	Tyrrhenian Sea									

\* data provided by Agenzia regionale del distretto idrografico della Sardegna, § Drought length was evaluated during the summer months (July - September) from 2006 and 2020 years

**TABLE 2.** Intra-population genetic diversity obtained by using mtDNA CR sequence analysis, PCR-RFLP analysis of *LDH-C1\** gene and 10 microsatellites genotyping on 20 wild brown trout Sardinian sampling sites, 2 reference samples from wild brown trout Corsican sampling sites and 2 reference populations for the brown trout Atlantic hatchery stock. LTT, CTT and HBT are Corsican sampling sites from Reynaud et al., 2011.

Location code	N	CR haplotypes (mtDNA)																	LDH-C1*		Microsatellites										
		A2	AD-Tyrrh1	AD-Tyrrh4	AD-Tyrrh7	AD-Tyrrh8	AD-Tyrrh9	AD-Tyrrh10	AD-Tyrrh11	AD-Tyrrh12	AD-Tyrrh13	AD-Tyrrh14	ADcs23	ADcs24	ADcs25	DaLa	Haplotype 1	Haplotype 2	Haplotype 3	Haplotype 4	AT-Tyrrh1	Alle	*90	*100	Ar	Ho	He	Fis	q (90% CI)	I	
Sardinia	COG	7	-	0.57	-	-	-	-	0.29	-	-	-	-	-	-	-	0.14	-	-	-	-	-	0.21	0.79	2.71	0.55	0.59	0.078	0.990 (0.933 - 1.000)	II	
	PAD	13	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.12	0.88	2.65	0.61	0.56	-0.097	0.987 (0.917 - 1.000)	II	
	POSa	7	-	0.86	-	-	-	-	-	0.14	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	2.83	0.50	0.56	0.118	0.955 (0.885 - 1.000)	III	
	POSB	18	-	0.74	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.26	0.36	0.64	3.07	0.58	0.61	0.038	0.974 (0.884 - 1.000)	III
	CED	30	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.02	0.98	2.06	0.50	0.52	0.048	0.993 (0.964 - 1.000)	II
	CDL	8	-	0.75	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.25	-	-	-	0.31	0.69	2.75	0.52	0.54	0.020	0.981 (0.891 - 1.000)	III
	FLUa	10	-	-	-	0.11	-	-	-	-	-	-	-	-	-	-	0.89	-	-	-	-	-	-	0.85	0.15	3.43	0.79	0.74	-0.071	0.012 (0.000 - 0.083)	IV
	FLUb	9	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	2.65	0.54	0.55	0.018	0.919 (0.828 - 1.000)	III	
	FLUc	11	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.99	0.49	0.45	-0.089	0.994 (0.967 - 1.000)	I	
	FMCa	8	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.13	0.88	2.83	0.52	0.65	0.221	0.992 (0.949 - 1.000)	II
	FMCb	12	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.67	0.33	-	-	-	-	0.83	0.17	3.37	0.72	0.72	-0.013	0.004 (0.000 - 0.019)	II
	PULa	12	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.77	0.30	0.54	0.475	0.970 (0.925 - 0.991)	II	
	PULb1	8	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.36	0.31	0.37	0.176	0.995 (0.978 - 1.000)	I	
	PULb2	23	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.28	0.33	0.35	0.027	0.998 (0.993 - 1.000)	I	
	FMPa	30	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.52	0.52	0.48	-0.086	0.997 (0.984 - 1.000)	I	
	FMPb	17	-	-	-	-	-	-	-	0.47	0.53	-	-	-	-	-	-	-	-	-	-	-	-	0.15	0.85	1.92	0.39	0.41	0.042	0.982 (0.912 - 1.000)	IV
TEM	6	-	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.33	0.67	1.87	0.45	0.42	-0.086	0.991 (0.941 - 1.000)	II	
RMN	10	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	0.77	0.22	3.30	0.65	0.72	0.107	0.875 (0.761 - 0.922)	III	
RMF	5	-	-	-	-	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	0.30	0.70	2.94	0.64	0.62	-0.036	0.992 (0.955 - 1.000)	II	
CIX	30	0.73	-	0.27	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.48	0.28	0.29	0.056	0.997 (0.987 - 1.000)	I		
Corse	LTT	5	-	-	-	-	-	-	-	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
	CTT	5	-	-	-	-	-	-	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
	HBT	5	-	-	-	-	-	-	-	-	-	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
	VES	19	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.82	0.43	0.51	0.081	0.998 (0.987 - 1.000)	I	
VIV	20	-	1.00	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	1.00	1.80	0.27	0.37	<b>0.283</b>	0.981 (0.944 - 1.000)	I		
Hac.	HATa	26	-	-	-	-	-	-	-	-	-	-	-	-	-	0.63	-	-	-	0.37	-	-	0.96	0.04	4.08	0.85	0.82	-0.044			
	HATb	20	-	-	-	-	-	-	-	-	-	-	-	-	-	0.13	0.74	0.13	-	-	-	-	1.00	-	4.06	0.75	0.81	0.075			

From left: location code; sample size (N); frequency of mtDNA Control Region haplotype(s) observed; *LDH-C1\** allele frequencies; Allelic richness (*Ar*); observed heterozygosity (*H<sub>o</sub>*); expected heterozygosity (*H<sub>e</sub>*); Fixation index (*F<sub>is</sub>*) with significant adjusted nominal level (5%) (*P* < 0.00021) given in bold; mean admixture coefficient (*q*) and 90% credible intervals (CI); Introgression rates (I, pure native trout; II, low introgressed trout; III, moderately introgressed trout; IV, non-native trout) based on admixture (*q*) values and their CIs, frequency of *LDH-C1\*90* allele and AT-DA haplotypes, see section 3.2 for more details.

**TABLE 3.** AMOVA hierarchical analysis examining the partitioning of genetic variance of mitochondrial (Control Region) and nuclear DNA (10 microsatellite loci) according to two hypothesized spatial structures: sites grouped by sea drainages and sites grouped by river basins (as defined in Table 1). The amount of variation (%) explained by differences among groups, among populations within groups and within populations, along with the p-value (statistically significant values are in bold) are provided.

No. of groups and group composition	Hierarchical level	Control Region		Microsatellites	
		Variation (%)	p	Variation (%)	p
<b>12 river basins</b>	among groups	83.37	<b>0.000</b>	16.49	<b>0.000</b>
COG / PAD / POSa+POSb / CED / CDL / FLUa+FLUb+FLUc / FMCa+FMCb / PULa+PULb1+PULb2 / FMPa+FMPb / TEM / RMN/ RMF / CIX	among populations within groups	4.64	<b>0.000</b>	29.22	<b>0.000</b>
	within populations	11.98	<b>0.000</b>	54.28	<b>0.000</b>
<b>4 sea drainages</b>	among groups	55.82	<b>0.000</b>	12.68	<b>0.000</b>
COG / PAD+POSa+POSb+CED+CDL+FLUa+FLUb+FLUc / FMCa+FMCb+PULa+PULb1+PULb2+CIX / FMPa+FMPb+TEM+ RMN + RMF	among populations within groups	33.56	<b>0.000</b>	34.44	<b>0.000</b>
	within populations	10.62	<b>0.006</b>	52.88	<b>0.000</b>

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**TABLE 4.** Effective population size estimates (Ne), with 95% confidence intervals based on linkage disequilibrium (NeEstimator, Ne1) and sibship approaches (Colony, Ne2), and tests of recent events of bottleneck based on Wilcoxon's test and using the allele frequency distribution mode-shift method for 19 wild Sardinian brown trout and two wild Corsican brown trout samples. In bold, the significant p-values ( $P < 0.05$ ) of the Wilcoxon tests.

	NeESTIMATOR (LD method)			COLONY (random mating method)			I.A.M Wilcoxon 1-way	T.P.M Wilcoxon 1-way	L-Shaped distribution
	Ne1	Lower 95% CI	Upper 95% CI	Ne2	Lower 95% CI	Upper 95% CI			
COG	∞	8.9	∞	56	16	∞	0.326	0.714	Shifted mode
PAD	∞	71.7	∞	∞	1	∞	0.752	0.997	Normal
POSa	7.4	2.2	162.6	42	12	∞	0.862	0.991	Normal
POSb	25.8	14.9	61.8	29	16	61	0.577	0.958	Normal
CED	42.6	16.5	∞	23	14	44	0.469	0.973	Normal
CDL	∞	9.4	∞	37	14	∞	0.934	0.998	Normal
FLUa	11.6	4.9	44.4	13	6	64	<b>0.001</b>	<b>0.005</b>	Shifted mode
FLUb	2.8	1.6	11.7	24	10	∞	0.385	0.754	Normal
FLUc	31.5	2.4	∞	28	12	315	0.629	0.987	Normal
FMCa	21.8	3.2	∞	28	11	∞	<b>0.001</b>	<b>0.002</b>	Normal
FMCb	5.6	2.9	10.2	16	7	50	<b>0.001</b>	<b>0.042</b>	Normal
PULa	2.6	0.5	∞	12	6	38	<b>0.008</b>	<b>0.040</b>	Shifted mode
PULb	9.9	1.2	∞	11	6	26	0.563	0.843	Shifted mode
FMPa	5.9	1.6	27.6	12	6	30	<b>0.016</b>	0.078	Normal
FMPb	∞	18	∞	20	10	43	0.500	0.898	Normal
TEM	∞	1.8	∞	∞	1	∞	0.980	0.989	Normal
RMN	16.5	6.7	170.8	23	10	299	<b>0.002</b>	0.215	Normal
RMF	∞	9.5	∞	20	6	∞	0.179	0.820	Shifted mode
CIX	1.6	0.8	3.7	10	5	28	0.422	0.781	Normal
VIV	10	3.2	30.9	25	14	52	0.629	0.980	Normal
VES	16	2.9	∞	15	7	31	<b>0.008</b>	0.055	Normal

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**TABLE 5** Pairwise  $F_{ST}$  based on 10 microsatellite loci between 19 wild Sardinian brown trout sampling sites (blue headers), 2 wild Corsican brown trout populations (orange headers) and 2 (yellow headers) Atlantic brown trout hatchery strains (below diagonal). p values (above diagonal) were obtained after 5060 permutations, indicative adjusted nominal level-5% for multiple comparisons is 0.000198. C G L =  $F_{ST}$  color gradient legend.

	COG	PAD	POSa	POsb	CED	CDL	FLUa	FLUb	FLUc	FMCa	FMCb	PULa	PULb1	PULb2	FMPa	FMPb	TEM	RMN	RMF	CIX	VES	VIV	HATa	HATb
COG		NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS
PAD	0.218		*	*	*	NS	*	*	NS	*	*	NS	NS	*	*	*	*	*	NS	*	*	*	*	*
POSa	0.191	0.176		*	*	NS	*	NS	NS	NS	*	NS	NS	*	*	*	NS	*	NS	*	*	*	*	*
POsb	0.174	0.151	0.108		*	*	*	*	NS	*	*	NS	NS	*	*	*	*	*	NS	*	*	*	*	*
CED	0.393	0.269	0.356	0.334		*	*	*	NS	*	*	NS	NS	*	*	*	*	*	NS	*	*	*	*	*
CDL	0.228	0.292	0.280	0.227	0.380		NS	NS	NS	NS	*	NS	NS	*	*	*	*	NS	NS	NS	*	NS	*	NS
FLUa	0.266	0.299	0.263	0.258	0.426	0.289		NS	NS	*	*	NS	NS	*	*	*	NS	*	NS	*	*	*	*	*
FLUb	0.277	0.287	0.271	0.248	0.407	0.322	0.284		NS	NS	*	NS	NS	*	*	*	NS	NS	NS	*	*	*	*	NS
FLUc	0.419	0.447	0.396	0.385	0.548	0.349	0.381	0.478		NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS
FMCa	0.219	0.269	0.221	0.210	0.420	0.270	0.227	0.232	0.397		*	NS	NS	*	*	*	NS	*	NS	*	*	*	*	*
FMCb	0.278	0.285	0.266	0.250	0.428	0.294	0.176	0.288	0.419	0.232		NS	NS	*	*	*	*	*	NS	*	*	*	*	*
PULa	0.379	0.440	0.370	0.367	0.558	0.421	0.357	0.404	0.555	0.365	0.429		NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS	NS
PULb1	0.473	0.480	0.407	0.394	0.563	0.524	0.454	0.445	0.635	0.431	0.479	0.213		NS	*	NS	NS	NS	NS	*	NS	*	*	NS
PULb2	0.607	0.572	0.537	0.489	0.625	0.621	0.572	0.565	0.696	0.559	0.591	0.232	0.273		*	*	*	*	*	*	*	*	*	*
FMPa	0.551	0.526	0.474	0.447	0.586	0.562	0.533	0.533	0.610	0.434	0.535	0.617	0.621	0.643		*	*	*	*	*	*	*	*	*
FMPb	0.447	0.455	0.393	0.370	0.517	0.463	0.443	0.423	0.545	0.363	0.403	0.553	0.569	0.625	0.550		*	*	NS	*	*	*	*	*
TEM	0.393	0.373	0.310	0.278	0.492	0.471	0.413	0.452	0.614	0.402	0.363	0.648	0.712	0.770	0.669	0.505		NS	NS	*	*	*	*	*
RMN	0.276	0.267	0.233	0.229	0.430	0.294	0.169	0.277	0.382	0.233	0.157	0.405	0.471	0.589	0.538	0.403	0.346		NS	*	*	*	*	*
RMF	0.257	0.246	0.218	0.209	0.397	0.284	0.271	0.248	0.423	0.214	0.261	0.431	0.491	0.619	0.531	0.388	0.394	0.211		NS	NS	*	*	NS
CIX	0.579	0.524	0.534	0.506	0.587	0.616	0.574	0.483	0.691	0.542	0.589	0.561	0.567	0.539	0.605	0.612	0.744	0.593	0.588		*	*	*	*
VES	0.454	0.446	0.468	0.395	0.540	0.421	0.471	0.498	0.527	0.463	0.486	0.585	0.654	0.705	0.652	0.583	0.613	0.448	0.473	0.697		*	*	*
VIV	0.514	0.524	0.490	0.437	0.586	0.512	0.478	0.532	0.593	0.479	0.493	0.619	0.673	0.726	0.645	0.605	0.650	0.458	0.519	0.713	0.584		*	*
HATa	0.232	0.254	0.219	0.216	0.370	0.256	0.093	0.234	0.333	0.162	0.075	0.327	0.381	0.468	0.425	0.327	0.320	0.109	0.211	0.479	0.408	0.409		*
HATb	0.261	0.254	0.229	0.220	0.377	0.278	0.101	0.251	0.352	0.178	0.085	0.363	0.407	0.506	0.456	0.355	0.338	0.094	0.205	0.510	0.421	0.420	0.026	
C G L	0.026	0.060	0.094	0.128	0.162	0.195	0.229	0.263	0.297	0.331	0.364	0.398	0.432	0.466	0.500	0.533	0.567	0.601	0.635	0.669	0.702	0.736	0.770	



## Figure Captions

**FIGURE 1** Map of the study area showing the brown trout sampling locations from investigated Sardinian and Corsican rivers. Solid lines mark boundaries of major drainage basins. Dashed line: coastline during the last glacial maximum (LGM); downloaded from Zickel et al. (2016) GIS dataset. Pie charts represent the geographic distribution and frequency of CR mtDNA haplotypes per sampling site. Pie chart size is proportional to the sampling site size.

**FIGURE 2** Calibrated chronogram of the genus *Salmo* created with an optimized relaxed clock in Beast2. Blue bars at the nodes represent 95% highest posterior density (hpd) intervals, only clade showing posterior probability greater than 0.9 are represented. Median node ages are shown as node labels and Beast/BI posterior probability greater than 0.5 are reported. Time estimates are given in millions of years. Calibration points are indicated by stars. Asterisk: the haplotype *AD-Tyrrh4* include also the haplotypes *AD-Tyrrh-9* and *I3* (see section 3.1).

**FIGURE 3** Parsimony network (95%) of CR *S. trutta* species complex and *S. orhidanus* haplotypes used in this study. In bold, the *S. trutta* CR haplotypes observed in this study. Pie charts indicate the frequency (circle sizes are proportional to observed haplotype frequencies) and distribution of haplotypes across basins (as indicated in Table 1). The white circles along the branches represent the mutational steps. The dashed box includes the CR Corso-Sardinian lineage haplotypes. Asterisk: the haplotype *AD-Tyrrh4* include also the haplotypes *AD-Tyrrh-9* and *I3* (see section 3.1).

**FIGURE 4** Hierarchical STRUCTURE analysis based on 10 microsatellites adopted to detect the genetic diversity of 273 wild brown trout from 20 sampling localities from 12 Sardinian river basins, 39 wild brown trout populations from 2 Corse populations and 46 specimens from 2 hatchery-reared Atlantic brown trout strains. Black lines separate sampling locations, whose codes (as in Table 2) are reported to the side of each bar plot. K outcomes obtained for each hierarchical round of STRUCTURE analysis are reported within the arrows positioned above the corresponding bar plot.

**FIGURE 5** Plots showing the two discriminant axes of a hierarchical discriminant analysis of principal components carried out on wild brown trout sampling sites from Sardinia and Corsica and two hatchery strains of Atlantic origin: A) all sampling sites included; B) all sampling sites, but PULa-b1-2, CIX, VIV and FMPa; C) all B step samples, but CED, VES and FMPb. Each trout is represented as a dot and the samples are represented as inertia ellipses.

**FIGURE S1** Second-order polynomial regressions between the frequency of the *LDH-C1\*90* allele and measures of per-site/hatchery genetic diversity: A, *Ar/LDH-C1\*90* allele frequency; B, *He/LDH-C1\*90* allele frequency.

**FIGURE S2** Plots of individual admixture coefficient ( $q$ ), including their 90% probability limits for individuals from 20 wild Sardinian brown trout. Sampling sites from the same river basin were plotted on the same plot. Location codes as in Table 1

## Appendices

**TABLE S1.** Control Region (CR) sequences used in this study. CR mtDNA lineage codes: ME = Mediterranean ; AD = Adriatic; MA = *marmoratus*; AT = Atlantic; DA = Danubian.

Haplotype	Lineage	Locality	Taxon	GenBank Accession number	Source
ADcs1	AD	Atlantic and West Mediterranean basin of Andalusia (Spain); Aegean basin (Balkans); Adriatic basin, Prespa (Albania, FYROM and Greece); Adige River (North Italy)	<i>S. trutta</i> , <i>S. carpio</i> , <i>S. peristericus</i> , <i>S. platycephalus</i>	AY836330	1; 2; 3; 4; 5
ADcs6	AD	West Mediterranean basin (Spain)	<i>S. trutta</i>	AY836335	1
ADcs7	AD	West Mediterranean basin (Spain)	<i>S. trutta</i>	AY836336	1
ADcs10	AD	West Mediterranean basin (Spain)	<i>S. trutta</i>	AY836339	1
ADcs11	AD	Adriatic basin (Greece; Albania; Montenegro; Serbia)	<i>S. trutta</i> , <i>S. dentex</i>	AY836340	1; 3; 6; 7
ADcs15	AD	Mediterranean basin (Corsica)	<i>S. trutta</i>	AY836344	1
ADcs16	AD	West Mediterranean basin (Spain)	<i>S. trutta</i>	AY836345	1
ADcs17	AD	West Mediterranean basin (Spain)	<i>S. trutta</i>	AY836346	1
ADcs18	AD	Atlantic basin-Andalusia (Spain)	<i>S. trutta</i>	AY836347	1
ADcs19	AD	Atlantic basin-Andalusia (Spain)	<i>S. trutta</i>	AY836348	1
ADcs20	AD	Adriatic and Aegean basins (Bulgaria, Greece)	<i>S. trutta</i>	AY836349	1; 2
ADrh1	AD	West Mediterranean basin Durance (France)	<i>S. trutta</i>	MK948035	8
ADporh1	AD	West Mediterranean basin Durance (France); Adriatic basins Pellice, Tanaro (North-West Italy)	<i>S. trutta</i>	MK948034	8
A_2	MA	Tyrrhenian basin (Sardinia)	<i>S. trutta</i>	KM216129	9; This study
AD-Tyrrh1	AD	Tyrrhenian basin (Corsica, Sardinia, Italy)	<i>S. trutta</i>	KX450257	9; This study
AD-Tyrrh2	AD	Tyrrhenian basin (Corsica and Italy)	<i>S. trutta</i>	KX450258	9
AD-Tyrrh3	AD	Tyrrhenian basin (Italy)	<i>S. trutta</i>	KX450259	9
AD-Tyrrh4	AD	Mediterranean and Tyrrhenian basins (Sardinia, Italy)	<i>S. trutta</i>	KX450260	9; This study
AD-Tyrrh5	AD	Tyrrhenian basin (Italy)	<i>S. trutta</i>	KX450261	9
AD-Tyrrh6	AD	Tyrrhenian basin (Italy)	<i>S. trutta</i>	KX450262	9
AD-Tyrrh7	AD	Tyrrhenian basin (Sardinia)	<i>S. trutta</i>	MT503201	10; This study
AD-Tyrrh8	AD	Tyrrhenian basin (Sardinia)	<i>S. trutta</i>		This study
AD-Tyrrh10	AD	Mediterranean basin (Sardinia)	<i>S. trutta</i>		This study
AD-Tyrrh11	AD	Tyrrhenian basin (Sardinia)	<i>S. trutta</i>		This study
AD-Tyrrh12	AD	Mediterranean basin (Sardinia)	<i>S. trutta</i>		This study
AD-Tyrrh14	AD	Mediterranean basin (Sardinia)	<i>S. trutta</i>		This study
<i>S. letnica</i> hap12	AD	Lake Ohrid (FYROM-Albania)	<i>S. letnica</i>	AY926570	11
<i>S. letnica</i> hap13	AD	Lake Ohrid (FYROM-Albania)	<i>S. letnica</i>	AY926573	11
<i>S. letnica</i> hap15	AD	Lake Ohrid (FYROM-Albania)	<i>S. letnica</i>	AY926572	11

MEcs2	ME	Western ME basin (Spain and France) AD basin (Albania and (North-West Italy), Krka River (Croatia)	<i>S. trutta</i>	AY83635 1	1; 3
MEcs3	ME	Western ME basin (Spain) Danube-Bistrica Ponto- Caspian basin (Slovenia)	<i>S. trutta</i>	AY83635 2	1
MEcs4	ME	Western ME basin (Spain).	<i>S. trutta</i>	AY83635 3	1
MEcs6	ME	Western ME basin (Spain).	<i>S. trutta</i>	AY83635 5	1
MEcs7	ME	Western ME basin (Spain).	<i>S. trutta</i>	AY83635 6	1
MEcs8	ME	Western ME basin (Spain).	<i>S. trutta</i>	AY83635 7	1
MAcs1	MA	Adriatic basin-Soca River (Slovenia); Adige and Po rivers (North Italy); Aegean basin (Greece)	<i>S. trutta</i>	AY83636 5	1; 2
Ma2a	MA	North Italy	<i>S. trutta</i>	DQ84118 9	5; 12
Ma2b	MA	North Italy	<i>S. trutta</i>	DQ84119 0	5; 8; 12
Ma2c	MA	North Italy	<i>S. trutta</i>	JQ58246 1	5; 8
Masl1	MA	North western Italy	<i>S. trutta</i>	MK9480 36	8
MAcs4	MA	North Italy	<i>S. trutta</i>	JN20802 2	13; 14
H1	AT	Denmark-Norway; Vistula, Elbe, Danube and Oder rivers (Central Europe); North Italy*	<i>S. trutta</i>	AF27308 6	5; 12; 17; 18; This study
H2	AT	Denmark-Norway; Vistula, Elbe, Danube and Oder rivers (Central Europe); North Italy*	<i>S. trutta</i>	AF27308 7	5; 12; 17; 18; This study
H3	AT	Denmark-Norway; Vistula, Elbe, Danube and Oder rivers (Central Europe); North Italy*	<i>S. trutta</i>	AF27457 4	5; 12; 17; 18; This study
H4	AT	Denmark-Norway; Vistula, Elbe, Danube and Oder rivers (Central Europe); North Italy*	<i>S. trutta</i>	AF27457 5	5; 12; 17; 18; This study
ATcs11	AT	Beherobentako (South France); Duero River (Spain)	<i>S. trutta</i>	AY83632 7	1
ATcs13	AT	Beherobentako (South France)	<i>S. trutta</i>	AY83632 9	1
At1e	AT	Adige River (Northern Italy)*	<i>S. trutta</i>	DQ84119 2	12; This study
ATSic	AT	Mediterranean basin (Sicily)	<i>S. trutta</i>	JF297974	14; 15
AT-Tyrrh1	AT	Tyrrhenian basin (Italy)	<i>S. trutta</i>	KX45026 3	9; This study
CloneJE1	AT	South European and African atlantic basin (Spain and Marocco) Mediterranean basin (Sicily)	<i>S. trutta</i>	AF25355 7	9; 16
Da1a	DA	Danube and Vistula basins (Central Europe, Bulgaria, Serbia); Adige River (Northern Italy)	<i>S. trutta</i>	AY18556 8	2; 12; 17; 18; This study
Da1b	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18556 9	17; 18
Da23a	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 4	17
Da23b	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 5	17
Da24	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 6	17
Da9	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 2	17
Da2	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 0	17; 18
Da3	DA	Danube basin (Austria)	<i>S. trutta</i>	AY18557 1	17
Da22	DA	Danube and Vistula basins (Central Europe), Balkans, Adige River (North Italy)	<i>S. trutta</i>	AY18557 3	12; 17; 18
<i>S. ohridanus</i> hap 3		Lake Ohrid	<i>S.ohridanus</i>	AY92656 8	11
<i>S. ohridanus</i> hap 4		Lake Ohrid	<i>S.ohridanus</i>	AY92656 1	11
<i>S.</i>		Lake Ohrid	<i>S.ohridanus</i>	AY92656	11

ohridanus hap 8				7	
S. ohridanus hap 9		Lake Ohrid	S.ohridanus	AY926565	11
S. ohridanus hap 10		Lake Ohrid	S.ohridanus	AY926562	11

**Table S1**  
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Repeat motif	Primers sequence (5' 3')
(CT) <sub>13</sub> ACCA(CT) <sub>3</sub>	F: CGG TGT GCT TGT CAG GTT TC

	R: GTC AAG TCA GCA AGC CTC AC
(GT) <sub>14</sub>	F: ACC CGC TCC TCA CTT AAT C R: AGG TGG GTC CTC CAA GCT AC
(TG) <sub>25</sub>	F: TTG TTC AGT GTA TAT GTG TCC CAT R: GAT CTT CAC TGC CAC CTT ATG ACC
(CA) <sub>4</sub> AA (CA) <sub>14</sub>	F: GCTGTGATTTCTCTCTG C R: AAAGGTGGGTCCAAGGAC
(GTTA) <sub>22</sub>	F: ATG TGG AGG TCA ACT AAC CAG CGT G R: CAT CAA TCA CAG AGT GAG GCA CTC G
(GTTA) <sub>25</sub>	F: GGCCAGACAGATAAAACAAACACGC R: GCCAACAGCAGCATCTACACCCAG
(GATA) <sub>19</sub>	F: AGA ATG CTA CTG GTG GCT GTA TTG TGA R: TCT GAA AGA CAG GTG GAT GGT TCC
(GATG) <sub>x</sub>	F: GGC ATT GGA GGTAAG GAC AC R: CCA GAC CAC TGA ACT TCT CAT C
(GACA) <sub>22</sub>	F: GGA AAA TAA TCA ATG CTG CTG GTT R: CTA CAA TCT GGA CTA TCT TCT TCA
(GACA) <sub>27</sub>	F: AAT GGA TTA CGG GTA CGT TAG ACA R: CTC TTG TGC AGG TTC TTC ATC TGT

## Table S2 references

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**Table S3.** Tmrca values for a time calibrated phylogeny of the *Salmo* genus. Clades showing posterior probability greater than 0.5 are reported.

Taxon/lineage	T <sub>MRCA</sub> [95% HPD ]	Posterior probability
<i>S. immigratus</i>	11.388 [10.093, 14.668]	1
<i>S. ohridanus</i>	1.659 [0.255, 4.672]	1
BT	3.829 [1.833, 8.536]	1
AT+DA	3.097 [1.206, 7.166]	0.51
AT	1.53 [0.367, 3.950]	1
DA	1.94 [0.547-4.731]	1
ME	1.263 [0.244-3.475]	1
MA	1.299 [0.213-3.601]	1
AD	2.515 [0.853-5.836]	1
Corso-Sardinian	1.051 [0.243-2.724]	1

**TABLE S4.** Definition of impassable barriers listed in Table 1

Ford	An impediment for stream crossing for fish passage, as they often combine many of the negative features of culverts and weirs. In particular, we have considered a ford impassable when it combines a downstream face with a steep drop exceeding 50 cm and shallow water over the ford.
Weir	Weirs combine several obstacles to upstream and downstream passage of fish, including fall heights that prevent swimming species from migrating upstream and crest shapes that may be challenging for climbing trout. We consider the weirs unsuitable for trout passage when they exceed a height of 1 meter.
Dam	Larger dams (average height of 42.5 ± 3 m) small dams with a height lower than 15 meters.
Waterfall	An abrupt change in water velocity, characterized by a vertical drop of at least 1 meter.