

The role of the south-western Alps as a unidirectional corridor for Mediterranean brown trout (Salmo trutta complex) lineages

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1 The Role of the south-western Alps as a unidirectional corridor for

2 Mediterranean brown trout (Salmo trutta complex) lineages

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- 22 **Running title:** The south-western Alps: a corridor for *Salmo trutta*
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31 Abstract

The role of south-western Alps as a corridor for Mediterranean trout (Salmo trutta complex Linnaeus, 1758) was evaluated in order to understand the influence of the last glacial events in shaping the spatial distribution of the genetic diversity of this salmonid. For this, the allochthonous hypothesis of a man-mediated French origin (19th century) of the Mediterranean trout inhabiting the Po tributaries in the Italian side of south-western Alps was tested. A total of 412 individuals were analyzed at the mitochondrial control region. The phylogenetic classification was carried out by using a Median-Joining Network analysis. Mismatch pair-wise analysis, molecular dating and Kernel density distribution analysis of the main mitochondrial lineages were evaluated to compare past demographic dynamics with the current spatial distribution of genetic diversity. The main outcomes resulted strongly in agreement with a biogeographic scenario where the south-western Alps acted as a unidirectional corridor that permitted the colonization of the upper Durance (Rhône River basin) by trout from the Po River basin. Therefore, the Mediterranean trout should be considered as native also along the Italian side of the south-western Alps and the allochthonous hypothesis should be rejected.

Key words: Salmo trutta complex, Alpine barrier, ice cover, conservation genetics, biological corridors

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61 INTRODUCTION

The study of the phylogeographic history of a species represents a fundamental step to understand the 62 factors producing genetic diversity both within and among species, with particular reference to the role 63 played by past environmental and climatic changes in shaping high levels of genetic complexity as well as 64 their current spatial distribution (Purvis & Hector, 2000). The reconstruction of the recent evolutionary 65 history of an organism is therefore essential to set-up concrete management and conservation actions aiming 66 at protecting its evolutionary potential (Losos et al., 2013). This is particularly true for those organisms 67 impacted by socio-economic interests, where conservation induces conflict with commercial interests 68 (Redpath et al., 2013). In addition, the tentative to state the natural occurrence of an organism on the basis of 69 70 phylogenetic studies could be made difficult by the long history of the human-mediated natural species range 71 alterations, that may act as a confounding factor that adds further complexity in the study of the already 72 complex evolutionary pathway of an organism (Fernández-García et al., 2014; Sanz, 2018).

73 Although, the human ability in shaping plant and animal distribution is rooted in ancient culture and 74 traditions (Larson & Fuller, 2014), nowadays, human-mediated transport beyond biogeographic barriers has 75 led to the introduction and establishment of alien species (sometimes invasive) in new regions worldwide 76 (Shackleton et al., 2019; Berrebi et al., 2020). About freshwater fish, the first written historical records date 77 back the major human-mediated introductions at Roman time and successively at medieval period (Sønstebø, Borgstrøm, & Heun, 2007; Miró & Ventura, 2013). Basically, the above ancient freshwater fish 78 79 translocations were promoted by food purposes, whereas, nowadays, freshwater fish represents one of the most important group of animals introduced for sport purposes worldwide (Leprieur et al., 2008). In 80 81 particular, brown trout (Salmo trutta complex) is listed as the 100 of the world's worst invasive alien species 82 (Global Invasive Species Database [2019], downloaded from http://193.206.192.138/gisd/100 worst.php on 83 13-03-2019). At the same time, its natural diversity is imperilled in much of its native range (Budy et al., 2013). 84

85 The Italian IUCN red list of vertebrates (Rondinini et al., 2013) classified its conservation status as near 86 threatened, and the French red list (IUCN, 2010) considers the trout status as low concern. It is probably 87 wildly optimistic statements considering that along the Apennine chain less than 3% of native populations were free of genetic introgressive hybridization with Atlantic genes of domestic origin (Splendiani et al., 88 2016a). Further, in the Alpine area, its conservation value appears at least chaotic. The brown trout 89 90 (regardless of its phylogenetic origin) is "on paper" considered as allochthonous in the Piemonte Region and 91 native in the neighbouring Lombardia Region (based on Regional lists of freshwater fishes). In addition, the 92 AIIAD (Italian Association of Freshwater Fish Ichthyologists) guideline for Italian salmonid management 93 proposes to adopt a passive conservation approach for the putative native populations of south-western Alps 94 (Zanetti, Nonnis Marzano & Lorenzoni 2013). Based on the opinions expressed in this document, the

95 Mediterranean trout populations recognizable in this region should be the consequence of historical humanmediated translocations from the Rhône River basin to the Po River basin. The reasons of these trout 96 translocations would be related with the well-known passion for trout fishing of the Queen Elena of Italy 97 98 (1873 – 1952) (e.g., Siccardi, 1996). Unfortunately, in the above AIIAD document, there is no citation of 99 historical records to sustain this hypothesis. On the contrary, bibliographic records collected in the 100 Geographic dictionary of the Sardinian State (Casalis, 1833, 1852), evidenced a widespread presence of 101 brown trout in the occidental Alps long before Elena's reign (1900-1946). Interesting, as far as it is known, 102 there are no pieces of evidence of trout translocations from France to Italy, taking into account the above historical literature. On the other hand, historical translocation of freshwater fish across the Alps have been 103 well documented, as the case of the historical (16th century) translocation of Salvelinus alpinus from Austria 104 105 to Trentino Alto Adige Region (Italy) (Tiberti & Splendiani, 2019).

106 Here, a comprehensive molecular data set is used to describe wild Mediterranean brown trout population genetic diversity encompassing from the Rhône River basin to the Po River basin to try to reach several 107 objectives. The first aim of the present study is to provide, for the first time, a focus on brown trout 108 mitochondrial DNA (mtDNA) genetic diversity distribution along a putative contact zone between several 109 110 lineages of this species complex. The second aim is to find solid hypotheses to correlate spatial distribution of genetic diversity with the main evolutionary forces that could have played a role in shaping the 111 Mediterranean brown trout lineage distribution observed in the study area, that are: i) effects of the last 112 113 glacial maximum (LGM, that are, ice cover extension, effects of ice flow pattern during deglaciation episodes, localization of glacial refuges, etc), and ii) the role of geomorphologic and hydrogeological 114 characteristics of the mountain relief in the south-western Alps in shaping genetic diversity. Finally, the third 115 aim is to test the hypothesis of non autochthony proposed by AIIAD and as a consequence provide 116 recommendations to preserve, in a rational manner, what remains of the native genetic diversity of brown 117 118 trout in south-western Alps watercourses.

119 MATERIAL AND METHODS

120 SAMPLING DESIGN

121 The S. trutta complex is characterized by a puzzling pattern of geographical forms probably underlying 122 the taxonomic inflation reported in literature with the description of nearly 50 Salmo species (Tougard et al., 2018). The description of the native trout taxa previously identified (e.g., Kottelat & Freyof, 2007) in the 123 study area have been summarised in Table S1. In this study, for practical reasons, the following terms will be 124 used hereafter: (1) "marble trout" for S. marmoratus specimens (generally fixed for MA haplotypes), (2) 125 "native brown trout" for the individuals showing both a fario phenotype and native haplotypes of the 126 127 Mediterranean area (that are, haplotypes belonging to the lineages AD, ME or MA), (3) "Lake Garda Carpione" to indicate S. carpio specimens and (4) "Atlantic brown trout" to indicate S. trutta specimens 128 129 hosting domestic and non-native haplotypes (AT lineage). The sampling efforts were mainly focused along

the south-western Alps, as for example, i) in the upper Durance, that belongs to the Rhône basin, but is located adjacent to the Po River basin along the French side of the south-western Alps, and ii) along the main sub-basins of the Po River basin, as regards the Italian side of the south-western Alps. For comparisons, further samples from other sites of the Rhône River basin and from neighbour minor Mediterranean independent rivers were also included, for example, the rivers Var, Loup, Roya and Sansobbia. The sampling size ranged from 4 to 24 with a mean value of 10 trout per sample. The total number of analysed *S. trutta* was 412 specimens that were collected from 42 sites (Figure 1, Table 1).

137 DNA EXTRACTION

From each fish, a small fin clip was removed and conserved in 95% ethanol until DNA extraction. Due to 138 the union of genetic data obtained from two different laboratories (i.e., from the ISEM, Université de 139 140 Montpellier and from the DiSVA, Università Politecnica delle Marche, hereafter, respectively, Lab.1 and Lab. 2), total genomic DNA was extracted using two methods. The first one (Lab. 1) consisted in a 141 Chelex/proteinase K-based protocol described by Estoup, Largiader, Perrot & Chourrot (1996). A small 142 piece of fin was incubated overnight at 56°C in 195 µl of 5% Chelex 100 Resin (Biorad) solution containing 143 144 50 mM of Tris-HCL (pH 7) and 500 µg/ml of proteinase K. Samples were then incubated at 95°C for 10 min 145 before centrifugation at 3500 g for 5 min. Supernatants were recovered and frozen at -20° C until required 146 for use. In the second method (Lab. 2), total genomic DNA was extracted using an automated DNA extractor 147 (MagCore® Automated Nucleic acid Extractor in combination with the Genomic DNA Tissue Kit 401).

In Lab. 1, the mtDNA control region (CR) was amplified by PCR using the PST (5'-148 CCCAAAGCTAAAATTCTAAAT-3') and FST (5'-GCTTTAGTTAAGCTACGC-3') primers (Cortey & 149 García-Marín, 2002). Each 50 µL reaction included 0.4 µM of each primer (Eurofins MWG Operon), dNTP 150 151 (2 mM each), 2 mM of MgCl2, 10 µL of 5 9 PCR buffer, 1 U of Taq polymerase (GoTaq® Promega), and about 50 ng of genomic DNA. The PCR conditions included initial denaturation (95°C, 5 min), followed by 152 153 30 cycles of strand denaturation (94° C, 1 min), primer annealing (52° C, 1 min), and DNA extension (72° C, 1 min) cycles, and then by a final extension (72° C, 5 min). All PCR amplifications were performed in 154 Eppendorf Mastercycler thermocyclers. The amplified DNA fragments were run on a 0.8% agarose gel to 155 verify the amplification efficiency. The amplified products were purified and sequenced in both directions to 156 confirm the polymorphic sites in an ABIPRISM 3130/xl/sequencer (Applied Biosystems). In Lab. 2, the 157 mtDNA CR was PCR-amplified according to Bernatchez & Danzmann (1993) (primer sequences: LN20, 5' 158 159 ACCACTAGCACCCAAAGCTA; HN20, 5' GTGTTATGCTTTAGTTAAAGC). Screening of mtDNA genetic variability was conducted through Single-Strand Conformation Polymorphism (SSCP) analysis. 160 161 Because shorter fragments are better suited for detection of mutations in SSCP gels (Hayashi, 1991), the CR 162 PCR products of c. 1000 bp were first digested with AluI restriction enzyme and then run on a nondenaturing polyacrylamide gel for 12-h at 5 W in a cool chamber. Finally, the non-digested segment of c. 163 1000 bp was sequenced in a sub-sample of individuals with the same SSCP profile (that is, three-four trout 164 165 per each SSCP morph detected).

166 POPULATION STRUCTURE, DEMOGRAPHIC HISTORY, AND MOLECULAR DATING

167 The mtDNA CR sequences were aligned using Clustal W (Larkin et al., 2007). In order to assign the sequence haplotypes observed in this study to each of the main brown trout mtDNA lineages, several 168 reference S. trutta CR sequences were downloaded from GenBank (belonging to the mtDNA lineages ME, 169 AD, MA and AT) (see Table S2 for more details). The genealogical relationship among haplotypes was 170 171 depicted using a Median-Joining Network (Bandelt, Forster, & Rohl, 1994) constructed using Network 5 (Fluxus Technology Ltd., www.fluxus-engineering.com), considering also gaps and missing nucleotides. The 172 173 ε parameter was set to zero. Historical demography inferences were drawn from three neutrality tests implemented in DnaSP 6 (Rozas et al., 2017): i) Fu's F_S (Fu, 1997), ii) Tajima's D (Tajima, 1989) and iii) R_2 174 175 (Ramos-Onsins & Rozas, 2002), and from mismatch distribution analysis by using Arlequin 3.5 (Excoffier & 176 Lischer, 2010). Briefly, a significantly negative Tajima's D and Fs, and a significantly positive R_2 indicate a 177 scenario of demographic expansion. In the mismatch analysis, a curve displaying the observed distribution of pair-wise differences within each lineage is compared to an expected curve under a model of population 178 179 growth-decline. Generally, a curve with a single peak associated with a low number of pair-wise differences indicates expansion, while a curve with two or multiple peaks indicates stability. Differences between 180 181 observed and expected pair-wise mismatch distribution were evaluated using the sum of squared deviations 182 (SSD) and the raggedness index (r) as implemented in Arlequin 3.5.

183 The estimation of time to the most recent ancestor (TMRCA) of the AD, ME and MA lineages was carried out with a Bayesian coalescent analysis using BEAST 1.10.4 (Suchard et al., 2018) under a HKI + I 184 185 model as inferred by using jModeltest (Posada, 2008). We adopted two fixed values with a normal prior 186 distribution (0.75 - 1%) of divergence rates (e.g., Sanz, 2018 and references therein), and taking into account that we were interested in the estimation of the separation time between the Mediterranean lineages AD, ME 187 188 and MA, we adopted the basic strict clock model, as implemented in BEAST, because, potentially 189 outperforming for trees with shallow roots (Brown & Yang, 2011). The strict molecular clock was used in combination with three coalescent models (constant size, exponential growth, expansion growth). To 190 191 determine the best fitting model of the data (Brandley, Schmitz, & Reeder, 2005), a modified Akaike 192 Information Criterion (AICM) as provided in TRACER 1.6 (Rambaut et al., 2018) was used. The models 193 were run five times for fifty million generation with a 10% burn-in stage. Markov chain convergence was 194 checked visually by the inspection of the traces, while the run stability was measured using the effective 195 sample size (ESS > 200 for all parameters) using Tracer. Results of the independent convergent runs were 196 combined with LogCombiner v 1.10.4 (auxiliary program implemented in the BEAST package) to estimate TMRCA and 95% highest probability density intervals (HPD). A consensus tree was then generated using 197 TreeAnnotator 1.10.4 (auxiliary program implemented in the BEAST package) with the following options: 198 maximum clade credibility and mean node heights. 199

Hierarchical analysis of molecular variance (AMOVA) was used to test how the effects of the last glacial
 events, occurred in the study area, could explain the current spatial genetic distribution. Groupings included:

i) Rhône samples *vs* Po samples and ii) Rhône samples *vs* Po and Durance samples. This latter grouping was
set-up to test the hypothesis of a recent post-glacial origin of brown trout samples from upper Durance
related to slope failure phenomena due to last deglaciation events. The above tests were carried out by using
Arlequin 3.5, using conventional F-statistics and testing the statistical significance of the tests with 5,000
permutations.

207 To depict the biogeographic scenarios underlying the observed haplotype spatial distribution, the mtDNA 208 lineage distribution observed in each population was both analysed by mapping pie charts geographically 209 and using Kernel density (KD) analysis along an elevation, longitudinal and latitudinal gradient. The KD analysis was conducted partitioning the samples as i) samples from the Rhône River basin and ii) samples 210 211 from the Po River basin. For these tests, brown trout samples were grouped in the following categories: i) 212 samples fixed for AD haplotypes, ii) samples fixed for ME haplotypes, iii) samples fixed for MA haplotypes 213 and mixed samples, iv) samples sharing AD and ME haplotypes, v) AD-MA, vi) ME-MA and vi) AD-ME-MA. The rational of the above partitioning was to verify if haplotype distribution can match with plausible 214 215 scenarios of extinction and recolonization events connected with the environmental changes occurred during the last glacial maximum in the south-western Alps. The KD analysis was carried out by using the density 216 function in R software (R Development Core Team, 2017). 217

218 RESULTS

219 HAPLOTYPE CLASSIFICATION

220 Before starting with sequence analyses, the PolyT region of the CR was considered of a constant length of 221 14bp in all sequences. In fact, such region is likely to be unstable and thus characterized by a high mutation 222 rate, showing frequent 14-T variants. In this circumstance, sequence stretch length identity, could be the mere consequence of homoplasy and would not represent a real phylogenetic signal. On the whole, an 223 224 alignment of 981 bp was obtained, from which 21 haplotypes emerged (Table 2). The observed haplotypes 225 belonged to four main mtDNA lineages: AD, ME, MA and AT (sensu Bernatchez et al., 1992 or Bernatchez, 2001). The AD lineage was represented by three haplotypes: ADporh-1 and ADrh-1, observed for the first 226 227 time in this study (GenBank accession numbers, respectively, MK948034 and MK948035) and ADcs-1 228 already described in literature (Cortey, Pla & García-Marín, 2004). As far for the ME lineage, seven 229 haplotypes were observed, two of which already detected in other studies, that are: MEcs-1 and MEcs-15 (e.g., Cortey et al., 2004) and the other five were detected for the first time in this study and named as 230 follow: MEcs-28 to MEcs-32 (from MK948029 to MK948033). The MA lineage displayed a new haplotype 231 232 named MAsl-1 (MK948036), and three mtDNA variants already detected in previous studies, namely Ma2a, Ma2b and Ma2c (Meraner, Baric, Pelster, & Dalla Via, 2007; Meraner, Gratton, Baraldi & Gandolfi, 2013). 233 234 Finally, the Atlantic lineage was represented by six haplotypes, five of them were already detected in 235 Mediterranean rivers and classified as haplotypes of hatchery origin, that are: haplotype 1 to haplotype 4 236 (Cortey & García-Marín, 2002) and the haplotype Atle, (Meraner et al., 2007), while one haplotype was

described for the first time in this study (haplotype 3b, MK948037). This latter haplotype should be also
considered of hatchery origin due to high similarity with the domestic variant haplotype 3 (Table 2, Figure
2).

240 HAPLOTYPE SPATIAL DISTRIBUTION

The ME lineage dominated in the Rhône River drainage and in the other minor French rivers included in this study. The most common ME haplotype was MEcs-1, the rest of the ME haplotypes (MEcs-28 – 32) was endemic of French Mediterranean rivers. On the other hand, within the Po River drainage and in the Sansobbia River, the ME lineage was represented by the haplotype MEcs-1.

245 The AD lineage was quite common in the Po River drainage, however, the ancestral haplotype of this lineage (ADcs-1, see Figure 2) was observed only in a sole trout (in sample GIU, Po River basin). However, 246 in other studies within the Po River drainage, the haplotype ADcs-1 was detected in both 19th century trout 247 (Splendiani et al 2017), as well as in recent samples (Gratton et al., 2014; Stefani, Anzani, & Marieni, 2019). 248 Toward the west part of the Po River drainage, the haplotype ADporh-1 was newly detected (one Mutational 249 250 Step from ADcs-1). This latter haplotype presented a spatial distribution confined to the upper reaches of 251 both upper Durance River (Rhône basin) and Po River along a transect of the south-western Alps extended 252 from the Cottian to the Maritime Alps (Figure 1 and Table 2, see also next paragraph). Finally, the haplotype ADrh-1 consistently with its position in the Median-Joining Network (one MS from the haplotype ADporh-253 1) was found only in the western part of the Rhône River drainage, in the Petit Buëch stream. 254

The MA haplotypes were found only within the Po River drainages. The most common haplotype detected for this lineage was Ma2b. This mtDNA variant occupied a central position within the MA lineage (Figure 2). In line with its haplotype network position, this haplotype was observed elsewhere within the Po River drainages in both modern (Meraner *et al.*, 2007, 2013) and museum specimens (from 19th century, *e.g.*, Splendiani *et al.*, 2017).

Finally, as expected, the AT haplotypes formed a separate cluster in the Median-Joining Network (Figure
2). These non-native haplotypes for the Mediterranean area showed an evident greater abundance within the
Italian samples (Table 2, Figure 1). Here, a mean value of AT haplotypes of 38% and a maximum value of
70% (in the locality MAIb) were observed. In four localities (TRO, RIP, FER and GES), the AT haplotypes
were not observed. In the French samples, the AT haplotypes were observed only in two localities out of 21,
GLU (60%) and ROY (20%).

266 MISMATCH ANALYSIS AND DIVERGENCE TIME ESTIMATES

Mismatch distribution analysis indicated (see Table S3 and Figure S1) a scenario consistent with a model of demographic expansion (Excoffier, 2004) for the brown trout mtDNA lineages ME and AD and a stable demographic trend for the lineage MA. 270 The AICM suggested that a strict clock under a constant size coalescent model best-fits our data. The 271 TMRCA estimations placed the origin of the AD lineage from 278,000 (95% HPD 170,000 - 391,000) to 272 212,000 (95% HPD 129,000 - 298,000) years ago by adopting, respectively, a substitution rate of 0.75 and 273 1%, the origin of the ME lineage from 267,000 (95% HPD 166,000-372,000) to 191,000 (95% HPD 274 122,000-265,000) years ago, and the origin of the MA lineage from 122,000 (95% HPD 172,000-205,000) to 275 117,000 (95% HPD 51,000-193,000) years ago (see Table S4). MA lineage appears so as the youngest one. 276 Finally, the origin of the AD branch composed by the haplotypes ADporh-1 and ADrh-1 was placed around 151,000 (95% HPD 11,000-99,000) and 120,000 (95% HPD 16,000-86,000) years. 277

278 AMOVA

With both two grouping options (Rhône samples *vs* Po Samples and Rhône samples *vs* Durance and Po samples), the AMOVA analyses showed that most of the genetic variation was explained at the within population level (53.26 and 51.62%, respectively) and among populations within group level (33.84 and 26.84%, respectively). However, the AMOVA analyses showed also that grouping the Durance samples within the Po River group explained much more genetic variation (21.54 %) than grouping the Durance within its main river basin (*i.e.*, the Rhône River basin) (14. 92 %). In both cases, the statistical significance of the source of variation represented among groups was highly significant (P = 0.0000).

286 KERNEL DENSITY (KD) BROWN TROUT LINEAGE DISTRIBUTION

Within the Rhône River drainage, the highest density of samples characterized by the sole presence of 287 ME haplotypes was found between 4.5 - 5.5 E longitudes (Figure 3A). Unfortunately, only one sample was 288 289 characterized by the sole presence of AD haplotypes (BUË), therefore, KD analysis was not applicable in 290 this case. On the contrary, the rest of the Rhône samples (i.e., samples from the Durance sub-basin) were composed by a mixture of AD and ME haplotypes. In this study, a total of 91 trouts originated from the 291 292 Rhône River basin in France, among them, 31 specimens showed the AD haplotypes. All these latter trouts came from the Durance sub-basin (samples BUË, BIA, CLA, UBA and GUI, see Table 2). The last two 293 294 rivers (UBA and GUI) are flowing in France directly from the France-Italy boundary. This kind of 295 populations peaked around 7.0 E longitude, corresponding with the upper part of the Durance River (Figure 296 3A). When KD was carried out to relate brown trout lineages distribution with elevation, a similar net 297 separation between different categories of samples was observed. For example, in the Rhône River, populations fixed for ME haplotypes were most abundant around 0 - 500 m (Figure 3B), whereas admixed 298 populations (AD – ME) showed higher values of probability between 1500 – 2000 m. Within the Po River 299 300 drainage, the different categories of populations defined based on mtDNA lineage composition, appeared clearly stratified along an altitudinal cline. Pure ME populations were detected only in one case (BAR, 301 Tanaro River, Ligurian Apennine). Admixed AD - ME populations were, however, the most common and 302 303 reached a density peak around 1600 – 2000 m. Pure AD populations peaked slightly lower, around 1400 m 304 (Figure 3C). Around this latter quote, peaked both pure brown trout samples fixed for MA haplotypes and

admixed MA – AD brown trout samples. Further downstream (c. 450 m), pure marble trout samples (MA) were abundant (data from Giuffra *et al.*, 1994). Finally, a pattern of brown trout mtDNA lineage density distribution along the south-western Alps was also evident along a latitudinal gradient (Figure 3D). For example, pure AD samples appeared most abundant around 44.0 - 44.5 N, roughly corresponding with the Maritime Alps (Italian side), while admixed populations (AD – ME) peaked around 45.0 N (*i.e.*, Cottian Alps).

311 DISCUSSION

In the following chapters of this study, a comprehensive phylogeographic analysis of brown trout 312 populations inhabiting adjacent tributaries of the Rhône and Po River basins is proposed for the first time. 313 Based on the biogeographic scenario reconstructed, the allochthonous hypothesis proposed by the Italian 314 Association of Freshwater Fish Ichthyologists (Zanetti et al., 2013) about Mediterranean brown trout 315 populations inhabiting the south-western Po tributaries was evaluated. Substantially, the shared haplotype 316 317 diversity (AD and ME haplotypes) observed along the two sides of the south-western Alps and, at the same time, the presence of an important ice cover along the western part of the Rhône basin (i.e., the Durance 318 319 glacier) during the glacial phases, incompatible with the presence of freshwater fish, suggest that native 320 brown trout survived the adverse phases of the upper Pleistocene just in the tributaries of the south-western 321 Po basin. Probably, the erosional events related with the deglaciation phenomena permitted the opening of a 322 biological corridor for brown trout from the Po basin toward the Rhône basin.

323 THE ORIGIN OF THE S. TRUTTA ME LINEAGE IN THE STUDY AREA

According to previous studies (Cortey *et al.*, 2004; Vera *et al.*, 2019), the Iberian Peninsula would represent the ideal candidate as the centre of the origin of the ME lineage (*e.g.*, Sanz, 2018). However, the ME haplotype diversity detected in the present study partially contrast with the above hypothesis. Within the Rhône River basin and neighbour rivers, five new ME haplotypes were detected. This fact appears in accordance with a western Mediterranean origin of the ME lineage as previously proposed (*e.g.*, Bernatchez, 2001; Cortey *et al.*, 2004; Oliver, 2014) but also suggests that the Rhône basin area acted as an important evolutionary centre for ME genetic diversity.

Although, the use of the sole mitochondrial control region could be taken into account with caution for inferring isolation time accurately (Schenekar, Lerceteau-Köhler, & Weiss, 2014), an attempt was, however, tried in this study. In this sense, it is interesting to indicate that the pre-defined divergence rates of 0.75 - 1% adopted in this study designed a time since expansion of the major Mediterranean lineages (ME, AD and MA) in accordance with their altitudinal distribution observed in the study area (see below).

The TMRCA analysis suggest a main expansion of the ME lineage around 191,000-267,000 years ago, roughly corresponding with the last (III) Mindel-Riss Interglacial, and resulting slightly more ancient respect previous estimations (Cortey *et al.*, 2004). Pleistocene Interglacial warming periods have been regarded as

339 phases of isolation in small headwaters for Mediterranean brown trout populations, thus promoting genetic 340 signatures within lineages (e.g., Sanz, 2018 and references therein). On the other hand, during glaciations, colder climate conditions may have triggered sea-ward migratory tactics in Mediterranean brown trout 341 342 populations, as highlighted by several paleontological evidences (Muñoz & Casadevall, 1997; Splendiani et 343 al., 2016b; Splendiani et al., 2020). Therefore, thanks to a seaward migratory route, the expansion of brown 344 trout in the Mediterranean area was possible. The colonisation of northern Corsica Island by the ME lineage 345 during the last glaciation is an example of this expansion (Gauthier & Berrebi, 2007). The spatial distribution 346 of the MEcs-1 haplotype suggests a potential eastward dispersion, from the Rhône River outlet to the Var 347 River (SAL, Gulf of Lion) where this haplotype was found and more eastern to the Sansobbia River (SAN, Ligurian Sea) (Figure 1, Table 2). Then, when the ME lineage reached the upper part of the Ligurian rivers, 348 349 the colonization of the Po River basin was likely possible thanks to river capture events occurred along the 350 Ligurian Apennine chain. This scenario could explain the finding of the MEcs-1 haplotype in the Rio 351 Baracca (BAR, Po River basin, see also Figure 1, Table 2). Interestingly, the role played by the 352 hydrographical captures of the upstream portions of Mediterranean rivers was also proposed in the literature 353 to explain, for example, the exchange of Duero haplotypes between rivers flowing along opposite slope of 354 the Cantabrian mountains (Iberian Peninsula) (Vera et al., 2015), as well as to explain the native occurrence 355 of the Danubian haplotypes within marble trout populations of the Sôca River (Berrebi, Jesenšek, & Crivelli, 356 2017). Concerning the study area, the phylogeographic history of the Italian vairone (*Telestes muticellus*) in 357 populations of west Liguria (Marchetto et al., 2010) could also be explained invoking the presence of 358 biological corridors opened by ancient Mediterranean river captures.

Once reached the Po River basin, it can be reasonably expected that the ME lineage tried to colonize available salmonid habitats. However, when the climate conditions went colder and the Alpine ice cover expanded, this lineage survived only in refuge areas as, for example, in south-western Alps. Milder conditions in this part of the Alpine chain and the absence of other brown trout lineages may have permitted the colonization of the upper reaches of the western Alps by the MEcs-1 haplotype (see the Kernel density analysis in the Results section and Figure 3).

365 THE ORIGIN OF THE S. TRUTTA AD LINEAGE IN THE STUDY AREA

The Adriatic-Balkan part of the Mediterranean basin is considered the centre of the origin of the AD 366 367 lineage (Sanz, 2018). The main expansion of the AD lineage seems to take place around 267,000 - 212,000 year ago and therefore nearly simultaneous with the last expansion proposed above for the ME lineage. 368 Although, as already stressed, divergence estimations should be interpreted with caution, mainly with regard 369 to the absolute date of expansion values, more reliable, on the contrary, appears the simultaneous time of 370 371 expansion of the AD and ME lineages observed in this study. The simultaneous expansion of these two 372 lineages fits well with both their similar peri-Mediterranean spatial distribution and with their phylogenetic 373 complexity (e.g., Sanz, 2018).

374 In north Italy, the central haplotype of the AD lineage (ADcs-1) was observed in two museum specimens (dating back to the end of 19th century) of Lakes Garda and Maggiore and in a modern sample of the Adige 375 Adriatic River (Meraner et al., 2013). According to Splendiani et al. (2016a, 2017), the spatial distribution of 376 377 the Mediterranean trout genetic diversity in north Italy represents a sort of "map" of the potential Alpine 378 peripheral refuges where brown trout survived during the extreme glacial phases. In addition, the role played 379 by the area of the Lake Garda as an important glacial refuge for the genus Salmo is also evidenced by the 380 detection of two endemic AD and MA haplotypes in Lake Garda Carpione, an endemic trout of this major 381 Italian lake (Gratton et al., 2014) (Figure 2). Further, the finding of a new AD haplotypes (ADporh-1), 382 endemic of the south-western Alps, suggests that also this area could have acted as both an important glacial 383 and interglacial refuge for brown trout. In the south-western Alps' part of the Po River basin, the haplotype 384 ADporh-1 was the sole AD haplotype observed. This haplotype was found as fixed in samples collected 385 around 1000 - 1500 m (Figure 3), that is slightly lower than the quote where AD-ME mixed populations peaked. Thus, the observed spatial distribution suggests that the ME lineage colonized first the headwaters of 386 387 the south-western Po River basin, whereas the AD lineage tried to do the same later. Based on molecular 388 dating analyses, both ME and AD lineages showed a similar divergence time and thus it is hard to explain 389 their different altitudinal distribution. However, the proximity of the south-western Alps to an important 390 centre of origin of the lineage ME, as can be considered the Rhône River basin, could explain why the ME 391 lineage reached the highest sites of the south-western part of the Po River basin first.

392 The peculiar AD haplotypes detected across the south-western Alps (*i.e.*, ADporh-1 and ADrh-1 393 haplotypes) probably split from the ADcs-1 ancestor when warmer climate condition promoted phases of 394 isolation. The estimated origin for this AD branch of c. 151,000 - 120,000 years ago, corresponding approximatively with the Riss-Würm Interglacial. During the warmer phases, brown trout population may 395 have survived in high altitude habitats of the south-western part of the Po River basin. This region would 396 397 also have been used as refuge during the colder phases (*i.e.*, the Younger Dryas stadial, c. 12,800 and 11,600 years BP) when the rest of the high-altitude Alpine streams was covered by a massive ice cover. Later, at the 398 399 beginning of the Holocene, the extreme erosional events produced by massive episodes of deglaciation 400 promoted the colonization of the upper Durance basin from the adjacent high altitude brown trout 401 populations survived in the south-western Po streams by river captures (see the next paragraph for more 402 information: The role of the south-western Alps as an asymmetrical biological corridor for brown trout 403 lineages).

The Italian side of the Maritime Alps represented also an isolated refuge. Interestingly, in this region, both the AD-Tyrrh-1, very common elsewhere within the Tyrrhenian watercourses (*e.g.*, Berrebi *et al.*, 2019) and ADcs-11, very common around the Adriatic Sea rivers (Sušnik *et al.*, 2007; Snoj *et al.*, 2009, 2010), haplotypes were not found. Based on the network haplotype topology (Figure 2), we can hypothesized that ADcs-1 colonized first the Po River, and then, during phases of geographic isolation within refugia, new haplotypes, as ADporh-1 within the Maritime Alps refuge and the *S. carpio* AD haplotypes (ScarAD-1 and

410 ScarAD-2), within the Lake Garda refuge, could split. According to Sanz (2018), the AD lineage was 411 characterized by multiple waves of expansions. Successive expansion opportunities were probably used by 412 individuals carrying haplotypes ADcs-11 (around the Adriatic Sea) and AD-Tyrrh-1 around the Tyrrhenian 413 Sea, these latter ones, however, were unable to colonize the Po River as it was already occupied by both 414 brown trout (showing the ADcs-1, ADporh-1 and MEcs-1 haplotypes) and marble trout (showing the Ma2a, 415 Ma2b, Ma2c and MAsl-1 haplotypes).

- 416

417 THE ORIGIN OF THE S. TRUTTA MA LINEAGE OBSERVED IN THE STUDY AREA

418 Within the Po River basin, the MA lineage showed an evolutionary pathway like that observed in the case of the AD lineage. For example, in the Lake Garda Carpione, Gratton et al. (2014) exhibited two endemic 419 420 MA haplotypes (named here ScarMA1 and ScarMA2, see also Figure 2). Further west, in brown trout 421 samples, a new MA haplotype (MAsl-1) was found, although in a sole specimen (Table 2). In this study, the 422 most diffuse MA haplotype detected in the south-western part of the Po River was Ma2b, also common 423 elsewhere in the Po River basin both in native brown trout and Lake Garda Carpione specimens (e.g., 424 Meraner et al., 2007; 2013). However, the most relevant result was represented by the spatial distribution of the MA lineage as detailed below. In this study, the native brown trout samples from the south-western 425 426 tributaries of the Po River basin, characterized by the sole presence of MA haplotypes, showed an altitudinal 427 range of distribution intermediate between marble trout (c. 450 m) and native brown trout populations characterized by a mix of AD and ME haplotypes (c. 1600 - 2000 m). Therefore, based on the altitudinal 428 429 distribution of the MA lineage it could be hypothesized that this lineage tried to reach the upstream thermal 430 refuges for last. This hypothesis seems to accord well with the younger origin of MA lineage respect to ME 431 and AD lineages emerged in this study and proposed also by Oliver (2014). In addition, the altitudinal 432 distribution of the MA lineage appear congruent with the stable or declining demographic scenario emerged 433 by mismatch analysis. The reduced habitat availability could have contrasted the demographic expansion of this lineage (e.g., Lavery et al., 1996; Bernatchez, 2001). Probably, this mtDNA lineage was fixed in marble 434 trout that inhabited the lower part of the Po River basin, then, when climate become warmer, this salmonid 435 tried to reach colder habitats at higher elevations. A similar palaeohistorical scenario was previously 436 proposed (Berrebi et al., 2000) to explain the spatial distribution of both brown trout and marble trout within 437 the Sôca River basin (Slovenia). The detection, in the present study, of only native brown trout phenotypes in 438 439 samples characterized by the sole presence of MA haplotypes suggests that the contact between brown trout and marble trout occurred within an ecological contact zone (i.e., ecotonal zone) where the parental 440 Mediterranean phenotype outcompeted (e.g., Arnold, 1997). Alternatively, the evolution of habitat 441 442 preference of the MA lineage for lower river sections could explain its altitudinal distribution observed in the 443 south-western Alps. However, elsewhere, as for example in the Adige River basin, the MA lineage was able to reach, accessible and formerly glaciated, high altitude sites (around 1000 - 1600 m) (Meraner *et al.*, 2007, 444 445 2010; Splendiani et al., 2016a).

446 THE ROLE OF THE SOUTH-WESTERN ALPS AS AN ASYMMETRICAL BIOLOGICAL447 CORRIDOR FOR BROWN TROUT LINEAGES

448 The comparison between the brown trout mtDNA genetic diversity observed along the two sides (east and 449 west) of the south-western Alps highlights the lack of a substantial genetic differentiation between the samples collected in the upper Durance River (Rhône River basin) and the upper reaches of the Po River. 450 451 Most samples from upper Durance and Po basins were composed by a mixture of the ADporh-1 and MEcs-1 452 haplotypes (Table 2). The similarity in haplotype composition between the native brown trout populations of 453 the two-opposite sides of the south-western Alps was also supported by the hierarchical analysis of molecular variance (AMOVA). In fact, when the samples of the upper Durance were grouped together with 454 455 the Po River group, the level of genetic variation explained between groups of populations increased from 14 456 to 21%, suggesting that the vicariant events between upper Durance and Po brown trout populations occurred 457 recently. In this regard, it is important to note that all the Durance collection sites extended in elevation from 1117 to 2077 m, that is, an altitudinal range occupied by the ice cover during the last glacial maximum (e.g., 458 459 Figure 1), which implies the arrival of trouts after this period. Interestingly, the private ADrh-1 haplotype, a 460 mtDNA variant distant of one MS only from the haplotype ADporh-1, was detected in the Petit Buëch stream, sited at the margin of the Durance glacier. Here, the milder climate condition of this part of Durance 461 basin could have allowed the maintenance of trout populations and represented a refuge for the haplotype 462 463 ADporh-1 and the centre of origin for the haplotype ADrh-1.

464 A possible explanation for the high genetic affinity observed between brown trout samples from the upper 465 Durance and Po River basins could be deducted when taking into account the effects, in terms of fish 466 exchange along the Alpine barrier, provided by the last deglaciation events occurred in the south-western 467 Alps. Likely, the first important factor that has drawn the present geographic structure of trout populations is 468 the spatial distribution of the ice cover during the LGM. During this period, the Durance paleo-glacier was 469 one of the most important Alpine glacier (Cossart, Braucher, Fort, Bourlès, & Carcaillet, 2008) (Figure 1). 470 On the other hand, along the Italian side of the south-western Alps, the ice cover appeared less extended (Hughes, Woodward, & Gibbard, 2006; Szövényi et al., 2009) (Figure 1). An explanation for the formation 471 472 of an unidirectional corridor between the two side of the south-western Alps could be related with the 473 formation of small ephemeral lakes and/or the swelling of connecting streams at the retreating edge of a glacier that may allow watershed crossing and drainage switching by freshwater fish (Waters et al., 2001). 474 475 This scenario was proposed to explain the colonization of the Lake Geneva (Rhône River basin) by bullhead 476 Cottus gobio migrants from the Rhine River basin during the last glacial retreat (Vonlanthen et al., 2007). 477 Further, also the spatial distribution of the genetic diversity of *Galaxias platei* in Patagonia along the Andes 478 also represents a similar example showing the role of glacial retreat events in promoting fish migrations 479 across watersheds (Zemlak et al., 2008; Habit et al., 2010). Therefore, a scenario can be suggested where, 480 first, during the colder phases, brown trout survived in the ice-free tributaries of the south-western Alps (*i.e.*, 481 Maritime and Cottian Alps), and second, during the erosional events related to the ice melting (early

Holocene), an unidirectional corridor opened and permitted the colonization of the empty habitats of the adjacent upper Durance river catchment from the Po watershed. Interestingly, along the south-western Alps, the haplotype MEcs-1 was recently detected (Splendiani *et al.*, 2017) in a museum specimen collected in 1876 in the Lake Mont Cenis (1974 m a.s.l.) (Figure 1), a former small Alpine Lake (since 1921 the lake was artificialized by the construction of a weir) of post-glacial origin belonging to the Dora Riparia River and located near the divide between the Po and Rhone catchments.

Finally, the above scenario could be also proposed to explain the spatial pattern of genetic diversity that has been observed in other freshwater organisms inhabiting the two sides of the south-western Alps. For example, as in the case of the high genetic similarity observed between adjacent populations of *Cottus gobio* (Šlechtová *et al.*, 2004), or similarly, the lack of genetic differentiation observed between adjacent populations of *Austrapotamobius pallipes* (Stefani *et al.*, 2011).

493 TAXONOMIC IMPLICATIONS

494 As suggested above, in material and methods section, the main aim of the present study was not related 495 with the attempt to solve the well-known problem of the S. trutta complex systematic (Splendiani et al., 496 2019b). However, the phylogeographic scenario of Mediterranean brown trout that emerged here represents 497 an opportunity to partially face with the above taxonomic issues. The trout from the two sides of the south-498 western Alps are traditionally classified into three-four nominal species that we have used here for practical 499 reasons: S. rhodanensis (Rhône River basin) a contested species (Berrebi et al., in prep.), S. cettii (a non-500 valid name when used to indicate trout from the Tyrrhenian and Ligurian Sea draining rivers) and S. 501 farioides or S. cenerinus (depending on the authors, Adriatic draining rivers) (Figure 1). At mtDNA level, 502 none of the above nominal species showed a genetic distinctiveness able to justify the recognition of 503 different species. For example, the two Ligurian samples (SAN, putative S. cettii and BAR, putative S. 504 farioides - S. cenerinus) collected from the two sides of the Apennine chain, were both fixed for the haplotype MEcs-1, that is a haplotype quite widespread in the study area, as well in the rest of the 505 Mediterranean rivers. More north, along the contact zone of the Rhône - Po River basins, the samples of the 506 Durance River (putative S. rhodanensis) showed a haplotype composition more similar to that observed 507 along the Italian side (putative S. farioides - S. cenerinus), respect to that shown by rest of the Rhône 508 samples as highlighted by the AMOVA analyses. Obviously, more sound conclusions should be drawn by 509 analysing also nuclear and morphological markers (see Ninua, Tarkhnishvili, & Gvazava, 2018 for similar 510 arumentations). These preliminary results however refute the traditional taxonomic position adopted until 511 now for the Mediterranean trout of the study area. 512

513 CONCLUSIONS

The main findings of this study highlight that brown trout should be considered native in the southwestern tributaries of the Po River basin. In this area, native brown trout survived the extreme climate phases of Pleistocene. In this respect, the biological value of the south-western Alps for the conservation of the last 15 wild native Mediterranean trout population should be considered of primary importance. As a consequence, the non-native statement and the non-intervention approach proposed by the Italian Association of Freshwater Fish Ichthyologists (Zanetti *et al.*, 2013), based on a human conjectural man-mediated origin of Italian slope trout from the Rhône River basin should be rejected. In addition, the weakness of the allochthonous hypothesis is also sustained by the lack of historical records describing the occurrence of such practices in the study area (*e.g.*, Splendiani *et al.*, 2019a).

In conclusion, caution must be exercised when planning conservation actions. For example, elsewhere, in 523 the Italian Alpine region, the massive introduction of domestic Mediterranean brown trout of Apennine 524 525 origin (AD, ME and MA haplotypes) started in the last five-ten years. In most cases, the outcomes of pivotal 526 genetic screening on local brown trout populations involved in these projects were not published, or even 527 never done. This probably occurred (and still occurs) in Italy because local administrations have transferred to sport fishing associations the fully management of these practices. In these circumstances, the rationale of 528 these putative conservation actions has not been evaluable by the scientific community. Paradoxically, 529 conservation plans can even represent a further threat for the protection of wild native trout. As far as we 530 531 know, Mediterranean trout hatchery managers in Italy have not published (even considering grey bibliography) the genetic description of their stocks. Recently, the genetic analysis of one of these putative 532 domestic Mediterranean stocks actually turned out as a mix of Mediterranean and Atlantic brown trout 533 (Splendiani et al., 2019b). The irrational planning of massive stocking activities, even if carried out with 534 535 native brown trout, can introduce further risks related to the potential deleterious genetic effects of supplementation programs (Fernández-Cebrián et al., 2014). These latter ones can result in the breakdown of 536 the delicate equilibrium persisting in the incipient parapatric speciation process subsisting between native 537 brown trout and marble trout, as for example in south-western Alps (Giuffra et al., 1994), and can affect the 538 adaptive genetic architecture of native genomes (Caputo, Giovannotti, Nisi Cerioni, Splendiani, & Olmo, 539 540 2009; Schenekar & Weiss, 2017). The situation is far better in the French side of the investigated area. In France, more and more administrative organizations, under the supervision of Ecology Ministry, adopted 541 542 "patrimonial" management during the last twenty years. For this, a large part of the trout populations have 543 been analysed and published in France (https://data.oreme.org/trout/home) with nuclear and mitochondrial 544 markers driving conservation and stocking.

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739 Figure captions740

741 Figure 1

- Map showing the sampling locations of brown trout throughout the Rhône and the western part of the Po River basins. Color pie charts indicate the mtDNA haplotype frequency distribution of the main brown trout lineages. Numbers in bold within pie charts represent the Atlantic haplotype frequency (%). The distribution of the Alpine ice cover during the last glacial maximum is represented by the light blue area. Box at the top right shows the range of the nominal Mediterranean trout species as reported in Kottelat & Freyof (2007).
- [†] The geographic range *of Salmo cettii* according to Kottellat & Freyhof (2007), # the revised geographic
 range of *S. cettii* according to Splendiani *et al.* (2019b and references therein). Locations code colored
 differentially based on the river basin of origin.
- 750 751 Figure 2
- Median-Joining Network showing the phylogenetic relationships subsisting between the 18 native brown trout haplotypes detected in this study (colored circles) and the brown trout haplotypes observed in previously published studies (grey circles, see also Table 1; the position of the haplotypes ADcs-11 and AD-Tyrrh-1 was reported because commented in the text). As regards the haplotypes observed in the study area, the size of each circle is proportional with the haplotype absolute frequency. Locations code colored differentially based on the river basin of origin.
- 759 Figure 3

Plot showing the probability density function (density) obtained comparing elevation, longitude and latitude with the mtDNA genetic composition of brown trout populations from the Rhône River basin (A and B) and from the Po River basin (C and D). The mtDNA lineage composition was represented by the following colored scheme: population characterized by the sole presence of ME haplotypes (Pure ME), in green; pure AD populations, in yellow; admixed AD-ME populations in orange; admixed AD-MA populations, in light blue and marble trout samples (from Giuffra *et al.*, 1994) in blue. To avoid confusion, other minor admixed population were not showed.

767



770 Fig. 2





Fig. 3

Table 1. Collection site information. Sampling code; Nominal taxon: *Salmo rhodanensis, S.rod; Salmo cenerinus, S.cen; Salmo fariodes, S. far.; Salmo cettii, S.cet.;* ?, no nominal proposed in literature; Sea drainage, Gulf of Lion, G; Ligurian, L, Adriatic, A; hierarchical description of the river network investigated (main-basin, sub-basin, stream); Sample size, No; Laboratories (Lab.) involved in genetic analyses, F = ISEM, Université de Montpellier (Lab.1) (Reynaud, Tougard & Berrebi 2011), and I = DiSVA,

	Tawar	Sea	Main-							Elevatio
Code	Taxon	drainage	basin	Sub-basin	Stream	No	Lab.	Lat.	Lon.	n (m)
MER	S. rod	G	Rhône	Ain	Merlue	5	F	46.51	5.64	433
MOU	S. rod	G	Rhône	Saône	Mouge	5	F	46.40	4.87	176
ALB	S. rod	G	Rhône	Ain	Albarine	5	F	45.94	5.38	258
ORC	S.cen/S.far	А	Ро	Orco		17	I.	45.43	7.42	700
VER	S.cen/S.far	А	Ро	Stura di Lanzo	Rio	18	I	45.38	7.28	1220
TET	S. rod	G	Rhône	Cance	Riotet	5	F	45.32	4.56	668
OVA	S.cen/S.far	А	Ро	Stura di Lanzo	Rio	20	I	45.24	7.27	1480
ARN	S.cen/S.far	А	Ро	Stura di Lanzo	Rio Arnas	10	I.	45.24	7.20	1370
VIU	S.cen/S.far	А	Ро	Stura di Lanzo	Stura di Viù	9	I	45.23	7.28	1540
RIP	S.cen/S.far	А	Ро	Dora Riparia	Ripa	24	I.	45.00	6.81	1900
PLA	S. rod	G	Rhône	Durance	Clarée	5	F	45.00	6.66	1484
TRO	S.cen/S.far	А	Ро	Pellice	Chisone	20	I.	44.95	6.95	1835
GER	S.cen/S.far	А	Ро	Pellice	Germanasca	9	I.	44.94	7.15	680
GERb	S.cen/S.far	А	Ро	Pellice	Germanasca	16	I.	44.92	7.27	750
CER	S. rod	G	Rhône	Durance	Cerveyrette	6	I.	44.87	6.78	2077
GLU	S. rod	G	Rhône	Eyrieux	Gluyère	5	F	44.81	4.48	568
GUIb	S. rod	G	Rhône	Durance	Guil	20	I.	44.77	6.97	1779
GHI	S.cen/S.far	А	Ро	Pellice	Ghiacciard	10	I.	44.76	7.09	1440
BIA	S. rod	G	Rhône	Durance	Biaysse	5	F	44.75	6.53	1200
GUI	S. rod	G	Rhône	Durance	Guil	5	F	44.73	6.84	1693
DRÔ	S. rod	G	Rhône	Drôme	Drôme	5	F	44.70	5.13	229
GIU	S.cen/S.far	А	Ро		Rio Giulian	10	I.	44.67	7.19	1120
ARD	S. rod	G	Rhône	Ardèche	Thines	5	F	44.64	4.39	217
BUË	S. rod	G	Rhône	Durance	Petit Buëch	5	F	44.55	5.88	1117
MAla	S.cen/S.far	А	Ро	Maira	Bedale di	10	I.	44.51	7.18	950
BAR	S.cen/S.far	А	Ро	Tanaro	Baracca	18	I.	44.50	8.65	570
MAIb	S.cen/S.far	А	Ро	Maira	Bedale	10	I	44.47	7.15	1180
GLE	S. rod	G	Rhône	Durance	Gleizolles	5	F	44.47	6.77	1319
UBA	S. rod	G	Rhône	Durance	Ubayette	15	I.	44.44	6.85	1953
SAN	S. cet	L	Sansobbi			20	I	44.43	8.50	660
FER	S.cen/S.far	А	Ро	Tanaro	Rio Ferriere	8	I	44.37	6.98	1480
FRE	S.cen/S.far	А	Ро	Tanaro	Rio Freddo	8	I.	44.24	7.17	1550
RAS	S.cen/S.far	А	Ро	Tanaro	Rio	14	I.	44.22	7.82	1100
OUV	S. rod	G	Rhône	Ouvèze	Ouvèze	5	F	44.22	5.11	222
SER	S.cen/S.far	А	Ро	Tanaro	Rio	13	I	44.21	7.67	1280
GES	S.cen/S.far	А	Ро	Tanaro	Gesso	4	I.	44.20	7.27	1450
BOU	S.cen/S.far	А	Ро	Tanaro	Bousset	13	I.	44.20	7.45	1170
SAL	?	G	Var	Tinée	Tinée	5	F	44.17	6.94	1930
FON	?	G	Roya	Roya	Roya	5	F	44.00	7.55	426
SUM	S. rod	G	Rhône	Doux	Sumène	5	F	43.98	3.72	203
VAU	S. rod	G	Rhône	Ouvèze	Sorgue	5	F	43.92	5.13	89
LOU	?	G	Loup	Loup	Loup	5	F	43.65	7.13	7

Table 2	Table 2. Haplotype frequency distribution at mtDNA control region in wild brown trout populations from the Rhône, Po and other neighbour Mediterranean rivers. Sampling code as in Table 1.																				
Code	Basin	MEcs1	MEcs15	MEcs28	MEcs29	MEcs30	MEcs31	MEcs32	ADporh-1	ADrh-1	ADcs-1	MA2b	Ma2a	MA2c	MAsl-1	hap1	hap2	hap3	hap4	At1e	hap3b
MER	Rhône	5																			
MOU	Rhône	5																			
ALB	Rhône	5																			
ORC	Ро												7	2			5				3
VER	Ро								6			6				6					
TET	Rhône	5																			
OVA	Ро											10	1				9				
ARN	Ро	3										2					1	4			
VIU	Ро											3	1		1		3	1			
PLA	Rhône	1							4												
RIP	Ро	19							5												
TRO	Ро	10							3			7									
GER	Ро	3							1			2						3			
GERb	Ро	5							5			2					2		1	1	
CER	Rhône	9							11												
GLU	Rhône			2														3			
GUIb	Rhône	18							2												
GHI	Ро	1							4									4	1		
BIA	Rhône						4		1												
GUI	Rhône	3							2												
DRÔ	Rhône	5																			
GIU	Ро										1		5				2	2			
ARD	Rhône	5																			
BUË	Rhône								1	4											
MAla	Ро											7						3			
BAR	Ро	11																7			
GIE	Rhône	1							4												
MAIb	Ро								3								1	5	1		
UBA	Rhône	8							13												
SAN	Sansob.	13															6	1			
FER	Ро								8												
FRE	Ро								8												
OUV	Rhône	5																			
RAS	Ро											10					4				
SER	Ро											12						0			
GES	Ро					L	L	L	4												
BOU	Ро								9								4				
SAL	Var	3			2																
FON	Roya							4										1			
SUM	Rhône		5																		
VAU	Rhône	5																			
LOU	Loup					5															

Table S1. Schematic summary of the Mediterranean taxa in the *Salmo trutta* complex of the study area. Reference to mtDNA lineages as follows: 1, Bernatchez 2001; 2, Berrebi et al. 2019; 3, Fabiani *et al.* 2017; 4, Fruciano *et al.* 2014; 5, Giuffra *et al.* 1994; 6, 1996; 7, Gratton *et al.* 2014; 8, Lerceteau-Köhler *et al.* 2013; Maric *et al.* (2017) 9, Meraner & Gandolfi 2018; 10, Sabatini *et al.* 2011; 11, Schöffmann *et al.* 2007; 12, Snoj *et al.* 2011; 13, Splendiani *et al.* 2006; 14, 2007; 15, 2017; 16, Zaccara *et al.* 2015; 17, Maric *et al.* 2017; 18, Tougard *et al.* 2018; 19, this study.

Taxon	Geographical range	mtDNA lineages	mtDNA references	Nomenclature change
Salmo rhodanensis Fowler, 1974	Rhône River basin	AD, ME	1, 19	Questioned the validity of specie rank in this study and in Berrebi <i>et al.</i> (in prep.).
S. cettii Rafinesque Schmaltz 1810	Described for the S.E. Sicily but extended to the Apennines (Tyrrhenian side) by Kottelat & Freyhof (2007)	AT (southern clade)	2, 4, 11, 18	Considered a senior synonym of <i>S. macrostigma</i> by Splendiani <i>et al.</i> (2019).
S. macrostigma (Dumeril, 1858)	Described for Algeria, but extended also to Apennines (Tyrrhenian side), Sicily, Sardinia (Italy) and Corsica (France) by Sommani (1951)	AD, MA, ME (According to Sommani, 1951) AT (southern clade) (according to Splendiani <i>et al.</i> , 2019)	1-4,7-13, 16	Considered a junior synonym of <i>S.cettii</i> by Splendiani <i>et al.</i> (2019).
<i>S. cenerinus</i> Chiereghini, 1847	Described for the Gulf of Trieste, but extended to the Apennines (Adriatic side) by Kottelat & Freyhof (2007)	AD, MA, ME (according to Kottelat & Freyhof, 2007) MA (according to Bianco & Delmastro (2011)	- 1, 2, 7, 9, 13-15	Considered by Bianco & Delmastro (2011) as a junior synonym for <i>Salmo marmoratus</i> Cuvier, 1829.
<i>S. farioides</i> Karaman, 1938	Described for south-western Balkans but extended to the Padano-Venetian district by Bianco & Delmastro (2011)	AD	17	Considered a valid name for the native trout of the Drin River basin (S.E. Balkans) by Marić <i>et</i> <i>al.</i> (2017) and extended by Bianco & Delmastro (2011) to the Adriatic slope of the Apennine.
<i>S. carpio</i> L., 1758	Lake Garda	AD, ME, MA	1, 5-7, 9, 13	Considered a subspecies of <i>S. trutta</i> by Tortonese (1970)
S. marmoratus Cuvier, 1829	Po River Basin and Balkan Peninsula	МА	1, 5-7, 9, 13-15	Considered a subspecies of <i>S. trutta</i> by Tortonese (1970)
Table S1 Bibliography Bernatchez L, 2001, The evolutions	ry history of brown trout (Salmo trutta I) inferred from phylogeog	canhic nested clade	and mismatch analyses of mitochondrial DNA
variation. Evolution 55:351–379.	ay instory of brown front (Sumo tratta E.	, mened nom phylogeogr	upine, nesteu clau	, and mismatch analyses of mitoefolidital DIVA

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Table S2: List of haplotypes retrieved from GenBank, with accession numbers and their distributions.

Haplotype	Accession number	Distribution – country/drainages					
		² Denmark (Skals), Norway (Bjornes Lake, Sima), Spain (hatchery					
haplotype 1 (ATcs1)	AF273086	stocks), ⁴ Spain (Garona), France (Gulf of Biscay), Iceland (Skorradalsvatn), British Isles (Coquet, Wear, Lune, Melvin), ¹¹ continental Italy (IT), Sardinia, Corsica					
		² Denmark (Skals, Karup), Norway (Guddal, Sima), Spain (hatchery					
haplotype 2 (ATcs2)AF273087stocks), ⁵ France (Gulf of Biscay), British Isles (Coquet Fowey, Teifi, Conwy, Loch Romoch), Russia (Nilin ¹¹ Sardinia, Corsica							
		² Denmark (Skals), Norway (Bjornes Lake, Guddal, Sima), Spain					
hanlotype 3 (ATcs3)	AF274574	(hatchery stocks), ⁴ Spain (Garona), France (Gulf of Biscay), British					
	1112/13/1	Isles (Coquet, Wear, Rother, Teifi, Conwy, Melvin), ¹¹ continental Italy, Sardinia, Calabria					
		² Denmark (Skals, Karup), Norway (Bjornes Lake, Guddal, Sima),					
haplotype 4 (ATcs4)	AF274575	Spain (hatchery stocks), ⁴ France (Gulf of Biscay), British Isles					
		(Lune), ¹¹ Sardinia, Calabria					
Hap3b	MK948037	²⁰ Italy (Po)					
Atle	DQ841192	⁸ Italy (Adige), ²⁰ Italy (Po)					
ADcs1	AY836330	³ Spain (Ter, Ebre, Túria, Segura), ⁶ Bulgaria (Struma, Mesta, Maritza), ^{1, 7} Macedonia (Prespa Lake, Vardar), ²⁰ Italy (Po)					
ADcs2	AY836331	³ Spain (Guadalfeo)					
ADcs3	AY836332	³ Spain (Ebre)					
ADcs4	AY836333	³ Spain (Ter)					
ADcs5	AY836334	³ Spain (Guadalfeo)					
ADcs6	AY836335	³ Spain (Ebre, Guadalfeo)					
ADcs7	AY836336	³ Spain (Ebre)					
ADcs8	AY836337	³ Spain (Ebre)					
ADcs9	AY836338	³ Spain (Turia)					
ADcs10	AY836339	³ Spain (Guadalfeo)					
ADes11	AV836340	³ Greece (Alfios), ¹⁰ Montenegro (Skadar Lake), ¹² Albania (Drin,					
	A1050540	Skumbini, Cermit), ¹³ Montengro (Zeta, Morača, Cijevna)					
ADcs15	AY836344	⁴ France - Corsica (Corsica stream)					
ADcs16	AY836345	³ Spain (Ebre)					
ADcs17	AY836346	³ Spain (Ebre)					
ADcs18	AY836347	³ Spain (Guadalquivir)					
ADcs19	AY836348	³ Spain (Guadalquivir)					
ADcs20	AY836349	³ Greece (Tripotamos), ¹³ Bulgaria (Maritza)					
ADporh-1	MK948034	²⁰ France (Rhône), Italy (Po)					
ADrh-1	MK948035	²⁰ France (Rhône)					
AD-Tyrrh1	KX450257	¹¹ Corsica, Sardinia, Calabria, continental Italy (Aniene River)					
AD-Tyrrh2	KX450258	¹¹ Corsica, continental Italy (Aniene River)					
AD-Tyrrh3	KX450259	¹¹ Calabria (Diga Giulia River)					

AD-Tyrrh4	KX450260	¹¹ Sardinia, continental Italy (Aniene River)
AD-Tyrrh5	KX450261	¹¹ continental Italy (Nera River)
AD-Tyrrh6	KX450262	¹¹ continental Italy (Nera River)
AD-zls-01	MG194729	¹⁵ Italy (Liri)
AD-zls-01	MG194729	¹⁵ Italy (Liri)
ScarAD-1 (C208)	KJ834848	¹⁶ Italy (Garda Lake)
ScarAD-2 (C021)	KJ834822	¹⁶ Italy (Garda Lake)
MEcs1	AY836350	 ³Spain (Ter, Llobregat, Ebre, Míjares, Palancia, Túria, Segura), ⁵Croatia (Krka), ²⁰France (Rhône, Var), Italy (Po, Sansobbia)
MEcs4	AY836353	³ Spain (Ter)
MEcs6	AY836355	³ Spain (Ebre)
MEcs7	AY836356	³ Spain (Ter)
MEcs8	AY836357	³ Spain (Túria)
MEcs9	AY836358	³ Spain (Túria)
MEcs10	AY836359	³ Spain (Ebre)
MEcs11	AY836360	³ Spain (Túria)
MEcs12	AY836361	³ Spain (Ter)
MEcs13	AY836362	³ Spain (Ebre)
MEcs14	AY836363	³ Spain (Ebre, Guadalfeo)
MEcs15	AY836364	³ Spain (Ebre), ²⁰ France (Rhône)
MEcs16	unpublished	¹⁷ Spain
MEcs17	unpublished	¹⁷ Spain
MEcs18	unpublished	¹⁷ Spain
MEcs20	unpublished	¹⁷ Spain
MEcs21	unpublished	¹⁷ Spain
MEcs22	unpublished	¹⁷ Spain
MEcs23	MG970273	¹⁸ Spain (Cardener)
MEcs25	MG970274	¹⁸ Spain (Cardener)
MEcs26	MG970275	¹⁸ Spain (Cardener)
MEcs27	MG970276	¹⁸ Spain (Cardener)
MEcs28	MK948029	²⁰ France (Rhône)
MEcs29	MK948030	²⁰ France (Var)
MEcs30	MK948031	²⁰ France (Loup)
MEcs31	MK948032	²⁰ France (Rhône)
MEcs32	MK948033	²⁰ France (Roya)
ME-nin63	MG194732	¹⁵ Italy (Sisto)
MAcs1	AY836365	³ Slovenia (Soča), ⁶ Greece (Aliakmon), ⁸ Italy (Adige), Italy (Po)
Ma2a	DQ841189	⁸ Italy (Adige), ²⁰ North Italy (Po)
Ma2b	DQ841190	⁸ Italy (Adige), ²⁰ North Italy (Po)
Ma2c	JQ582461	⁹ Italy (Adige), ²⁰ North Italy (Po)
ScarMA1 (C201)	KJ834841	¹⁶ Italy (Garda Lake)

ScarMA2 (C024)	KJ834825	¹⁶ Italy (Garda Lake)
marm1	KJ834770	¹⁶ North Italy (Po, Adige), Slovenia (Soca River)
Mak1	JX846931	¹⁹ Croatia (Krka)
MAsl-1	MK948036	²⁰ North Italy (Po)

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- ²⁰ Present study (new haplotypes)

Table S3. Demographic indices calculated for three brown trout mtDNA lineages based on control region sequence analysis.

		Neutrality to	est	Mismatc	Model					
	Fs	D	R ₂	SSD	r					
ME	-2.874*	-1.897*	0.124***	0.027**	0.066*	Expansion				
AD	-2.126	-2.147*	0.121***	0.023**	0.076**	Expansion				
MA	-1.019	-1.070	0.188***	0.039	0.111	Stable				
Fs = Fu's	Fs = Fu's F statistic, D = Tajima's D statistic, R2 = Ramos-Osisns and Rozas statistic, SSD = sum of									
standard deviations of mismatch distribution, r = raggedness index of mismatch distribution.										
***p<0.0	001, **p<0.01	L <i>,</i> *p<0.05.								

Substitution rate (%) Node TMRCA (MY) Upper 95% HPD (MY) Lower 95% HPD (MY) ADporh1-ADrh1 0.151 0.011 0.099 0.278 AD 0.170 0.391 0.75 ME 0.267 0.166 0.372 MA 0.122 0.172 0.205 ADporh1-ADrh1 0.120 0.016 0.086 AD 0.212 0.129 0.298 1 ME 0.191 0.122 0.265 MA 0.117 0.193 0.051

Table S4. Time to the most common ancestor (TMRCA) estimates for the Mediterranean brown trout lineages AD, ME and MA and for the sub-clade ADporh-1, ADrh-1 with 95% highest probability density (HPD) intervals.