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Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/90494> since 2015-12-01T13:40:48Z

Published version:

DOI:10.1007/s10342-011-0584-3

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by P. Belletti, D. Ferrazzini, A. Piotti, I. Monteleone, F. Ducci

[European Journal of Forest Research, 131 (4): 1127-1138, 2012, DOI 10.1007/s10342-011-0584-3]

*The definitive version is available at:
La versione definitiva è disponibile alla URL:*

[<http://link.springer.com/journal/10342/131/4/page/1>]

26 **Genetic variation and divergence in Scots pine (*Pinus sylvestris* L.) within its natural range in Italy**

27

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37

38 **Abstract**

39 Twenty-one populations of Scots pine sampled over the entire Italian range of the species were
40 analysed for genetic variation scored at 9 nuclear SSR markers. The main aim of the work was to find
41 genetic features useful for conservation management, namely allelic composition, gene diversity and
42 differentiation.

43 High levels of intra-population variability were scored. The only population sampled in the
44 Apennines gave the lowest values, confirming the genetic erosion undergone in the Scots pine remnants in
45 this area. A low level of genetic variability was also scored for populations from the Po valley and hills of
46 Piedmont. Most genetic diversity was found within-populations, while only a small amount occurred among
47 them ($F_{ST} = 0.058$). Both Bayesian clustering and sPCA analysis showed a East-West subdivision,
48 notwithstanding the unclear position of populations from the Po valley. The population from the Apennines
49 was always clearly separated from the others.

50 The results are discussed in terms of post-glacial recolonisation, as well as for defining the regions of
51 provenance of Scots pine in Italy. The management of genetic resources could benefit from the identification
52 of genetically homogeneous regions, thereby avoiding the use of non-local reproductive material for
53 plantations, which is well known as one of the most important reason for failure of reforestation.

54

55 **Key words**

56 Genetic differentiation, genetic variation, glacial refugia, regions of provenance, Scots pine, SSR markers

57

58 **Introduction**

59 Genetic erosion is one of the most serious threats for the survival of forest ecosystems worldwide. A
60 high level of variability is essential to supply populations with strong adaptability. This is particularly
61 important for forest species, which consist of individuals with long life-cycles and no possibility of migrating
62 to more favourable sites (Palmberg-Lerche 2001, Toro and Caballero 2005). Genetic erosion can be strongly
63 enhanced by habitat fragmentation and marginality in the species' range (Jump and Peñuelas 2006, Eckert et
64 al. 2008). However, the expected 'genetic signal' depending on increasing isolation and decreasing
65 population size is often undetected, as fragmentation may have occurred relatively recently and fragments
66 may contain large remnant populations and the longevity of many tree species can delay the loss of genetic
67 variability (Kramer et al. 2008). In addition, the historically underestimated long distance dispersal capability
68 of forest trees can also maintain a high connectivity among widely isolated stands (Robledo-Arnuncio and
69 Gil 2005, Nathan 2006, Williams 2010). Conversely, climate warming can strengthen the loss of genetic
70 variability due to fragmentation, causing a marked decline in growth and survival of marginal southern
71 populations of temperate tree species, as demonstrated in European *Pinus sylvestris* populations by Rubiales
72 et al. (2008) and Reich and Oleksyn (2008).

73 Knowledge of the level and distribution of genetic variation is of the utmost importance in providing
74 information for the conservation and the management of genetic resources. Furthermore, genetic analysis
75 based on molecular markers can increase our understanding of the historical processes that led to the present
76 distribution of a species (Petit et al. 2003). These markers can provide us with appropriate means to obtain
77 information on the genetic structure of populations, as well as to analyse the distribution of within-species
78 variability (Pautasso 2009). Such data provide important insights for preservation and restoration programs,
79 indicating areas of high genetic diversity and geographic limits for seed collection, and delimiting the scale
80 at which conservation should be planned (Escudero et al. 2003).

81 Scots pine (*Pinus sylvestris* L.) is the most widespread European conifer tree, and its natural range
82 extends from the arctic circle in Scandinavia down to southern Spain and central Italy, and from western
83 Scotland to eastern Siberia. In southern Europe and Asia Minor, isolated occurrences are confined to the

84 mountain zone (up to 2200 m in altitude in the Balkans and Spain, and 2700 m in the Caucasus). In Italy the
85 species is widely spread throughout the Alps, and some relic populations can be found in the northern
86 Apennines, in the hilly areas of Piedmont (north-western Italy) and in the upper part of the Po valley
87 (Pignatti 1982). The present distribution is highly influenced by human activities. In the Alps Scots pine has
88 often been substituted by other species, namely Norway spruce and black pine, while the other populations
89 are nowadays regressing, mainly due to recruitment limitation, competition with other forest trees and as a
90 consequence of rural depopulation and the abandonment of wood management (Camerano et al. 2008).

91 The complex biogeographic history of Scots pine in Europe has been extensively studied (Sinclair et
92 al. 1999, Soranzo et al. 2000, Cheddadi et al. 2006). Recent work has shown the presence of a previously
93 unknown glacial refugium in northern Europe during the last glaciation (Naydenov et al. 2007, Pyhäjärvi et
94 al. 2008), and of several small putative refugial areas in Southern Europe that gave rise to geographically
95 limited “interglacial refugia” (Cheddadi et al. 2006). Among the southernmost populations the Spanish,
96 Balkan and Turkish ones have usually been considered in broad scale phylogeographic studies (Sinclair et al.
97 1999, Soranzo et al. 2000, Naydenov et al. 2007, Pyhäjärvi et al. 2008), whereas Alpine and Apennine
98 populations have received less attention (Scalfi et al. 2009). In the few studies where they were extensively
99 sampled it was demonstrated that Italian populations shared a common mitochondrial haplotype, and that
100 they are different from the surrounding Austrian, Swiss, and French alpine populations (Cheddadi et al.
101 2006). Cheddadi et al. therefore hypothesised a common origin for Italian Scots pine populations from a
102 refugial area in Southern Italy, even though the results by Puglisi and Attolico (2000), Labra et al. (2006) and
103 Scalfi et al. (2009), obtained with more polymorphic genetic markers (respectively allozymes, ISSRs and
104 SSRs), showed a marked differentiation between Alpine and Apennine populations, suggesting different
105 recolonisation routes for the two mountain chains. The preservation of genetic resources of highly
106 fragmented Apennine populations therefore appears to be a high priority challenge.

107 At the population level genetic structure and gene flow patterns have been studied in Spanish
108 populations from the northern Meseta by Robledo-Arnuncio and Gil (2005) and Robledo-Arnuncio et al.
109 (2005), and show high within-populations genetic diversity and extremely high levels of pollen flow over
110 long distances (5% longer than 30 km). On the other hand, small relic Scots pine populations from the
111 Apennines had significantly lower genetic diversity, and are differentiated from the alpine ones, possibly as a

112 consequence of progressive isolation since the early Holocene and their origin from different glacial refugia
113 (Labra et al. 2006, Scalfi et al. 2009). Scalfi et al. (2009) however, hypothesised a possible different role of
114 gene flow via-pollen and seed in determining this genetic differentiation.

115 In this study we surveyed the genetic variability of 21 Scots pine populations throughout the species
116 distribution in the Italian peninsula, using 9 highly informative nuclear microsatellite (nSSR) markers. Our
117 main aim was to assess the levels and distribution of genetic variability of this conifer by intensively
118 sampling the entire Italian range of the species, and to investigate the presence of any cryptic genetic
119 structure shaped by postglacial recolonisation which went undetected in previous studies based on organellar
120 markers. We also discussed our results in the light of the European Directive 105/1999, with particular
121 emphasis on the preservation and restoration of Scots pine genetic resources in the Alps and Apennines.

122

123 **Material and Methods**

124 *Plant material*

125 Twenty-one native populations of Scots pine were chosen within the natural range of diffusion of the
126 species in Italy (Figure 1). Four of them (BOS, VEZ, PAS and CAS) are located in the hilly areas of
127 Piedmont, two (TIC and OLG) grow in the upper part of Po valley and another (CRO) is found in the
128 northern Apennines. All the others are distributed along the entire Italian Alpine region. Table 1 summarises
129 names and locations of the populations analysed. Most of the populations belong to mixed forests, with the
130 exception of populations CAR, FEN and SAV which are pure stands of Scots pine. In other locations
131 accompanying species vary according to altitude and latitude: English oak, hornbeam and wild cherry (TIC),
132 pubescent oak, flowering ash and juniper (CRO), sessile oak and European ash (OLG, MAS, GAR), sessile
133 oak, black locust, elm and maples (BOS, VEZ, PAS, CAS), black pine (DOG), Norway spruce (VAL, SIU,
134 BRU, COR, CLA), silver fir (TOC), larch (SAR) and mountain pine (BRI). Among the populations sampled,
135 CRO, CAR, FEN, OLG, VAL, SIU and BRU are registered in the Italian National Book of Seed Stands for
136 Scots pine, selected for their phenotypic characteristics and health status (Morandini and Magini 1975).

137 Twenty-four adult non-adjacent trees were chosen at random in each population. Since some
138 individuals did not show reliable nSSRs banding patterns, they were excluded from the analysis. The number
139 of these individuals varied among populations: from zero up to six. Consequently, the total number of

140 analysed individuals was 449. Needles collected from the trees were stored at -20°C until DNA extraction
141 was carried out.

142

143 *Molecular analysis*

144 Frozen needles (100 mg of tissue) were powdered in liquid nitrogen and genomic DNA was
145 extracted using the QIAGEN[®] DNeasy plant mini kit, according to the manufacturer's protocol. A total
146 concentration of 20 ng/ μl was obtained for each sample.

147 Twelve simple sequence repeat markers (SSR) were selected according to the literature, and tested
148 on our plant material (Table 2). While the SPAC and SPAG series consisted of primers specific for *P.*
149 *sylvestris* (Soranzo et al. 1998), the PtTX series included primers originally designed for *Pinus taeda*, but
150 they also proved to be as useful as the markers selected for *P. sylvestris* (Elsik et al. 2000, Auckland et al.
151 2002, González-Martínez et al. 2004).

152 Polymerase Chain Reaction (PCR) amplifications were performed using a Perkin Elmer GeneAmp[®]
153 PCR System 9600 thermal cycler. The protocol was slightly modified, according to the presence of a
154 fluorochrome (IR-Dye 700 and IR-Dye 800) attached to each forward primer. Each amplification reaction
155 contained 1X reaction buffer (Promega), 2.5 mM MgCl_2 , 0.2 mM dNTPs, 0.5 μM of each primer, 0.65 U of
156 GoTaq DNA Polymerase (Promega), approximately 10 ng genomic DNA, and deionised water to a total
157 volume of 13 μl .

158 The PCR profiles varied for different primers, and were adjusted for the presence of the forward
159 labelled ones: for *P. sylvestris* specific primers the profiles included an initial step of 3 min at 94°C , followed
160 by 5 cycles of touchdown consisting of 94°C for 30 s, 65°C for 30 s $\Delta\downarrow 1^{\circ}\text{C}$ (SPAC 11.4 and SPAC 11.5) or
161 60°C for 30 s $\Delta\downarrow 1^{\circ}\text{C}$ (SPAC 11.8, SPAC 12.5, SPAG 3.7 and SPAG 7.14), 72°C for 1 min, and subsequent
162 25 cycles of amplification consisting of 94°C for 30 s, 60°C for 30 s, 72°C for 1 min, and a final extension of
163 72°C for 10 min. Touchdown PCR was not necessary for SPAC 11.6, where the protocol consisted of a first
164 step at 95°C for 5 min, and 35 cycles (denaturation at 94°C for 1.5 min, annealing at 55°C for 1.5 min and
165 elongation at 72°C for 1.5 min), followed by 10 min at 72°C . For *P. taeda* primers PCR profiles consisted of
166 an initial step of 5 min at 94°C , followed by 20 cycles of touchdown [94°C for 1 min, 59°C for 30 s $\Delta\downarrow$
167 $0,5^{\circ}\text{C}$ (PtTX 3032) or 55°C for 30 s $\Delta\downarrow 0,5^{\circ}\text{C}$ (PtTX 3116) or 60°C for 30 s $\Delta\downarrow 0,5^{\circ}\text{C}$ (PtTX 3107, PtTX

168 4001 and PtTX 4011), 72°C for 1 min)], followed by 20 cycles of amplification at 94°C for 1 min, annealing
169 temperature for 1 min, 72°C for 1 min, and a final extension of 72°C for 3 min.

170 The forward sequence of each primer pair was labelled with a fluorescent dye at its 5' end: IR-Dye
171 800 for SPAC 11.6, SPAC 12.5, SPAG 7.14, PtTX 3107 and PtTX 4001, IR-Dye 700 for SPAC 11.4, SPAC
172 11.8, SPAC 11.5, SPAG 3.7, PtTX 3032, PtTX 3116 and PtTX 4011.

173 Electrophoresis and detection of PCR products were carried on a 6%, 25 cm long, 0.25 mm thick,
174 denaturing polyacrylamide gel using a sequencer (model DNA 4200 Sequencer LI-COR Biotechnology).
175 Gels were run at 2000 V in TBE 1X buffer, for 1 to 3 h, depending on the product sizes. Determination of
176 polymorphism was obtained using two different marked standards of known molecular weight (50–350 bp,
177 and 50–700 bp). Data were collected by e-Seq software (DNA Sequencing and Analysis Software), and all
178 the size scores were visually checked.

179

180 *Data processing*

181 Allele frequencies and within-population genetic diversity parameters (mean number of alleles per
182 locus, A ; mean number of private alleles per locus, P_a ; effective number of alleles per locus, N_e ; observed and
183 expected heterozygosity, H_O and H_E , respectively) were estimated using GenAEx v.6 software (Peakall and
184 Smouse 2006). Allelic richness, based on a minimum sample size of 16 gene copies (A_{r16}), was calculated
185 using FSTAT (Goudet 1995). Genotypic disequilibrium between pairs of loci was tested at the single
186 population level and across all populations, with Fisher's exact test using Arlequin software (Excoffier et al.
187 2005).

188 Fisher's exact test using the Markov Chain algorithm (Guo and Thompson 1992) was used to assess
189 deviations from Hardy-Weinberg equilibrium for each population and each locus, and where significant
190 deficiencies of heterozygotes from Hardy-Weinberg expectations were found the presence of a relatively
191 high frequency of null alleles was suspected (Pemberton et al. 1995). Loci with high frequencies of null
192 alleles were identified by estimating null allele frequencies for each locus and each population, using the
193 software Micro-Checker (Van Oosterout et al. 2004). In further analysis we eliminated problematical loci
194 with high null allele frequency from the dataset, using only loci with < 0.19 null allele frequencies. This
195 value has been considered as a threshold over which significant underestimate of H_E due to null alleles can

196 be found (Chapuis et al. 2008). Where possible, we employed robust analyses for the presence of null alleles
197 (Chapuis and Estoup 2007, Chapuis et al. 2008), including STRUCTURE (Pritchard et al. 2000, Falush et al.
198 2003), ordination methods (sPCA, Jombart et al. 2008), and the Mantel test on chord distance (DC, Cavalli-
199 Sforza and Edwards 1967). In particular the inbreeding coefficients (F_{IS}) were calculated taking into account
200 the estimated null allele frequencies using the program INEst, and running the individual inbreeding model
201 (IIM) with a Gibbs sampler of 10^5 iterations (Chybicki and Burczyk 2009).

202 FreeNA was used to compute the value of Weir's (1996) estimators of F-statistics to analyse the
203 population genetic structure of the overall samples. In particular, F_{ST} was calculated in order to estimate the
204 proportion of the total genetic variation due to differentiation among populations. Genetic differentiation
205 between populations was estimated computing a pairwise F_{ST} . FreeNA applies the ENA correction method to
206 efficiently correct for the positive bias induced by the presence of null alleles on the F_{ST} estimation (Chapuis
207 and Estoup 2007).

208 The genetic structure of the populations was explored using Bayesian clustering and spatial principal
209 components analysis (sPCA). Bayesian clustering was performed with the software STRUCTURE (Pritchard
210 et al. 2000). The program uses a Markov chain Monte Carlo (MCMC) algorithm to cluster individuals into
211 populations on the basis of multilocus genotypic data. Individual multilocus genotypes are first assigned
212 probabilistically to genetic clusters (K) without considering sampling origins. Admixed or hybrid individuals
213 can be identified as they will have a fraction of their alleles derived from each genetic cluster. The program
214 was run setting a burn-in period of 10^5 followed by 5×10^5 iterations, and using the admixture ancestry model
215 and the correlated allele frequency model, given the low F_{ST} and the high genetic connectivity typical of
216 forest tree populations. Posterior probabilities of K were calculated from the means of 20 runs for each value
217 of $K \in \{1, \dots, 10\}$, and the optimum K determined using the method of Evanno et al. (2005).

218 The sPCA is a spatially explicit multivariate method recently developed by Jombart et al. (2008) to
219 investigate the spatial pattern of genetic variability using allelic frequency data of individuals or populations.
220 It takes spatial information directly into account as a component of the adjusted model, focusing on the part
221 of the variability that is spatially structured. This analysis does not require data to meet the Hardy–Weinberg
222 expectations, or linkage equilibrium to exist between loci. The sPCA yields scores summarising both the
223 genetic variability and the spatial structure among individuals (or populations). Global structures (patches,

224 clines and intermediates) are disentangled from local ones (strong genetic differences between neighbours)
225 and from random noise. Neighbouring sites were defined by building a connection network based on
226 Delaunay triangulation. The existence of global and local structuring was tested using the multivariate Monte
227 Carlo tests implemented, as the sPCA procedure, in the *adegenet* package for R (Jombart 2008; R
228 Development Core Team, 2009).

229 A Mantel (1967) test was applied to the matrices of pairwise chord distance and log-transformed
230 geographical distance between populations (natural logarithm scale) to assess isolation-by-distance, namely
231 the model under which genetic differentiation between populations is the result of drift. Chord distance
232 (Cavalli-Sforza and Edwards 1967) for each pair of populations was calculated using the INA correction
233 described in Chapuis and Estoup (2007) with FreeNA. The test of significance for Mantel test was carried
234 out on 9,999 permutations of the data. Mantel test was executed on the entire dataset and on clusters detected
235 by STRUCTURE and sPCA analyses.

236 Finally, we used the program Bottleneck v.1.2.02 (Piry et al. 1999) to test for recent population
237 bottlenecks. A Wilcoxon's sign rank test was used to compare expected heterozygosity from Hardy-
238 Weinberg equilibrium with predicted heterozygosity at mutation-drift equilibrium, on the basis of the
239 observed allele number (Piry et al. 1999). The program was run under a two-phase model of mutation (TPM)
240 that generally fits microsatellite evolution better than either pure stepwise or infinite allele models (Di
241 Rienzo et al. 1994). One thousand simulations were performed for each sample based on a TPM consisting
242 of 90% single-step mutations and 10% multistep changes.

243

244 **Results**

245 *Allelic diversity of microsatellite loci*

246 Ten out of the 12 tested microsatellites (SPAC 11.4, SPAC 11.6, SPAC 11.8, SPAC 12.5, SPAG
247 7.14, PtTX3032, PtTX3107, PtTX3116, PtTX4001 and PtTX4011) showed reliable banding patterns with
248 clear and repeatable bands. On the contrary, SPAC 11.5 did not amplify in any sample, and SPAC 3.7 failed
249 to amplify the DNA in the majority of samples. The latter two markers were therefore excluded from the
250 analysis. SPAC 11.8 was also excluded due to the high frequency of possible null alleles (0.57). The

251 presence of null alleles was also suspected for 6 out of the remaining 9 loci, with lower frequencies ranging
252 from 0.07 (PtTX3032) to 0.19 (SPAC 11.6).

253 The 9 selected microsatellites were highly polymorphic, and generated a total of 532 alleles (range
254 21 to 151), with a mean number of ~48 alleles per locus. It was also possible to detect 137 private alleles, i.e.
255 present only in one population. The population frequency of private alleles was on average 0.036 and ranged
256 between 0.021 and 0.208. The distribution of these alleles among loci ranged from 43 (PtTX3032) to 1
257 (PtTX3107). PAS (17 alleles) was the population with the highest number of private alleles, followed by
258 TOC (14 alleles) and COR (10 alleles), and three populations (BOS, CLA and GAR) presented only two
259 private alleles each.

260

261 *Genetic variation within populations*

262 Statistics on the genetic diversity within populations are given in Table 3. A high level of intra-
263 population variability was found, since on average more than 14 alleles per locus were observed ($A = 14.45$).
264 CRO (the only populations from the Apennines) showed the lowest value of genetic diversity ($A = 10.4$, $N_e =$
265 5.4 , $A_{r16} = 6.781$, and $H_e = 0.754$). The populations belonging to the Po valley (TIC and OLG) also showed a
266 low variability, as well as the populations from the hills of Piedmont (BOS, VEZ, PAS, and CAS), even
267 though the genetic impoverishment was less pronounced. Except for these particular populations (which, in
268 most cases, are small, isolated, and/or on the edge of the main range) only BRI, GAR, and CLA showed a
269 more or less marked reduction in genetic diversity. This pattern was confirmed for all genetic diversity
270 parameters considered.

271 Despite the fact that some of the investigated populations are characterised by a low genetic
272 diversity, no evidence for recent bottlenecks was found. In fact, for all the populations HW heterozygosity
273 and expected gene diversity at mutation drift equilibrium did not differ significantly.

274 The probability that two randomly sampled alleles in a given population were not the same was
275 higher than 84% (mean $H_e = 0.847$), whereas the observed heterozygosity (mean $H_o = 0.670$) was lower than
276 expected. The difference, that determines a significant positive value for mean inbreeding coefficient, is
277 mainly due to non-random mating and null alleles. Recalculating the inbreeding coefficients, taking into
278 account the frequencies of null alleles, we found that deviations from the Hardy-Weinberg equilibrium were

279 low (F_{IS} ranging from 0.052 in FEN to 0.016 in TIC, with a mean of 0.033). None of estimated F_{IS} was
280 significantly different from zero (confidence interval calculated through INEst estimates of F_{IS} overlapped
281 zero in all populations).

282

283 *Genetic differentiation among populations*

284 Most of the genetic diversity was found within populations, while a small amount of the variability
285 occurred among populations ($F_{ST} = 0.058$, CI: 0.037-0.081). The F_{ST} values per locus ranged from 0.024
286 (SPAC 11.6) to 0.12 (PtTX 4011), and there were no obvious differences between the *P. sylvestris* and the *P.*
287 *taeda* sets of markers. The genetic divergence between populations was further investigated by computing a
288 pairwise F_{ST} matrix. Multilocus F_{ST} values varied between 0.015 (CAR and SAR) and 0.141 (CRO and
289 VEZ). The population from the Apennines (CRO) was always clearly separated by the others. Almost all
290 pairwise F_{ST} values were significantly greater than zero, confirming the presence of a slight, although
291 significant, amount of population structuring in Italian Scots pine (results not shown).

292 Following the method of Evanno et al. (2005), the Bayesian clustering results obtained with
293 STRUCTURE indicate that $K=2$ clusters represents the most informative representation of the overall
294 genetic structure that we analysed (Figure 2). We found that most individuals from western populations
295 (VEZ, PAS, CAS, SAV, and TOC) clearly belong to cluster 2, whereas eastern populations and CRO (the
296 Apennine population) are primarily composed by individuals from cluster 1, with the exception of some
297 admixed populations (COR, SIU, and VAL). Although populations belonging to the pedo-climatic region of
298 the Po valley (TIC and OLG) are closer to western populations, their individuals are predominantly assigned
299 to cluster 1. This East-West subdivision was confirmed by sPCA analysis (Figure 3). The existence of such
300 ‘global structure’ (*sensu* Jombart et al., 2008) was demonstrated using both Delaunay triangulation ($t_{max} =$
301 0.0631, $P < 0.05$) and Gabriel graph ($t_{max} = 0.0672$, $P < 0.05$) for building the connection network.

302 The correlation between genetic diversity, expressed as Cavalli-Sforza and Edwards (1967) chord
303 distance for pairs of populations, and the logarithm of distances expressed in km, did not show the typical
304 pattern of isolation by distance, and did not suggest a strong relationship between the two factors. Mantel’s
305 test was not significant when performed on the entire dataset ($P = 0.094$), as well as when performed

306 separately on the two clusters detected by sPCA analysis ($P = 0.669$ in eastern populations, and $P = 0.482$ in
307 western populations excluding CRO).

308

309 **Discussion and Conclusion**

310 The aim of this study was to assess the level and the distribution of genetic variation of Scots pine
311 throughout its natural range in Italy, in order to get fundamental knowledge that can be applied for plant
312 propagation and genetic resources conservation. The results could be slightly biased due to the limited
313 number of individuals sampled per populations. Although a larger database (more than 50 individuals per
314 population) would be preferred to obtain stronger data on genetic variation at the level of polymorphic loci
315 (Nei 1978), phylogeographic studies using nSSR markers often are based on less than 30 individuals per
316 populations (e.g. Williams et al. 2007, Ferrazzini et al. 2008, Bagnoli et al. 2009, Scalfi et al. 2009, Bai et al.
317 2010). Kalinowski (2005) demonstrated that some genetic distances, such as F_{ST} , showed limited sampling
318 variance at loci characterised by a high mutation. He also showed that increasing the number of loci, instead
319 of increasing the number of individuals, is an effective way to improve the precision of measures of genetic
320 differentiation. Miyamoto et al. (2008) showed by resampling simulations, that an accurate estimate of
321 genetic diversity (H_e) can also be achieved with small samples (less than 30 plants per population) genotyped
322 at nSSRs. On the other hand, larger sample sizes are needed in order to obtain more accurate estimates of
323 allelic richness, although the statistical technique of rarefaction can compensate for sampling disparities and
324 allow for meaningful comparisons among populations. In general, in the present paper we used population
325 genetic indexes and techniques that minimize possible estimation biases caused by small sample size.

326 The populations analysed showed a considerable amount of genetic diversity, as estimated by means
327 of variation scored at nine nuclear microsatellite loci. The observed number of alleles per locus (A) ranged
328 from 10.4 to 18.4, with an average per population of 14.45, and the average gene diversity (H_e) was as high
329 as 0.847. The high degree of observed diversity is not surprising since it has been recognised for a long time
330 as a peculiar characteristic of woody plants (Hamrick et al. 1992). Furthermore, species such as Scots pine,
331 which do not have a strong habitat specificity, and are almost continuously distributed, are expected to have
332 more within-population diversity than those with strong habitat preference and a scattered distribution. Data
333 from this analysis are coherent with those reported in the literature: for example Robledo-Arnuncio et al.

334 (2005) estimated nSSR loci values of 23.0 and 0.923 for A and H_e respectively. However, this Authors used
335 only three loci, but if we reference our data with these markers we obtained values of 19.4 (A) and 0.885
336 (H_e).

337 The Apennine population (CRO) showed lower values for all the calculated genetic variation indices.
338 This confirms the genetic erosion undergone by this population, most likely as a consequence of isolation
339 and limited population size. Lower values of genetic diversity in Apennine populations, compared to Alpine
340 ones, were already recorded by Scalfi et al. (2009) at nSSR loci, and by Puglisi and Attolico (2000) and
341 Labra et al. (2006), using different genetic markers, respectively isozymes and ISSRs. Similar patterns were
342 also observed with reference to populations from the Po valley (TIC and OLG). As for CRO, habitat
343 fragmentation could be the main cause of the genetic diversity reduction detected for these populations. TIC
344 and, to a lesser extent, OLG can be considered as relic populations, especially after the reduction of the Scots
345 pine range in the Po valley due to intensive use of land for both agriculture and urbanisation purposes. These
346 non-alpine populations are nowadays regressing, mainly due to problems of seed dispersal, competition with
347 other forest trees and as a consequence of rural depopulation and abandonment of wood management
348 (Camerano 2008, Regione Piemonte and Regione Valle d'Aosta 2008). Their preservation is therefore a
349 primary goal: it is needed to maintain genetic diversity as well as the forest landscape. Particular attention
350 should be addressed to the population PAS. It is a very small population, where only a few dozens of
351 individuals still survive, threatened by the competition with other species (namely black locust). This
352 population showed the highest number of private alleles (17) confirming an ongoing genetic erosion process
353 which could cause the local extinction of the population over a short period of time.

354
355 The overall level of genetic diversity arising from the differentiation between populations found in
356 this study ($F_{ST} = 0.058$) is moderate, but higher than that previously observed in the same species in other
357 European countries (Müller-Starck et al. 1992). In the Scandinavian region, for instance, values of $F_{ST} \leq 0.02$
358 were found in populations of Scots pine studied with different markers (Karhu et al. 1996). Allozymes gave a
359 $F_{ST} = 0.03$ between Sweden and Siberian populations (Wang et al. 1991), and a $G_{ST} = 0.021 \div 0.046$ among
360 many European stands (Prus-Glowacki et al. 1993, Prus-Glowacki et al. 2003). The F_{ST} value found in our
361 study is interesting, especially if we consider the relatively small geographic distances between the Italian

362 populations. A low level of genetic differentiation among populations is common in conifers. They maintain
363 most of their variation within populations (Hamrick et al. 1992), which can be explained by the mainly
364 allogamous mating system, and by the high gene flow rate favoured by their dispersal strategy and
365 widespread diffusion (Petit and Hampe 2006, Piotti et al. 2009, Williams 2010). Mantel's test, performed in
366 order to check the presence of IBD between the populations studied was not significant, and thus IBD was
367 apparently not a mechanism shaping the present distribution of genetic variability.

368 The analysis of genetic differentiation, and the most likely population clustering according to
369 STRUCTURE and sPCA analyses indicate that the global structure detected separate gene pools for the
370 eastern and the western Alps. Despite the higher similarity of the Apennine population with the eastern Alps
371 populations, rather than with closer populations from the western Alps, we found a generally high
372 differentiation between populations from the two mountain chains. These two results seem to exclude a
373 common postglacial origin of Italian *P. sylvestris* populations based on the shared RFLP and *nad* 1 intron
374 mitotypes (Sinclair et al. 1999, Cheddadi et al. 2006, Labra et al. 2006). Our results support the hypotheses
375 of an Apennine glacial refugium (see for instance Puglisi and Attolico 2000), but with no evidence of any
376 expansion from Apennines into the southern slope of the Alps, as hypothesised by Cheddadi et al. (2006).
377 The *P. sylvestris* Apennine population (CRO) is genetically distinct from those in the western Alps, although
378 a certain level of admixture exists within the BOS and CAR populations. Similar results have been recently
379 obtained by Piovani et al. (2010), studying *Abies alba* populations. These authors also found that Apennine
380 populations are genetically different from populations from the western Alps. Recent studies based on the
381 analysis of stratigraphic records of pollen, stomata, and macrofossils in northern Italy showed that, although
382 the southern slope of the Alps was extensively glaciated during the last glacial maximum (LGM), conifer and
383 several broad-leaved tree species survived in the Po plain and along the southeastern Alpine border (Vescovi
384 et al. 2007). It has also been recently demonstrated that *P. sylvestris* survived the LGM in the Euganean Hills
385 (north-eastern Italy), a hilly area 50-60 km south of the maximum extent of the last Alpine glaciation
386 (Kaltenrieder et al. 2009). An early presence (ca 15000 BP) of *P. sylvestris* after LGM was also signalled by
387 Finsinger et al. (2006) in the western Alps, at Lago piccolo di Avigliana (Piedmont). Our results depict a
388 scenario where populations from the western Alps, eastern Alps and the Apennines originated from at least
389 three different refugia, with possible contact zones between the western and eastern Alps,

390 colonisation/expansion routes at the latitude of the TIC and TOC populations, and between the western Alps
391 and the Apennine populations in southern Piedmont (BOS and CAR populations). In addition, our study
392 presents further evidence of the possible past genetic connection of western Alps and Apennine populations
393 through the Po plain, a vast area (ca 200 km large) where *P. sylvestris* was widespread during the early
394 Holocene (Labra et al. 2006, Scalfi et al. 2009).

395 The management of genetic resources may benefit from the identification of genetically
396 homogeneous regions, since genetic pollution of local genetic stocks by plantations of non-local origin
397 material can be greatly reduced by the use and transfer of suitable propagation material. Assuming local
398 populations to be optimal as basic material for local uses, genetic zones would have relevance for defining
399 variation in potential commercial and adaptive traits; and they would provide the genetic background
400 required in order to establish the number and location of primary gene-pool reserves for the species.
401 Moreover, the definition of genetic parameters based on molecular markers for each genetic zone can be
402 useful for certification of seedlots and breeding programmes (Bucci and Vendramin 2000). Since 1999, the
403 European Council Directive 1999/105/CE regulates the forest reproductive material market and transfer in
404 Europe. The Italian Government has implemented this directive, with the Decree No. 386/2003. One of the
405 most important features of the acts is the definition of region of provenance as “the area or group of areas
406 subjected to sufficiently uniform ecological conditions in which stands or seed sources showing similar
407 phenotypic or genetic characters are found”. The identification of these areas plays a basic role for a rational
408 management of activities linked with forest tree propagation, including afforestation and in situ genetic
409 preservation.

410 The results of our study contribute to a better understanding of our knowledge on genetic variation of
411 Scots pine in Italy. The information is of basic importance for the definition of regions of provenance for
412 Scots pine, although a deeper knowledge of the ecological characteristics of the areas of the study, such as
413 vegetational and phytogeographical data, is also needed.

414

415 **Acknowledgments**

416 The research was supported by the project “RISELV.ITALIA” (task 1.1, Biodiversity and
417 production of forest reproductive material) from the Italian Ministry for Agriculture, Food and Forestry

418 Policy. Thanks are also due to the Italian Ministry of Education, Universities and Research, which partially
419 granted the work (local research funds). Thanks are also due to colleagues from “Translations Group” for the
420 language revision of the manuscript (www.translation-group.com/).

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648 FIGURE LEGENDS

649

650 Figure 1. Geographical distribution of the 21 populations of Scots pine analysed in this study.

651

652 Figure 2. STRUCTURE results. On the top the relationship between K (number of inferred clusters) and
653 Ln(K) and DeltaK, respectively. On the bottom, the probability of belonging to each of the two inferred
654 clusters, according to the method by Evanno et al. (2005) for each of the individuals.

655

656 Figure 3. On the left the geographic representation of the connection matrix based on the Delaunay
657 triangulation. On the right the geographic distribution of the first positive sPCA scores.

658 TABLE TITLES

659

660 Table 1. Details of site characteristics of Scots pine populations from Italy which were sampled for the study.
661 Data on annual and summer rainfall, and average annual temperature were inferred for a period of at least
662 five years.

663

664 Table 2. Descriptive statistics for the twelve microsatellite loci considered for the study.

665

666 Table 3. Statistics of genetic variation within Scots pine populations at nine microsatellite loci. A , mean
667 number of alleles per locus; N_e , effective number of alleles per locus; A_{r16} Allelic richness based on a
668 minimum sample size of 16 gene copies; P_a mean number of private alleles per locus; H_o , average observed
669 heterozygosity; H_E , average gene diversity or expected heterozygosity; F_{IS} , average inbreeding coefficient,
670 calculated taking into account the estimated null allele frequencies. Values in parenthesis are standard errors.

Code	Population	Province	Location	Average elevation (m a.s.l.)	Annual (and summer) rainfall (mm)
CRO	Vezzano sul Crostolo	Reggio Emilia	44° 31' N, 10° 31' E	450	945 (174)
CAR	Carpe	Imperia	43° 55' N, 7° 47' E	1000	1100 (260)
BRI	Carnino Briga Alta	Cuneo	44° 18' N, 7° 43' E	1200	1170 (378)
BOS	Bossolasco	Cuneo	44° 32' N, 8° 03' E	750	877 (145)
VEZ	Veza d'Alba	Cuneo	44° 45' N, 8° 00' E	300	660 (125)
FEN	Fenestrelle	Torino	45° 02' N, 7° 03' E	1450	900 (300)
SAV	Savoulx	Torino	45° 05' N, 6° 40' E	1300	396 (102)
PAS	Passerano Marmorito	Asti	45° 05' N, 8° 05' E	225	554 (156)
CAS	Casalborgone	Torino	45° 10' N, 8° 00' E	500	673 (142)
SAR	Sarre	Aosta	45° 43' N, 7° 15' E	1300	698 (174)
TOC	Toceno	Verbania	46° 10' N, 8° 32' E	1100	698 (459)
TIC	Ticino	Novara	45° 30' N, 8° 39' E	250	1020 (234)
OLG	Olgelasca	Como	45° 44' N, 9° 11' E	350	1635 (587)
MAS	Val Masino	Sondrio	46° 09' N, 9° 34' E	300	458 (165)
GAR	Valvestino Garda	Brescia	45° 45' N, 10° 35' E	800	685 (263)
VAL	Valda	Trento	46° 12' N, 11° 16' E	950	750 (350)
SIU	Alpe di Siusi	Bolzano	46° 33' N, 11° 33' E	1200	800 (400)
BRU	Brunico	Bolzano	46° 48' N, 11° 56' E	1100	692 (360)
COR	Cortina	Belluno	46° 32' N, 12° 18' E	1200	1100 (383)
CLA	Claut	Pordenone	46° 16' N, 12° 31' E	600	1186 (436)
DOG	Val Dogna	Udine	46° 26' N, 13° 19' E	1000	908 (365)

Locus	Repeat motif	Primer sequences (5' → 3')	Number of alleles	Molecular weight range (bp)
SPAC 11.4 ^a	(AT) ₅ (GT) ₁₉	TCACAAAACACGTGATTCCACA GAAAATAGCCCTGTGTGAGACA	38	130-170
SPAC 11.5 ^a	(AT) ₈ (GT) ₁₉ -(TA) ₁₁	TGGAGTGGAAAGTTTGAGAAGC TTGGGTTACGATACAGACGATG	no amplification	
SPAC 11.6 ^a	(CA) ₂₉ (TA) ₇	CTTCACAGGACTGATGTTCA TTACAGCGGTTGGTAAATG	76	103-220
SPAC 11.8 ^a	(TG) ₁₆	AGGGAGATCAATAGATCATGG CAGCCAAGACATCAAAAATG	25	123-181
SPAC 12.5 ^a	(GT) ₂₀ (GA) ₁₀	CTTCTTCACTAGTTTCCTTTGG TTGGTTATAGGCATAGATTGC	62	116-202
SPAG 3.7 ^a	(TC) ₄₅	GTTAAAGAAAATAATGACGTCTC AATACATTTACCTAGAATACGTCA	no scorable bands	
SPAG 7.14 ^a	(TG) ₁₇ (AG) ₂₁	TTCGTAGGACTAAAAATGTGTG CAAAGTGGATTTTGACCG	64	174-252
PtTX 3032 ^b	(GAT) ₃₅ (GAC) ₃ GAT(GAC) ₈ -(GAC) ₆ AAT(GAT) ₆	CTGCCACACTACCAACC AACATTAAGATCTCATTTCAA	151	254-572
PtTX 3107 ^c	(CAT) ₁₄	AAACAAGCCCACATCGTCAATC TCCCCTGGATCTGAGGA	22	144-175
PtTX 3116 ^c	(TTG) ₇ -(TTG) ₅	CCTCCCAAAGCCTAAAGAAT CATAACAAGGCCTTATCTTACAGAA	67	100-276
PtTX 4001 ^d	(CA) ₁₅	CTATTTGAGTTAAGAAGGGAGTC CTGTGGGTAGCATCATC	31	197-231
PtTX 4011 ^d	(CA) ₂₀	GGTAACATTGGGAAAACACTCA TTAACCATCTATGCCAATCACTT	21	230-284

675

676 ^a Soranzo et al. 1998

677 ^b Elsik et al. 2000

678 ^c Elsik and Williams 2001

679 ^d Zhou et al. 2002

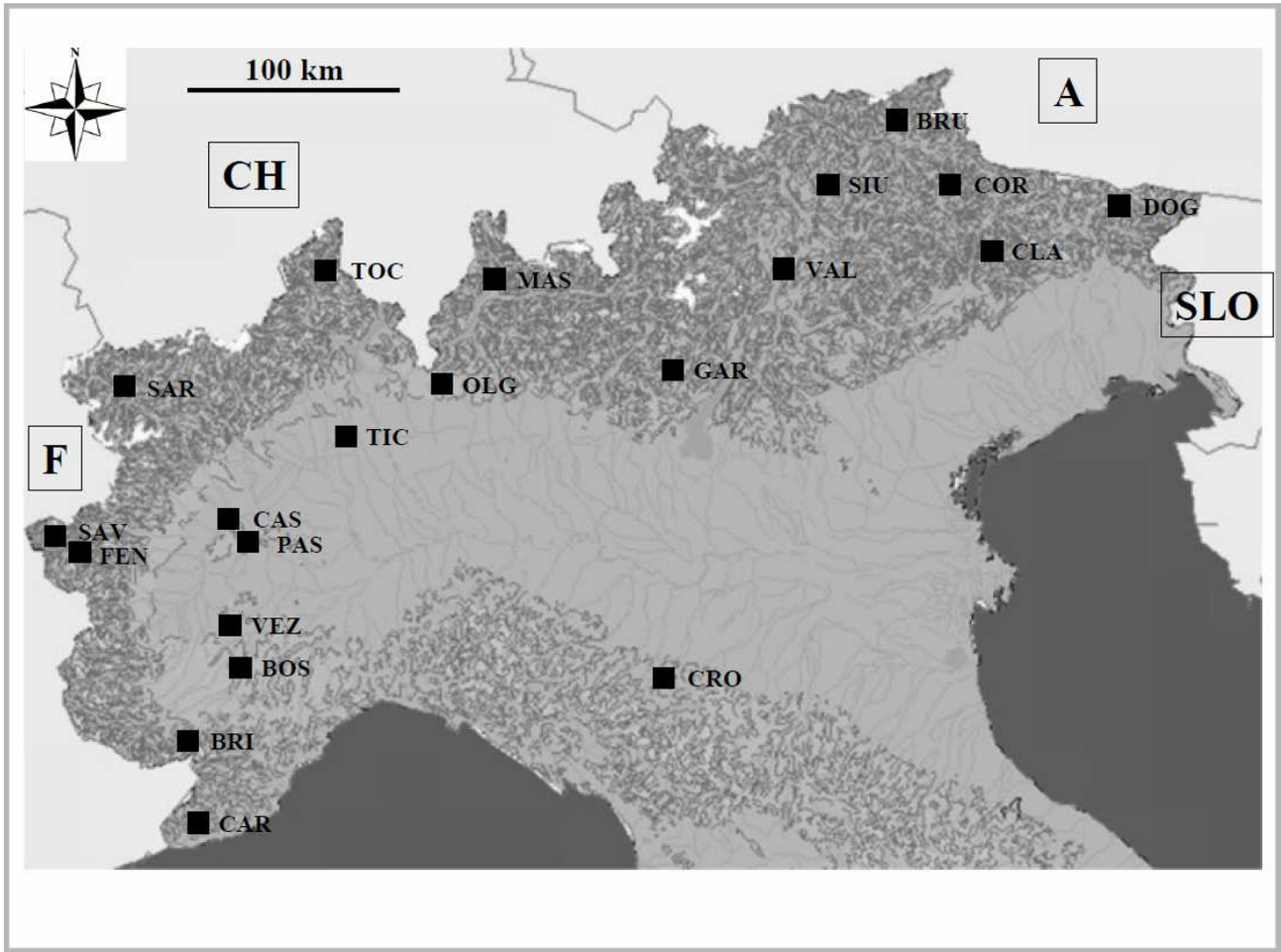
680 Table 3
681

Population	A	N_e	A_{r16}	P_a	H_o	H_e	F_{IS}
CRO	10.4	5.4	6.781	0.333	0.603	0.754	0.037
CAR	15.9	10.4	9.525	1	0.714	0.886	0.030
BRI	12.3	9.3	8.636	0.333	0.651	0.846	0.036
BOS	12.4	8.0	8.353	0.222	0.724	0.829	0.022
VEZ	14.6	9.4	8.939	0.778	0.670	0.852	0.032
FEN	16.0	10.2	9.391	0.556	0.664	0.845	0.052
SAV	18.4	12.2	10.003	1	0.665	0.877	0.023
PAS	14.1	10.1	9.830	1.889	0.576	0.874	0.049
CAS	13.9	8.7	8.601	0.778	0.645	0.858	0.031
SAR	16.4	10.9	9.641	0.889	0.698	0.876	0.029
TOC	17.6	11.0	9.637	1.556	0.641	0.850	0.025
TIC	12.9	6.9	7.713	0.778	0.703	0.822	0.016
OLG	11.2	7.2	8.049	0.444	0.621	0.811	0.047
MAS	16.3	10.6	9.363	0.889	0.703	0.855	0.049
GAR	13.1	8.2	8.398	0.222	0.661	0.808	0.028
VAL	15.3	10.6	9.793	0.333	0.772	0.872	0.022
SIU	15.7	9.7	9.137	0.889	0.676	0.854	0.028
BRU	16.0	9.8	9.326	0.444	0.630	0.865	0.021
COR	14.0	10.0	9.635	1.111	0.724	0.872	0.024
CLA	12.8	8.1	8.771	0.222	0.684	0.828	0.044
DOG	14.0	9.8	9.304	0.556	0.642	0.863	0.042
Overall mean	14.45 (0.46)	9.37 (0.41)	8.992	0.724	0.670 (0.012)	0.847 (0.007)	0.033 (0.001)

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683 Figure 1

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