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1 **Genetic variation in Italian wild cherry (*Prunus avium* L.) as characterised by nSSR**
2 **markers**

3

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8

9 **Abstract**

10 The main aim of our research was to describe the level and distribution of genetic variability of wild
11 cherry (*Prunus avium* L.) in Italy, using eight nuclear microsatellite markers. The sampled plants were
12 grouped in 11 internally homogeneous breeding zones, defined according to their ecological and
13 vegetational conditions. The mean observed heterozygosity (H_o) was 0.573, and the expected (H_e)
14 0.698. Significant departures from Hardy-Weinberg equilibrium at each locus were found for all
15 breeding zones ($P < 0.01$). The mean fixation index, calculated taking into account the estimated null
16 allele frequencies, was 0.075, showing a slight excess of homozygotes. F_{ST} (departure of genotype
17 frequencies within populations from Hardy-Weinberg expectations, commonly used as estimator of
18 genetic differentiation among populations), showed a mean value of 0.046, indicating a slight,
19 although significant, differentiation among breeding zones. However, in general, it was not possible to
20 observe a structuring linked to the geographical location of the breeding zones. The results of the
21 study contribute to a better understanding of our knowledge of the wild cherry genetic variation in
22 Italy, thus making for more efficient programs aimed at the preservation of biodiversity and for more
23 rational planning of the management of reproductive material. Since our results do not show a clear
24 structuring of genetic variability within the Italian diffusion area of wild cherry, it is not possible to
25 draw any indications on Regions of Provenance delimitation based only on genetic data, and the
26 identification of the latter should be based mainly on ecological and vegetational features.

27

28 **Key words**

29 wild cherry, genetic variability, breeding zones, Regions of Provenance.

30

31 **Introduction**

32 Wild cherry (*Prunus avium* L.) is an important forest tree, usually characterised by a scattered
33 distribution. It is used for high quality wood production, and is of great ecological and naturalistic
34 importance, being an essential component of sensitive and threatened ecosystems, such as mixed
35 deciduous forest. The species has hermaphroditic flowers, usually pollinated by a range of insects,
36 mainly honeybees and bumblebees. Outcrossing is the principal mating system, with self-fertilisation
37 usually being prevented by gametophytic incompatibility (Vaughan et al. 2008). Seeds are dispersed
38 by gravity, and birds also play an important role in their dissemination. The species can also propagate
39 through vegetative reproduction via root suckering, involving up to 65% of individuals in the
40 population (Vaughan et al. 2007).

41 The species has a widespread distribution, extending from the Black Sea to Ireland and Spain,
42 and from southern Scandinavia to northern Africa. In Italy it is mainly present in the Po Valley, in the
43 bottom of Alpine valleys and along the Apennines. As we move to the south the species becomes even
44 more and more scattered, and only a few isolated individuals can be found (Ducci 2005). The effects
45 of the past civilisations, as well as the traditional silviculture coppice systems have played a role in
46 determining this situation (Ducci and Proietti 1997).

47 Forest trees are static long-lived organisms which grow under environmental conditions that
48 are heterogeneous in time and space. Moreover, they are exposed to many stress factors, most of
49 which are due to human activities: pollution, climate change and habitat fragmentation. In order to
50 survive these threats, and to persist over time, a high adaptive potential is needed, which is mainly
51 determined by the within-species genetic diversity (Boshier and Amaral, 2004). Programmes aimed at
52 the conservation of forest genetic resources should address the issue of maintenance of this diversity
53 (Palmberg-Lerche 2001). To this end, knowledge of genetic variation, as well as information on
54 mating system and pollen and seed dispersal, are of the utmost importance. These data provide
55 important insights for preservation and restoration programmes, indicating areas of high genetic

56 diversity and geographic limits for seed collection, helping breeders to take decisions about crosses
57 and germplasm management (Marchese et al. 2007), and delimiting the scale at which conservation
58 actions should be planned (Escudero et al. 2003). Furthermore, genetic analysis can increase
59 understanding of the historical processes that led to the present distribution of a species, while the
60 preservation of germplasm is extremely important in order to meet future climatic, abiotic and biotic
61 change (Marchese et al. 2007).

62 Molecular markers are now available which can provide us with the relevant means to acquire
63 information on the genetic structure of populations, and to study the pattern of distribution of within-
64 species variability. In particular, simple sequence repeats (SSRs, also known as microsatellites) are
65 commonly used in genetic studies of plant populations. SSRs are tandem repeats of short DNA
66 sequences (1 to 6 base pairs); they are highly polymorphic, widely distributed throughout the genome
67 and codominant. Allelic variation can be detected quickly using the DNA-polymerase chain reaction
68 (PCR) technique (Beckmann and Soller 1990).

69 Microsatellites have been widely used for genetic studies of cherry, although most studies
70 involved the sweet cultivated forms. Research has included genetic diversity analysis (Dirlewanger et
71 al. 2002, Wünsch and Hormaza 2002, Vaughan and Russell 2004, Ohta et al 2005, Guarino et al. 2009,
72 Laciš et al. 2009, Ercisli et al. 2011), cultivar identification and fingerprinting (Cantini et al. 2001,
73 Schüller et al. 2003, Gisbert et al. 2008, Clarke and Tobutt 2009, Wünsch 2009, Xuan et al. 2009,
74 Avramidou et al. 2010, Ganopoulos et al. 2010, Gulen et al. 2010), self-incompatibility and population
75 genetic structure evaluation and recruitment (Struss et al. 2003, De Cuyper et al. 2005, Kaçar et al.
76 2005, Schüller et al. 2006, Stoeckel et al. 2006, Vaughan et al. 2006, Marchese et al. 2007, Vaughan et
77 al. 2007), and mapping genetic linkage (Olmstead et al. 2008).

78 In Italy the genetic diversity of wild cherry populations is also under threat due to destruction
79 and restriction of habitat, intercross with sweet cherry, pests and diseases, low natural regeneration
80 and competition with other species, deforestation, pollution, and climate change. Genetic diversity in
81 wild cherries has been extensively studied because of its role as a potential gene pool for improving
82 sweet cherry cultivars (Ducci 2005). Research has been conducted to assess genetic diversity and the
83 level of adaptability of different clones (Ducci and Proietti 1997, Santi et al. 1998, Curnel et al. 2003).

84 In this study we surveyed the genetic variability of wild cherry throughout the main area of
85 species distribution in northern and central Italy, using eight highly informative nuclear microsatellite
86 (nSSR) markers. Our main aim was to describe the levels and distribution of genetic variability of the
87 species in Italy, and to investigate the genetic structure of plant groups growing at the southern border
88 of the area where the species is naturally widespread, with particular emphasis on the conservation of
89 the species evolutionary processes and the valorisation of the adaptive potential of its breeding
90 resources. In fact, it is well known that geographically peripheral populations are often genetically
91 differentiated from central populations, due to smaller size, fragmentation and as response to different
92 biotic and abiotic conditions (Lesica and Allendorf 1995). We also discuss our results in the light of
93 the European Directive 105/1999, emphasising the preservation and restoration of wild cherry genetic
94 resources in Italy.

95

96 **Material and Methods**

97 *Plant materials*

98 A total of 278 plants were sampled within the natural range of dispersion of the species in central and
99 northern Italy. They were chosen at random, but in order to minimize the sampling of close relatives a
100 minimum distance among them of about 50 m was adopted. Since in case of wild cherry, due to the
101 low density and scattered distribution of plants, it was not considered valid to define populations *sensu*
102 *stricto*, and the sampled plants were therefore grouped into 11 internally homogeneous “*breeding*
103 *zones*”, defined according to their ecological and vegetational conditions (Figure 1). Each breeding
104 zone was intended to be a separate breeding population, with genetic improvement being limited to
105 these zones to take advantage of naturally evolved adaptation (Silen and Wheat 2003). The boundaries
106 of these breeding zones were based on physiographic, climatic and economic factors. The number of
107 individuals per breeding zone varied from 14 to 39 (Table 2) and no substantial differences in
108 individual density among them were present.

109

110 *Molecular analysis*

111 Young leaves or buds (100 mg of tissue) were powdered in liquid nitrogen, and genomic DNA
112 was extracted using the QIAGEN® DNeasy plant mini kit, according to the manufacturer's protocol. A
113 total concentration of 20 ng μ l⁻¹ was obtained for each sample, and DNA quality was evaluated with an
114 Eppendorf biophotometer.

115 Ten simple sequence repeat markers (SSR) were selected according to the literature, and tested
116 on our plant material (Table 1). While most of the primers were specific for wild cherry (Vaughan and
117 Russell 2004), EMPaO04, EMPaO05 and EMPaO15 were originally designed for sweet cherry
118 (Clarke and Tobutt 2003).

119 Polymerase Chain Reaction (PCR) amplifications were carried out using a DNA Engine (PTC-
120 200) Thermal Cycler, according to the protocols developed by the authors (listed in Table 1). Each
121 amplification reaction contained 1x reaction buffer, 2.5 mM MgCl₂, 0.2 mM dNTPs, 6 μ l Ready Mix
122 Taq, 0.2 μ M of each primer, 4.9 μ l bidistilled water, and 2.5 ng μ l⁻¹ of genomic DNA. Each forward
123 primer was labelled with 5'-fluorescence dye Cy5.

124 The PCR conditions varied for different primers, and were adjusted according to the literature
125 (Vaughan and Russell 2004, Clarke and Tobutt 2003). PCR amplification products were separated in
126 polyacrylamide gel using an ALFexpress II Sequencer (Amersham) laser sequencer. The
127 electrophoresis was carried out, according to the instrument protocol, on 8-10 μ l working solution (3
128 μ l loading dye, 1-2 μ l internal standard sizers (100 - 300), 1-2 μ l external standard sizers 50-500 and 3
129 μ l buffer) added to 2-3 μ l PCR product, under the following conditions: voltage 1500 V, temperature
130 55°C, power 30W, running time 500 min. Fragments sizes were estimated using the Allele Locator
131 software (Amersham Pharmacia Biotech 1999).

132

133 *Data processing*

134 At each locus alleles were characterised exclusively by the length of the DNA fragments
135 generated by the amplification process. The percentage of polymorphic loci (P), mean number of
136 alleles per locus (i. e. present only in one breeding zone, A), effective numbers of alleles (N_e), allele
137 frequencies, mean number of private alleles per locus, observed heterozygosity (H_o), gene diversity
138 (expected heterozygosity, H_e) and the inbreeding coefficient F_{IS} were calculated using EXCEL and the

139 GENALEX 6 software (Peakall and Smouse 2006). The same software was used to compute the
140 population genetic structure of the overall samples for each locus with Wright's F-statistics (Wright
141 1946, 1951, 1965), and to compute a pairwise F_{ST} matrix among breeding zones. For the analyses of
142 departures from Hardy–Weinberg equilibrium (HWE), and to evaluate the significance of inbreeding
143 that occurred in each locus, in each population and in overall loci, we used ARLEQUIN 3.5 (Excoffier
144 et al. 2005, Excoffier and Lisher 2010). The same programme was used to perform tests for genotypic
145 linkage disequilibrium (LD).

146 Where significant deficiencies of heterozygotes from HWE were observed the presence of null
147 alleles was suspected (Pemberton et al 1995). Loci with high frequencies of null alleles were identified
148 by estimating the presence of the latter using the software MICRO-CHECKER (Van Oosterhout et al
149 2004). In subsequent analysis, loci with high null allele frequency were eliminated and only those with
150 a frequency on null alleles lower than 0.19 were used. This value has been considered a threshold over
151 which underestimation of H_e due to null alleles becomes significant (Chapuis et al. 2008). Where
152 possible, analyses with robustness prerogative in presence of null alleles were used (Chapuis and
153 Estoup 2007), including STRUCTURE (Pritchard et al. 2000). The inbreeding coefficients (F_{IS}) were
154 calculated taking into account the estimated null frequencies using the programme INEST and running
155 the individual inbreeding model (IIM) with a Gibbs sampler of 105 iterations (Chybicki and Burczyk
156 2009).

157 Unbiased genetic distances among populations were estimated according to Nei (1978), using
158 GENALEX. This matrix constituted the base for investigating the relationships among groups, by an
159 unweighted pair group method using the arithmetic means (UPGMA) dendrogram developed by
160 SAHN software in the NTSYSpc package (Rohlf 2005). The cophenetic values matrix was calculated
161 from the tree matrix using the program COPH. The cophenetic matrix was used to evaluate goodness
162 of fit for the cluster analysis by comparing it to genetic distances matrix (MXCOMP of NTSYS).

163 In addition, the genetic structure was analysed using a Bayesian clustering approach with
164 software STRUCTURE version 2.2.3 (Pritchard et al. 2000). The data was explored without
165 considering prior classification within the sample, by performing 10 replicates of each simulation from
166 $K = 2$ to $K = 12$, with a burn-in of 10,000 steps followed by 10,000 Markov chain Monte Carlo

167 (MCMC) iterations under the admixture model, and the assumption of correlated allele frequencies
168 among populations. We used the program Structure Harvester (Earl and von Holdt 2012) that applies
169 the Evanno's method to assess the optimal level of K (Evanno et al. 2005).

170

171 **Results**

172 All 10 primer pairs analysed produced polymorphic amplification products, which could be
173 easily identified (Table 1). However, two markers were excluded from the analysis due to the high
174 frequency of possible null alleles: respectively 0.23 (EMPaS11) and 0.20 (EMPaO15). The presence
175 of null alleles was also suspected for 3 out of the remaining 8 loci, with frequencies of 0.10
176 (EMPaS06), 0.12 (EMPaS10) and 0.13 (EMPaS12). In total, 124 distinct alleles were scored in the
177 278 studied individuals. The number of alleles per locus ranged from 4 (EMPaS14) to 25 (EMPaS10),
178 with an average value of 15.5. It was possible to detect 24 private alleles. The frequency of these
179 alleles was always low, ranging from 0.013 to 0.048. The distribution of private alleles among
180 microsatellites was unbiased, ranging from one (EMPaO04, EMPaS02 and EMPaS14) to six
181 (EMPaO05). The breeding zone which displayed the highest number of such alleles was TSE (8
182 private alleles), while no private alleles were detected in EMR.

183 Sampling of individuals belonging to the same clone could be excluded since plants with
184 identical genotypes were never detected. Statistics on the genetic diversity within breeding zones are
185 given in Table 2. A relatively high level of intra-breeding zones variability was found, since on
186 average more than 7 alleles per locus were observed ($A = 7.53$). The probability that two randomly
187 sampled alleles in a given breeding zone were not the same was almost 70% ($H_e = 0.698$), whereas the
188 observed heterozygosity was lower than expected (mean $H_o = 0.573$). Since the difference, that
189 determines a significant positive value for mean inbreeding coefficient, could be due to the presence of
190 null alleles the inbreeding coefficients were recalculated. Taking into account the frequencies of null
191 alleles, we found that deviations from the Hardy-Weinberg equilibrium were low (F_{IS} ranging from
192 0.031 in FON to 0.220 in HPD with a mean of 0.075). Significant per locus departure from Hardy-
193 Weinberg equilibrium were however found in all breeding zones ($P < 0.01$).

194 The genetic divergence between breeding zones was investigated by computing a Nei's
195 genetic distance matrix (Nei 1978), and by a pairwise F_{ST} matrix (Table 3). Most of the diversity was
196 found within breeding zones, with only a small amount among breeding zones: the average F_{ST} value
197 was 0.046 (SE = 0.003) and ranged from 0.034 (EMPaS14) to 0.059 (EMPaS12). Almost all pairwise
198 F_{ST} values were significantly greater than zero. The Nei's distances values varied between 0.053 (TSE
199 and TSW) and 0.223 (VNE and FVG). The UPGMA dendrogram confirmed the presence of
200 differentiation between breeding zones, although only in rare cases it was possible to observe a
201 structuring linked to their geographical location. The cophenetic correlation indicated a fair fit of the
202 cluster analysis to data ($r = 0.605$, $P < 0.01$). In particular, VNE and FVG, which are geographically
203 close, were identical; another group consisted of FON and LOM, which are located in the same
204 Region, while PDA showed the highest amount of genetic differentiation (Figure 2).

205 Following the method of Evanno et al. (2005), the Bayesian clustering results obtained with
206 STRUCTURE indicate that $K=11$ clusters represents the most likely representation of the overall
207 genetic structure that we analysed (Figure 3). Breeding zones showed a high level of admixture and no
208 general trends were detected.

209

210 **Discussion**

211 The main aim of our research was to describe the level and distribution of genetic variability
212 of wild cherry in northern and central Italy, which, together with the southern part of the country
213 (where the presence of the specie is extremely scattered) represents the southern border of the species
214 diffusion area.

215 In general, our values are higher than those found in studies of sweet cherry: this most likely
216 reflects the fact that the wild cherry conserves a wider basis of genetic diversity compared to the
217 cultivated forms. The number of alleles in this study (range 4-25 per locus) is high, when compared
218 with data reported for other wild cherry populations: Vaughan et al. (2007) detected a range of 4–14
219 alleles ($N_a = 7.77$) in 551 plants, Stoeckel et al. (2006) detected 4–21 alleles in 350 plants (among
220 which, 247 were from the same population), while Guarino et al. (2009) observed a number of alleles
221 ranging between 2 and 14 in 50 plants from three different countries. Our results are consistent with

222 the fact that Italy was probably a centre of dispersion of wild cherry in Europe after the last glacial
223 period, where a higher number of alleles were conserved than in other countries. The size ranges of
224 alleles overlapped with reported values, when compared with results of previous works (Clarke and
225 Tobutt 2003, Vaughan and Russell 2004, Guarino et al. 2009).

226 We found neither low differences between H_e and H_o , nor the excess of heterozygotes that was
227 reported in previous research (for sweet cherry see Dirlewanger et al. 2002, Wünsch and Hormaza
228 2002, Marchese et al. 2007, Ganopoulos et al. 2010; for wild and sweet cherry see Guarino et al. 2009;
229 for wild cherry see Clarke and Tobutt 2003, Schüller et al. 2003, Vaughan and Russell 2004, Stoeckel
230 et al. 2006, Avramidou et al. 2010). On the contrary, our F_{IS} always gave positive values, indicating an
231 excess of homozygotes. This could be due to the presence of null alleles for some of SSRs markers
232 that increase the number of homozygotes. For instance, in analysis of a pool of 50 wild cherry trees,
233 compared with near 80 sweet cherry cultivars, Guarino et al. (2009) found different frequencies of null
234 alleles, ranging from 0.011 (EMPaS01) to 0.327 (EMPaS02). However, our F_{IS} values remained
235 positive notwithstanding the correction applied by MICRO-CHECKER for null alleles and the
236 consequent exclusion from the analysis of two markers where the presence of null alleles exceeded the
237 threshold commonly adopted in literature.

238 To explain the positive value of inbreeding coefficient, it is also possible to assume the
239 presence of a Wahlund effect, that is the structuring of the populations in subunits within which
240 mating is more probable. This does not seem to be the case of our sampling, since we collected
241 material from plants not closer than 50 m from each other and sampling density was homogeneous
242 within each breeding zone.

243 Nevertheless, to limit the impact of the number of homozygotes on the interpretation of
244 inbreeding indices, it is important to notice that an excess of homozygotes was also found for Italian
245 populations with isozymes (Ducci 2005), where H_o and H_e values were lower than those found in the
246 present study, but F_{IS} values were relatively similar in size. In our work, the higher values of F_{IS} were
247 found in HPD, LOM, TSW and VNW. Some of these areas are producers of sweet cherry in Northern
248 Italy, and it is possible that some of our samples consisted of individuals derived from seed produced
249 by cultivated trees widespread in the wild or as results of intercross between wild and sweet forms.

250 This level of inbreeding contrasts with the fact that cherry is a species characterised by a well-known
251 self-incompatibility system that prevents self-fertilisation (Sonnenveld et al. 2003, Granger 2004,
252 Schüler et al. 2006, Sonnenveld et al. 2006, Marchese et al. 2007, Vaughan et al. 2008). The system is
253 based upon a single multiallelic locus with gametophytic action that controls the ribonucleases in the
254 style of flowers and arriving pollen grains. When the allele of the haploid genotype of the pollen is
255 identical to one of the diploid genotype of the style, fertilisation does not occur, while crosses between
256 individuals sharing only one S-allele can occur, although seed production is limited. Many studies
257 have paid attention to these kind of alleles, that are particularly important in cultivation of sweet
258 cherry (Wiersma et al. 2001, Sonnenveld et al. 2006), but also for the maintenance of wild cherry in
259 the wild (De Cuyper et al. 2005, Vaughan et al. 2006). Since individual trees are often widely
260 scattered, the number and proximity of compatible pollen donors have a large impact on seed set and
261 the genetic diversity of the progeny.

262 However, self-incompatibility can be overcome under particular conditions, for instance when a
263 population goes through a bottleneck (Reinartz and Les 1994, Gigord et al. 1998).

264 Another explanation is suggested by the evidence that in natural populations of *Prunus avium*
265 individuals with a high suckering ability can produce more gametes, thereby having a higher
266 probability of undergoing somatic mutations at the gametophytic incompatibility S-locus, as already
267 observed by Lewis and Crowe (1954).

268 Self-incompatibility could be bypassed by other circumstances. In *Sorbus domestica* L., a
269 predominantly self-incompatible species belonging to the *Rosaceae* family, Kamm et al. (2011) found
270 that self-fertilisation is probably due to a break-down of the incompatibility system occurring as
271 flowers grow old without being pollinated, or under cold weather conditions, while there was no
272 selfing occurred with controlled pollinations of newly opened flowers. Holderegger et al. (2008),
273 studying 15 populations in Switzerland of another self-incompatible *Rosaceae* (*Pyrus pyraster*
274 Burgsd.), with both S-alleles and nuclear SSRs, found positive values of F_{IS} (0.021-0.164). They
275 explained this fact by the spatially restricted pollen dispersal that can cause the production of seeds
276 from pollination by a limited number of compatible trees in the neighbourhood. The genetically
277 related offspring could therefore produce a progeny with elevated F_{IS} index. The questions posed by our

278 samples can only be answered by S-allele analysis in the future. Reduced heterozygosity at nuclear
279 microsatellite loci, representative of whole-genome inbreeding, is likely to be associated with reduced
280 individual fitness and impaired potential of the population to adapt to a changing environment, and it
281 is therefore important to better identify the way in which our populations share genetic variability.

282 Population structure in wild cherry has been characterised by relatively few studies: many
283 researches having focused on the cultivated sweet cherry, employing many different markers (Granger
284 et al. 1993, Gerlach and Stösser 1997, Struss et al. 2001, Wünsch and Hormaza 2002, Xhou et al.
285 2002, Struss et al. 2003). Studies with chloroplast DNA (Mohanty et al. 2001a, 2001b) found a low
286 level of differentiation among populations spread in different European regions, and Panda et al.
287 (2003) found a higher cpDNA diversity in wild cherry with respect to a group of common sweet
288 cultivars.

289 In Italy, genetic differentiation measured among the eleven zones was relatively high, and
290 only a small number of alleles per locus are in common among populations (only 1 allele for
291 EMPaS12; 2 alleles for EMPaS14; 3 alleles for EMPaS01, S06, S10, O04, O05; 4 alleles for
292 EMPaS02). We found a $F_{ST} = 0.046$, comparable with other studies developed in France (Frascaria et
293 al. 1993, Mariette et al. 1997), where G_{ST} values among populations were 0.05 and 0.06 respectively,
294 or in Georgia ($F_{ST} = 0.057$) (Santi and Dufour 2010).

295 The value of differentiation is compatible with the reproductive system. Wild cherry plants are
296 insect pollinated, especially by bees, which use the nectar and pollen as an important nutrient source.
297 Pollen can be transferred by various modes, resulting in largely variable dispersal distances. “*Short*
298 *pollen dispersal*” is generally the most common, due to insect-mediated “*tree-to-tree*” transfer, that
299 tends to cause a differentiation in plants populations. Larger distances of transfer could potentially be
300 realised by other processes, such as “*insect-to-insect*” transfer within the colonies of social insects.
301 When a colony uses different nutrient sources it can accumulate a vast number of pollen mixtures,
302 from wide surrounding areas, in its hive (Free and Williams 1972, De Grandi-Hoffman et al. 1984,
303 1986). However, at lower frequencies, long distance foraging has also been observed (Zurbuchen et al.
304 2010). As a typical pioneer tree species wild cherry colonises early forest successional stages as a
305 result of forest disturbances, but it is generally replaced by climax tree species during subsequent

306 succession of the local disturbance, so it is difficult to identify pure, extensive and continuous
307 populations, where insects could share pollen exchange. In the Mediterranean environment we find
308 small groups, and their suckers, or isolated trees, so pollen transfer among distant groups become more
309 difficult, since it is known that smaller or less dense populations attract fewer pollinators.

310 From UPGMA we observe a general lack of geographic pattern. This could also be due to
311 different management strategies of wild cherry within the considered area. For instance, in the eastern
312 regions wild cherry is traditionally cultivated for wood production, and populations are larger and
313 widespread in larger areas, but there is also an extensive cultivation of sweet cherry for fruit
314 production. “Bosco Fontana”, in Lombardy, is a particular situation. It’s a wood situated in a
315 biogenetic reserve of 235 ha, established during the 70’s of the 20th century on an ancient hunting
316 reserve, which preserves a relict of the ancient plain forest in the Po Valley, rich in woody species and
317 managed by the local “National Forest Service”, and where coppicing has been avoided for 60 years.
318 In the Apennines there are small populations, isolated in the valley bottoms, with few individuals, very
319 distant from each other. Furthermore, the silvicultural practices that privilege the vegetative
320 propagation contributes to shaping the genetic variation in these zones, and generally wild cherry has
321 been managed as other species, with coppicing.

322 At the individual level, STRUCTURE analysis using Evanno’s method suggests that wild
323 cherry is constituted from such a high number of groups, that the entire studied gene pool can be
324 considered as an unstructured population. This result is not so far from that obtained by Mariette et al.
325 (2010) in a collection of 211 wild cherry trees sampled in France, and also to those revealed in Italian
326 populations of walnut (*Juglans regia* L.) (Ferrazzini et al. 2007b), that share with wild cherry the
327 anthropic effect due to cultivation for wood and fruit production.

328 Jordano et al. (2007) found that in *Prunus mahaleb* L. seed dispersal due to different types of
329 frugivores resulted in distinct contributions of different distance classes and microhabitats, with only a
330 few species responsible for long-distance dispersal events. Small-sized birds accounted for most short-
331 distance dispersal, and larger frugivores (both birds and mammals) accounted for most long-distance
332 dispersal. They pointed out that three components are particularly important when we want to
333 understand genetic distribution by seed dispersal: the frugivore abundance, their feeding and post-

334 feeding behaviour and the structure of landscape. While these factors could partially explain the lower
335 genetic distance among VNE and FVG as well as among FON and LOM, that are relatively close from
336 the geographic point of view (with distances compatible with the animal impact on seed dispersal), the
337 human influence on wild cherry distribution seems to be one of the most important causes of the lack
338 of a clear pattern in general distribution of the genetic diversity. As for most European forests, the
339 studied areas have experienced human activities and management for generations. The traditional
340 management method (coppice-with-standards) has probably favoured the maintenance of clonal
341 lineages through time, and due to the edible flesh of fruits, long distance seed dispersal by humans
342 appears as a source of gene flow among populations.

343 Foresters began to use wild cherry for reforestation and afforestation in the 1970s, usually
344 with material of unknown origin. Frequently, this material originated not only from wild cherry, but
345 also from sweet cherry or even from sour cherry. For this reason, it is not so unlikely that many
346 individuals originated from the introduction of domesticated trees into the wild woods, or from the
347 intercross among wild and sweet forms. In Italy, since 1988, reproductive material for plantations has
348 been collected from physiographic seed areas (Ducci et al. 1988).

349

350 **Conclusions**

351 The results of the study contribute to a better understanding of our knowledge of wild cherry
352 genetic variation in Italy, thus making for more efficient programs aimed at the preservation of the
353 biodiversity. Furthermore, our results give useful indications on how to act for more rational planning
354 of the management of reproductive material. In the light of the European Council Directive 105/1999,
355 forest reproductive material falling into the “*source-identified*” and “*selected*” categories should be
356 used only within the Region of Provenance where it originated. The transfer of material from one
357 Region to another should be avoided, due to possible problems associated with adaptability. The
358 identification of Regions of Provenance is therefore a basic aspect for a rational management of
359 activities linked with forest tree propagation, including afforestation and *in situ* genetic preservation.
360 In the case of wild cherry, in Italy, we cannot describe populations *sensu stricto*, but use breeding
361 zones, where trees grow with a scattered dispersion. Since our results do not show a clear structuring

362 of genetic variability within the Italian diffusion area of wild cherry, it is not possible to draw any
363 indications on Regions of Provenance delimitation based only on genetic data, contrary to the results
364 of similar studies carried on in Italy on different species (common ash and Scots pine), where genetic
365 data proved to be useful for Regions of Provenance identification (Ferrazzini et al. 2007a, Belletti et
366 al. 2012). Therefore, the identification of Regions of Provenance for wild cherry in Italy should be
367 based mainly on ecological and vegetation features. Results similar to ours were obtained also in
368 France, where it has been suggested to consider a single Region of Provenance (Anonymous 2003).
369 Furthermore, the development of seed orchards where it should be possible to collect clones from
370 different sites should be encouraged, since they would allow the preservation of as much as possible of
371 the scattered genetic diversity of wild cherry, and to simplify the operation of finding reproductive
372 material for forest breeders.

373

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378

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581 **Figure legends**

- 582 • Figure 1. Geographical location of the breeding zones where individuals of wild cherry analysed
583 in this study were sampled.
- 584 • Figure 2. Dendrogram constructed from UPGMA cluster analysis of 11 breeding zones of wild
585 *Prunus avium* in northern and central Italy, based on Nei's genetic distance for SSRs produced by
586 eight primers.
- 587 • Figure 3. Probability of assignment of 278 plants of wild cherry from Italy to the three genetic
588 clusters identified by hierarchical STRUCTURE analysis. Each vertical bar corresponds with a
589 distinct genotype and different colours indicate the part of its genome assigned to each cluster.

590

591 **Table titles**

- 592 • Table 1. Allelic diversity of the ten microsatellite loci considered for the study.
- 593 • Table 2. Statistics of genetic variation within Italian wild cherry at eight microsatellite loci. N,
594 sample size; A, mean number of alleles per locus; N_e , effective number of alleles per locus; P_a
595 mean number of private alleles per locus; H_o , average observed heterozygosity; H_e , average gene
596 diversity or expected heterozygosity; F_{IS} , average inbreeding coefficient calculated taking into
597 account the estimated null allele frequencies. Values in parenthesis are standard errors.
- 598 • Table 3. Pairwise F_{ST} (above the diagonal) and genetic distances (according to Nei (1978), below
599 the diagonal) among the 11 breeding zones considered in the study.

600 Table 1

Locus	Reference	Number of alleles	Molecular weight range (bp)
EMPaS01	Vaughan and Russell 2004	8	225-254
EMPaS02	Vaughan and Russell 2004	16	133-148
EMPaS06	Vaughan and Russell 2004	24	200-230
EMPaS10	Vaughan and Russell 2004	25	151-185
EMPaS11*	Vaughan and Russell 2004	11	81-109
EMPaS12	Vaughan and Russell 2004	23	121-152
EMPaS14	Vaughan and Russell 2004	4	197-213
EMPaO04	Clarke and Tobutt, 2003	9	177-195
EMPaO05	Clarke and Tobutt, 2003	15	230-262
EMPaO15*	Clarke and Tobutt, 2003	24	202-300

601

602 * markers excluded from the analysis due to the high frequency of possible null alleles.

603 Table 2

Breeding zone	Code	N	A	N_e	P_a	H_o	H_e	F_{IS}
Inner hills of Piedmont	HPD	27	8.5	5.2	0.8	0.490	0.734	0.229
Western Alps of Piedmont	PDA	22	7.1	4.6	0.4	0.523	0.687	0.075
Fontana Forest	FON	23	6.4	3.8	0.1	0.596	0.666	0.031
Prealps of Lombardy	LOM	30	9.1	5.1	0.9	0.540	0.687	0.111
Eastern Veneto	VNE	22	7.1	4.4	0.1	0.585	0.730	0.033
Western Veneto	VNW	37	8.4	4.7	0.3	0.560	0.718	0.085
Friuli Venetia Giulia	FVG	14	6.3	3.9	0.4	0.644	0.660	0.042
Emilia Romagna	EMR	16	6.3	4.2	0.0	0.617	0.683	0.034
Western Tuscany	TSW	39	8.8	4.7	0.8	0.583	0.731	0.103
Eastern Tuscany	TSE	31	9.1	5.0	1.0	0.602	0.733	0.041
Marche	MAR	15	5.9	3.8	0.3	0.558	0.648	0.041
Overall mean		24.9	7.53 (0.410)	4.48 (0.259)	0.44	0.573 (0.019)	0.698 (0.019)	0.075 (0.047)

604 Table 3

	HPD	PDA	FON	LOM	VNE	VNW	FVG	EMR	TSW	TSE	MAR
HPD	0.000	0.019	0.028	0.023	0.022	0.025	0.025	0.020	0.021	0.029	0.035
PDA	0.105	0.000	0.023	0.021	0.025	0.021	0.031	0.020	0.013	0.015	0.018
FON	0.136	0.108	0.000	0.039	0.026	0.025	0.035	0.038	0.026	0.024	0.027
LOM	0.125	0.110	0.187	0.000	0.032	0.023	0.035	0.019	0.024	0.028	0.030
VNE	0.136	0.132	0.127	0.174	0.000	0.021	0.042	0.019	0.018	0.022	0.035
VNW	0.140	0.113	0.128	0.118	0.120	0.000	0.036	0.020	0.020	0.016	0.024
FVG	0.126	0.166	0.170	0.176	0.223	0.187	0.000	0.036	0.039	0.040	0.039
EMR	0.117	0.104	0.182	0.098	0.088	0.097	0.192	0.000	0.021	0.029	0.031
TSW	0.124	0.065	0.124	0.135	0.109	0.115	0.205	0.104	0.000	0.010	0.020
TSE	0.162	0.067	0.113	0.148	0.129	0.092	0.202	0.136	0.053	0.000	0.022
MAR	0.180	0.084	0.126	0.151	0.168	0.116	0.194	0.142	0.088	0.094	0.000

605