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# Ascoma genotyping and mating type analyses of mycorrhizas and soil mycelia of Tuber borchii in a truffle orchard established by mycelial inoculated plants

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1	Ascoma genotyping and mating type analyses of mycorrhizas and soil mycelia of
2	Tuber borchii in a truffle orchard established by mycelial inoculated plants
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#### 15 Significance/originality statement

16 The maternal and paternal genotypes in the first truffle orchard realized with plants inoculated with 17 the mycelia of the heterothallic fungus *T. borchii* were analyzed. The inoculated mycelia persisted in 18 soil 9 years after plantation and the paternal genotypes were mostly recombinants of the inoculated 19 mycelia, corresponding to their offspring. The study confirms that meiospores are the most likely 20 paternal candidate for truffle fertilization and improves our knowledge of sexual truffle reproduction.

#### 21 Abstract

22 *Tuber borchii* (the Bianchetto truffle) is a heterothallic Ascomycete living in symbiotic association 23 with trees and shrubs. Maternal and paternal genotype dynamics have already been studied for the black truffles Tuber melanosporum and Tuber aestivum but not yet for T. borchii. In this study we 24 25 analyzed maternal and paternal genotypes in the first truffle orchard realized with plants inoculated 26 with five different T. borchii mycelia. Our aims were to test the persistence of the inoculated mycelia, 27 if maternal and/or paternal genotypes correspond to inoculated mycelia and to assess the 28 hermaphroditism of *T. borchii*. The mating type of each isolate as well as those of mycorrhizas, 29 ascomata and extraradical soil mycelia was determined. Moreover, simple sequence repeat (SSR) 30 profiles of maternal and paternal genotypes were assessed in 18 fruiting bodies to investigate the 31 sexual behavior of this truffle.

The maternal genotypes of the fruiting bodies corresponded to those of the inoculated mycelia with only two exceptions. This confirmed that the inoculated mycelia persisted 9 years after plantation. As regards paternal partner, only two had the same genotype as those of the inoculated mycelia, suggesting hermaphroditism. Most of the new paternal genotypes originated from a recombination of those of inoculated mycelia.

37 Key words: mating type, SSR, truffle reproduction, meiospores, mycelial cultures

38

# 39 Introduction

40 Ectomycorrhizal fungi assist plants in their growth, therefore playing key roles in forest ecosystem 41 functioning. In addition, some of them produce edible fructifications representing income 42 opportunities for farmers and foresters. True truffles are hypogeous fungi belonging to the genus 43 *Tuber*, which live in ectomycorrhizal association with a wide range of shrubs and trees (Zambonelli 44 et al., 2016). The genus Tuber comprises around 200 species but only a few of them have a 45 considerable value, such as the European species *Tuber magnatum* Picco (Italian white truffle), *Tuber* 46 melanosporum Vittad. (black truffle), Tuber aestivum Vittad. (summer truffle) and Tuber borchii 47 Vittad. (bianchetto truffle) (Bonito et al., 2010). In recent years, important milestones have been 48 reached, allowing a better understanding of the truffle life cycle. In 2006, Paolocci and colleagues 49 found that T. magnatum is heterothallic and its ectomycorrhizas are formed by primary 50 (homokaryotic) mycelia. This condition was also confirmed in T. melanosporum (Riccioni et al., 51 2008; Rubini et al., 2011a) and T. borchii (Belfiori et al., 2016), indicating that heterothallic life style 52 is common within the *Tuber* genus and fruiting body production depends on the mating between 53 mycelia harboring different mating types. Mycelia of both mating types can act as maternal partner, 54 indicating hermaphroditism (Selosse et al., 2017). The haploid maternal genotype of truffles forms 55 the sterile tissues of the fruiting body, and it is fed from the surrounding mycorhizas throughout the maturation time (Rubini et al., 2011a; Murat et al., 2013; Le Tacon et al., 2013). On the contrary, the 56 57 paternal genes are confined to the asci and, indeed, the paternal genotypes were rarely found to form 58 mycorrhizas (Taschen et al., 2016; De la Varga et al., 2017).

In addition to the characterization of mating type idiomorphs, the sequencing of the *T. melanosporum* genome (Martin *et al.*, 2010) has allowed large-scale screening of SSR markers which are being implemented on a small scale in field trials in order to unravel the interaction between maternal and paternal genotypes. In the field, *T. melanosporum* displays a strong genetic isolation with strains of opposite mating types spatially confined in separate patches in which a few dominant genotypes can persist throughout the years (Rubini *et al.*, 2011b; Taschen *et al.*, 2016; De la Varga *et al.*, 2017).
Due to the important turnover and small spatial size of the paternal genotypes, it has been suggested
that in most cases germinating ascospores could act as paternal partner (Selosse *et al.*, 2013; Le Tacon *et al.*, 2016; Taschen *et al.*, 2016; De la Varga *et al.*, 2017; Selosse *et al.*, 2017). However, this
hypothesis remains to be demonstrated and the nature of the paternal genotype is still unclear.

69 Tuber borchii was long used as a model species within the Tuber genus for transcriptomic and 70 functional analyses. *Tuber borchii*, together with *T. aestivum*, is the species with the widest diffusion 71 in Europe, adapting to different climatic and soil conditions (Hall et al., 2007; Zambonelli et al., 72 2002; Gardin, 2005; Lancellotti *et al.*, 2016). For its gastronomic value and adaptability to different 73 environmental conditions, T. borchii cultivation was introduced in European Mediterranean countries 74 (Italy, Portugal and Spain) and in non-European countries (Hall et al., 2017; Zambonelli et al., 2015). 75 The cultivation of T. borchii, similarly to that of other truffles, is achieved by synthetizing 76 mycorrhizal plants in specialized nurseries by spore inoculum and planting them in suitable places 77 (Chevalier and Grente, 1978; Hall et al., 2007). Mycelial inoculum was successfully tested in the 78 second half of the last century to obtain Tuber mycorrhizal plants (Palenzona et al., 1972; Chevalier, 79 1973) but, later, it was applied only for experimental purposes due to the difficulties of obtaining 80 large quantities of mycelium for large-scale inoculation purposes (Iotti et al., 2002; Giomaro et al., 81 2005; Iotti et al., 2012a). Also, once reliable methods for growing T. borchii mycelium had been 82 perfected, mycelial inoculation was no longer used because it was commonly thought that the 83 obtained plants would not have been able to produce fruiting bodies due to heterothallic sexual 84 reproduction of truffles (Zambonelli et al., 2008).

85

Recently, the first truffle orchard realized with plants inoculated by *T. borchii* mycelia began to produce (Iotti *et al.*, 2016). This truffle orchard was established in Cadriano (Bologna) inside an intensive agricultural area where *T. borchii* had not been found for at least 30 years (Zambonelli and Morara, 1984; Morara *et al.*, 2009). Moreover, the presence of large mycophagus mammals like wild boars, which could introduce truffle spores in the orchard from far sites (Piattoni *et al.*, 2014), was excluded. The plants were inoculated with five different strains singly and in mixture. Truffle production was first assessed in February 2016, 9 years after planting. We hypothesized that the strains used for inoculation had different mating types and that fertilization had occurred between them (Iotti *et al.*, 2016). This plantation represents a unique opportunity to investigate *T. borchii* sexual reproduction.

96 The aims of this study were to address the following questions: 1) were mycelium genotypes able to 97 be perennial for 9 years in the root system? 2) did maternal genotypes correspond to mycelium used 98 for plant inoculation? 3) did paternal genotypes correspond to offspring of original genotypes or was 99 hermaphroditism more frequent in T. borchii than T. melanosporum? In order to address these 100 questions, polymorphic simple sequence repeat (SSR) markers were identified in the T. borchii 101 genome (Murat et al., 2018). These SSRs were applied to genotype the inoculated strains used for 102 plant colonization, as well as maternal and paternal genotypes of harvested ascomata. Mating type 103 distribution in the orchard was also determined for inoculated mycelia, mycorrhizas, ascomata and 104 soil samples.

# 105 **Results**

# 106 SSR characterization

More than one thousand SSRs (1,111 in total) were found in the *T. borchii* genome. Among the 31 SSR primer pairs selected *in silico* analysis, 14 were used in subsequent analyses since produced a single band per sample and showed polymorphisms. They were used to analyze the polymorphism of 50 ascomata harvested in Italy and Hungary as well as the samples from the Cadriano truffle orchards. MLGsim analysis showed that all the 50 ascomata from Italy and Hungary were different from each other, resulting in 50 multilocus genotypes (MLGs) (genotypic diversity = 1; Supplementary Table S1). The number of alleles ranged from 3 to 10 and the He from 0.208 to 0.739 (Table 1 and 114 Supplementary Table S1). Most of the SSRs are localized in intergenic regions with the exception of

115 Tb151, Tb293, and Tb46 that are localized in gene models.

116 Table 1

117 Genotyping of the inoculated mycelia

The SSRs and mating type analyses made it possible to genotype the five *T. borchii* strains used to inoculate truffle seedlings. The strain 2364 was assigned to C-MLG\_13, the strain Tb98 to C-MLG\_8, the strain 1Bo to C-MLG\_6, the strain 2292 to C-MLG\_18 and the strain 2352 to the C-MLG\_14. Two strains (2292 and 1Bo) have the MAT 1-2-1 gene coding for the HMG-domain protein and the other three strains (Tb98, 2352 and 2364) have the MAT 1-1-1 gene coding for the alpha-domain

protein (Table 2). Although the arrangement of the groups of plants inoculated with the same strain(s)in four rows along the plantation was casual at the time of plantation, the distribution of the two

125 mating types proved to be spatially alternated across the truffle ground (Fig. 1).

126 Figs 1 and 2

127 Genotyping of the maternal tissue (gleba) and mating characterization of ECM and soil mycelium

The maternal tissue of 18 ascomata out of 33 collected in the truffle orchard was successfully genotyped using both SSR and mating type primers (Table 2 and 3). At least two ascomata for each plant group inoculated with the same strain were genotyped. The other 15 ascomata were only mating typed and not considered for SSR analyses because the respective paternal genotypes failed to be characterized (Table 2).

133 Table 2 and 3

134 Amplifications of the maternal tissue with both mating type and SSR primers gave single and specific

amplicons. The 18 ascomata were grouped into 6 different genotypes (C-MLGs): 4 with MAT 1-1-1

136 (C-MLG\_8, C-MLG\_13, C-MLG\_10, C-MLG\_14) and 2 with MAT 1-2-1 (C-MLG\_3, C-MLG\_18)

137 (Table 2 and Supplementary Table S2). Most of the analyzed ascomata (15/18) have the same C-

138 MLG of the strains used to inoculate the plants where they were collected (C-MLG\_8, C-MLG\_13,

139 C-MLG\_18, C-MLG\_14). Three female genotypes (C-MLG\_8, C-MLG\_13, C-MLG\_14) showed a

140 significant P sex value that supports the hypothesis that the ascomata of each of these genotypes are 141 part of the same clone (Supplementary Table S3). One ascoma (4658), found in the group of plants 142 inoculated with the strain 1Bo (C-MLG 6) had the same genotype (C-MLG 14) of the neighboring 143 strain 2352 (about 5 m far) (Figs. 1 and 2). Only two ascomata (4601, 4633) had female genotypes (C-MLG 3 and C-MLG 10) different from those used to inoculate the plants (Table 2). In particular, 144 C-MLG 10 showed a SSR profile which differs from that of the inoculated strain 2364 (C-MLG 13) 145 146 for the presence at locus Tb1 of a different allele which is present in the strains Tb98 and 2292. On 147 the contrary, C-MLG 3 had two loci (Tb151 and Tb46) with alleles not present in any inoculated 148 strain (Supplementary Table S1). The genotype C-MLG 6 corresponding to the inoculated strain 1Bo 149 was never found as maternal genotype in the truffle ground. 150 The ascomata found in the plants inoculated with a mixture of strains showed the maternal genotypes

151 C-MLG\_8 and C-MLG\_14, corresponding to the strains Tb98 and 2352, respectively.

The persistence of the inoculated strains in the areas where the plants were originally planted was also confirmed by mating type analyses of the other 15 ascomata. All these ascomata had the same mating type as the strain used to inoculate the plants where they were collected (Table 2). Nine samples were identified as MAT 1-1-1 and 6 as MAT 1-2-1. In the case of plants inoculated with the mixture of strains, PCRs revealed the presence of both mating types although spatially isolated in different plants: MAT 1-2-1 under the plants 15 and 17 (ascomata 4639 and 4660) and MAT 1-1-1 under the plants 51, 52 (ascomata 4640, 4657, 4664) and 13 (ascoma 4675) (Fig. 1, Table 2).

A total of 23 root samples taken under 23 of the collected ascomata were processed to characterize mating types of ectomycorrhizas (ECMs) (Fig. 1, Table 2). Molecular analyses with *T. borchii* species-specific primers confirmed the identity of the ECMs morphotyped in each root sample. In total, 115 ECMs identified as *T. borchii* (5 for each collection point) were analyzed.

Remarkably, each ECM amplified a single and specific mating type amplicon. All ECMs had the same mating type as the inoculated strain and as the ascoma under which they were collected (Table

165 2).

Soil analysis also confirmed the persistence of the inoculated mating types. In almost all soil samples, the mating type corresponded to that of the inoculated strain and of the ascoma maternal tissue and ECMs collected in the same position, with only two exceptions: soils collected under ascomata 4677 and 4642. They were found in the same group of plants where the genotypes of the two ascomata (4601 and 4633) differed from the one used to inoculate the plants (Table 2, Fig. 2).

171

#### 172 *Genotyping of paternal tissue (spores)*

The SSRs analysis performed on the spores was successful only for 18 of the 33 analyzed ascomata.
Sixteen paternal genotypes were found on the 18 ascomata characterized by SSRs (Table 2 and 4).
The percentage of success of spore DNA extraction was only 55% probably due to the difficulties in breaking *T. borchii* spores or the too low quantity of extracted DNA.

177 Table 4

178 In zygotes, the  $F_{IS}$  was 0.058 with a p-value of 0.11, indicating a non-significant departure from 0; 179 consequently, even if the  $F_{IS}$  is positive there is no heterozygote deficit.

180 Most of the paternal C-MLGs were different from the female C-MLGs and were characterized by the 181 recombination of the alleles of the inoculated strains (Fig. 3). Four paternal MLGs (C-MLG 1; C-MLG 2; C-MLG 4, C-MLG 16, ascomata 4600, 4652-4599, 4651 and 4635 respectively) showed 182 183 new alleles (loci Tb151 and/or Tb46) never detected in the inoculated mycelia, although they were 184 detected in the maternal tissue of ascoma 4601 (Supplementary Table S1). Two paternal C-MLGs had new alleles not present in any maternal genotype in the loci Tb17 (C-MLG 9) and Tb293 (C-185 MLG 9 and C-MLG 16) (Fig. 3, Supplementary Table S2). Only two MLGs (C-MLG 14 and C-186 187 MLG 18) were detected as both maternal and paternal genotypes but only one (C-MLG 14) 188 according to the P-Sex value can be considered as a true hermaphroditic strain (Table 4 and 189 Supplementary Table S3). This hermaphroditic C-MLG 14 was found as male and female in adjacent 190 areas of the experimental field (Fig. 2). Only one C-MLG found only as male (C-MLG 2) for two

191 samples was supported by a significant P sex value, indicating that this genotype is a clone192 (Supplementary Table S3).

193 Fig. 3

194 **Discussion** 

195 In this study, truffles harvested in a T. borchii orchard were investigated by using mating type and 196 SSR markers for the first time. Mating type primers were recently designed and tested only on T. 197 borchii pure cultures and not applied to environmental samples (Belfiori et al., 2016). On the other 198 hand, SSR analyses were carried out on other truffle species (T. aestivum and T. melanosporum) but 199 never on T. borchii (Rubini et al., 2005; Riccioni et al., 2008; Murat et al., 2013; Molinier et al., 200 2015, 2016a, 2016b; Taschen, et al., 2016; De la Varga et al., 2017; Schneider-Maunory et al., 2018). 201 In this study, T. borchii-specific SSR loci were selected, tested for their polymorphisms on 50 202 ascomata of different origin and used to assess the genetic structure of T. borchii population in the 203 studied truffle orchard. The level of polymorphism of the developed SSR is highly variable (He 204 ranges from 0.208 to 0.739) as already observed for other *Tuber* spp. (Murat *et al.*, 2011; Molinier *et* 205 al., 2013).

Most of the studies on the genetic structure of truffle population have been conducted in human-made or natural truffle grounds where the identity of the strains colonizing roots was not known (Murat *et al.*, 2013; Taschen, *et al.*, 2016; Molinier *et al.*, 2016b; De la Varga *et al.*, 2017). On the contrary, the *T. borchii* plantation under investigation was established with plants inoculated with five different mycelial strains without knowing their mating type. As supposed by Iotti *et al.* (2016), the inoculated strains were found to belong to different mating types which is the necessary condition for fruiting in a heterotallic fungus like *T. borchii* (Belfiori *et al.*, 2016).

213

214 *Clonal structure suggests the persistence of the inoculated strains* 

215 Ascoma SSR analysis demonstrated that the maternal C-MLGs of the fruiting bodies was the same 216 as the corresponding inoculating strains, as also supported by the significant Psex value of three 217 female genotypes corresponding to the inoculated strains. Several works demonstrated that in *Tuber* 218 species the maternal tissue of ascomata and surrounding ECMs have the same genotype (Rubini et 219 al., 2011b, Murat et al., 2013; Molinier et al., 2016b; Taschen et al. 2016; De la Varga et al., 2017). 220 Moreover, a physical and nutritional link between these fungal structures has been proved (Le Tacon 221 et al., 2013; Deveau et al., 2019). For these reasons, our results suggest the persistence of the 222 inoculated mycelia 9 years after plantation on the root systems. A similar result was already found 223 for seedlings inoculated with the basidiomycete Laccaria bicolor for which the introduced strain was 224 still present in the inoculated plots 10 years after plantation (Selosse et al., 1998).

The maternal genotypes of ascomata 4601 and 4633 (C-MLG\_3 and C-MLG\_10, respectively) found in adjacent groups of plants at the southern edge of the truffle orchard differed from those of the inoculated mycelia. Ascomata 4601 had two alleles not present in the inoculated strains (loci Tb151 and Tb46) and this suggests an introgression of a foreign genotype on the host roots inoculated with 1Bo. Ascoma 4633 showed a different combination of the alleles found in the inoculated strains and thus could originate from ECMs colonized by a germinating spore corresponding to an offspring of inoculated mycelia.

232 Mating type of ECMs and almost all ascomata and soil mycelia was the same as the inoculated strain, 233 confirming the results obtained by SSR analyses. The only exception was found in the group of plants 234 inoculated with the strain 1Bo where a different mating type (MAT 1-1-1) was found in the soil (Fig. 235 2). Since the strain 2352 located in the adjacent group of plants has MAT 1-1-1, its mycelium was 236 probably able to migrate towards the plants inoculated with strain 1Bo. This migration could also 237 have been favored by the death of one uninoculated hazel separating the group of plants inoculated 238 with the strains 1Bo and 2352. Moreover, the strain 1Bo did probably not establish itself in the truffle 239 ground because of the low level of root colonization at the planting time (data not shown). Indeed 240 1Bo genotype was not found as maternal genotype in these or other plants and its specific alleles of 241 the loci Tb155 and Tb46 were also never found in any paternal genotype. The 1Bo mycelium was, in fact, isolated in 1997 and repeated subcultures are known to affect the viability and infectivity of a 242 243 *Tuber* strain (Piattoni *et al.*, 2017). Both the adjacent strain 2352 and the foreign genotype C-MLG 3 found as female in the ascomata 4658 and 4601, respectively, could have taken advantage of the 244 245 scarce colonization of strain 1Bo (Fig. 2). However, we cannot exclude the possibility that the 246 mycelium and ECMs of strain 1Bo were still present in the soil and the production of the ascoma of 247 this strain could have occurred.

248

249 The plants inoculated with the mixture of strains showed that MAT 1-1-1 and MAT 1-2-1 dominated 250 in separate areas either considering ECMs, soil mycelium or maternal tissue of ascomata. In order to better understand the dynamics of the mating types, we analyzed another 29 ascomata, and the 251 252 respective ECMs and soils collected in the following year (February-March 2017). These additional 253 analyses confirmed that, the ascomata, mycorrhizas and soil mycelium had the same mating type of the inoculated strain (Table and Fig. S5). As in 2016, the only exceptions were the two sample sets 254 255 collected under plants inoculated with the strain 1Bo. Fourteen of these 29 ascomata were collected 256 under the plants inoculated with the mixture of strains and only MAT 1-1-1 was found in these new 257 samples (Table and Fig. S5). This result confirms that also in *T. borchii* a single mating type tends to 258 dominate in a single soil patch, as found for T. melanosporum and T. aestivum in natural and cultivated truffle ground, and confirms that in field conditions only one strain or multiple strains of 259 260 the same mating type tend to colonize one single plant (Rubini et al., 2011b; Linde and Selmes, 2012; 261 Zampieri et al., 2012; Murat et al., 2013; Rubini et al., 2014; Molinier et al., 2016b; De la Varga et 262 al., 2017). This can be explained by considering that a vegetative incompatibility (VI) system exists 263 in T. borchii which prevents hyphal interactions between hyphae of different strains (Sbrana et al., 2007). Although the sets of genes which regulated VI in filamentous ascomycetes was not found in 264

*T. melanosporum* genome (Iotti *et al.*, 2012b), other molecular mechanisms controlling self/nonself
 recognition are involved in segregation between strains of different mating types in the field (Rubini
 *et al.*, 2011b; Selosse *et al.*, 2013) and prevent hyphal fusion in axenic conditions (Iotti *et al.*, 2016).

268

# 269 Meiospores are the most important origin of male genotype

270 Most of the male C-MLGs are new genotypes and seem to be originated from spores, confirming the 271 hypothesis made for T. melanosporum that the locally dispersed spores are the major source of male 272 genotypes (Selosse et al., 2013; Taschen et al., 2016; Le Tacon et al., 2016; De la Varga et al., 2017). 273 In fact, the male C-MLGs showed an allele recombination of the alleles present in the maternal 274 genotypes which are mostly (5 out of 6) represented by the inoculated strains. The numbers of 275 generations to account for the observed diversity of paternal recombinant genotypes should be at least 276 two. In fact, for example, C-MLG 15 could derive from a first mating between C-MLG 8 and C-277 MLG 18 and then the mycelium/conidia originated from the spores of the formed ascoma could have 278 fertilized strain 2352 or 2364. Thus the ascoma production started before our first survey with trained dogs in the truffle ground (Iotti et al., 2016). Only three male genotypes presented new alleles and 279 perhaps we could suppose that they originated from spores coming from other sites. Although it is 280 281 not possible to exclude the arrival of spores from the natural T. borchii grounds which are located 282 several kilometers away (Iotti et al., 2016), likely they could also have come from cultivated T. 283 borchii orchards in the area. Tuber borchii cultivation has become very popular in Italy in the last few years after the first results obtained by spore inoculation (Zambonelli et al., 2000). 284

The role of spores in fertilization could explain the increase in truffle production obtained by inoculating spores in the field (Murat *et al.*, 2016) and the decrease in production attributed to ascoma overharvesting in natural truffle grounds. Moreover, it stresses the importance of animals in spore dispersal (Piattoni *et al.*, 2014; Zambonelli et al., 2017; Ori *et al.*, 2018) not only to promote truffle colonization of new areas but also to favor truffle fertilization in non-productive plants.

#### 290

291 Only one strain (strain 2352; MAT 1-1-1; C-MLG 14) displayed actual evidence for 292 hermaphroditism. This strain was found as male in one ascoma (4601) collected in the adjacent plants 293 inoculated with another strain (1Bo, MAT 1-2-1). This ascoma was located under the same group of 294 plants where both mating types were found in the soil and the C-MLG 14 was also found as female 295 in one ascoma (4658). Although not statistically supported, another C-MLG (C-MLG 18) was found 296 in adjacent groups of plants as male and female, supporting the possibility that fertilization can occur 297 also between mycelia. These results suggest that hermaphroditism is not frequent in *T. borchii*, as 298 already reported for T. melanosporum (De la Varga et al., 2017).

299

# Consideration on the truffle life cycle 300

301 Although this study gives new important insights into truffle biology, it was not able to completely 302 resolve the mystery of the reproduction strategy in truffles. How are the ascospores able to fertilize 303 the mycelium of different mating types? We can suppose that the ascospores germinate and originate 304 a mycelium that, directly or by conidia formation, fertilizes other compatible mycelia in the soil or 305 on the roots. As suggested for *T. melanosporum*, competition and/or vegetative incompatibility events 306 can prevent the growth of this new mycelium in the root system extensively colonized with the 307 mycelium of the opposite mating type (De la Varga *et al.*, 2017). However, when host roots are not 308 colonized by other truffle strains, the new genotype could be able to establish on them and function 309 as a new female (maternal) genotype. T. melanosporum and T. magnatum presented a significant 310 heterozygote deficit with high levels of inbreeding (Paolocci et al., 2006; Riccioni et al., 2008; 311 Taschen et al. 2016; De la Varga et al., 2017). In T. borchii we did not find such inbreeding since Fis 312 was not significantly different from 0, suggesting no departure from panmixia in the Cadriano 313 population. This means that all opposite genotypes have the same probability to breed. Could it be 314 explained by the possibility of T. borchii to form conidia? Indeed, the conidia formation in T.

*melanosporum* and *T. magnatum* was never observed in contrast to *T. borchii* and other species in the Puberulum clade (Urban *et al.*, 2004; Healy *et al.*, 2012; Ian Hall personal communication). Additional studies will be necessary to clarify these aspects of truffle life cycle, focusing on field experiments to detect conidia in the Cadriano population as well as in the laboratory to induce conidia production and ascospore germination.

320

# 321 Experimental Procedures

### 322 Plantation and ascoma sampling

323 The study was carried out in an experimental truffle orchard in Cadriano (Bologna, Italy) established 324 in autumn 2007-2008, planting seedlings of *Pinus pinea* L., *Quercus pubescens* Willd., *Quercus* 325 robur L. and Corvlus avellana L. The seedlings were inoculated with 5 different T. borchii pure 326 cultures (strains Tb98, 2352, 2292, 1Bo, 2364) separately and together as described by Iotti et al. 327 (2016). Seedlings inoculated with the same strain(s) were grouped along four rows and separated by non-inoculated guard plants of C. avellana (Fig. 1). Ascoma production was firstly verified in 2016 328 329 when 99 ascomata (total weight 722.2 g) were collected through February and March (Iotti et al., 330 2016). Each ascoma was weighed and fragments were either freeze-dried at -65 °C for three days in 331 a Virtis Benchtop 2 K lyophilizer (SP Industries) and then stored at -20 °C (gleba) or fixed in FAA 332 (gleba and peridium) for molecular and morphological analyses, respectively. The remaining portion 333 of each ascoma was dried and deposited in the Mycological Herbarium of Hypogeous Fungi of the 334 Bologna University (CMI-UNIBO). Fruiting position and metadata of 33 truffles processed in this 335 study are reported in Fig. 1 and Table 3.

336

#### 337 Mycelial strains

338 The 5 strains of *T. borchii* used for the seedling inoculation were preserved at 4 °C in 15 ml tubes

containing 6 ml of Potato Dextrose Agar half strength (hsPDA) (Difco) in the culture collection of
the Mycological Center of Bologna University (CMI-UNIBO). The cultures were renewed every year
on fresh hsPDA.

For their genetic characterization, the cultures were transferred on fresh modified woody plant medium (mWPM) (Iotti *et al.*, 2005) without agar addition and incubated in the dark at  $23 \pm 1$  °C for 60 days.

345

#### 346 Soil and root sampling

Soil and ECMs were sampled under 23 ascomata collected in February 2016 (Table 2) during truffle
surveys. A 20-cm-long soil core was taken under each ascoma by using a 6-cm-diameter soil corer.
ECMs were carefully separated from soil and washed in sterile water while any root fragment, stone
or organic debris was removed under a stereomicroscope (× 12) from the remaining soil.

ECMs were examined under a stereomicroscope (× 40) and those of *T. borchii* were identified based on their morphological features (Zambonelli *et al.*, 1993). *Tuber borchii* ECMs were vortexed in a 1.5 ml tube for 30 s, spun for 2 min at 17,000 g to remove soil particles from the mantle and then stored in sterile water at -80 °C pending further molecular characterization.

Soil samples were freeze-dried at - 65 °C for three days and then pulverized and homogenized by
mortar and pestle. Three 15 ml tubes containing 5 g of soil were prepared for each sample and then
stored at - 20°C until DNA extraction.

358

# 359 Molecular assays

# 360 DNA extraction

361 The complexity of the genetic analyses carried out in this study and, in particular, the necessity to
362 differentially target the maternal and paternal tissues of ascomata have implicated the selection of

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363 different strategies to isolate and/or amplify DNAs.

A direct PCR strategy avoiding DNA isolation (Iotti and Zambonelli, 2006) was applied to confirm
the identity of ECMs and to characterize the mating type of mycelia, ascomata and ECMs.

366 Soil DNA was extracted using the CTAB-based protocol described by Iotti et al. (2012c) adapted for

367 1 g of soil. Crude DNA solutions were then purified using the Nucleospin Plant II kit (Macherey-

368 Nagel, Düren, Germany) following the manufacturer's instructions. Total DNAs were quantified by

a NanoDrop ND-1000 Spectrophotometer (Thermo Scientific) and their quality evaluated with
optical density (OD) 260/280 nm and 260/230.

371 DNA extraction for analyses of SSR polymorphisms was performed using three different protocols: 372 1) DNeasy® Plant Mini Kit (Qiagen, Hilden, Germany) was used to isolate DNA from 50 mg of 373 lyophilized mycelia by applying the manufacturer's instructions; 2) Nucleospin Plant II kit 374 (Macherey-Nagel, Düren, Germany) was used to isolate DNA from 50 mg of frozen gleba 375 (corresponding to the female tissue) by applying the manufacturer's instructions; 3) spore DNA was 376 isolated by placing thin slices of lyophilized gleba in 1 ml of sterile water within a petri plate; after 377 precipitation from the floating gleba slice to the bottom of the plate, about 200 spores were transferred 378 in a 1.5 ml tube and their DNA isolated according to De la Varga et al. (2017). DNA extraction from 379 each spore sample was repeated up to three times if the amplification of paternal genotypes failed.

380 DNA extracts were stored at -20°C until processed.

381 Tuber borchii identification

The species-specific primer pair TboI-TboII (Amicucci *et al.*, 1998) were used to confirm the identity of the morphotyped *T. borchii* ECMs by direct PCR and to detect the presence of *T. borchii* extraradical mycelium in DNA soil extracts. For PCRs we used 1 µl of a 1:10 dilution of DNA (10–50 ng DNA) in a reaction volume of 10 µl. The REDTaq DNA polymerase and REDTaq PCR reaction buffer (1.1 mM MgCl<sub>2</sub> final concentration) were used according to the protocol of SIGMA, with 200  $\mu$ M of each dNTP, 0.2  $\mu$ M of each primer, and 75  $\mu$ g BSA (only for ECMs). PCR reactions were performed in a BioRad thermalcycler with the following conditions: 6 min at 94 °C followed by 34 cycles of denaturation at 94 °C for 30 s, annealing at 50 °C for 30 s, extension at 72 °C for 30 s, and a final extension at 72 °C for 5 min. PCR products were run on 2% agarose gel and visualized by staining with ethidium bromide.

392

# 393 *Mating type identification*

Mating type identification of pure cultures, ascomata, ECMs (five from each soil sample), and extraradical soil mycelium was performed by applying a multiplex PCR, using both the specific primer pairs B1-B3 and B23-B33 (Belfiori *et al.*, 2016) designed to identify the *T. borchii* MAT 1-1-1 and MAT 1-2-1 genes, respectively.

398 Multiplex PCRs of mating type genes were performed in a total volume of 25 µl consisting of 1× Ex-399 Taq Buffer (TaKaRa), 400 nM for each dNTP, 40 µg of bovine serum albumin, 400 nM for each 400 primer, 0.75 U of Ex Tag® DNA polymerase (TaKaRa). Few aerial hyphae, or small portions of 401 gleba (sterile veins) and ECM mantle were transferred directly to the PCR tubes in place of the 402 extracted DNA. The direct PCRs were performed with the following conditions: 6 min at 94 °C followed by 34 cycles of denaturation at 94 °C for 30 s, annealing at 60 °C for 30 s, extension at 72 403 404 °C for 30 s, and a final extension at 72 °C for 5 min. PCR products were run on 2% agarose gel. The 405 same conditions were applied to identify the mating type of extra-radical soil mycelium by adding 30 406 ng of soil DNA extracts to the PCR mixtures and avoiding BSA addition.

407

# 408 Selection of polymorphic SSR primers

409 MISA program (http://pgrc. Ipk gatersleben.de/misa/download/misa.pl) was used to search for SSR 410 markers in the recently released *T. borchii* genome (Murat *et al.*, 2018). Tri-, tetra-, penta-, and 411 hexanucleotide satellites with at least 6 repetitions and excluding composite SSRs were selected as 412 search parameter. Subsequently, primer pairs were designed based on the sequences flanking the 413 selected SSRs using Primer 3 software (Rozen and Skaletsky, 2000). The functionality of the 414 designed primers was tested using AmplifX software (http://crn2m.univ-mrs.fr/pub/amplifx-dist). 415 After *in silico* analyses, a total of 31 primer pairs were designed and tested in PCR and visualized in 416 4% agarose gel. Among then 14 were retained since they produced only one band per sample and 417 showed polymorphisms. They were used to search for polymorphisms on 50 ascomata from Italy and 418 Hungary (Supplementary Tables S1 and S4). The genotyping was realized in the Gentyane platform 419 (INRA, Clermont-Ferrand, France). The mating type of these ascomata was also determined as 420 described above.

421

### 422 SSR identification

The extracted DNA from mycelia, gleba and spores was amplified using a set of primer pairs amplifying the 14 polymorphic SSR loci (Table 1 and Supplementary Table S4). The amplification of maternal genotypes (gleba) was carried out only for the ascomata on which the amplification of paternal genotypes (spores) was successful.

427 PCRs were performed in a total volume of 10  $\mu$ l consisting of 1× REDTaq Buffer (1.1 mM MgCl<sub>2</sub> 428 final concentration), 200  $\mu$ M for each dNTP, 0.2  $\mu$ M of each primer and 1U DNA polymerase (RED

429 Taq Sigma-Aldrich) and 1 μl template DNA diluted 10 times (10–50 ng DNA).

430 The PCR reactions were performed with the following conditions: 4 min at 94 °C followed by 34

431 cycles of denaturation at 94 °C for 30 s, annealing at 55-60-65°C for 30 s depending on the primer,

432 extension at 72 °C for 30s, and a final extension at 72 °C for 5 min. PCR products were run on 4%

- 433 agarose gel.
- 434 Genomic DNA was analyzed using an ABI 3730XL sequencer (Applied Biosystems, Foster City,
- 435 California, USA) from "Plateforme de Génotypage GENTYANE" (Clermont-Ferrand, France). The
- 436 size of the alleles was analyzed with the Peak scanner software v.1.0.

- 437 SSR analyses were carried out only on ascomata because, when the genome sequencing of *T. borchii*
- 438 (Murat et al., 2018) made possible to select the SSR markers, the mycorrhizas were already analyzed
- 439 for mating type characterization by using direct PCR and successively disrupted (see above).

#### 440 Data analyses

Gene diversity or expected heterozygosity (He) was calculated using GenAlex v6.51.b2 (Peakall and Smouse, 2006) in Excel 2011. The He is a measure of the probability that two copies of the same gene chosen randomly in a population will have different alleles (Nei, 1973). This estimator is based on the allele frequency and can be calculated as:  $He = 1\sum Rpi^2$ . The genotypic diversity was calculated using Multilocus 1.3 (Agapow and Burt, 2001). It corresponds to the probability that two individuals taken at random have different genotypes. In other words, this value is 0 if every individual has the same genotype, and 1 if every individual has a different genotype.

The fixation index  $F_{IS}$  was calculated for zygote (maternal+paternal genotypes) in the Cadriano population using GenAlex v6.51.b2. In this analysis only the nine polymorphic SSR in this population were used. The significant departure from  $F_{IS} = 0$  was estimated using the R script developed by Taschen *et al.* (2016); script available at http://dx.doi.org/10.5061/dryad.vm11r.

MLG analysis of the inoculated mycelia, and the maternal (gleba) and paternal (spores) ascoma tissues harvested in Cadriano was realized using the 14 polymorphic SSR markers combined with the mating type gene. MLGsim 2.0 (Stenberg *et al.*, 2003) was used for MLG identification and the calculation of the likelihood (PSex) that copies of MLGs result from sexual reproduction or clonal spread. The threshold value (< 0.05) for testing the significance of the PSex for each genotype was estimated using 1000 simulations. When the PSex values fell below the threshold value, it was concluded that identical genotypes originated from clonal multiplication.

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460

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#### 467 **References**

468 Agapow, P.M., and Burt A. (2001) Indices of multilocus linkage disequilibrium. Mol Ecol
469 Notes 1:101–102.

Amicucci, A., Zambonelli, A., Giomaro, G., Potenza, L., and Stocchi V. (1998) Identification
of ectomycorrhizal fungi of the genus *Tuber* by species-specific ITS primers. Mol Ecol 7: 273-277.

Belfiori, B., Riccioni, C., Paolocci, F., and Rubini, A. (2016) Characterization of the
reproductive mode and life cycle of the whitish truffle *Tuber borchii*. Mycorrhiza 26: 515–527.

Bonito, G.M., Gryganskyi, A.P., Trappe, J.M., and Vilgalys, R. (2010) A global meta-analysis
of *Tuber* ITS rDNA sequences: species diversity, host associations and long-distance dispersal. Mol
Ecol 19: 4994-5008.

477 Chevalier, G. (1973) Synthese axenique des mycorhizes de *Tuber brumale* Vitt. a partir de
478 cultures puresdu champignon. Ann Phytopathol 5: 163-182.

479 Chevalier, G., and Grente, J. (1978) Application practique de la symbiose ectomycorrhizienne:
480 production a grande echelle de plants mycorrhizes par la truffe (*Tuber melanosporum* Vitt.).
481 Mushroom Science 10: 483– 505.

482 De la Varga, H., Le Tacon, F., Lagoguet, M., Todesco, F., Varga, T., Miquel, I., et al. (2017)

483 Five years investigation of female and male genotypes in Périgord black truffle (*Tuber melanosporum* 

484 Vittad.) revealed contrasted reproduction strategies. Environ Microbiol 19: 2604-2615.

485	Deveau, A., Clowez, P., Petit, F., Maurice, JP., Todesco, F., Murat, C., et al. (2019). New							
486	insights into black truffle biology: discovery of the potential connecting structure between a Tuber							
487	aestivum ascocarp and its host root. Mycorrhiza 29: 219–226.							
488	Gardin, L. (2005) I tartufi minori in Toscana. Gli ambienti di crescita dei tartufi marzuolo e							
489	scorzone. Firenze: Quaderno ARSIA [WWW document] URL							
490	http://www.provincia.pisa.it/uploads/2012_08_28_17_46_12.pdf							
491	Giomaro, G., Sisti, D., and Zambonelli, A. (2005) Cultivation of edible ectomycorrhizal fungi							
492	by in vitro mycorrhizal synthesis. In In vitro Culture of Mycorrhizas. Declerck, S., Strullu, D.G., and							
493	Fortin, J.A. (eds). Soil Biology series, vol.4. Berlin Heidelberg: Springer-Verlag, pp. 253–267.							
494	Hall, I., Brown, G., and Zambonelli, A. (2007) Taming the Truffle. The History, Lore, and							
495	Science of the Ultimate Mushroom. Portland: Timber Press.							
496	Hall, I., Fitzpatrick, N., Miros, P., and Zambonelli, A. (2017) Counter-season cultivation of							
497	truffles in the Southern Hemisphere: an update. Italian Journal of Mycology 46: 21-36.							
498	Healy, R.A., Smith, M.E., Bonito, G.M., Pfister, D.H., Ge Z.W., Guevara, G.G., et al., (2012)							
499	High diversity and widespread occurrence of mitotic spore mats in ectomycorrhizal Pezizales. Mol							
500	Ecol 22: 1717–1732.							
501	Iotti, M., Amicucci, A., Stocchi, V., Zambonelli, A. (2002) Morphological and molecular							
502	characterization of mycelia of some <i>Tuber</i> species in pure culture. New Phytol 155: 499-505.							
503	Iotti, M., Barbieri, E., Stocchi, V., and Zambonelli, A. (2005) Morphological and molecular							
504	characterization of mycelia of ectomycorrhizal fungi in pure culture. Fungal Divers 19: 51-68.							
505	Iotti, M., and Zambonelli, A. (2006) A quick and precise technique for identifying							
506	ectomycorrhizas by PCR. Mycol Res 110: 60 – 65.							
507	Iotti, M., Piattoni, F., and Zambonelli, A. (2012a) Techniques for host plant inoculation with							
508	truffles and other edible ectomycorrhizal mushrooms. In Edible ectomycorrhizal mushrooms, current							

knowledge and future prospects. Zambonelli, A., and Bonito, G.M. (eds). Soil Biology series, vol.34.
Berlin Heidelberg: Springer-Verlag, pp. 145 – 161.

511 Iotti, M., Rubini, A., Tisserant, E., Kholer, A., Paolocci, F., and Zambonelli, A. (2012b) 512 Self/nonself recognition in *Tuber melanosporum* is not mediated by a heterokaryon incompatibility 513 system. Fungal Biol 116: 261-275.

- Iotti, M., Leonardi, M., Oddis, M., Salerni, E., Baraldi, E., and Zambonelli A. (2012c)
  Development and validation of a real-time PCR assay for detection and quantification of *Tuber magnatum* in soil. BMC Microbiol 12: 93.
- 517 Iotti, M., Piattoni, F., Leonardi, P., Hall, I.R., and Zambonelli, A. (2016) First evidence for 518 truffle production from plants inoculated with mycelial pure cultures. Mycorrhiza 26: 793–798.
- Lancellotti, E., Iotti, M., Zambonelli, A., and Franceschini, A. (2016) The Puberulum group sensul lato (whitish truffles). In True truffle (*Tuber* spp.) in the world. Zambonelli, A., Murat, C., and lotti, M. (eds). Soil Biology, vol. 47. Berlin Heidelberg: Springer-Verlag, pp. 105-124.
- Le Tacon, F., Zeller, B., Plain, C., Hossann, C., Bréchet, C., and Robin, C. (2013) Carbon transfer from the host to *Tuber melanosporum* mycorrhizas and ascocarps followed using a 13C pulse-labeling technique. PLoS One 8: e64626.
- Le Tacon, F., Rubini, A., Murat, C., Riccioni, C., Robin, C., Belfiori, B., *et al.* (2016) Certainties and uncertainties about the life cycle of the Périgord black truffle (*Tuber melanosporum* Vittad.) Ann For Sci 73: 105–117.
- Linde, C.C., and Selmes, H. (2012) Genetic diversity and mating type distribution of *Tuber melanosporum* and their significance to truffle cultivation in artificially planted truffieres in Australia. Appl Environ Microbiol 78: 6534–6539.

symbiosis. Nature
5

Molinier, V., Murat, C., Morin, E., Gollotte, A., Wipf, D., and Martin F (2013) First Identification of polymorphic microsatellite markers in the burgundy truffle, *Tuber aestivum* (Tuberaceae). Appl Plant Sci 1: 1200220

Molinier, V., Murat, C., Peter, M., Gollotte, A., De la Varga, H., Meier, B., *et al.* (2015) SSRbased identification of genetic groups within European populations of *Tuber aestivum* Vittad.
Mycorrhiza. 26: 99-110.

Molinier, V., Murat, C., Peter, M., Gollotte, A., De la Varga, H., Meier, B., *et al.* (2016a) SSRbased identification of genetic groups within European populations of *Tuber aestivum* Vittad.
Mycorrhiza 26: 99–110.

Molinier, V., Murat, C., Baltensweiler, A., Büntgen, U., Martin, F., Meier, B., *et al.* (2016b)
Fine-scale genetic structure of natural *Tuber aestivum* sites in southern Germany. Mycorrhiza 26:
895–907.

Morara, M., Saitta, A., Venturella, G., Gargano, M.L., and Zambonelli, A. (2009) I funghi
ipogei dell'Emilia Romagna e della Sicilia a confronto. In: Annales Confederationis Europaeae
Mycologiae Mediterraneensis. Venturella, G. and Gargano M.L. (eds) XVII Giornate Micologiche
della CEMM, 14–21 Novembre 2009, Cefalù, Italia, pp 53–68

Murat, C., Riccioni, C., Belfiori, B., Cichocki, N., Labbé, J., Morin, E. *et al.* (2011) Mining
microsatellites in the Perigord black truffle genome and identification of new molecular markers.
Fungal Genet Biol 48: 592–601.

553	Murat, C., Rubini, A., Riccioni, C., De la Varga, H., Akroume, E., Belfiori, B., et al. (2013)
554	Fine-scale spatial genetic structure of the black truffle (Tuber melanosporum) investigated with
555	neutral microsatellites and functional mating type genes. New Phytol 199: 176-187.
556	Murat, C., Bonneau, L., De la Varga, H., Olivier, J.M., Sandrine, F. and Le Tacon, F. (2016)
557	Trapping truffle production in holes: a promising technique for improving production and unravelling
558	truffle life cycle. Italian Journal of Mycology 45: 47-53.
559	Murat, C., Kuo, A., Barry, W.W., Clum, A., Dockter, R.B., Fauchery, L., et al. (2018) Draft
560	genome sequence of <i>Tuber borchii</i> Vittad., a whitish edible truffle. Genome Announc 6: e00537-18.
561	Nei, M. (1973) Analysis of gene diversity in subdivided populations. PNAS 70: 3321–3323.
562	Ori, F., Trappe, J., Leonardi, M., Iotti, M., and Pacioni, G. (2018) Crested purcopines (Hystrix
563	<i>cristata</i> ): mycophagist spore disperser of the ectomycorrhizal truffle <i>Tuber aestivum</i> . Mycorrhiza 28:
564	561-565.
565	Palenzona, M., Chevalier, G., and Fontana, A. (1972) Sintesi micorrizica tra i miceli in coltura
566	pura di Tuber brumale, T. melanosporum, T. rufum e semenzali di conifere e latifoglie. Allionia 18:
567	41–52.
568	Paolocci, F., Rubini, A., Riccioni, C., and Arcioni, S. (2006) Reevaluation of the life cycle of
569	Tuber magnatum. Appl Environ Microbiol 72: 2390–2393.
570	Peakall, R., and Smouse, P.E. (2006) GenAlEx 6: genetic analysis in Excel. Population genetic
571	software for teaching and research. Mol Ecol Notes 6: 288–295.
572	Piattoni, F., Amicucci, A., Iotti, M., Ori, F., Stocchi, V., and Zambonelli, A. (2014) Viability
573	and morphology of <i>Tuber aestivum</i> spores after passage through the gut of <i>Sus scrofa</i> . Fungal Ecol
574	9: 52–60.
575	Piattoni, F., Leonardi, P., Boutahir, S., Iotti, M., and Zambonelli, A. (2017) Viability and
576	infectivity of <i>Tuber borchii</i> after cryopreservation. CryoLetters 38: 58-64.

577	Riccioni, C., Belfiori, B., Rubini, A., Passeri, V., Arcioni, S., and Paolocci, F. (2008) Tuber
578	melanosporum outcrosses: analysis of the genetic diversity within and among its natural populations
579	under this new scenario. New Phytol 180: 466–478.
580	Rozen, S., and Skaletsky, H. (2000) Primer3 on the WWW for general users and for biologist
581	programmers. In Bioinformatics Methods and Protocols. Misener, S., and Krawetz, S.A. (eds).
582	Methods in Molecular Biology <sup>™</sup> , vol 132. Totowa, NJ: Humana Press, pp. 365–386.
583	Rubini, A., Paolocci, F., Riccioni, C., Vendramin, G.G., and Arcioni, S. (2005) Genetic and
584	phylogeographic structures of the symbiotic fungus <i>Tuber magnatum</i> . Appl Environ Microbiol 71:
585	6584-6589.
586	Rubini, A., Belfiori, B., Riccioni, C., Tisserant, E., Arcioni, S., Martin, F., and Paolocci, F.
587	(2011a) Isolation and characterization of MAT genes in the symbiotic ascomycete Tuber
588	melanosporum. New Phytol 189: 710-722.
589	Rubini, A., Belfiori, B., Riccioni, C., Arcioni, S., Martin, F., and Paolocci, F. (2011b) Tuber
590	melanosporum: mating type distribution in a natural plantation and dynamics of strains of different
591	mating types on the roots of nursery-inoculated host plants. New Phytol 189: 723-735.
592	Rubini, A., Riccioni, C., Belfiori, B., and Paolocci, F. (2014) Impact of the competition between
593	mating types on the cultivation of Tuber melanosporum: Romeo and Juliet and the matter of space
594	and time. Mycorrhiza 24:19–27.
595	Sbrana, C., Nuti, M.P., and Giovanetti, M. (2007) Self-anastomosing ability and vegetative
596	incompatibility of Tuber borchii isolates. Mycorrhiza 17: 667–675.
597	Schneider-Maunoury, L., Clément, C., Coves, H., Lambourdière, J., Leclercq, S., Richard, F.,
598	Selosse, M.A., and Taschen, E. (2018) Is Tuber melanosporum colonizing the roots of herbaceous,
599	non-ectomycorrhizal plants? Fungal Ecol 31: 59-68.

600	Selosse, M.A., Jacquot, D., Bouchard, D., Martin, F., and Le Tacon, F. (1998) Temporal
601	persistence and spatial distribution of an American inoculant strain of the ectomycorrhizal
602	basidiomycete Laccaria bicolor in a French forest plantation. Mol Ecol 7: 561–573.
603	Selosse, M.A., Taschen, E., and Giraud, T. (2013) Do black truffles avoid sexual harassment
604	by linking mating type and vegetative incompatibility? New Phytol 199: 10-13.
605	Selosse, M.A., Schneider-Maunoury, L., Taschen, E., Rousset, F., and Richard, F. (2017) Black
606	truffle, a hermaphrodite with forced unisexual behavior. Trends Microbiol 25: 784-787:
607	Stenberg, P., Lundmark, M., and Saura, A. (2003) MLGsim: a program for detecting clones
608	using a simulation approach. Mol Ecol Notes 3: 329–331.
609	Taschen, E., Rousset, F., Sauve, M., Benoit, L., Dubois, M., Richard, F., and Selosse, M. (2016)
610	How the truffle got its mate: insights from genetic structure in spontaneous and planted Mediterranean
	$N_{\rm el} = 125 \cdot 5(11 \cdot 5(27$
611	populations of <i>Tuber melanosporum</i> . Mol Ecol 25: 5611-5627.
611 612	Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular
<ul><li>611</li><li>612</li><li>613</li></ul>	Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–
<ul><li>611</li><li>612</li><li>613</li><li>614</li></ul>	<ul> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular</li> <li>studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–</li> <li>758.</li> </ul>
<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> </ul>	<ul> <li>populations of <i>Tuber melanosporum</i>. Mol Ecol 25: 3611-3627.</li> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e</li> </ul>
<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> <li>616</li> </ul>	<ul> <li>populations of <i>Tuber melanosporum</i>. Mol Ecol 25: 3611-3627.</li> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e micromorfologica delle micorrize di <i>Tuber</i> spp. su <i>Quercus pubescens</i> Willd. Micol Ital 3: 73-90.</li> </ul>
<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> <li>616</li> <li>617</li> </ul>	<ul> <li>populations of <i>Tuber metanosporum</i>. Mol Ecol 25: 5611-5627.</li> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e micromorfologica delle micorrize di <i>Tuber</i> spp. su <i>Quercus pubescens</i> Willd. Micol Ital 3: 73-90.</li> <li>Zambonelli, A., Iotti, M., Rossi, I., and Hall, I. (2000) Interaction between <i>Tuber borchii</i> and</li> </ul>
<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> <li>616</li> <li>617</li> <li>618</li> </ul>	<ul> <li>populations of <i>Tuber melanosporum</i>. Mol Ecol 25: 5611-5627.</li> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e micromorfologica delle micorrize di <i>Tuber</i> spp. su <i>Quercus pubescens</i> Willd. Micol Ital 3: 73-90.</li> <li>Zambonelli, A., Iotti, M., Rossi, I., and Hall, I. (2000) Interaction between <i>Tuber borchii</i> and other ectomycorrhizal fungi in a field plantation. Mycol Res 104: 698-702.</li> </ul>
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<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> <li>616</li> <li>617</li> <li>618</li> <li>619</li> <li>620</li> </ul>	<ul> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e micromorfologica delle micorrize di <i>Tuber</i> spp. su <i>Quercus pubescens</i> Willd. Micol Ital 3: 73-90.</li> <li>Zambonelli, A., Iotti, M., Rossi, I., and Hall, I. (2000) Interaction between <i>Tuber borchii</i> and other ectomycorrhizal fungi in a field plantation. Mycol Res 104: 698-702.</li> <li>Zambonelli, A., Iotti, M., Giomaro, G., Hall, I., and Stocchi, V. (2002) <i>T. borchii</i> cultivation: an interesting perspective. In Edible mycorrhizal mushrooms. Proceedings of 2nd international</li> </ul>
<ul> <li>611</li> <li>612</li> <li>613</li> <li>614</li> <li>615</li> <li>616</li> <li>617</li> <li>618</li> <li>619</li> <li>620</li> <li>621</li> </ul>	<ul> <li>Dippulations of <i>Tuber metanosporum</i>. Mol Ecol 25: 5611-5627.</li> <li>Urban, A., Neuner-Plattner, I., Krisai-Greilhuber, I., Haselwandter, K. (2004) Molecular studies on terricolous microfungi reveal novel anamorphs of two <i>Tuber</i> species. Mycol Res 108: 749–758.</li> <li>Zambonelli, A., Salomoni, S., and Pisi, A. (1993) Caratterizzazione anatomo-morfologica e micromorfologica delle micorrize di <i>Tuber</i> spp. su <i>Quercus pubescens</i> Willd. Micol Ital 3: 73-90.</li> <li>Zambonelli, A., Iotti, M., Rossi, I., and Hall, I. (2000) Interaction between <i>Tuber borchii</i> and other ectomycorrhizal fungi in a field plantation. Mycol Res 104: 698-702.</li> <li>Zambonelli, A., Iotti, M., Giomaro, G., Hall, I., and Stocchi, V. (2002) <i>T. borchii</i> cultivation: an interesting perspective. In Edible mycorrhizal mushrooms. Proceedings of 2nd international workshop on edible ectomycorrhizal mushrooms. Hall, I., Wang, Y., Danell, E., and Zambonelli, A.</li> </ul>

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623	Zambonelli, A., Iotti, M., and Piattoni, F. (2008) Problems and perspectives in the production
624	of Tuber infected plants. In Mushroom biology and mushroom products. Proceeding of the sixth
625	International conference on Mushroom Biology and Mushroom products. Lelley, J.I., and Buswell,
626	J.A. (eds). Krefeld: GAMU GmbH, pp. 263 – 271.
627	Zambonelli, A., Iotti, M., and Hall, I. (2015) Current status of truffle cultivation: recent results
628	and future perspectives. Italian Journal of Mycology 44: 31–40.
629	Zambonelli, A., and Morara, M. (1984) Le specie di Tuberales dell'Emilia Romagna, ecologia
630	e distribuzione. Natura e Montagna 4:9–32.
631	Zambonelli, A., Murat, C., and Iotti, M. (2016) True truffle (Tuber spp.) in the world. Soil
632	Biology, vol. 47. Berlin Heidelberg: Springer-Verlag.
633	Zambonelli, A., Ori, F., and Hall, I. (2017) Mycophagy and spore dispersal by vertebrates. In
634	The fungal community. Its organization and role in ecosystems, 4th edn. Dighton, J., and White, J.F.
635	(eds). Boca Raton, FL: CRC Press Taylor and Francis Group, pp. 347-358.
636	Zampieri, E., Rizzello, R., Bonfante, P., Mello, A. (2012) The detection of mating type genes
637	of <i>Tuber melanosporum</i> in productive and non productive soils. Applied soil Ecology 57:9-15.
638	
639	Legends of tables
640	Table 1 – Characteristics of the 14 SSRs used to characterize the <i>Tuber borchii</i> clones involved in
641	this study.
642	Table 2 – Genotyping of the inoculated strain and of the ascoma, ectomycorrhizas and mycelium at
643	the points where the samples were collected.
644	Table 3 – Metadata of the <i>T. borchii</i> ascomata of Cadriano truffle ground used in this study.
645	Table 4 - Genetic diversity for maternal, paternal genotypes.
646	Figure legends

- 647 Fig. 1 Scheme of the experimental plantation realized with mycelial inoculated seedlings. The
- 648 circles show the codes of inoculated strains or their mixture (Mix). Different circle types indicate
- 649 different host plants. Triangles and asterisks indicate the position of samples and types of analyses
- applied to them. The dotted rectangle indicates the area of the plantation enlarged in Fig. 2.
- Fig. 2- Area of the plantation where a true hermaphrodite strain was found (C-MLG\_14). This
- strain was found as female in ascomata 4651, 4652 and 4658 and as male in ascoma 4601 located in
- the adjacent plants. This ascoma presents a new female C-MLG (C-MLG\_3).
- Fig. 3 Characteristics of the maternal (a) and paternal (b) genotypes of the 18 analyzed ascomata.

# 655 Supplemental material

- 656 Table S1 Multilocus genotypes (MLG) found in the 50 ascomata analyzed from different
- 657 populations.
- Table S2 Multilocus genotypes (C-MLG) found in the Cadriano plantation.
- Table S3- Genotype description, probability of genotype occurrence resulting from distinct sexual
- 660 events (PSex) for all samples harvested in the plantations.
- 661 Table S4- Tested microsatellite primers. In bold the selected polymorphic microsatellites.
- Table S4- Tested microsatellite primers. In bold the selected polymorphic microsatellites.
- 663 Table and Fig. S5- Mating type analysis of the ascomata, ectomycorrhizas and soil mycelia of the
- samples collected in 2017. In the figure triangles indicate the position of samples analyzed.

SSR name	Repeat motif	Genome localization <sup>a</sup>	Na <sup>b</sup>	Hec
Tb244	(GAGGGA) <sub>6</sub>	Intergenic	5	0.562
Tb1	(TATTTT) <sub>10</sub>	Intergenic	3	0.626
Tb83	(GACT) <sub>8</sub>	Intergenic	3	0.263
Tb11	(AGGC) <sub>8</sub>	Intergenic	4	0.319
Tb151	(AAC) <sub>8</sub>	Gene model (UTR)	8	0.472
Tb155	(GGA) <sub>12</sub>	Intergenic	7	0.649
Tb156	(GAG) <sub>8</sub>	Intergenic	5	0.432
Tb17	(TTTAGA) <sub>5</sub>	Intergenic	3	0.266
Tb206	(CCTT) <sub>8</sub>	Intergenic	3	0.617
Tb293	(AGAAGG)5	Gene model (intron)	10	0.739
Tb43	(CTTTT) <sub>5</sub>	Intergenic	4	0.255
Tb704	(AAAG) <sub>8</sub>	Intergenic	6	0.595
Tb43bis	(TACC) <sub>8</sub>	Intergenic	4	0.208
Tb46	(AGA) <sub>9</sub>	Gene model (CDS)	6	0.456
<sup>a</sup> UTR = untra	slated region, CDS = er of allele observed	coding sequence		

	sample	inoculated strain		strain ascoma genotype		genotypes			
plant n.	n.	n.	strain	ascoma	ECM	soil		maternal	paternal
1	4597	2292	2	2	2	2	C-MLG_18	nd	nd
46	4598	Tb 98	1	1	nd	nd	C-MLG_8	C-MLG_8	C-MLG_7
10	4599	Tb 98	1		nd	nd	C-MLG_8	C-MLG_8	C-MLG_2
69	4600	2364	1	1	nd	nd	C-MLG_13	C-MLG_13	C-MLG_1
63	4601	1Bo 🗲	2	2	nd	nd	C-MLG_6	C-MLG_3	C-MLG_14
47	4602	Tb 98	1	1	nd	nd	C-MLG_8	C-MLG_8	C-MLG_18
33	4633	2364 🗲	1	1	1	1	C-MLG_13	C-MLG_10	C-MLG_20
39	4634	2292	2	2	nd	nd	C-MLG_18	C-MLG_18	C-MLG_5
68	4635	2364	1	1	nd	nd	C-MLG_13	C-MLG_13	C-MLG_16
15	4639	MIX	na	2	2	2	na	nd	nd
51	4640	MIX	na	1	1	1	na	nd	nd
44	4641	Tb 98	1	1	1	1	C-MLG_8	nd	nd
65	4642	1Bo	2	2	2	1	C-MLG_6	nd	nd
6	4644	Tb 98	1	1	1	1	C-MLG_8	nd	nd
47	4647	Tb 98	1	1	1	1	C-MLG_8	nd	nd
10	4648	Tb 98	1	1	nd	nd	C-MLG_8	C-MLG_8	C-MLG_15

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44	4649	Tb 98	1	1	nd	nd	C-MLG_8	C-MLG_8	C-MLG_19
23	4650	2352	1	1	1	1	C-MLG_14	nd	nd
59	4651	2352	1	1	1	1	C-MLG_14	C-MLG_14	C-MLG_4
61	4652	2352	1	1	1	1	C-MLG_14	C-MLG_14	C-MLG_2
68	4653	2364	1	1	1	1	C-MLG_13	C-MLG_13	C-MLG_9
45	4654	TB98	1	1	1	1	C-MLG_8	nd	nd
41	4655	2292	20	2	2	2	C-MLG_18	nd	nd
44	4656	TB98	1	1	1	1	C-MLG_8	C-MLG_8	C-MLG_7
52	4657	MIX	na		1	1	na	C-MLG_14	C-MLG_17
65	4658	1 Bo 🗲	2	1	nd	nd	C-MLG_6	$C-MLG_14^+$	C-MLG_11
6	4659	TB98	1	1		1	C-MLG_8	nd	nd
17	4660	MIX	na	2	2	2	na	nd	nd
27	4662	2352	1	1	1	1	C-MLG_14	nd	nd
52	4664	MIX	na	1	1	1	na	C-MLG_8	C-MLG_21
13	4675	MIX	na	1	1	1	na	C-MLG_8	C-MLG_12
65	4677	1Bo	2	2	2	1	C-MLG_6	nd	nd
13	4699	MIX	na	1	1	1	na	nd	nd

\*1 correspond to the mating type MAT 1-1-1, 2 correspond to MAT 1-2-1.  $^\dagger$  C-MLG of the strain 2352

The arrow indicate the samples having a maternal C-MLG (in bold) different from the inoculated strain

nd = not determined

na = not applicable

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ascoma	sampling date	weight (g)	plant	tree species	inoculated strain
4597	08/02/2016	35	1	Pinus pinea	2292
4598	08/02/2016	34	46	Pinus pinea	Tb98
4599	08/02/2016	13.9	10	Quercus pubescens	Tb98
4600	08/02/2016	54.7	69	Corylus avellana	2364
4601	08/02/2016	20.9	63	Quercus pubescens	1Bo
4602	08/02/2016	2	47	Pinus pinea	Tb98
4633	12/02/2016	0.59	33	Quercus robur	2364
4634	12/02/2016	1	39	Quercus pubescens	2292
4635	12/02/2016	2.28	68	Quercus robur	2364
4639	19/02/2016	5.53	15	Pinus pinea	MIX
4640	19/02/2016	9.73	51	Quercus pubescens	MIX
4641	19/02/2016	6.84	44	Quercus pubescens	Tb98
4642	19/02/2016	3.42	65	Pinus pinea	1BO
4644	19/02/2016	0.27	6	Quercus pubescens	TB98
4647	19/02/2016	27.73	47	Pinus pinea	TB98
4648	25/02/2016	9.94	10	Quercus pubescens	Tb98
4649	25/02/2016	0.53	44	Quercus pubescens	Tb98
4650	25/02/2016	6.18	23	Pinus pinea	2352
4651	25/02/2016	0.38	59	Pinus pinea	2352
4652	25/02/2016	17.23	61	Pinus pinea	2352
4653	25/02/2016	7.39	68	Quercus robur	2364
4654	25/02/2016	2.73	45	Quercus pubescens	TB98
4655	25/02/2016	5.17	41	Quercus pubescens	2292
4656	25/02/2016	1.19	44	Quercus pubescens	Tb98
4657	25/02/2016	12.79	52	Quercus pubescens	MIX
4658	25/02/2016	6.66	65	Pinus pinea	1Bo
4659	02/03/2016	4.67	6	Quercus pubescens	Tb98
4660	02/03/2016	6.33	17	Pinus pinea	MIX
4662	02/03/2016	19.24	27	Quercus pubescens	2352
4664	02/03/2016	6.96	52	Quercus pubescens	MIX
4675	18/03/2016	11.24	13	Pinus pinea	MIX
4677	18/03/2016	2.06	65	Pinus pinea	1Bo
4699	06/04/2016	5.87	13	Pinus pinea	MIX

Number of sampled ascocarps	18
Number of inocula genotyped	5
Maternal genotypes	
Number of samples genotyped	18
Number of genotypes	6
Number of genotypes with Psex < 0.05	3
Maximum number of ascocarps per genotype	8
Number of genotype represented by a single ascocarp	3
Paternal genotypes	
number genotyped zigotes	18
Number of homozygosus zygotes	0
Zigotes Fis	1
Number of genotypes	16
Number of genotypes with Psex < 0.05	1
Maximum number of ascocarps per genotype	2
Number of male genotypes represented by a single ascocarp	12
Hermaphodite genotypes	
Total number (number of genotypes whit Psex < 0.05)	1

x < 0.05) 1



Fig. 1 - Scheme of the experimental plantation realized with mycelial inoculated seedlings. The circles show the codes of inoculated strains or their mixture (Mix). Different circle types indicate different host plants. Triangles and asterisks indicate the position of samples and types of analyses applied to them. The dotted rectangle indicates the area of the plantation enlarged in Fig. 2.

190x274mm (300 x 300 DPI)



Fig. 2- Area of the plantation where a true hermaphrodite strain was found (C-MLG\_14). This strain was found as female in ascomata 4651, 4652 and 4658 and as male in ascoma 4601 located in the adjacent plot This ascoma presents a new female C-MLG (C-MLG\_3).

190x274mm (300 x 300 DPI)



Fig. 3 - Characteristics of the maternal (a) and paternal (b) genotypes of the 18 analyzed ascomata.