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## Local epidemiology of the wood decay agent Laetiporus sulphureus in carob stands in Sicily

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## UNIVERSITÀ DEGLI STUDI DI TORINO

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16	Local epidemiology of the wood decay agent <i>Laetiporus sulphureus</i> in carob stands
17	in Sicily
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## 28 Summary

## 29

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30 The basidiomycete Laetiporus sulphureus (Bull) Murrill is a forest pathogen causing brown cubical 31 heart rot in a broad range of host trees. Despite its wide distribution and importance, studies aimed at 32 understanding the epidemiology of the fungus in specific areas or hosts are lacking. In this study, an 33 incidence of L. sulphureus as high as 34% was determined through molecular analysis of wood 34 samples collected from 70 carob (Ceratonia siliqua L.) trees in the South West of Sicily, Italy. A 35 phylogenetic analysis of Internal Transcribed Spacer (ITS) sequences indicated that all carob isolates 36 belonged to the cluster E of Laetiporus taxonomy. Ten molecular markers based on Single Sequence 37 Repeats (SSRs) designed on the L. sulphureus genome were developed and isolates were genotyped 38 through High Resolution Melting (HRM) analysis. High gene diversity (0.581), no correlation 39 between fungal genotype and host tree species and significant correlation between spatial and genetic

40 distance were observed, suggesting an important role of basidiospores in the epidemiology of the 41 fungus and a risk of transmission from a host tree species to the others. Finally, from a prognostic 42 perspective, significant differences among isolates in terms of mycelial growth suggest that in 43 addition to an accurate identification, a phenotypic characterization of isolates affecting trees may 44 also be important.

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## 47 Introduction

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49 The genus Laetiporus Murril (Polyporales) includes wood-rotting basidiomycetes growing on a wide variety of broadleaf and conifer trees (Bernicchia, 2005). The origin and biogeography of the genus 50 51 has been resolved only recently (Song & Cui, 2017). Currently, it is recognized that the genus 52 Laetiporus comprises eleven species and four undescribed worldwide distributed taxa (Gilbertson & 53 Ryvarden, 1986; Song & Cui, 2017), including L. ailaoshanensis B.K. Cui & J. Song, L. caribensis 54 Banik & D.L. Lindner, L. cincinnatus (Morgan) Burds., Banik & T.J. Volk, L. cremeiporus Y. Ota & T. Hatt., L. conifericola Burds. & Banik, L. gilbertsonii Burds, L. huroniensis Burds. & Banik, L. 55 montanus Černý ex Tomšovský & Jankovský, L. sulphureus (Bull) Murrill, L. versisporus (Lloyd) 56 57 Imazeki, and L. zonatus B.K. Cui & J. Song (Song & Cui, 2017).

L. sulphureus is one of the most studied species of the genus. The edible shelf-shaped, pink-orange 58 59 to yellow fruiting bodies of L. sulphureus are indeed a rich source of antioxidant and antimicrobial 60 compounds, including polysaccharides, hemolectins and laetiporic acids used in pharmaceutical and 61 industrial processes (Turkoglu et al., 2007; Petrović et al., 2013). In addition, fruiting bodies also 62 contain  $\alpha$ -1-3-glucans, which are used as inducers of bacterial mutanases, i.e. enzymes involved in 63 the degradation of biofilms formed by the etiologic agents of dental caries (Wiater et al., 2012). The nutritional and pharmaceutical values of L. sulphureus fruiting bodies (Petrović et al., 2014) led to 64 65 significant efforts for the development of cultivation methods of this fungus at the industrial scale 66 (Pleszczyńska et al., 2013).

67 However, L. sulphureus is also a forest pathogen (Schwarze, Engels, & Mattheck, 2000; Dai et al. 68 2007). As an important wood decay agent, it infects a broad range of hosts both in forests and urban 69 areas, including Castanea sativa L., Eucalyptus spp., Fagus spp., Quercus spp., Populus spp., Prunus 70 spp., Pyrus spp., Robinia spp., Salix spp., and occasionally Tilia spp. (Schwarze, Engels, & Mattheck, 71 2000; Bernicchia, 2005; Giordano et al., 2015). Infection is hypothesized to occur by means of 72 basidiospores germinating on injured bark of stems or roots (Schwarze, Engels, & Mattheck, 2000). 73 The fungus colonizes the wood through the libriform fibers by breaking down the cellulose 74 (Schwarze, Engels, & Mattheck, 2000). The stem decay columns can be 5-8 meters long and are 75 generally associated with heartwood (Bernicchia, 2005). Sapwood may also be invaded, but only in 76 the root systems (Schwarze, Engels, & Mattheck, 2000). After the establishment into the tree, sexual 77 reproduction may occur in fruiting bodies emerging outside the trunk (Bernicchia, 2005). 78 Furthermore, being a necrotrophic parasite, L. sulphureus is able to colonize dead trees and stumps 79 as a saprotroph (Schwarze, Engels, & Mattheck, 2000).

80 Research on L. sulphureus was mostly focused on the development of protocols to cultivate the 81 mycelia (Pleszczyńska et al., 2013), on the characterization of the plethora of compounds in the 82 fruiting bodies (Petrović et al., 2013), and on resolving the puzzling phylogenesis of the species and 83 related taxa (Song & Cui, 2017). For example, somatic incompatibility assays, allozyme and, more 84 recently, multi-locus phylogenetic analyses allowed mapping the different Laetiporus taxa in Europe, 85 North America and Asia (Banik & Burdsall Jr, 2000; Lindner & Banik, 2008, 2011; Vasaitis et al., 86 2009; Song et al., 2014). Comparative phenotypic assays in vitro further supported and confirmed the 87 differentiation among species (Vasaitis et al., 2009).

However, studies aimed at understanding the epidemiology of *L. sulphureus* in specific areas or hosts are lacking, with a very few exceptions (Rogers, Holdenrieder, & Sieber, 1999). The main issue is that, although *L. sulphureus* can be easily found in natural ecosystems in North America and Europe (Vasaitis et al., 2009), sites where fungal fruiting bodies are abundant are rare. In South West of Sicily

92 (Italy), L. sulphureus appears to be associated with carob trees (Ceratonia siliquia L.) (Bernicchia, 93 2005). In the past, carob was considered a profitable tree species in Sicily, since its pods were 94 consumed as food and used to feed livestock (Barbagallo et al., 1997). Currently, carob is regaining 95 importance as ornamental and forest tree in marginal areas and because of the industrial use of its pods (La Malfa et al., 2014). In the complex phytosanitary conditions of carob, which can be affected 96 97 by several plant pathogens and pests (Ramon-Laca & Mabberley, 2004; Vannini et al., 2017), a 98 relevant role is played by stem decays caused by L. sulphureus (Battle & Tous, 1997). The abundance 99 of L. sulphureus fruiting bodies on carob trees in the area of Modica and Ragusa, South West of Sicily, 100 makes this area an excellent sampling site for epidemiological studies. The aims of this work were: 101 I) to determine the incidence of L. sulphureus on carob trees in a selected area in Sicily, II) to develop 102 molecular markers to elucidate the genotypic and allelic diversity of L. sulphureus in this area, III) to 103 perform an analysis of the phenotypic diversity of L. sulphureus, through in vitro growth assays, and 104 IV) to explore the relationship between *in vitro* growth levels of mycelia and genotypic diversity.

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## 106 Materials and methods

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## 108 Sampling sites, wood and fungal materials

109 To estimate the incidence of L. sulphureus in carob trees, the sampling area comprised about 270 km<sup>2</sup> 110 in Modica Municipality (South West of Sicily, Italy) (Figure 1). A total of 70 randomly selected 111 standing carob trees were sampled in December 2015. Trees were about 80 years old and 112 characterized by a Diameter at Breast Height (DBH) ranging from 55 to 80 cm. Twenty-four trees 113 showed symptoms of wood decay, i.e. discoloration of exposed wood in pruning wounds or cracks, 114 while 46 trees did not show any mechanical damage, nor visible wood decay or crown symptoms. 115 These 46 trees were regarded as asymptomatic. Laetiporus fruiting bodies were neither observed on the sampled trees nor on the neighbouring trees. Each tree was drilled four times (drillings at 90° 116 117 from one another) 50 cm aboveground according to a previously described sampling protocol 118 (Guglielmo et al., 2010). Wood chips generated from different drillings of the same tree were pooled 119 together as a single sample in a 90-mm Petri dish and lyophilized overnight (Guglielmo et al., 2010). 120 For phylogenetic, genetic and phenotypic analyses, L. sulphureus fruiting bodies were collected 121 during Summer 2016 (June-August 2016) in a broader area in South West of Sicily (Figure 1). In detail, 24 fruiting bodies were collected from carob trees and one from an almond tree (Prunus 122 123 amygdalus Batsch). Distance between the trees from which fruiting bodies were collected ranged 124 between 0.5 to 35.4 km. Four additional fruiting bodies collected in Northern Italy were included in 125 the genetic analysis as outgroups, three of these originating from European chestnut (Castanea sativa 126 L.) and one from oak (Quercus sp.). Altogether 29 fruiting body samples were processed (Table 1). 127 Isolates were obtained from fruiting bodies by placing fragments of tissues (approximately 2 x 2 x 128 1.5 cm in size) excised from the context in 90-mm Petri dishes containing Malt Extract Agar and 129 citric acid (MEA; malt extract agar 33.6 g/L, citric acid 0.5 g/L). Isolates were incubated at 25°C±2°C 130 for one week and subsequently sub-cultured in 250-mL flasks, filled with Malt Extract liquid medium 131 (2% w/v) in the dark at  $25^{\circ}\text{C}\pm2^{\circ}\text{C}$  for a week before being harvested for DNA extraction.

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## 133 Molecular detection of *L. sulphureus* in wood samples

134 About 200 mg of lyophilized wood chips per sample were homogenized with the aid of glass beads 135 (3 mm and 5 mm) in a FastPrep FP120 Cell Disrupter (Qbiogene, Irvine, CA, USA). DNA extraction 136 from wood samples was performed by using the E.Z.N.A.<sup>TM</sup> Stool DNA Isolation Kit (Omega Bio-137 Tek, Doraville, CA, USA). Taxon specific primers developed by Guglielmo et al. (2007) were used in PCR to identify samples positive to L. sulphureus s.l. PCR assays were performed in a 25 µl volume 138 139 containing 5x of PCR buffer, 1.5 mM of MgCl<sub>2</sub>, 0.2 mM of dNTPs mix, 0.5 µM each of the taxon 140 specific primers, 0.025 U/µl of GoTaq® polymerase (Promega, Madison, WI, USA) and 6.25 µl of 141 the 50-fold dilution of the DNA extracted from wood (Guglielmo et al., 2007). PCR reactions were performed using an initial denaturation at 94°C for 3 minutes, followed by 35 cycles with each cycle 142 143 consisting of a denaturation at 94°C for 30 seconds, an annealing at 60°C for 45 seconds, an extension

at 72°C for 1 minute, and one final extension cycle at 72°C for 10 minutes. The presence of amplicons
in samples were checked, after electrophoretic migration (6 V/cm) on a gel containing 1.5% (w/v) of
agarose, through the use of the Image Lab<sup>™</sup> Software (Bio-Rad Laboratories, Hercules, CA, USA).

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## 148 Phylogenetic analysis of *L. sulphureus* isolates

149 Mycelia from the 29 isolates were dried overnight with a lyophilizer equipped with a vacuum pump. 150 About 200 mg of dried mycelia per sample were homogenized by using glass beads (diameter 2 mm 151 and 4 mm) in a FastPrep FP120 Cell Disrupter (Qbiogene, Irvine, CA, USA). Total DNA extraction was performed using DNeasy Plant Mini Kit (Qiagen Inc., Valencia, CA, USA), following 152 153 manufacturer instructions. The DNA samples were used as templates for PCRs with the primer pair 154 ITS1F/ITS4, which amplifies the Internal Transcribed Spacer (ITS) regions of the fungal nuclear 155 ribosomal RNA gene (nrDNA; Gardes & Bruns, 1993). The PCR mix included 6.25 µL of DNA, 0.75 156 U of GoTaq polymerase (Promega, Madison, WI, USA), 0.5 µM of each primer, 200 µM of each 157 dNTP, and 5  $\mu$ L of the 5x buffer in a total volume of 25  $\mu$ L. The PCR protocol was as follows: an 158 initial denaturation step at 94°C for 3 minutes, followed by 35 cycles at 94°C for 30 seconds, 54°C 159 for 30 seconds, and 72°C for 45 seconds, and a final elongation step at 72°C for 10 minutes. 160 Amplicons were visualized on 1% (w/v) agarose gel after electrophoresis at 6 V/cm for 30 minutes. PCR products were purified by using ExoSAP-IT (Affymetrix, Santa Clara, CA, USA) at 37°C for 161 162 15 minutes followed by 80°C for 15 minutes. The purified PCR products were sequenced at BMR 163 Genomics S.r.l. (Padua, Italy). A multiple sequence alignment was built using the ClustalW algorithm inside MEGA version 6 (Tamura et al., 2013). Phylogenetic trees were constructed using the 164 165 Maximum Likehood (ML) method (Jukes-Cantor model), with the MEGA v. 6 software. Bootstrap analyses were carried out on the basis of 100 re-samplings of the sequence alignment. A total of 78 166 167 Laetiporus spp. nucleotide sequences deposited in GenBank from previous phylogenetic studies by Vasaitis et al. (2009) and by Song & Cui (2017) were also included in the phylogenetic analysis. 168 169 Representative ITS nucleotide sequences from the present study were deposited in the GenBank

170 database under the accession numbers MG386383- MG386385.

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## 172 Development of SSR-based markers, HRM genotyping and analysis of genetic diversity of 173 isolates

MSDB 2.4.2 (Microsatellite Search and Database) (Du et al., 2013) was used to scan the entire L. 174 sulphureus genome (Laetiporus sulphureus var. sulphureus v1.0; Nagy et al., 2015) to search for 175 perfect Single Sequence Repeats (SSRs), using the "perfect" search mode. Six classes of 176 177 microsatellites were detected as follows: mono-, di-, tri-, tetra-, penta- and hexa-nucleotide SSR 178 motifs with minimum repeat number of 12, 7, 5, 4, 4, 4, respectively. Five mono-, four di-, nine tri-, 179 one tetra- and one hexa-nucleotide SSRs were selected as candidate marker regions, based on length 180 of SSR motifs (minimum length 10 repeats) and GC-content above 60% in the flanking regions, this 181 totalling 20 SSR markers. The design of specific primer sets to amplify SSR loci was performed by 182 using Primer3Plus (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi/). Each primer 183 pair was tested in PCR on DNA of the isolates VALLERE and C2 (Table 1), and the presence of 184 amplification products was checked by agarose electrophoresis.

185 All the isolates were subjected to PCR with the verified primer pairs, coupled with High Resolution 186 Melting (HRM) analysis, to identify polymorphic loci and to determine isolate specific allelic 187 variation at such loci. The DNA of each sample was quantified by using the NanoDrop (Thermo 188 Scientific, Wilmington, DE, USA) in order to use a standard concentration of template DNA in PCR 189 reactions. The PCR for the HRM analysis was carried out with Connect<sup>™</sup> Real-Time PCR Detection 190 System (Bio-Rad Laboratories, Hercules, CA, USA). Each PCR reaction was conducted on a total 191 volume of 10 µl, containing 1 µl of appropriately diluted DNA (20 ng/µl), 5 µl Sso Fast Eva Green 192 Supermix (Bio-Rad Laboratories, Hercules, CA, USA), 0.3 µl of each primer (3 µM) and 3.4 µl of 193 water, using a 96 well plate. The following PCR programme, which included the calculation of a 194 melting curve, was used: 98°C for 2 minutes, followed by 45 cycles of 98°C for 10 seconds and 59°C for 20 seconds, and terminated by ramp from 65°C to 95°C with a temperature increment of 0.1°C 195

196 and a plate read every 10 seconds. Melting curves were analysed by using the Precision Melt Analysis<sup>™</sup> Software from Bio-rad, setting the Tm difference threshold at 0.15 and the Melt curve 197 198 shape sensitivity at 50. This software allowed to group the melting curves in different clusters, 199 representing different alleles of SSR loci. By analyzing the melting curves, alleles were assigned to 200 all isolates for each SSR locus and a matrix including all the allelic data obtained was prepared. 201 Number of observed alleles *per* locus (Na), number of effective alleles based on their frequencies 202 (Ne) and gene diversity (h) were estimated from the data matrix using the software GenAlEx version 203 6.5 (Peakall & Smouse, 2012). The software was also used for calculating the overall observed and 204 expected levels of heterozygosity based on the number of melting curve peaks and effective number 205 of alleles detected per locus, respectively.

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## 207 In vitro growth assays and phenotypic diversity of isolates

Growth assays were performed by inoculating a plug of mycelium (6 mm in diameter) obtained from the margin of actively growing cultures of each Sicilian isolate on the centre of 90-mm Petri dishes containing MEA. Five replicates for each isolate were prepared and Petri dishes were incubated at  $25^{\circ}C\pm2^{\circ}C$  in the dark. The extent of fungal colonization in mm was measured every 48 hours until the mycelia reached the edges of the Petri dish. Two perpendicular measures of radial growth for each replicate were taken. Observations on the colony pigmentation were conducted on 8-days old pure cultures of Sicilian isolates grown at  $25^{\circ}C$  on Petri dishes filled with MEA.

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## 216 Statistical analyses

The software GeneAlEx version 6.5 was used to perform the Principal Coordinates Analysis (PCoA) on both genetic (matrix of genetic distances between isolates) and phenotypic (matrix obtained from pairwise comparison of growth curves) data matrices. The same program was used to assess the minimum number of SSR loci needed to discriminate all genotypes, through a genotype accumulation curve of multi-locus match probability for increasing combinations of loci. The XLSTAT<sup>©</sup> software

222	package (Addinsoft, Paris, France) was used to assess the association between spatial and genetic
223	distances, by performing Spearman's rank correlation test. The XLSTAT <sup>©</sup> software package was also
224	used to assess the association between genetic distance and growth coefficients pairwise by
225	performing the Spearman's rank correlation test. Significant ( $p < 0.05$ ) differences among isolates in
226	terms of growth in vitro were assessed by using the compareGrowthCurves function
227	(http://bioinf.wehi.edu.au/software/compareCurves/) from the R Statistical Modeling package
228	statmod. This function performed permutation tests (1000 permutations) of the differences between
229	groups in growth rate.

- 230
- 231 **Results**
- 232
- 233 Incidence of *L. sulphureus* in carob trees

The use of taxon-specific primers detected *L. sulphureus* in 24 samples out of 70, corresponding to a 34.28% incidence. The positive samples, showing a typical band of 146 bp, were obtained from 11 trees showing symptoms of decay and from 13 asymptomatic trees.

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### 238 Phylogenetic analysis of *L. sulphureus* isolates

Phylogenetic analysis based on ITS sequences of Laetiporus spp. included the now generated ITS 239 sequences and ITS sequences of reference Laetiporus spp. from different geographic origins and 240 241 different hosts. In the first phylogenetic tree, the 25 Sicilian isolates clustered inside the cluster E: 17 242 grouped in the sub-cluster E1, and seven in the sub-cluster E2 (Figure 2). The ITS sequence of isolate 243 C1 showed five additional SNPs compared to the other isolates and was considered as belonging to 244 E1. The four additional Italian isolates grouped in the sub-cluster E2 (Figure 2). In the second phylogenetic tree, which included only isolates of L. sulphureus, the Italian isolates grouped in the 245 246 cluster E, which contained isolates obtained from different host trees, i.e. Eucalyptus sp., Fraxinus sp., *Prunus* sp. and *Quercus* sp., in addition to those from carob (Figure 3).

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# Development of SSR-based markers, HRM genotyping and analysis of genetic diversity of isolates

Eleven out of 20 tested SSR markers were amplifiable through PCR and hence were used in the HRM genotyping. One locus turned out to be monomorphic as it showed no variation in melting curves between the samples and hence it was excluded from analysis. The remaining ten SSR markers were polymorphic based on HRM analysis, showing a minimum of two alleles (locus Ls\_GGA\_174) and a maximum of ten alleles (locus Ls\_ATGCCC\_111) (Figure 4). Primers developed and used to amplify the polymorphic SSR markers are reported in Table 2.

257 Melting curve difference plots for the ten polymorphic loci are visualized in Figure 4. Number of 258 alleles and number of effective alleles based on allele frequencies for each locus are reported in Table 259 2. All SSR loci analysed in this study were homozygous in all isolates, with the exception of locus 260 Ls AC 322 for which isolates F3 and F6 showed a typical double peak in the melting curve plots, 261 suggesting a heterozygous condition (Figure S1) at this locus. For this locus, these two isolates were 262 grouped in a separate cluster/allele in the subsequent HRM analysis. The observed and expected levels of heterozygosity were 0.007 and 0.581, respectively. Average gene diversity (h) was 0.581 ( $\pm$ 263 0.079). The HRM genotyping identified 29 genotypes out of 29 isolates. Multilocus matches by locus 264 265 for increasing combinations of all the ten loci showed that the minimum number of SSR markers allowing distinction between all isolates was nine. 266

The PCoA based on genetic data did not show distinct groups (Figure 5). The percentage of variation explained by the first two axes was 34.68%. A slight distinction between isolates belonging to E1 and E2 sub-clusters was observed, with the exception of isolate C1 (Figure 5). The isolates obtained from chestnut (GAIOLA, Em-a, Em-b), oak (VALLERE) and almond (G5) grouped together with the other isolates (Figure 5). The Spearman's rank correlation test between pairwise genetic and geographical distances within the 25 Sicilian isolates showed a significant positive correlation (0.142, *p*-value = 273 0.014).

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## 275 Growth assay *in vitro* and analysis of phenotypic diversity

276 Mycelial growth assessed in vitro showed that six isolates (C2, D4, D5, O2, O3 and F5) were able to fully colonize the Petri dish in 6 days only (Table 3). Isolates C1, C4 and F1 did not reach the edge 277 278 of the Petri dish after 8 days (Table 3). PCoA on mycelial growth data allowed to distinguish four 279 different groups: one group representing the fastest growing isolate O3, two intermediate groups 280 including 10 and 12 isolates, respectively, and a group including the isolates C1 and F1 characterized 281 by low growth (Figure 6). The percentage of variation explained by the first two axes was 66.41%. 282 Statistical analysis performed with *compareGrowthCurves* function showed significant differences 283 among the four clusters identified (p-value < 0.05). The Spearman's rank correlation test between 284 pairwise genetic distances and pairwise mycelial growth coefficients showed no significant 285 correlation (p-value > 0.05).

A slight difference in pigmentation between Sicilian isolates was observed. In particular, 11 isolates (C1, C3, D3, D5, F1, F4, F5, G4, G5, O1 and O3) showed white mycelia, while the remaining 14 isolates (C2, C4, D1, D4, F2, F3, F6, G1, G2, G3, G6, G7, I1 and O2) showed yellow-to-orange mycelia.

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## 291 Discussion

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This epidemiological study on *L. sulphureus* associated with carob trees has provided insights into the presence and the distribution of this wood decay agent in the South West of Sicily. In this work, 34% of the sampled trees were found to be infected by *L. sulphureus*, based on molecular detection. The high occurrence frequency of this species, generally diagnosed based on the inspection of visible fruiting bodies, implies that the prevalence of this species is largely underestimated as previously observed in surveys conducted in urban areas (Giordano et al., 2015). None of the sampled carob 299 trees showed visible fruiting bodies, which may be partly due to the period of sampling (i.e. 300 December). The majority of trees (66%) did not show any external symptoms of wood decay either. 301 From an ecological perspective, L. sulphureus has long been regarded as belonging to the group of 302 true heart rots colonizing heartwood exposed by natural injuries (Vasiliauskas, Sunhede, & Stenlid, 303 2003; Vasaitis, 2013). The now observed prevalent incidence of L. sulphureus in asymptomatic carob 304 trees might suggest that the fungus could be considered as a true heart rot of intact tree stems, when 305 following the classification of Vasaitis (2013). However, it should be noted that unravelling the 306 ecological strategies of L. sulphureus was beyond the scope of our study. Further research with an 307 experimental design tailored for the purpose is needed to understand whether L. sulphureus may 308 belong to the category of true heart rots of intact tree stems, which would imply an ability to colonize 309 trees without wounding.

310 The widespread presence of L. sulphureus on carob trees may have been favoured by the 311 abandonment of cultivation of this tree species in the area. In the recent past, carob pods prices 312 dramatically decreased, making the cultivation of carob trees no longer economically profitable 313 compared to other crops, such as vineyards (Barbera & Cullotta, 2012). The cultivation of this woody 314 crop has indeed radically decreased from about 70,000 ha during the 60's to less than 30,000 ha by 315 early 21st Century (Massa & La Mantia, 2007). As a consequence, the abandonment of agricultural 316 practices on Sicilian carob trees, such as removal of decayed wood residuals, may have favoured the 317 establishment and spread of L. sulphureus in this area.

The prevalent presence of *L. sulphureus* fruiting bodies on several trees in the Modica municipality allowed to perform a genetic diversity analysis of this pathogen in the area. Previous phylogenetic analyses on ITS region of *Laetiporus* isolates from different countries have showed that the species *L. sulphureus* includes at least two distinct clusters, C and E (Vasaitis et al., 2009; Song et al., 2014). The cluster C included European isolates only, while cluster E comprised isolates originating from different European, South American, and North American countries (Vasaitis et al., 2009). Our phylogenetic analysis showed that all Italian isolates grouped within the cluster E. Seventeen isolates 325 from South Western Sicily were grouped in the sub-cluster E1, while the remaining 12 isolates, 326 including isolates from other Italian regions, i.e. Tuscany and Piedmont, clustered in the sub-cluster 327 E2. These two sub-clusters were previously defined on the basis of phenotypical observations, i.e. 328 colours of the fruiting body pores (white or yellow) (Burdsall Jr & Banik, 2001; Song et al., 2014). Our phylogenetic analysis on ITS regions confirmed the absence of a link between cluster and host 329 330 species, as previously observed (Vasaitis et al., 2009). Isolates obtained from fruiting bodies collected 331 from carob trees, European chestnuts, oak and almond clustered together with other European isolates 332 collected from different host plants, such as ashes and eucalypts.

333 In order to increase the resolution of genetic characterization, novel molecular markers specific for 334 L. sulphureus were developed, by using large scale genomic data (Nagy et al., 2015). SSR-based 335 markers are widely used and popular due to their high reproducibility and multiallelic nature, and 336 their power for genetic characterization of populations of wood decay fungi has been demonstrated 337 (Franzen et al., 2007; Travadon et al., 2012; Maurice et al., 2014; Gonthier et al., 2015). The analysis 338 of SSRs coupled with HRM is a robust and reproducible method (Ganopoulos, Argiriou, & Tsaftaris, 339 2011), as it has been successfully used in genotyping of plants (Xanthopoulou et al., 2014; DiStefano 340 et al., 2012), and, more recently, of fungal pathogens (Zambounis et al., 2016; Sillo et al., 2017). The 341 HRM genotyping of the ten SSR markers allowed to distinguish all isolates from one another. In 342 addition, this analysis allowed detection of heterozygous allelic conditions, often observed in 343 heterokaryotic basidiomycetes (Nazrul & YinBing, 2011), yet uncommon in the L. sulphureus isolates 344 analysed in this study. The relatively high number of observed alleles per locus and the high gene 345 diversity (0.581) are not in disagreement with the hypothesis that L. sulphureus retains high allelic 346 recombination due to sexual reproduction (Milgroom, 1996), although the fungus is also known to 347 develop an asexual stage (Stalpers, 1978; Bernicchia, 2005). However, the observed level of heterozygosity was substantially lower than the expected one, which might indicate that L. 348 349 sulphureus, or at least the studied population, is homothallic, a condition previously suggested for 350 other Laetiporus taxa, but not for L. sulphureus (Banik & Burdsall Jr, 2000; Banik et al., 2010).

351 The PCoA of genetic data did not divide the samples into distinct groups, but there was a slight 352 differentiation between isolates from sub-cluster E1 and E2. Interestingly, the genotype C1, belonging 353 to sub-cluster E1, was more similar to genotypes of sub-cluster E2. This genotype also showed the 354 lower mycelial growth in the *in vitro* assay. It could be hypothesized that this genotype may putatively belong to an additional sub-cluster, since it showed a) several polymorphisms in the ITS region 355 356 compared to the other Sicilian isolates, b) a peculiar SSR profile, and c) a phenotype different from the others. PCoA also showed the absence of correlation between L. sulphureus genotypes and host 357 358 plants. These findings confirm the results of the single-locus phylogenetic analysis based on ITS and 359 support the hypothesis that L. sulphureus genotypes may infect a broad spectrum of host plants.

The Spearman's rank correlation test between genetic and geographical pairwise distances showed a significant positive correlation (*p*-value < 0.05). This outcome, along with the fact that high gene diversity was observed within isolates, may provide evidence that *L. sulphureus* spreads and infects trees through basidiospores, as previously hypothesized (Schwarze, Engels, & Mattheck, 2000) and as documented for other wood decay fungi (Gonthier et al., 2012; Travadon et al., 2012; Sillo et al., 2016).

366 The phenotypic characterization of isolates through the *in vitro* growth assay allowed to distinguish 367 at least four different groups within Sicilian isolates, based on the differences of mycelial growth. 368 Vasaitis et al. (2009) reported that L. sulphureus isolates belonging to the cluster E were faster in 369 growth than isolates in cluster C (Vasaitis et al., 2009). In this work, cluster E showed isolate specific 370 variation in mycelial growth rate, suggesting a high phenotypic diversity. The phenotypic diversity 371 within the cluster E was also displayed by differences in the pigmentation of pure cultures (Figure 372 S2). In fact, although no clear differences in the pigmentation among isolates belonging to the two 373 sub-clusters E1 and E2 were observed, pure cultures of Sicilian isolates as a whole ranged from white 374 to yellow-to-orange in colour. The results of the *in vitro* growth assays may mirror the saprotrophic ability of the fungus on wood, as suggested by preliminary results of a growth assay of selected 375 376 Sicilian isolates on wood substrates (Gianchino, 2017). Interestingly, a significant correlation 377 between growth on artificial media and on wood substrates has been recently documented for 378 *Armillaria ostoyae* (Romagn.) Herink (Labbé et al., 2017). Phenotypic assays *in vitro* may thus be 379 useful to determine the potential decaying ability of *L. sulphureus* isolates inside the trees, an aspect 380 that may be useful when considering the most appropriate management measures.

381 No correlation was observed between genetic and phenotypic data in terms of mycelial growth rates 382 *in vitro* (Spearman's rank correlation test, *p*-value > 0.05). The phenotypic diversity of isolates may 383 be the result of adaptive evolution to the environment mediated by epigenetic changes, as recently 384 demonstrated for other plant pathogens (Dubey & Jeon, 2016). Alternatively, it could be due to 385 genetic polymorphisms affecting loci different from those analysed in this study.

386 In conclusion, this work allowed us to determine the incidence of L. sulphureus on carob in selected 387 areas of Sicily, to develop and use ten SSR markers, specific for L. sulphureus, for a precise 388 genotyping of isolates through HRM analysis, and to assess the phenotypic diversity through simple 389 in vitro assays. Results showed not only that over one third of carob trees are affected by L. sulphureus 390 in the area, but also suggest a risk of transmission of the pathogen from one tree species to another 391 through basidiospores as inferred from data of genetic diversity and on the lack of correlation between 392 fungal genotype and host trees. In addition, the high phenotypic variability observed in vitro may 393 reflect a similar variation in natural environment, highlighting the importance of a phenotypic 394 characterization of isolates affecting the trees. From a practical perspective, the pruning/removal of 395 decayed wood, which can serve as substrate for L. sulphureus and a source for the emergence of 396 fruiting bodies, could minimize the spread of this wood decay agent in carob orchards.

397

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401

## 402 **References**

404

405	of Laetiporus sulphureus sensu lato. Mycologia, 92, 649–655.
406	
407	Banik, M. T., Lindner, D. L., Ota, Y., & Hattori, T. (2010). Relationships among North American and
408	Japanese Laetiporus isolates inferred from molecular phylogenetics and single-spore incompatibility
409	reactions. Mycologia, 102, 911–917.
410	
411	Barbagallo, M. G., Di Lorenzo, R., Meli, R., & Crescimanno, F. G. (1997). Characterization of carob
412	germplasm (Ceratonia siliqua L.) in Sicily. Journal of Horticultural Science, 72, 537–543.

Banik, M. T., & Burdsall Jr, H. H. (2000). Incompatibility groups among North American populations

413

Barbera, G., & Cullotta, S. (2012). An inventory approach to the assessment of main traditional
landscapes in Sicily (Central Mediterranean Basin). *Landscape Research*, *37*, 539–569.

416

Battle, I., & Tous, J. (1997). Carob tree: *Ceratonia siliqua* L. – Promoting the conservation and use
of underutilized and neglected crops. 17. Institute of Plant Genetics and Crop Plant Research,

419 Gatersleben/International Plant Genetic Resources Institute, Rome, Italy.

420

421 Bernicchia, A. (2005). Polyporaceae s.l. Ed. Candusso, Alassio, Italy.

422

Burdsall Jr, H. H., & Banik, M. T. (2001). The genus *Laetiporus* in North America. *Harvard Papers in Botany*, 6, 43–55.

425

426 Dai, Y. C., Cui, B. K., Yuan, H. S., & Li, B. D. (2007). Pathogenic wood–decaying fungi in China.
427 *Forest Pathology*, *37*, 105–120.

429	Distefano, G., Caruso, M., La Malfa, S., Gentile, A., & Wu, S. B. (2012). High resolution melting
430	analysis is a more sensitive and effective alternative to gel-based platforms in analysis of SSR-an
431	example in citrus. <i>PloS ONE</i> , 7, e44202.

Du, L., Li, Y., Zhang, X., & Yue, B. (2013). MSDB: a user–friendly program for reporting distribution
and building databases of microsatellites from genome sequences. *Journal of Heredity*, *104*, 154–
157.

- 437 Dubey, A., & Jeon, J. (2016). Epigenetic regulation of development and pathogenesis in fungal plant
  438 pathogens. *Molecular Plant Pathology*, *18*, 887–898.
- 439
- 440 Franzen, I., Vasaitis, R., Penttilä, R., & Stenlid, J. (2007). Population genetics of the wood–decay
  441 fungus *Phlebia centrifuga* P. Karst. in fragmented and continuous habitats. *Molecular Ecology*, *16*,
  442 3326–3333.
- 443
- Ganopoulos, I., Argiriou, A., & Tsaftaris, A. (2011). Microsatellite high resolution melting (SSR–
  HRM) analysis for authenticity testing of protected designation of origin (PDO) sweet cherry
  products. *Food Control*, 22, 532–541.
- 447
- Gardes, M., & Bruns, T. D. (1993). ITS primers with enhanced specificity for basidiomycetes
  application to the identification of mycorrhizae and rusts. *Molecular Ecology*, *2*, 113–118.
- 450
- 451 Gianchino, C. (2017). Incidenza e diversità genetica e fenotipica del fungo lignivoro *Laetiporus*452 *sulphureus* in carrubeti della Sicilia. Master thesis, University of Bologna, Bologna, Italy.
- 453
- 454 Gilbertson, R. L., & Ryvarden, L. (1986). North American polypores 1. Fungiflora, Oslo, Norway.

- Giordano, L., Sillo, F., Guglielmo, F., & Gonthier, P. (2015). Comparing visual inspection of trees
  and molecular analysis of internal wood tissues for the diagnosis of wood decay fungi. *Forestry: An International Journal of Forest Research*, 88, 465–470.
- 459
- Gonthier, P., Lione, G., Giordano, L., & Garbelotto, M. (2012). The American forest pathogen *Heterobasidion irregulare* colonizes unexpected habitats after its introduction in Italy. *Ecological Applications*, 22, 2135–2143.
- 463
- Gonthier, P., Sillo, F., Lagostina, E., Roccotelli, A., Cacciola, S. O., Stenlid, J., & Garbelotto, M.
  (2015). Selection processes in simple sequence repeats suggest a correlation with their genomic
  location: insights from a fungal model system. *BMC Genomics*, *16*, 1107.
- 467
- Guglielmo, F., Bergemann, S. E., Gonthier, P., Nicolotti, G., & Garbelotto, M. (2007). A multiplex
  PCR-based method for the detection and early identification of wood rotting fungi in standing trees. *Journal of Applied Microbiology*, *103*, 1490–1507.
- 471
- 472 Guglielmo, F., Gonthier, P., Garbelotto, M., & Nicolotti, G. (2010). Optimization of sampling
  473 procedures for DNA-based diagnosis of wood decay fungi in standing trees. *Letters in Applied*474 *Microbiology*, *51*, 90–97.
- 475
- 476 La Malfa, S., Currò, S., Douglas, A. B., Brugaletta, M., Caruso, M., & Gentile, A. (2014). Genetic
  477 diversity revealed by EST–SSR markers in carob tree (*Ceratonia siliqua* L.). *Biochemical Systematics*478 *and Ecology*, 55, 205–211.
- 479
- 480 Labbé, F., Lung-Escarmant, B., Fievet, V., Soularue, J. P., Laurent, C., Robin, C., & Dutech, C.

- 481 (2017). Variation in traits associated with parasitism and saprotrophism in a fungal root–rot pathogen
  482 invading intensive pine plantations. *Fungal Ecology*, *26*, 99–108.
- 483
- Lindner, D. L., & Banik, M. T. (2008). Molecular phylogeny of *Laetiporus* and other brown rot
  polypore genera in North America. *Mycologia*, *100*, 417–430.
- 486
- 487 Lindner, D. L., & Banik, M. T. (2011). Intragenomic variation in the ITS rDNA region obscures
  488 phylogenetic relationships and inflates estimates of operational taxonomic units in genus *Laetiporus*.
  489 *Mycologia*, 103, 731–740.
- 490
- Massa, B., & La Mantia, T. (2007). Forestry, pasture, agriculture and fauna correlated to recent
  changes in Sicily. *Forest@–Journal of Silviculture and Forest Ecology*, *4*, 418.
- 493
- Maurice, S., Skrede, I., LeFloch, G., Barbier, G., & Kauserud, H. (2014). Population structure of *Serpula lacrymans* in Europe with an outlook to the French population. *Mycologia*, *106*, 889–895.
- 496
- Milgroom, M. G. (1996). Recombination and the multilocus structure of fungal populations. *Annual Review of Phytopathology*, *34*, 457–477.
- 499
- 500 Nagy, L. G., Riley, R., Tritt, A., Adam, C., Daum, C., Floudas, D., Sun, H., Yadav, J. S., Pangilinan,
- 501 J., Larsson, K. H., Matsuura, K., Barry, K., Labutti, K., Kuo, R., Ohm, R. A., Bhattacharya, S. S.,
- 502 Shirouzu, T., Yoshinaga, Y., Martin, F. M., Grigoriev, I. V., & Hibbett, D. S. (2015). Comparative
- 503 genomics of early-diverging mushroom-forming fungi provides insights into the origins of
- 504 lignocellulose decay capabilities. *Molecular Biology and Evolution*, 33, 959–970.
- 505
- 506 Nazrul, M. I., & YinBing, B. (2011). Differentiation of homokaryons and heterokaryons of Agaricus

507 *bisporus* with inter-simple sequence repeat markers. *Microbiological research*, 166, 226–236.

508

509 Peakall, P. E., & Smouse, R. (2012). GenAlEx 6.5: genetic analysis in Excel. Population genetic
510 software for teaching and research—an update. *Bioinformatics*, 28, 2537–2539.

- Petrović, J., Glamočlija, J., Stojković, D. S., Ćirić, A., Nikolić, M., Bukvički, D., Guerzoni, M. E., &
  Soković, M. D. (2013). *Laetiporus sulphureus*, edible mushroom from Serbia: Investigation on
  volatile compounds, *in vitro* antimicrobial activity and *in situ* control of *Aspergillus flavus* in tomato
  paste. *Food and Chemical Toxicology*, *59*, 297–302.
- 516
- 517 Petrović, J., Stojković, D., Reis, F. S., Barros, L., Glamočlija, J., Ćirić, A., Ferreira, I. C., & Soković
  518 M. (2014). Study on chemical, bioactive and food preserving properties of *Laetiporus sulphureus*519 (Bull.: Fr.) Murr. *Food & Function*, *5*, 1441–1451.
- 520
- 521 Pleszczyńska, M., Wiater, A., Siwulski, M., & Szczodrak, J. (2013). Successful large–scale
  522 production of fruiting bodies of *Laetiporus sulphureus* (Bull.: Fr.) Murrill on an artificial substrate.
  523 *World Journal of Microbiology and Biotechnology*, 29, 753–758.
- 524
- Ramón-Laca, L., & Mabberley, D. J. (2004). The ecological status of the carob-tree (*Ceratonia siliqua*, Leguminosae) in the Mediterranean. *Botanical Journal of the Linnean Society*, 144, 431–436.
- 527
- Rogers, S. O., Holdenrieder, O., & Sieber, T. N. (1999). Intraspecific comparisons of *Laetiporus sulphureus* isolates from broadleaf and coniferous trees in Europe. *Mycological Research*, *103*, 1245–
  1251.
- 531
- 532 Schwarze, F. W., Engels, J., & Mattheck, C. (2000). Fungal strategies of wood decay in trees.

533 Springer–Verlag Berlin Heidelberg, Heidelberg, Germany.

Sillo, F., Savino, E., Giordano, L., Girometta, C., Astegiano, D., Picco, A. M., & Gonthier, P. (2016).
Analysis of genotypic diversity provides a first glimpse on the patterns of spread of the wood decay
fungus *Perenniporia fraxinea* in an urban park in Northern Italy. *Journal of Plant Pathology*, *98*,
617–624.

- 539
- Sillo, F., Giordano, L., Zampieri, E., Lione, G., De Cesare, S., & Gonthier, P. (2017). HRM analysis
  provides insights on the reproduction mode and the population structure of *Gnomoniopsis castaneae*in Europe. *Plant Pathology*, *66*, 293–303.
- 543
- Song, J., & Cui, B. K. (2017). Phylogeny, divergence time and historical biogeography of *Laetiporus*(Basidiomycota, Polyporales). *BMC Evolutionary Biology*, *17*, 102.
- 546
- Song, J., Chen, Y., Cui, B., Liu, H., & Wang, Y. (2014). Morphological and molecular evidence for
  two new species of *Laetiporus* (Basidiomycota, Polyporales) from southwestern China. *Mycologia*, *106*, 1039–1050.
- 550
- Stalpers, J. A. (1978). Identification of wood–inhabiting Aphyllophorales in pure culture. *Studies in Mycology*, *16*, 1–248.
- 553
- Tamura, K., Stecher, G., Peterson, D., Filipski, A., & Kumar, S. (2013). MEGA6: molecular
  evolutionary genetics analysis version 6.0. *Molecular Biology and Evolution*, *30*, 2725–2729.
- 556
- 557 Travadon, R., Smith, M. E., Fujiyoshi, P., Douhan, G. W., Rizzo, D. M., & Baumgartner, K. (2012).
- 558 Inferring dispersal patterns of the generalist root fungus Armillaria mellea. New Phytologist, 193,

559 959–969.

560

Turkoglu, A., Duru, M. E., Mercan, N., Kivrak, I., & Gezer, K. (2007). Antioxidant and antimicrobial
activities of *Laetiporus sulphureus* (Bull.) Murrill. *Food Chemistry*, *101*, 267–273.

563

Vannini, A., Contarini, M., Faccoli, M., Valle, M. D., Rodriguez, C. M., Mazzetto, T., Guarneri, D.,
Vettraino, A. M., & Speranza, S. (2017). First report of the ambrosia beetle *Xylosandrus compactus*and associated fungi in the Mediterranean maquis in Italy, and new host–pest associations. *EPPO Bulletin*, 47, 100–103.

568

- Vasaitis, R., Menkis, A., Lim, Y. W., Seok, S., Tomsovsky, M., Jankovsky, L., Lygis, V., Slippers, B.,
  & Stenlid, J. (2009). Genetic variation and relationships in *Laetiporus sulphureus* s. lat., as
  determined by ITS rDNA sequences and in vitro growth rate. *Mycological Research*, *113*, 326–336.
- Vasaitis, R. (2013). Heart rots, sap rots and canker rots. In: *Infectious forest diseases*. Ed. by Gonthier,
  P. & Nicolotti, G. Wallingford: CAB International, pp. 197–229.
- 575
- Vasiliauskas, R., Sunhede, S., & Stenlid, J. (2003). Distribution, status and biology of oak polypores
  in Baltic Sea region. In: *Forest Health Problems in Older Forest Stands. Proceedings of the Nordic/Baltic Forest Pathology Meeting, Denmark, September 2002.* Ed. I.M., Thomsen. Danish
  Forest and Landscape Research Institute, Horsholm, pp. 61–66.

580

581 Wiater, A., Pleszczyńska, M., Szczodrak, J., & Janusz, G. (2012). Comparative studies on the 582 induction of *Trichoderma harzianum* mutanase by  $\alpha$ -(1 $\rightarrow$ 3)–glucan–rich fruiting bodies and mycelia 583 of *Laetiporus sulphureus*. *International Journal of Molecular Sciences*, *13*, 9584–9598.

586	Madesis, P. (2014). Microsatellite high-resolution melting (SSR-HRM) analysis for genotyping and
587	molecular characterization of an Olea europaea germplasm collection. Plant Genetic Resources, 12,
588	273–277.
589	
590	Zambounis, A., Xanthopoulou, A., Karaoglanidis, G., Tsaftaris, A., & Madesis, P. (2016). A new
591	accurate genotyping HRM method for Alternaria species related to fruit rot diseases of apple and
592	pomegranate. International Journal of Phytopathology, 4, 159–165.
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601	Figure legends
602	
603	FIGURE 1. Map of the sampling area. Gray triangles represent sampled carob trees. Black circles
604	represent locations where fungal fruiting bodies were collected. Map was created and customized by
605	using TileMill ( <u>https://tilemill-project.github.io/tilemill/</u> ).
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Xanthopoulou, A., Ganopoulos, I., Koubouris, G., Tsaftaris, A., Sergendani, C., Kalivas, A., &

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FIGURE 2. Phylogenetic tree including ITS sequences of different *Laetiporus* taxa. The sequences
were aligned using ClustalW in MEGA v. 6 (Tamura et al., 2013) and the tree (unrooted) was

constructed using the Maximum Likehood (ML) method. Numbers indicate bootstrap values. Only
taxonomically described *Laetiporus* species were represented. The four undescribed taxa (*Laetiporus*sp.) from South Africa, Hawaii, Costa Rica and Argentina, corresponding to cluster H, I, L and M,
respectively, were not included in the tree.

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FIGURE 3. Phylogenetic tree including ITS sequences of *Laetiporus sulphureus*. The sequences were aligned using ClustalW in MEGA v. 6 (Tamura et al., 2013) and the tree (unrooted) was constructed using the Maximum Likehood (ML) method. Numbers indicate bootstrap values. For each sequence, information on geographic origin and host tree are provided. Abbreviations: AUS, Austria; CAN, Canada; CZE, Czech Republic; DEN, Denmark; ITA, Italy; LAT, Latvia; LIT, Lithuania; SPA, Spain; SWE, Sweden; URU, Uruguay.

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FIGURE 4. Melting curve difference plots after normalization and overlay for the ten polymorphic
SSR loci. Differences in terms of relative fluorescence were obtained using the Precision Melt
Analysis<sup>™</sup> Software. Each curve represent a different allele. (a) Ls\_A\_345; (b) Ls\_AC\_322; (c)
Ls\_ATGCCC11; (d) Ls\_CAG\_633; (e) Ls\_CAG\_159; (f) Ls\_G\_413; (g) Ls\_GAG\_238; (h)
Ls\_GCA\_174; (i) Ls\_GGA\_479; (l) Ls\_TTC\_500.

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FIGURE 5. Results of PCoA on genetic data generated by the SSR-HRM genotyping. Gray squares
 represent isolates belonging to *L. sulphureus* sub-cluster E1, while black diamond represent isolates
 belonging to sub-cluster E2.

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631 FIGURE 6. Results of PCoA on phenotypic data obtained from the in vitro growth assay.
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Isolate ID Code	Host	Geographic origins	Latitude	Longitude
C1	Ceratonia siliqua	Sicily (Italy)	36.89825	14.86479
C2	Ceratonia siliqua	Sicily (Italy)	36.92324	14.74267
C3	Ceratonia siliqua	Sicily (Italy)	36.89288	14.88271
C4	Ceratonia siliqua	Sicily (Italy)	36.94353	14.87465
D1	Ceratonia siliqua	Sicily (Italy)	36.92010	14.82380
D3	Ceratonia siliqua	Sicily (Italy)	36.94248	14.88063
D4	Ceratonia siliqua	Sicily (Italy)	36.91627	14.74451
D5	Ceratonia siliqua	Sicily (Italy)	37.02037	14.71584
F1	Ceratonia siliqua	Sicily (Italy)	36.91437	14.86937
F2	Ceratonia siliqua	Sicily (Italy)	36.81274	14.84336
F3	Ceratonia siliqua	Sicily (Italy)	36.94810	14.88038
F4	Ceratonia siliqua	Sicily (Italy)	37.04736	14.78326
F5	Ceratonia siliqua	Sicily (Italy)	36.92972	14.74760
F6	Ceratonia siliqua	Sicily (Italy)	37.13120	14.85319
Gl	Ceratonia siliqua	Sicily (Italy)	36.92437	14.83428
G2	Ceratonia siliqua	Sicily (Italy)	36.92519	14.84561
G3	Ceratonia siliqua	Sicily (Italy)	36.94364	14.92788
G4	Ceratonia siliqua	Sicily (Italy)	36.93788	14.73893
G5	Prunus amygdalus	Sicily (Italy)	36.91716	14.74974
G6	Ceratonia siliqua	Sicily (Italy)	36.95976	14.83583
G7	Ceratonia siliqua	Sicily (Italy)	36.97077	14.84270
I1	Ceratonia siliqua	Sicily (Italy)	36.92116	14.87124
01	Ceratonia siliqua	Sicily (Italy)	36.98445	14.83350
02	Ceratonia siliqua	Sicily (Italy)	36.92574	14.81369
03	Ceratonia siliqua	Sicily (Italy)	36.95476	14.88278
Em-A	Castanea sativa	Tuscany (Italy)	-	-
Em-B	Castanea sativa	Tuscany (Italy)	-	-
GAIOLA	Castanea sativa	Piedmont (Italy)	-	-
VALLERE	Quercus sp.	Piedmont (Italy)	-	-
L		1		

 Table 1. List of L. sulphureus isolates used in this study.

Table 2. SSR markers developed in the study with related allelic diversity. Sequence and ID of
primers, number of alleles (Na), number of effective alleles based on their frequencies (Ne) and gene
diversity (*h*) per SSR locus are shown.

Locus	Sequence of the primer	ID primer	Na	Ne	h
Ls_GAG_238	GCAAGGCTAAGGTGTGTCCA	laets_GAG_238_f	3	2.129	0.530
	TCCTCTTCCTCTGCCAATTC	laets_GAG_238_r			
Ls CAG 633	ATATGCCCTCCAATGAGCAG	laets_GAG_238_r	5	3.461	0.711
	ATGAACGATCCGTTCTGCAA	laets_CAG_633_f			
Ls GGA 174	TCACGGAGAGCATGAGACTG	laets_GGA_174_f	2	1.071	0.067
	GTGTCCATCTCGTCCAGGTT	laets_GGA_174_r			
Ls CGA 159	TCCGCCATTCAACTTAACAA	laets_CGA_159_f	7	3.948	0.747
	TGACGTTGTACTCGGATGGA	laets_CGA_159_r			
Ls AC 322	TCAGGTGCACTTTCTGTCCTT	laets_AC_322_f	6	3.267	0.694
	GGCTGCTCATCCTGTAGGG	laets_AC_322_r			
Ls_A_345	TGAGACTCAGGGGAAGGAGA	laets_A_345_f	7	4.918	0.797
	GATTTCCTGAACCCGATAACC	laets_A_345_r			
Ls_G_413	TTGAGCAACCTGTTGAGTGG	laets_G_413_f	4	3.461	0.711
	GCTCTCTGTTCCGTGTCTCC	laets_G_413_r			
Ls_GGA_174 Ls_CGA_159 Ls_AC_322 Ls_A_345 Ls_G_413 Ls_GGA_479 Ls_TTC_500 Ls_ATGCCC_111	TGAACGTGAGGCAGATCAAG	laets_GGA_479_f	3	1.532	0.347
	GCGGTCTCAGCATAAAGTCC	laets_GGA_479_r	ts_GGA_479_r		
Ls TTC 500	CCCTCAAACGTCTCCACATT	laets_TTC_500_f	6	1.566	0.361
	AGAGCGTCAGCAAGGAAGAC	laets_TTC_500_r			
Ls_ATGCCC_111	CACGCTCTGCACTACACCAT	laets_ATGCCC_111_f	10	6.622	0.849
	GAGGAGTCCGAGTCATACGAA	laets_ATGCCC_111_r			
			Na	Ne	h
		Average	5.300	3.198	0.581
		Standard Error	0.761	0.543	0.079

Table 3. Mycelial radial growth expressed in mm of colonization of agar medium in Petri dishes of
Sicilian isolates at different times (days after inoculation). For each isolate, average of measures of
the five replicates and standard deviation (SD) are shown.

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Isolate ID Code	Radius of fungal colony (mm) ± SD				
	After 2 days	After 4 days	After 6 days	After 8 days	
C1	4.75±2.24	20.70±2.30	33.60±1.30	38.00±1.22	
C2	7.05±2.19	29.50±5.24	38.50±1.29	40.00±0.00*	
C3	7.35±1.20	24.90±2.17	39.10±0.90	40.00±0.00*	
C4	6.65±4.28	25.90±0.84	37.10±2.26	38.30±1.27	
D1	7.30±3.42	32.30±8.96	39.53±0.51	40.00±0.00*	
D3	8.10±2.17	28.60±2.17	36.50±2.24	40.00±0.00*	
D4	8.25±1.50	30.20±1.52	40.00±0.00*	40.00±0.00*	
D5	9.35±1.15	30.20±4.77	40.00±0.00*	40.00±0.00*	
F1	5.75±1.46	22.80±1.52	31.60±2.28	37.00±2.55	
F2	7.50±1.32	30.20±1.67	40.00±0.00*	40.00±0.00*	
F3	5.75±1.00	27.10±1.64	37.90±1.64	40.00±0.00*	
F4	6.00±1.06	27.70±2.88	38.70±1.30	40.00±0.00*	
F5	7.10±1.52	30.70±4.16	40.00±0.00*	40.00±0.00*	
F6	6.35±1.40	27.50±2.92	38.10±3.42	40.00±0.00*	
G1	7.85±5.92	28.40±6.30	38.50±1.00	40.00±0.00*	
G2	6.05±1.43	27.00±1.22	37.30±2.30	40.00±0.00*	
G3	6.75±1.12	26.50±1.58	37.40±2.42	40.00±0.00*	
G4	6.65±0.57	27.90±3.03	38.70±0.55	40.00±0.00*	
G5	6.55±1.29	25.70±1.52	36.90±0.84	40.00±0.00*	
G6	7.85±2.11	27.20±2.70	36.00±1.00	40.00±0.00*	
G7	7.50±1.77	30.30±3.78	39.10±0.89	40.00±0.00*	
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5.05±2.36	27.90±3.27	38.30±1.34	40.00±0.00*	
7.15±1.44	26.35±0.45	37.00±2.55	40.00±0.00*	
9.80±2.27	31.60±1.92	40.00±0.00*	40.00±0.00*	
11.05±0.55	38.30±2.00	40.00±0.00*	40.00±0.00*	
	5.05±2.36 7.15±1.44 9.80±2.27 11.05±0.55	$5.05\pm2.36$ $27.90\pm3.27$ $7.15\pm1.44$ $26.35\pm0.45$ $9.80\pm2.27$ $31.60\pm1.92$ $11.05\pm0.55$ $38.30\pm2.00$	$5.05\pm2.36$ $27.90\pm3.27$ $38.30\pm1.34$ $7.15\pm1.44$ $26.35\pm0.45$ $37.00\pm2.55$ $9.80\pm2.27$ $31.60\pm1.92$ $40.00\pm0.00*$ $11.05\pm0.55$ $38.30\pm2.00$ $40.00\pm0.00*$	$5.05\pm2.36$ $27.90\pm3.27$ $38.30\pm1.34$ $40.00\pm0.00*$ $7.15\pm1.44$ $26.35\pm0.45$ $37.00\pm2.55$ $40.00\pm0.00*$ $9.80\pm2.27$ $31.60\pm1.92$ $40.00\pm0.00*$ $40.00\pm0.00*$ $11.05\pm0.55$ $38.30\pm2.00$ $40.00\pm0.00*$ $40.00\pm0.00*$

647 \*fully-grown culture













