1	Species diversity and community composition of native arbuscular mycorrhizal fungi in apple roots are affected
2	by site and orchard management
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14	Abstract
15	Arbuscular mycorrhizal fungi (AMF) are beneficial microrganisms which establish mutualistic symbioses with the roots
16	of most food crops, improving plant performance, nutrient uptake and tolerance to biotic and abiotic stresses. A better
17	understanding of the factors affecting AMF occurrence and diversity is fundamental to implement sustainable
18	agricultural managements effectively profiting from beneficial plant symbionts. Here, we investigated AMF occurrence,
19	diversity and community composition in the roots of apple trees from 21 orchards in South Tyrol, as affected by
20	location, management (organic vs integrated) and altitude, by PCR cloning and sequencing and PCR-DGGE of partial
21	18S rRNA gene. The screening of 448 clones from 21 clone libraries allowed the identification of 6 native AMF at the
22	species level: Glomus indicum, Sclerocystis sinuosa, Funneliformis mosseae, Rhizoglomus irregulare, Septoglomus
23	constrictus and Claroideoglomus lamellosum. The most abundant genera were represented by Glomus (29.7 % of the
24	sequences), Paraglomus (19.4 %), Claroideoglomus (17.2 %), Sclerocystis (16.1 %) and Rhizoglomus (12.3 %).
25	Septoglomus, Diversispora and Funneliformis sequences corresponded to less than 4 % of total sequences. Although the
26	degree of root colonization was unaffected by treatments, ANOSIM analysis of PCR-DGGE clusters revealed
27	significant differences in apple root AMF diversity between sites and agricultural managements. Species richness was
28	significantly higher in organically managed orchards than in integrated ones. Our findings provide insights into
29	important factors affecting native AMF communities of apple trees, which could be exploited in sustainable fruit
30	production systems, where beneficial soil biota boost biogeochemical cycles, energy fluxes and crop productivity.
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Keywords: Apple trees; Native AMF communities; Mycorrhizal colonization; Small Ribosomal Subunit (SSU rDNA);
 Clone libraries; PCR-DGGE

34

35 1. Introduction

36 Soil microorganisms are essential elements of soil health, fertility and productivity in sustainable and organic farming 37 systems, playing key roles in the completion of biogeochemical cycles and availability of mineral nutrients, carbon 38 sequestration and soil structure improvement, pest and disease control (Pimentel et al., 1997). Arbuscular mycorrhizal 39 (AM) fungi (AMF, Phylum Glomeromycota) are beneficial microorganisms establishing mutualistic symbioses with the 40 roots of most food crops, including cereals, legumes and fruit plants, and improving plant performance, nutrient uptake 41 and tolerance to biotic and abiotic stresses (Smith and Read, 2008). AMF are obligate biotrophs, obtaining sugars from 42 their host plants while providing soil mineral nutrients - such as P, N, S, K, Ca, Fe, Cu and Zn - absorbed and 43 translocated by means of large and ramified extraradical hyphal networks that spread from plant roots into the 44 surrounding soil (Giovannetti et al., 2015). Moreover, AMF deliver important agroecosystem services, such as soil 45 aggregation and carbon sequestration (Gianinazzi et al., 2010), and are considered important soil biological indicators 46 (Creamer et al., 2016; Stone et al., 2016). AMF enhance plant performance and fitness also through the synergistic 47 action of beneficial mycorrhizosphere bacteria, *i.e.* strictly associated to mycorrhizal hyphae, colonised roots and 48 spores, which display multifunctional activities, from antibiotic, siderophores and indole acetic acid production to P-49 solubilisation, phytate mineralization and N-fixation (Barea et al., 2002; Philippott et al., 2013; Agnolucci et al., 2015). 50 Thus, AMF represent environmentally-friendly biofertilisers and biostimulants, as they reduce the need of chemical 51 fertilizers and pesticides in sustainable food production systems (Rouphael et al., 2015). In addition, they contribute to 52 the production of safe and high-quality food, positively affecting the synthesis of health-promoting secondary 53 metabolites in food crops (Battini et al., 2016a, b).

54 In spite of all the potential benefits of the symbiosis, AMF role has been often marginalised in agriculture. In 55 field crops, several agronomic practices, such as continuous monocultures, deep ploughing, intensive fertiliser and 56 pesticide use can decrease AMF occurrence, activity and diversity, often leading to a reduction of AMF benefits to crop 57 production and soil quality (Douds and Millner, 1999; Jansa et al., 2003; Oehl et al., 2004, 2005b; Brito et al., 2012; 58 Avio et al., 2013). Moreover, in fruit production the soil is usually replanted with the same tree species short after the 59 preceding trees are removed. In organic farming systems, thanks to the fact that synthetic fertilizers, herbicides and 60 pesticides are not applied, AMF diversity, activity and abundance have been reported to be higher than in conventional 61 systems, supporting the view that AMF play a fundamental functional role in the maintenance of soil fertility and crop

production in low-input agroecosystems, compensating for the reduced use of agrochemicals (Mäder et al., 2002; Oehl
et al., 2003, 2004; Gosling et al., 2006; Ryan and Tibbet, 2008; Mazzoncini et al., 2010; Verbruggen et al., 2010).

64 Different strategies have been devised to increase the mycorrhizal potential of soils in sustainable agriculture: 65 among them, the inoculation with non-native AMF (Jeffries et al., 2003; Gianinazzi and Vosatka, 2004) has proved 66 effective in improving root colonization, plant biomass production and P uptake (Lekberg and Koide, 2005). However, 67 such an approach involves high costs of production and application when utilised on a large scale, and raises concerns 68 about potential negative impacts of non-native invasive AMF inoculants on the composition and structure of native 69 AMF communities, possibly leading to biodiversity losses (Schwartz et al., 2006). A different approach focuses on the 70 enhancement of native AMF by means of mycotrophic crops able to maintain or increase native mycorrhizal potential 71 of soils and root colonization and growth of the subsequent crops (Kabir and Koide, 2002; Karasawa and Takebe, 2012; 72 Lehman et al., 2012; Njeru et al., 2015, 2014).

73 Little information is available on the impact of cultural practices on AMF occurrence, species richness and 74 composition in apple crop production systems. Different orchard floor managements create different levels of soil 75 disturbance and may therefore promote or depress AMF occurrence. Organic farming guidelines, for example, 76 differently from conventional or integrated fruit production techniques (Kelderer, 2004), do not allow the use of 77 chemical herbicides, but recommend, instead, the use of alternative techniques to manage the orchard floor by either 78 superficial soil tillage, grass mowing or mulching. Similarly, while conventional and integrated management techniques 79 allow the use of synthetic mineral fertilisers, organic farming guidelines recommend increasing soil fertility by 80 enhancing soil organic matter. The use of straw mulches and compost have been reported to increase apple root 81 colonisation, which was positively linked with soil pH and availability of P, K, Zn, Mn and C, suggesting that the 82 relevant organic orchard floor management practices promoted functional AMF associations more effectively than 83 conventional practices (Meyer et al., 2015). A recent work carried out using molecular methods showed that soil 84 characteristics and farming systems affect AMF diversity and community composition in the roots of cultivated apple 85 trees. In particular, a low soil available P content was associated with a higher AMF diversity in organically managed 86 orchards, as compared with conventionally managed ones (Van Geel et al., 2015). In such a study, the native AMF 87 colonizing apple roots were identified by 454-pyrosequencing of small subunit rRNA gene amplicons, with a taxonomic 88 resolution at the family level. So far, the characterization of apple AMF communities at the species level has been 89 performed only by conventional morphological methods, which show several shortfalls, mainly due to the use of spores 90 extracted from the soil, whose integrity and quality may induce species misidentification (Cavallazzi et al., 2007; Miller 91 et al., 1985; Purin et al., 2006).

92 Here we used two molecular methods, PCR-cloning and sequencing and PCR denaturating gradient gel 93 electrophoresis (PCR-DGGE) of partial 18S rRNA gene to characterize AMF communities of apple roots at genus and 94 species levels. We investigated AMF occurrence and diversity in the root systems of apple trees cultivated under 95 organic and integrated management from 21 orchards in the South Tyrol province (Italy), one of the most important 96 apple production districts in Europe, with more than 18,000 ha acreage and approx. 1 million tons apples harvested 97 yearly (Dalla Via and Mantinger, 2012). Our data provide knowledge of the factors which affect AMF colonization and 98 shape the native AMF community composition in apple roots, allowing the implementation of effective sustainable 99 management strategies that take advantage from beneficial plant symbionts.

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101 2. Materials and Methods

102 2.1. Soil and orchard management parameters

The research was carried out in South Tyrol (Northern Italy). In May 2013, 21 commercial apple (*Malus domestica* Borkh) orchards located either in the municipalities of Terlano (46°31' 59" N, 11°14' 47" E) or in that of Lagundo (46° 41' 0" N, 11° 8' 0" E) were randomly selected from a list provided by the local advisory service (The South Tyrolean Advisory Service for Fruit and Grape growing, Lana, Italy). This list included drip-irrigated apple orchards planted between 2007 and 2009, managed since planting following either the integrated fruit production (hereafter referred to as "integrated") or the organic farming (hereafter referred to as "organic") guidelines.

In the municipality of Lagundo 3 organic and 3 integrated orchards were selected at low (298-334 m a.s.l.) and at high altitude (591-660 m a.s.l.) while in the municipality of Terlano 6 organic and 3 integrated orchards were selected at a low altitude (ranging from 243 to 299 m a.s.l.). Orchards were classified according to the area (Te, Terano; La, Lagundo), management (O, Organic; I, Integrated) and altitude (L, Low altitude; H, High altitude) (Fig. 1).

The climate in area is warm temperate according to the Koppen-Geiger classification (Kottek et al., 2006). Average (1993 to 2013) climatic parameters are reported in Table 1. Average annual and maximum temperatures at high elevation were approx. 2 °C lower than at low elevation, while average minimum temperature was approx. 1 °C lower. Soils in the area are mainly sandy loam or loamy sand and are classified as Calcaric Cambisol according to the FAO Soil Taxonomy (IUSS Working Group WRB 2015) (Table 2).

The trees were always grafted on the same clonal rootstock (M9, T337 strain) and belonged to one of the following varieties: Gala, Golden Delicious, Pinova, Modì and Red Delicious. Orchards were uniform in terms of training systems (spindle bush) and planting distances (approx. 3 m between rows and 1 m between trees along the row). Orchard floor management always included the presence of grassed alleys between rows (2.0 to 2.3 m wide) that were mowed 3-4 times per year, while in the 0.7-1.0 m soil strip along the tree row, soil management differed between organic and integrated orchards (see below). The organic and integrated orchards differed both for the type of protection against aboveground pests and for soil management. In particular, in the organic orchards weed control was carried out through mechanical removal of weeds (approximately five times per year) and soil fertility was maintained only by the addition of organic fertilisers. In the integrated orchards, weed control in the soil strip under the trees was performed by spring and autumn application of glyphosate; mineral fertilizer supply was carried out according to the nutrient budget, considering expected yields of approximately 60 t ha⁻¹ and soil fertility, in the following ranges, 30-80 kg N, 19-28 kg P, 71-155 kg K.

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131 2.2. Root, leaf and soil sampling

132 In each orchard, root and leaf were samples from 6 randomly chosen trees from different rows, all located in the central 133 part of the plot. Sampling was carried out between May 15th and June 6th, 2013. Six soil cores were collected at a depth 134 of 10-30 cm and at approx. 20 cm distance from the trunk, using a split tube sample of 5 cm-diameter (Eijkelkamp 135 Agrisearch Equipment, BV). Roots were gently cleaned from the soil and approx. 20 g of fine roots (<2 mm diam) per 136 tree were collected and stored at 4 °C in polyethylene bags, to be successively analysed for mycorrhizal colonization. A 137 single root sample per orchard was prepared for molecular analyses (100-125 mg each) pooling together root 138 subsamples from the individual trees. Samples were stored at -20 °C, until processed. Soil samples were oven-dried at 139 65 °C until constant weight and stored at room temperature until analysed. Ten leaves per tree (without petiols) were 140 sampled from the 6th or the 7th nodes of one year-old shoots, oven-dried at 65 °C until constant weight and stored until 141 analysed.

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143 2.3. Mycorrhizal colonization

Mycorrhizal colonization was determined on 10 g thoroughly washed root samples. Roots were cleared with 10% KOH
in a 80 °C water bath for 15 min and stained with Trypan blue in lactic acid (0.05 %) after 10 min in 2 % aqueous HCl.
The percentage of AMF colonization was calculated from root samples from individual trees from each orchard using a
dissecting microscope at x25 or x40 magnification and the gridline intersect method (Giovannetti and Mosse, 1980).

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149 **2.4.** Soil and leaf analysis

With the exception of soil N and C concentration (performed by an elemental analyzer, Thermo Scientific), soil parameters were determined according to the methods of the Verband Deutscher Landwirtschaftlicher Untersuchungsund Forschungsanstalten (VDLUFA, 1991). Leaf N concentration was determined by Elemental Analyser (Flash 2000, Thermo Scientific), while leaf K, Ca, P and magnesium (Mg) concentrations were determined by inductive coupled plasma-optical emission spectrometry (ICP-OES, Spectro Ametek, Arcos and Spectro Ciros CCD) after microwave
digestion (Milestone UltraWAVE) with 69 % of ultrapure nitric acid.

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157 2.5. Statistical analyses of soil, leaves and mycorrhizal colonization

Data of mycorrhizal colonization (after angular transformation) and leaf mineral nutrient concentrations were analysed by two-way-ANOVA. The sub-dataset from Lagundo was used to assess the interactive effects of orchard management (organic vs. integrated) and orchard altitude (low vs. high). The sub-dataset obtained at low altitude was used to assess the interactive effects of orchard management (organic vs. integrated) and site (Terlano vs. Lagundo). Mycorrhizal colonization data were linearly correlated with soil parameters using the entire dataset. The coefficient of mycorrhizal colonization data variation (CV) within each farm was also calculated. The statistical analyses were conducted using the dedicated software SigmaPlot Centurion XV (StatPoint Technologies, Inc).

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166 **2.6.** Molecular analyses

167 2.6.1. DNA extraction from roots

Genomic DNA was isolated from root material by grinding with mortar and pestle in liquid nitrogen and subsequently using DNeasy Plant Mini Kit (Qiagen Milan, Italy), before performing cloning and sequencing analyses. Moreover, in order to obtain DNA containing less inhibitors possibly interfering with the amplification procedure of the PCR-DGGE technique, we extracted DNA from roots using the PowerSoilTM DNA Isolation Kit (MO BIO Laboratories, Solana beach, CA, USA). The isolated DNA was stored at -20 °C for subsequent analyses.

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174 2.6.2. Cloning and sequencing

DNA (10-20 ng) extracted from roots was used as template. Partial small subunit (SSU) ribosomal RNA gene fragments were amplified in volumes of 25 μ l with 0.125 U of GoTaq Flexi DNA Polymerase (Promega, Milan, Italy), 0.4 μ M of each primer (NS31 /AML2, Simon et al., 1992; Lee et al., 2008), 0.2 mM of each dNTP, 1.5 mM of MgCl₂ and 1x manufacturer's reaction buffer. The thermal cycler (Eppendorf Mastercycler personal, Eppendorf, Milan, Italy) was programmed as follows: a manual "hot start" at 94 °C for 3 min, 30 cycles at 94 °C for 30 sec, 58 °C for 40 sec, 72 °C for 55 sec and a final extension step at 72 °C for 10 min. Reactions yields were estimated using a 1 % (*w/v*) agarose I (Euroclone, Milan, Italy) in TBE 1X buffer (Euroclone) gels stained with ethidium bromide (0.5 μ g ml⁻¹).

182 NS31/AML2 amplicons from apple root samples were purified using Wizard® SV Gel and PCR Clean-up
183 system (Promega), then ligated into pGem-T Easy vector (Promega) to transform XL10-Gold Ultracompetent
184 *Escherichia coli* cells (Stratagene, La Jolla, CA, USA).

The structure and composition of the AM fungal communities were determined using PCR-RFLP screening of clone libraries. Putative positive clones were amplified by using standard SP6/T7 amplifications, followed by a nested PCR using NS31/AML2 primers as described above. Twenty-five amplicons per clone library were digested by HinfI and Hsp92II restriction enzymes (Promega) and run on 2 % agarose at costant 50 V for 2 h. A 100 bp DNA ladder (Promega) was used as a molecular weight marker. DNA profiles were visualized under UV illumination and captured as TIFF format file by Liscap program for Image Master VDS System (Pharmacia Biotech).

A total of 448 clones were examined. Recombinant plasmids of representative clones of each RFLP pattern
 were purified by Wizard® Plus SV Minipreps (Promega) and sequenced using T7 vector primers at BMR Genomics
 s.r.l. (University of Padova, Italy).

Sixty-nine unique sequences of the clones generated in this study have been deposited in EMBL Nucleotide
 Sequence Database (www.ebi.ac.uk/embl/) under the accession numbers from LT600783 to LT600851.

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197 2.6.3. PCR-DGGE analysis

198 To analyse AMF communities by PCR-DGGE, a semi-nested PCR approach was used. Initially a 550 bp fragment of 199 the 18S rRNA gene was amplified using the primer NS31 (Simon et al., 1992) in combination with the primer AM1 200 (Helgason et al., 1998). Amplification reactions were performed in a final volume of 50 µl, using 10-20 ng of DNA, 5 201 µl of 10X Ex Taq Buffer (Takara Biotechnology, Milan, Italy), 1.25U of TaKaRa Ex Taq (Takara Biotechnology), 0.2 202 mM of each dNTP (Takara Biotechnology), 0.5 µM of each primer (Eurofins genomics, Milan, Italy) and an aliquot of 203 $0.3 \,\mu g \,\mu l^{-1}$ acetylated bovine serum albumin (BSA, Promega). PCR amplifications were carried out using an iCycler-iQ 204 Multicolor Real-Time PCR Detection System (Bio-Rad, Milan, Italy) with the following conditions: 94 °C initial 205 denaturation for 1 min; 35 amplification cycles of 30 s at 94 °C, 30 s at 66 °C, 30 s at 72 °C; final extension at 72 °C for 206 5 min. The presence of amplicons was confirmed by electrophoresis in 1.5 % (w/v) agarose gels in TBE 1X buffer 207 (Tris-borate-EDTA, pH 8.0), stained with $0.5 \,\mu g \,m L^{-1}$ ethidium bromide.

For the DGGE analysis, 20 µl_of the PCR products plus 20 µl of buffer 2X made with 70 % glycerol, 0.05 %
xylene cyanol and 0.05 % bromophenol blue, were separated in a 8 % polyacrylamide-bisacrilamide (37.5:1) gel with a
35-56 % urea-formamide gradient, using the DCodeTM Universal Mutation Detection System (Bio-Rad). A composite

216 mix of AMF 18S rRNA gene fragments from *Funneliformis mossae* AZ225C, *Funneliformis coronatum* IMA3 and 217 *Rhizoglomus intraradices* IMA5 was added in the middle and at both ends of each gel as DGGE marker (M). Gels were 218 run and profiles were visualized as described in Agnolucci et al. (2013).

DGGE bands were excised from polyacrylamide gels and DNA was extracted by eluting in 50 µl ddH₂O at 4°C overnight. One µL of the supernatant diluted 1:100 was used to re-amplify the 230 bp DNA fragment of the 18S rRNA gene according to the PCR protocol described above, except that the primer NS31 was used without GC clamp.
PCR products were then purified by UltraClean PCR CleanUp Kit (MO-BIO Laboratories) according to the manufacturers' protocol, quantified and 5' sequenced at the BMR Genomics (Padova, Italy).

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225 2.7. Bioinformatics

Sequences from *E. coli* libraries and DGGE bands were edited in MEGA 6.0 and their similarities were determined using the Basic Local Alignment Search Tool (BLASTn) provided by NCBI. The detection of chimeric sequences was performed using USEARCH 6.0 (http://fungene.cme.msu.edu/FunGenePipeline/chimera_check/form.spr). Sequences were aligned with those corresponding to the closest matches from GenBank as well as with sequences from major clades of Glomeromycota using MUSCLE as implemented in MEGA6. Phylogenetic trees were inferred by Neighbourjoining analysis. The evolutionary distances were computed using the Maximum Composite Likelihood method. The confidence of branching was assessed using 1000 bootstrap resamplings.

233 Richness and composition of AMF communities obtained from clone libraries were evaluated for each 234 treatment. We determined the rarefaction curves with Past software to estimate whether the number of screened 235 sequenced were sufficient to capture AMF diversity of each host. Estimates of community diversity were determined as 236 Richness (S), bias-corrected Chao1 richness, Shannon-Weaver (Hs) and dominance index of Simpson (D). The indices 237 were calculated using PAST 3.0 and 1000 bootstraps were used to determine confidence intervals. Non parametric 238 Kruskal Wallis test was used to determine differences in the diversity indices among AMF communities in the roots of 239 apple plants cultivated in the two sites (Terlano vs Lagundo), under different managements (organic vs integrated) and 240 at different altitudes (low and high). AMF communities were also evaluated by permutational multivariate analysis of 241 variance (PERMANOVA), to test the effect of the management (organic vs integrated), site and altitude. Non 242 parametric tests and PERMANOVA were performed in SPSS version 20 software (IBM Corp., Armon, NY Inc, USA) 243 and PAST 3.0, respectively.

DGGE profiles were digitally processed with BioNumerics software version 7.5 (Applied Maths, St-Martens-Latem, Belgium) following the manufacturer's instructions. The lanes were normalized to contain the same amount of total signal after background subtraction and the gel images were straightened and aligned to give a densitometric curve. DGGE markers were used for digital gel normalization to allow comparison between gels. Bands were assigned using the auto search bands option and then checked manually and their positions were converted to Rf % values. All profiles were compared using the band matching tool with a position tolerance and optimization of 0.5 and 0 % respectively.

251 Similarities between DGGE patterns were calculated by determining Dice's similarity coefficients for the total 252 number of lane patterns from the DGGE gel. The similarity coefficients were then used to generate the dendrogram 253 utilizing the clustering method UPGMA (Unweighted Pair Group Method Using Arithmetic Average). DGGE profiles 254 were also analysed using non-metric multidimensional scaling analysis (NMDS) performed from a data matrix based on 255 presence/absence of bands (Bray-Curtis coefficient). The significance of data was assessed by the two way ANOSIM 256 method (analysis of similarities; 999 permutations) (PAST 3.0) with geographical site (Terlano vs Lagundo) and 257 management (organic vs integrated) as variability factors. Note that the stability index R describes the extent of 258 similarity between each pair in the ANOSIM, with values close to unity indicating that the two groups are entirely 259 separate and a value of zero indicating that there is no difference between the groups.

260 DGGE banding data were used to estimate four different indices treating each band as an individual 261 operational taxonomic unit (OTU). Richness (S) indicates the number of OTUs present in a sample and was determined 262 from the number of fragments. The overall diversity index of Shannon-Weaver (Hs) and the dominance index of 263 Simpson (D) were calculated using the equations Hs = $-\sum (P_i \times \ln P_i)$ and D = $\sum P_i^2$ respectively, where the relative 264 importance of each OTU is $P_i = n_i N^{-1}$, and n_i is the peak intensity of a band and N is the sum of all peak intensities in a 265 lane. Evenness index (E), which allows the identification of dominant OTUs, was calculated as $E = H (lnS)^{-1}$. Two-way 266 ANOVA was applied to diversity indices with site (Terlano vs Lagundo) and management (organic vs integrated) as 267 variability factors, after checking the normal distribution of data. One-way ANOVA was used for Lagundo diversity 268 indices with management as the variability factor. The means were compared by the Tukey's test (P < 0.05). Analyses 269 were carried out with the SPSS version 20 software.

- 270
- 271 **3. Results**

272 **3.1.** Soil fertility and mycorrhizal colonization

Soils texture in the Lagundo area had either a silty loam or a sandy loam texture, while those in the Terlano area were either loamy or sandy loam. In general, as indicated by the standard error values (Table 2), there was a relatively low variability of N concentration among soils belonging to different treatments as well as within each treatment. All soils were well endowed with P and Mg, while soil K availability ranged from moderate-low to normal values. In Lagundo only, soils managed according to organic guidelines had higher pH (P=0.02), organic matter (OM) (P=0.01) and C 278 concentration (P=0.03) than those managed according to the integrated guidelines (Table 2). Soils at higher elevation 279 also had higher soil OM (P=0.05) and soil C (P=0.06) than those at lower elevation (Table 2). When only data 280 belonging to low elevation were analyzed, no significant differences in soil characteristics between the two orchard 281 managements were detected.

282 No significant effect of altitude and management was recorded on leaf nutrient concentration (Table 3). Leaf P
283 and K were significantly higher (P=0.01 and 0.001, respectively) and leaf Mg significantly lower (P=0.02) in orchards
284 located in Terlano than in Lagundo (Table 3).

In all sampled orchards, apple roots were well colonized by AMF (Table 3). The percentage of colonized root length, ranging from 41 to 60 %, was not significantly affected by site, management and altitude. We did not find significant effects of treatments on the degree of mycorrhizal colonization variation among and within orchards (Table S1). The level of mycorrhizal colonization was not significantly correlated with most measured soil parameters, but a negative linear correlation with soil organic matter (r=-0.49, P=0.03) was found.

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3.2. Identification of native AMF colonizing apple roots

The DNA extracted from apple roots was successfully amplified using the primer pair NS31/AML2, obtaining a fragment of the expected size (~550 bp). A total of 448 clones from the 21 clone libraries were screened by PCR-RFLP analysis, obtaining 19 different RFLP patterns. For each RFLP group several clones originating from different libraries were sequenced, giving 17 Glomeromycota sequences. All non-redundant sequences from the 21 clone libraries (69 out of 212) and 27 references from GenBank were used for neighbour-joining phylogenetic analyses (Fig. 2). After RFLPs, BLASTn and phylogenetic analyses the sequences were grouped into 17 OTUs supported by a bootstrap value >87 %.

298 Among the 17 OTUs, we retrieved sequences belonging to 8 out of the 19 Glomeromycota genera (Redecker et 299 al., 2013). In particular all the genera of Glomeraceae were found (Glomus, Septoglomus, Rhizoglomus, Sclerocystis, 300 Funneliformis) together with the genera Claroideoglomus, Diversispora and Paraglomus, belonging to 301 Claroideoglomeraceae, Diversisporaceae and Paraglomeraceae, respectively. The most abundant genera were 302 represented by Glomus, accounting for 29.7 % of the sequences, followed by Paraglomus (19.4 %), Claroideoglomus 303 (17.2 %), Sclerocystis (16.1 %) and Rhizoglomus (12.3 %). Septoglomus, Diversispora and Funneliformis sequences 304 corresponded to less than 4 % of total sequences. Twelve OTUs were ascribed to Glomeraceae family: within the genus 305 Glomus we retrieved seven OTUs, of which only one (Glo7, 5.8 % of total sequences) was identified as Glomus 306 indicum; the remaining six OTUs (Glo1 to Glo6) represented sequences of uncultured Glomus species (Table 4). In the 307 genera Sclerocystis, Funneliformis and Rhizoglomus, OTUs named Scle, Fun and Rhi were identified as Sclerocystis 308 sinuosa, Funneliformis mosseae and Rhizoglomus irregulare (synonym Rhizophagus irregularis, formerly known as

309 Glomus irregulare), respectively. OTUs Sept1 and Sept2 and Cl1, Cl2 and Cl3 assigned to Sept0glomus and 310 Claroideoglomus, respectively, matched to sequences of either known (Sept2, Septoglomus constrictus and Cl1, 311 Claroideoglomus lamellosum), or unknown species (Septoglomus sp., Claroideoglomus sp.), while OTUs belonging to 312 Paraglomus (Par) and Diversispora (Div) matched only unknown Glomeromycota. MaarjAM database 313 (http://maarjam.botany.ut.ee/, accessed on March/2016) was used to confirm the assignment of our OTUs to sequences 314 of Glomeromycota (Table 4). Rarefaction analyses indicated that the number of analysed sequences was generally 315 sufficient to capture the AMF diversity in the roots of most orchards, since the curves almost reached the asymptote 316 (Fig. S1).

- The bands of interest from the DGGE profiles, excised and sequenced to determine their affiliation, identified the same AMF species as those described above, except Claroideoglomeraceae and Paraglomeraceae (data not shown).
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320 **3.3.** AMF community diversity in apple roots as affected by site, management and altitude

321 3.3.1. Analysis of clone libraries

322 The cloning and sequencing method allowed the identification of 14 and 15 OTUs in Terlano and Lagundo orchards, 323 respectively, 12 of which shared between the two sites (97.5 and 88.8 % of the total sequences of the two sites, 324 respectively). Fun (F. mosseae) and Glo6 were found only in Terlano apple roots, while Sept2 (S. constrictus), Glo2 325 and Cl3 were retrieved only in Lagundo. Two ways PERMANOVA analyses revealed differences in AMF community 326 composition between the two sites (P=0.02) and between organic and integrated managements (P=0.02). Moreover, in 327 Lagundo, where orchards were located at two different altitudes (300 m and 600 m) two ways PERMANOVA analysis 328 showed differences among AMF communities in relation to management (P=0.003) and altitude (P=0.003). In total, 329 orchards under organic management hosted 13 and 6 OTUs at low and high altitude, respectively. In organic orchards 330 growing at higher elevation the most common species (Par, Glo7, Rhi, Scle) present at both sites and under different 331 managements were detected together with Cl1 and Div (species found in all organic orchards at low altitude) (Fig. 3). 332 By contrast, in integrated orchards 10 and 9 OTUs in total were found at low and high altitude, respectively. At high 333 altitude, together with the most common species, a particular OTU (Cl3, 31 % of the sequences) was retrieved (Fig. 3).

Among diversity indices, S and Chao-1, indicating observed and estimated AMF richness, were significantly higher in organic orchards (6.33 ± 0.55 and 7.14 ± 0.88 , respectively) than in integrated ones (4.00 ± 0.01 and 4.00 ± 0.01 , respectively) in Terlano (P=0.027 and P=0.034, respectively). In Terlano 13 OTUs were retrieved in organic management, in contrast with only 6 OTUs found in integrated ones. No differences among diversity indices were detected among orchards in Lagundo.

340 3.3.2. Analysis of PCR-DGGE profiles

AMF 18S rDNA fragments of approx. 230 bp were successfully amplified from all the samples, by the semi-nested
 PCR approach. DGGE analyses of PCR products showed profiles characterized by a high number of intense and clearly
 defined fragments.

AMF community composition was assessed by cluster analysis of DGGE profiles (Fig. 4). The dendrogram showed two main subclusters with a low similarity, 20 %. In particular, the first subcluster included all Terlano samples (except one), while the other one included all Lagundo samples (except one). The latter was formed by two subclusters in which organic and integrated samples were separated with a similarity of 36 %. The analysis of DGGE profiles did not separate samples originating from orchards growing at different altitudes (Fig. 4). Terlano samples showed a more complex clustering, with no clearcut separation by management.

These findings were confirmed by the NMDS analysis of the DGGE AMF community profiles (Fig. 5). ANOSIM revealed significant differences in AMF community composition of Lagundo samples compared with that of Terlano (R=0.786, P=0.0001) and between samples from organic and integrated sites (R=0.227, P=0.05).

AMF DGGE profiles were also analysed to assess S, Hs, D and E diversity indices. Richness (S) and Hs indices were significantly different between the two geographical sites, while a significant interaction was observed for Hs and D (Table 5). The relevant one-way ANOVA revealed significant differences between organic and integrated samples in Lagundo, where S and Hs were significantly higher in the organic management (P=0.049 and P=0.029 respectively) (Table 6).

358

359 4. Discussion

Our data showed that, in South Tyrol orchards, geographical area, altitude and management shaped AMF species community composition in apple roots, which however always maintained high levels of mycorrhizal colonization. Here, for the first time, we identified at the species and genus level native AMF colonizing apple roots and characterized their diversity and community composition utilizing two molecular methods - PCR cloning and sequencing and PCR-DGGE.

365

366 4.1. Mycorrhizal colonization of apple roots

The percentage of colonized root length of apple trees ranged from 41 to 60 %, comparing well with previous data on mycorrhizal colonization of apple seedlings, that ranged from 40 to 60 % after inoculation with *F. mosseae* and *Glomus macrocarpum*, respectively (Miller et al., 1989) and with findings obtained by Meyer et al. (2015) in conventionally and organically-managed apple orchards in South Africa. Other authors reported more variable values, ranging from 24 to 68 % in micropropagated apple trees, depending on soil pH and the identity of the inoculated AMF species (Cavallazzi
et al., 2007). In our work, mycorrhizal colonization of apple roots was uncoupled from soil parameters, except for
organic matter, confirming previous findings on the decrease of mycorrhizal colonization levels in an organic apple
orchard after a straw mulch floor management treatment (Meyer et al., 2015).

Overall, mycorrhizal root length was not significantly affected by site, altitude and management in the apple orchards investigated. As to site, Miller et al. (1985) in a survey of 18 apple rootstock plantings in USA found a high variability in mycorrhizal colonization, ranging from 5 to 75 %, with the lowest levels observed in California and Washington and the highest in Virginia, Pennsylvania, Georgia and Iowa. Such data are expected, given the different range of geographical distances of the apple orchards analysed, entailing the most divergent environmental conditions, at the subcontinental level.

Most studies investigating the effects of altitude on mycorrhizal colonization have been carried out at very high elevation gradients (from 1,500 to 5,300 m), in alpine habitats of Tibet, Andes, Rocky Mountains, Alps and Mount Fuji (Read and Haselwandter, 1981; Wu et al., 2007; Lugo et al., 2008; Schmidt et al., 2008; Gai et al., 2012). Overall, these works reported an adverse effect of increasing altitude on AMF colonization across a large altitude gradient. In our study, we assessed the effects of elevation in a single site, Lagundo, from apple orchards cultivated at 300 and 600 m, an altitudinal range probably too low to produce significant effects on AMF ability to colonize apple roots, which, indeed, showed high percentages of mycorrhizal root length, 35-52 %.

388 The type of farming practices has long been known to affect mycorrhizal colonization in different crops, such 389 as wheat (Ryan et al., 1994; Mäder et al., 2000), vetch-rye and grass-clover (Mäder et al., 2000), onion (Galván et al., 390 2009), maize (Douds et al., 1993; Bedini et al., 2013), soybean (Douds et al., 1993). Very few studies investigated the 391 effects of organic vs. conventional management regimes on the establishment of mycorrhizal symbiosis in apple roots. 392 The absence of differences in the percentage of mycorrhizal root length in our experimental apple orchards may be 393 ascribed to the uniformity of the orchard floor managements, which included the presence of grassed alleys between 394 rows. AMF-host weed species may have contributed to the maintenance of wide and infective extraradical mycelial 395 networks establishing linkages among the roots of cover plants and apple trees (Atkinson, 1983; Giovannetti et al., 396 2004; Njeru et al., 2014). The application of glyphosate in the small soil strip along the tree rows and that of mineral 397 fertilizer supply in the integrated orchards did not have a significant impact on the ability of AMF to colonize apple 398 roots. This finding is consistent with previous studies reporting variable and unpredictable effects of such chemicals on 399 AMF symbioses in different environments and geographical locations (Gosling et al., 2006).

400

401 4.2. Identification of native AMF colonizing apple roots

402 The screening of 448 clones from 21 clone libraries allowed us to affiliate DNA sequences with 8 out of the 19
403 Glomeromycota genera, which encompassed all the genera of the family Glomeraceae described so far: *Glomus*,
404 *Septoglomus*, *Rhizophagus*, *Sclerocystis*, *Funneliformis*. In addition, members of the families Claroideoglomeraceae,
405 Diversisporaceae and Paraglomeraceae were found.

406 Overall, the most common species belonged to the genus Glomus, in agreement with previous findings 407 obtained by morphological description of AMF spores in a survey of 18 apple rootstock plantings in USA (Miller et al., 408 1985). Some of the species retrieved in the quoted study occurred also in our apple roots, such as F. mosseae and S. 409 constrictus (formerly Glomus mosseae and Glomus constrictum, respectively) and S. sinuosa, while no species of the 410 genus Gigaspora, common in the USA apple plantings, were found. Our data differ from those obtained in a survey of 411 apple orchards in Santa Catarina, south region of Brazil, where 15 species of the genus Acaulospora were retrieved and 412 described after spore sievings from the soil, using morphological methods (Purin et al., 2006). Only few AMF species 413 were in common, belonging to F. mosseae, Claroideoglomus spp. and S. sinuosa (formerly Glomus sinuosum). Another 414 interesting work taxonomically characterized AMF species by morphological identification of spores retrieved from soil 415 sieving and trap cultures in Brazil (Cavallazzi et al., 2007): the species with the highest number of spores was 416 Acaulospora mellea, followed by Scutellospora heterogama, Gigaspora decipiens and Acaulospora spinosa, none of 417 which occurred in apple roots of South Tyrol orchards. A molecular study, utilising 454-pyrosequencing of small 418 subunit rRNA gene amplicons, identified, at the family level, the AMF colonizing roots of cultivated apple in central 419 Belgium (Van Geel et al., 2015). In the work, 73 % and 19 % of OTUs were affiliated with the Glomeraceae and 420 Claroideoglomeraceae, in agreement with our findings, 70 % and 18 %, respectively. In addition, the authors detected 421 only a few OTUs belonging to the families Gigasporaceae, Diversisporaceae and Acaulosporaceae, while the 422 percentage of OTUs affiliated with Paraglomeraceae was much lower in Belgian apple orchards, compared with South 423 Tyrol ones, 1 % vs. 19 %. A recent work, carried out in an experimental apple orchard near Sint-Truiden, Belgium, 424 confirmed the previous data, with 72 % of OTUs belonging to Glomeraceae, 26 % to Claroideoglomeraceae and 2 % to 425 Paraglomeraceae (Van Geel et al., 2016).

The absence or rarity of specific taxa, such as Acaulosporaceae and Gigasporaceae, from the three European apple orchards investigated so far, compared with the frequency of their retrieval from USA and Brazil, may be ascribed to a number of complex and interacting factors - soil, environment, climate, agronomic practices, history of the sites etc. - beyond geographical position. Although the number of experimental data on apple from the two continents is still too small to deduce general trends, recent studies at the global scale revealed that differences in AMF communities diversity do occur among different continents and climatic zones (Kivlin et al., 2011; Öpik et al., 2013; Davison et al., 2015). It is interesting to note that a survey of AMF species distributed all over the world in protected areas reported a general predominance of Acaulosporaceae and Gigasporaceae in tropical or subtropical forests in Brazil and Argentina
(Turrini and Giovannetti, 2012).

In our study we found sequences (uncultured *Clareidoglomus*-Cl2, *Paraglomus*-Par and *Glomus*-Glo1 sequences, corresponding to 18 % of total OTUs), matching those retrieved in apple roots from German orchards (JN644447, unpublished), in roots of other fruit tree plants, such as *Pyrus pyrifolia* (AB695049) (Yoshimura et al., 2013) and *Citrus* rootstocks (JQ350797) (Wang and Wang, 2014). It is tempting to speculate that fruit trees may show a preference for specific fungi, possibly recruiting AMF species in relation to their functional significance.

440

441 4.3. AMF community diversity in apple roots as affected by site, altitude and management

Present results show that root AMF community composition of apple trees cultivated in two production sites, Terlano and Lagundo, differed significantly and were affected by agricultural management, organic vs. integrated, and altitude. Such findings are supported by consistent data obtained by combining two molecular methods, cloning and sequencing and PCR-DGGE, and utilizing two different primer pairs to amplify the same region of the partial 18S rRNA gene. To the best of our knowledge, this is the first application of such a comprehensive approach for the characterization of root AMF diversity of fruit trees, allowing the differentiation of AMF communities colonizing apple roots in different agricultural conditions.

449 The differences detected in root AMF community composition between the two apple production sites, as 450 revealed by cloning and sequencing analysis, were ascribed to specific taxa occurring only in Terlano (Fun and Glo6) or 451 Lagundo (Sept2, Glo2 and Cl3), to their relative abundance and to the different distribution of shared OTUs between 452 the two sites. Such site differences were consistently detected by PCR-DGGE community profiles analysis, which 453 separated the two relevant clusters with a very high dissimilarity (80 %), and confirmed by NMDS analysis. Our data 454 are in agreement with a previous work, carried out at the regional scale, investigating the distribution of 6 AMF species 455 in 154 agricultural soils across Switzerland, which showed that AMF communities were strongly affected by 456 geographical distance (max. 294 km) (Jansa et al., 2014). Consistent findings were also reported by van der Gast et al. 457 (2011), who investigated AMF community diversity across England (max. 250 km). Although the maximum distances 458 among the sampled apple orchards were much lower than those described above (max. 27 km), our work identified 459 different AMF species and genera in the two sites, Terlano and Lagundo. Such distribution patterns may be the result of 460 biogeographic history, reflecting the dispersal of AMF taxa over time, and the variable climatic and environmental 461 conditions (Morton et al., 1995). Recent molecular works (Hazard et al., 2013; De Beenhouwer et al., 2015; Van Geel et 462 al., 2015) suggested a key role of soil environment in shaping AMF diversity and distribution patterns. Further in-depth 463 and comprehensive studies should be performed in order to separate geographical distance effects from those due to the

464 characteristics of each sampling site, i.e. soil properties, use of pesticides and chemical fertilizers, environs, even when 465 comparing sites at the regional and local scale. In addition, beyond geographical, soil and environmental variables, host 466 plant identity should be taken into account, given its essential role played in the selection of AMF symbionts 467 functionally established in the roots (Helgason et al., 2002; Gollotte et al., 2004; Sýkorová et al., 2007). Actually, at 468 local scale, a marked host preference has been recently found in maize plants, which hosted completely different AMF 469 communities, as compared with those of the preceding cover crops (Turrini et al., 2016).

470 No differences in AMF community composition were found by PCR-DGGE cluster analysis between orchards 471 located at 300 m and 600 m altitude in Lagundo, in contrast with cloning and sequencing, that allowed the detection of 472 a particular OTU (Cl3-Claroideoglomus sp., 31 % of the sequences) only in the 600 m high orchards. Such divergent 473 data may be ascribed to the different primers utilized during DNA amplification by the two diverse molecular 474 techniques: indeed, Glo1 primer, used in PCR-DGGE, did not allow the amplification of Claroideoglomeraceae, as 475 shown by band sequencing. The OTU Cl3 is infrequent, probably fitting better to the colder environment and to the 476 higher levels of organic matter and soil C found in the 600 m Lagundo orchards. Indeed, the occurrence of infrequent 477 AMF species at high altitude was previously reported (Liu et al., 2011; Sýkorová et al., 2007), while several new 478 species have been described in soils from Swiss Alps (Oehl and Sieverding, 2004; Oehl et al., 2005a, 2006, 2011).

479 In this work, the relative abundance of single AMF species colonizing apple roots showed significant 480 differences between organic and integrated management, as assessed by PERMANOVA, both in Terlano and in 481 Lagundo orchards. Differences were found also by the analysis of PCR-DGGE AMF community profiles, carried out by 482 NMDS and ANOSIM. Diversity indices showed higher AMF richness in organically managed apples compared with 483 integrated ones, using both molecular approaches. Although scarce information is available on AMF diversity in apple 484 orchards, as affected by management, our findings compare well with those obtained by Purin et al. (2006), who, by a 485 morphological approach, found a higher AMF richness in organic compared with conventional orchards in Brazil. A 486 few other studies, performed on plant species other than apple, reported the effects of organic farming on the 487 composition of AMF communities. For example, Oehl et al. (2004), in the DOC field experiment in Switzerland found 488 that organic management enhanced AMF spore diversity and abundance, while Bedini et al. (2013) showed a 489 progressive increase of AMF richness and composition during the transition from conventional to organic agriculture. 490 Other authors, using molecular techniques, showed that the diversity of AMF species composition in maize and potato 491 roots was higher in organic sites, compared with conventional ones, in agricultural fields throughout the Netherlands 492 (Verbruggen et al., 2010).

493 The experimental data collected so far on the influence of organic and conventional managements on AMF 494 diversity, either in the soil or in the roots, using either morphological or molecular approaches, are limited and do not

495 allow us to infer consistent response patterns. Indeed, the words "organic" and "conventional" may have different 496 meanings in the different experimental studies, as the diverse farming systems often encompass heterogeneous local soil 497 properties, environmental conditions and host plants. Thus, AMF diversity or community composition may be affected 498 by a number of uncontrolled variables, such as the use of weed cover, soil tillage, cover crops, manure, quality and 499 quantity of herbicides, pesticides and fertilizers. Whereas floor management in the orchard alleys was similar in organic 500 and integrated orchards, the two management systems differed mainly by the weed control and fertilizer supply to the 501 soil in the strip centres on the tree row: we speculate that the use of glyphosate and mineral fertilizers in integrated 502 managed orchards may have selectively modulated the abundance and composition of AMF taxa able to tolerate such 503 chemicals.

504

505 5. Conclusions

506 In this work, we utilized a multimodal approach to study AMF communities living in symbiosis with apple roots in 507 South Tyrol orchards, under different geographical and environmental conditions. High levels of mycorrhizal 508 colonization were detected across the different variables, while AMF diversity and community composition were 509 affected by geographical area, altitude and farming system management, as detected by PCR cloning and sequencing 510 and PCR-DGGE. In particular, species richness was significantly higher in organically managed orchards than in 511 integrated ones. We identified, for the first time, the native AMF communities of apple roots at the species and genus 512 level, detected infrequent taxa and retrieved some environmental sequences matching those obtained from other fruit 513 plant species. Our findings provide insights into factors affecting native AMF communities of apple trees, which could 514 be exploited to implement sustainable fruit production systems, where beneficial soil biota can boost biogeochemical 515 processes fundamental for energy fluxes, ecosystem functioning and crop productivity.

516

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- 527 molecular and data analyses. M.A., M.G. and A.T. wrote the paper. F.S., M.T. and E.T. provided agronomic expertise
- 528 and apple root samples. M.A., M.G., F.S., M.T. and A.T. contributed reagents/ materials/analysis tools.
- 529

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734 Main climatic parameters of the studied sites

		Altitude						
Parameter	Unit	Low	High					
average annual temperature	°C	11.7	10.3					
average minimum temperature (January)	°C	-3.8	- 5.1					
average maximum temperature (July)	°C	29.2	27.5					
annual rainfall	mm	714	522					
Data are from the meteorological stations (average 1971-2013) of the Province of Bolzano-Bozen								

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739 Main soil characteristics in the selected orchards according to their geographical site, management and altitude

Area	Management	Altitude	Organic matter (g kg ⁻¹)	pH (H ₂ O)	C (g kg ⁻¹)	N (g kg ⁻¹)	P (mg kg ⁻¹)	K (mg kg ⁻¹)	Mg (mg kg ⁻¹)
	Organic	Low	58.7 ± 8.7	7.3 ± 0.1	34.1 ± 1.6	3.2 ± 0.1	57 ± 24	43 ± 21	380 ± 45
Lagundo	C	High	81.7 ± 10.7	7.2 ± 0.1	34.1 ± 1.0	3.2 ± 0.1	108 ± 19	56 ± 23	293 ± 18
C	Integrated	Low	38.7 ± 4.1	6.9 ± 0.2	25.1 ± 1.1	2.8 ± 0.3	79 ± 5	44 ± 13	360 ± 105
	-	High	48.7 ± 1.8	6.2 ± 0.5	31.1 ± 1.8	3.2 ± 0.1	61 ± 8	103 ± 37	187 ± 26
Terlano	Organic	Low	40.0 ± 6.7	7.3 ± 0.1	32.1 ± 4.7	2.6 ± 0.3	88 ± 16	86 ± 18	293 ± 14
	Integrated	Low	36.0 ± 5.5	6.4 ± 1.0	28.7 ± 2.9	3.0 ± 0.2	43 ± 8	70 ± 17	283 ± 38

740 Data are averages \pm s.e. For clarity, statistics is reported only in the text.

743 Percentage of root mycorrhizal colonisation and leaf nutrient concentrations in the selected orchards according to their geographical site, management and altitude

Area	rea Management Altitude My		Mycorrhizal colonization (%) N(%)		P(%)	K (%)	Mg (%)	Ca (%)	
	Organic	Low	41.44 ± 12.23	3.64 ± 0.70	0.22 ± 0.01	0.95 ± 0.05	0.34 ± 0.01	1.23 ± 0.16	
Lagundo	Organie	High	34.97 ± 15.69	2.94 ± 0.29	0.21 ± 0.01	0.97 ± 0.05	0.29 ± 0.01	1.07 ± 0.04	
	Integrated	Low	52.02 ± 14.33	3.21 ± 0.39	0.21 ± 0.01	1.02 ± 0.08	0.36 ± 0.05	1.17 ± 0.12	
	Integrated	High	48.19 ± 11.11	2.42 ± 0.09	0.21 ± 0.01	1.15 ± 0.06	0.25 ± 0.02	1.18 ± 0.10	
Terlano	Organic	Low	60.01 ± 13.76	2.82 ± 0.10	0.24 ± 0.01	1.26 ± 0.04	0.3 ± 0.01	1.12 ± 0.06	
	Integrated	Low	54.71 ± 23.41	3.01 ± 0.24	0.24 ± 0.01	1.2 ± 0.05	0.27 ± 0.03	0.9 ± 0.15	

744 Data are averages \pm s.e. For clarity, statistics is reported only in the text.

⁷⁵⁴ grown in South Tyrol orchards

OTU name	Identity NCBI database (%)	Identity Maarjam database (%)	Maarjam database virtual taxa	Taxonomic affiliation
Cl1 [†]	AJ276087 (99)	FN869808 (99)	VTX00057	Claroideoglomus lamellosum
C12	JN644447 (99)	HE615004 (99)	VTX00056	Claroideoglomus sp.
C13	KF290671 (99)	KF290671 (99)	VTX00225	Claroideoglomus sp.
Div	FJ831643 (100)	FJ831643 (100)	VTX00062	Diversispora sp.
Fun	AJ306438 (99)	AY635833 (99)	VTX00067	Funneliformis mosseae
Glo1	JQ350797 (99)	KF386274 (99)	VTX00214	Glomus sp.
Glo2	KF467269 (99)	KF467269 (99)	VTX00135	Glomus sp.
Glo3	H4380136 (99)	HG004495	VTX00153	Glomus sp.
Glo4	JX144121 (99)	KC579423	VTX00304	Glomus sp.
Glo5	JN009364 (99)	JN009364 (99)	VTX00151	Glomus sp.
Glo6	JX144133 (99)	HG004465	VTX00301	Glomus sp.
Glo7	GU059539 (99)	GU059539 (99)	VTX00222	Glomus indicum
Par	AB695049 (100)	-	-	Uncultured Glomeromycota sp.
Scle	AJ33706 (99)	AJ33706 (98)	VTX00069	Sclerocystis sinuosa
Sept1	KF386332 (99)	KF386332 (99)	VTX00063	Septoglomus sp.
Sept2	FR750212 (99)	FJ831626 (99)	VTX00064	Septoglomus constrictus
Rhi	FJ009618 (99)	FJ009617 (99)	VTX00114	Rhizoglomus irregulare
[†] Names	denote the most similar	AM fungal species	of sequenced clones	: Cl1, Claroideoglomus sp.1; Cl2,

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756 Claroideoglomus sp.2; Cl3, Claroideoglomus sp.3; Div, Diversispora sp.; Fun, Funneliformis mosseae; Glo1, Glomus

757 sp.1; Glo2, Glomus sp.2; Glo3, Glomus sp.3; Glo4, Glomus sp.4; Glo5, Glomus sp.5; Glo6, Glomus sp.6; Glo7, Glomus

758 sp.7; Par, Paraglomus sp.; Scle, Sclerocystis sinuosa; Sept1, Speptoglomus sp.1; Sept2, Speptoglomus sp.2; Rhi,

759 Rhizoglomus irregulare.

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⁷⁵³ Sequence types of arbuscular mycorrhizal fungi, identified using NS31-AML2 primers pair, in the roots of apple plants

763 Richness (S), Shannon-Weaver (Hs), Simpson (D) and Evenness (E) indices calculated from AMF DGGE profile

associated with apple plant roots growing in orchards cultivated in two geographical sites (Terlano vs Lagundo) and

~ /	~ г	1		•		/ 1 1	~
11	ነካ	under two managements	(organic	vs integrated	1)	(mean + standard error	r)
<i>'</i> '	10	under two managements	(Of game	vs mugiatet	•)	(mean ± standard ento.	1)

Geographical area	Management	S	Hs	D	Е
Lagundo	Organic	9.83 ± 0.87	2.19 ± 0.08	0.12 ± 0.01	0.97 ± 0.00
	Integrated	6.83 ± 1.01	1.72 ± 0.16	0.20 ± 0.03	0.93 ± 0.02
Terlano	Organic	10.5 ± 1.63	2.18 ± 0.17	0.13 ± 0.02	0.96 ± 0.01
	Integrated	12.3 ± 1.33	2.38 ± 0.08	0.09 ± 0.01	0.95 ± 0.01
Analysis of variance (P values)				
Geographical area		0.033	0.049	0.069	0.734
Management		0.666	0.392	0.252	0.126
Geographical area x M	Ianagement	0.086	0.044	0.033	0.171

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770 Richness (S), Shannon-Weaver (Hs), Simpson (D) and Evenness (E) indices calculated from AMF DGGE profile

- associated with apple plant roots growing in orchards cultivated in Lagundo under two managements (organic vs
- 772 integrated) (mean ± standard error)
- 773

	Geographical area	Management	S	Hs	D	E
-	Lagundo	Organic	9.83 ± 0.87	2.19 ± 0.08	0.12 ± 0.01	0.97 ± 0.00
		Integrated	6.83 ± 1.01	1.72 ± 0.16	0.20 ± 0.03	0.93 ± 0.02
-	Analysis of variance (P values)					
	Management		0.029	0.028	0.049	0.069
774						
775						

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778 Captions

Fig. 1. a) Maps showing the location of apple orchards in South Tyrol, Italy. Integrated managed orchards at high altitude, Organically managed orchards at high altitude, Integrated managed orchards at low altitude, Organically managed orchards at low altitude. Source of the air photo on the right: Google earth V 7.1.2.2041 image
2016 DigitalGlobe http://www.earth.google.com. Pictures of organically b) and integrated c) managed apple trees in the study site.

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Fig. 2. Neighbor-Joining phylogenetic tree of glomeromycotan sequences derived from apple roots of orchards growing in South Tyrol (Italy). Bootstrap values are shown when they exceed 75 % (1,000 replications). The analysis is based on partial nuclear small subunit ribosomal RNA gene sequences (SSU; ~ 550bp; NS31/AML2 fragment) and involved 96 nucleotide sequences. Different sequence types are indicated in brackets and names are reported in Table 1. AMF family are also reported. Sequences obtained in the present study are shown in bold and their accession numbers are prefixed with site/management/altitude clone identifiers (Te, Terlano; La, Lagundo; O, Organic; I, Integrated; L, Low altitude; H, High altitude). The tree is rooted with a reference sequence of *Corallochytrium lymacisporum* (L42528).

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Fig. 3. Relative abundance (%) of AMF phylotypes detected in the roots of the different orchards cultivated in two
geographical sites (Lagundo and Terlano), under two managements (organic and integrated) and at two altitudes (Low,
300 m and High, 600m).

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Fig. 4. Dendrogram obtained by UPGMA (Unweighted Pair Group Method Using Arithmetic Average) based on AMF
DGGE profiles obtained from apple plant roots growing in orchards cultivated in two geographical sites (Terlano and
Lagundo) and under two managements (Organic, Integrated) (Te, Terlano; La, Lagundo; O, Organic; I, Integrated;
L, Low altitude; H, High altitude).

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Fig. 5. Non-metric multidimentional scaling plot of DGGE analysis. Each point on the plot represents the AMF
community composition associated with apple plant roots growing in orchards cultivated in Terlano area (blue symbols)
or Lagundo area (orange symbols). The stress value is 0.2, the ANOSIM values (R) indicates significant differences
between Lagundo and Terlano (0.786, P=0.0001) and between organic and integrated (0.227, P=0.05) (Te, Terlano; La,
Lagundo; O, Organic; I, Integrated; L, Low altitude; H, High altitude).











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