



AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Impact of an arbuscular mycorrhizal fungus versus a mixed microbial inoculum on the transcriptome reprogramming of grapevine roots

This is the author's manuscript

Original Citation:

Availability:

This version is available http://hdl.handle.net/2318/1622433 since 2017-11-14T15:28:21Z

Published version:

DOI:10.1007/s00572-016-0754-8

Terms of use:

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)



UNIVERSITÀ DEGLI STUDI DI TORINO

This is an author version of the contribution published on:

Questa è la versione dell'autore dell'opera: [Mycorrhiza, 2017, DOI: 10.1007/s00572-016-0754-8]

The definitive version is available at:

La versione definitiva è disponibile alla URL: [http://link.springer.com/article/10.1007%2Fs00572-016-0754-8] 1 Raffaella Balestrini^{1*}, Alessandra Salvioli², Alessandra Dal Molin³, Mara Novero², Giovanni

2 Gabelli², Eleonora Paparelli^{4, 5}, Fabio Marroni^{4, 5}, Paola Bonfante²

Impact of an arbuscular mycorrhizal fungus *versus* a mixed microbial inoculum on the transcriptome reprogramming of grapevine roots

¹Istituto per la Protezione Sostenibile delle Piante del CNR, SS Torino, Viale P.A. Mattioli 25,
10125 Torino, Italy; ²Dipartimento di Scienze della Vita e Biologia dei Sistemi, Università degli
Studi di Torino, Viale P.A. Mattioli 25, 10125 Torino, Italy; ³Centro di Genomica Funzionale
dell'Università di Verona, Strada le Grazie 15, 37134 Verona, Italy; ⁴Dipartimento di Scienze
Agroalimentari, Ambientali e Animali (DI4A), Università degli Studi di Udine, Viale delle Scienze
208, 33100 Udine, Italy; ⁵Istituto di Genomica Applicata (IGA), Via J. Linussio 51, 33100 Udine,

12

- 13 *Corresponding author:
- 14 Raffaella Balestrini
- 15 <u>raffaella.balestrini@ipsp.cnr.it</u>
- 16 Phone: 00390116502927
- 17 Fax: 00390116705962
- 18
- 19
- 20
- 21
- 22

23 Acknowledgments

This work was supported by the VIT-INNOVA project (Val D'Aosta) to PB. The authors thank all the partners for the collaboration, the Centro di Genomica Funzionale dell'Università di Verona for the support in the bioinformatics analyses, Matteo Chialva for the KEGG pathways, Walter Chitarra for Figure 3 preparation and Samuele Voyron for the Bioanalyzer analysis.

28 Abstract

Grapevine, cultivated for both fruit and beverage production, represents one of the most economically important fruit crops worldwide. With the aim of better understanding how grape roots respond to beneficial microbes, a transcriptome sequencing experiment has been performed to evaluate the impact of a single arbuscular mycorrhizal (AM) fungal species (Funneliformis mosseae) versus a mixed inoculum containing a bacterial and fungal consortium, including different AM species, on Richter 110 rootstock. Results showed that the impact of a single AM fungus and of a complex microbial inoculum on the grapevine transcriptome differed. After three months, roots exclusively were colonized after the F. mosseae treatment, and several AM marker genes were found to be up-regulated. The mixed inoculum led only to traces of colonization by AM fungi, but elicited an important transcriptional regulation. Additionally, the expression of genes belonging to categories such as nutrient transport, transcription factors, and cell wall-related genes was significantly altered in both treatments, but the exact genes affected differed in the two conditions. These findings advance our understanding about the impact of soil beneficial microbes on the root system of a woody plant, also offering the basis for novel approaches in grapevine cultivation.

44 Keywords

			• •	•		
15	A M gymbiogig.	microhial	inoculum.	aranevine.	root transerir	tome protile
45	AIVI SVIIIUIUSIS.	microular	moculum,	grapevine,	100t transcrip	
	,		,	<u> </u>		



56 Introduction

Grapevine, cultivated since the dawn of civilization for the production of fruit, juice and wine, represents one of the most economically important fruit crops worldwide, with widespread cultivation (77,181,122.00 Mt produced in 2013; <u>http://faostat.fao.org</u>) and high commercial value (Vivier and Pretorius 2002). It has become a model organism for fruit trees, as mirrored by the two genome-sequencing projects developed on grape (Jaillon et al. 2007; Velasco et al. 2007). The molecular regulation occurring during berry development has been investigated using several

high-throughput technologies (Zenoni et al. 2010; Fasoli et al. 2016). Additionally, in the last years,
several studies investigated different aspects related to water transport and water deficit impact
(Perrone et al. 2012; Chitarra et al. 2014; Tombesi et al. 2015; Corso et al. 2015) as well as to the
interactions with pathogens (Milli et al. 2011; Dal Santo et al. 2013; Vitali et al. 2013; Pantaleo et
al. 2016).

Only a few papers so far have been published on transcriptomics in *Vitis spp.* roots. Du et al. (2014) recently studied the root transcriptome, using the Affymetrix *V. vinifera* genome array, to verify the impact of phylloxera attack in a resistant rootstock (140Ru) and in the susceptible cultivar "Crimson Seedless". Because of the ongoing climate change in wine-growing regions, the selection of rootstocks tolerant to several biotic and abiotic stresses is considered a crucial factor for developing sustainable agriculture. As a consequence, next generation viticulture is aimed to select appropriate rootstocks bred from several *Vitis* species (Corso and Bonghi 2014; Flexas et al. 2009).

75 Grapevine is highly responsive to local environmental conditions and vineyard management 76 practices. In this context, Anesi and colleagues (2015) have suggested a genome plasticity in relation to environment, overall known as terroir, that characterizes a specific vineyard and impacts 77 grape and wine quality. Soil qualities, rootstocks, location, climatic factors, and soil management 78 have been reported to influence grape development and fruit and wine quality (Koundouras et al. 79 2006; de Andrés-de-Prado et al. 2007; Marè et al. 2013). The transcriptome variation in relation to 80 81 different soils and rootstocks recently has been studied in leaves of the scion cv. Pinot noir through 82 a microarray approach, suggesting a link among soil composition, rootstock and gene expression (Marè et al. 2013). Today, viticulturists aim to produce high quality wine, increasing profit from the 83 land and reducing agronomic inputs, through encouraging natural soil beneficial organisms 84 (Trouvelot et al. 2015). 85

In this changing context, many Italian wines are now labelled as "organic wines" and are produced by introducing commercially available microbial inoculants to the soil. These inoculants include bacteria belonging to the genera *Bacillus*, *Pseudomonas*, *Streptomyces* and biocontrol fungi such as *Trichoderma spp.* and/or arbuscular mycorrhizal (AM) fungi (Pinto and Gomes 2016). The latter

microbes are among the most relevant soil organisms that colonize the roots of most land plants, 90 where they facilitate mineral nutrient uptake from the soil in exchange for plant-assimilated carbon 91 92 (Bonfante and Genre 2010). It is already known that vineyard soils support indigenous AM fungi 93 (AMF) and it is well established that grapevine roots are colonized by native AMF (Balestrini et al. 2010; Trouvelot et al. 2015). The use of molecular approaches, including metagenomics and/or 94 Next Generation Sequencing (NGS) techniques, has offered new information about the AMF 95 assemblages that live in symbiosis with this important, typical Mediterranean fruit crop (Schreiner 96 and Mihara 2009; Balestrini et al. 2010; Lumini et al., 2010; Holland et al. 2014). By contrast, the 97 molecular basis underlying the interactions between grapevine and AM fungi still has to be 98 99 elucidated. While RNAseq techniques have been used to study transcriptome profiles in AMcolonized roots from herbaceous plants such as tomato, rice and Lotus japonicus (Ruzicka et al. 100 2012; Fiorilli et al. 2015; Handa et al. 2015), information is scarce on woody crops. Recently, 101 102 transcriptome data have been obtained from litchi (Litchi chinensis Sonn.) roots, identifying 103 transcripts involved in the interaction with AM fungi under carbohydrate starvation (Shu et al. 2016). 104

With the aim to better understand how grape roots respond to beneficial microbes, a transcriptome
 experiment has been performed to evaluate the impact of a single AM fungal species (*Funneliformis mosseae*) versus a mixed inoculum containing a bacterial and fungal consortium on Richter 110
 rootstock.

109

110 Materials and Methods

111 Plant material and growth conditions

112 Pinot noir plants grafted on 110 Richter rootstock have been prepared at Roero Viti Vivai 113 (http://www.roerovitivivai.it/). In detail, after that rootstock developed roots, grafted plants were 114 grown in pots with a natural soil collected in vineyards of Val d'Aosta, Italy (characterized by 81%) sand, 16% loam and 3% clay; pH 7.70; organic matter content 3.73 g/100g; cation exchange 115 116 capacity 15.47 meq/100g), previously sterilised. The same sterilised soil was supplemented with an inoculum of Funneliformis mosseae (formerly Glomus mosseae) Gerd. & Trappe (BEG 12) 117 118 purchased from MycAgro Lab for single species inoculation (FMOS; 30% inoculum/soil for each 119 plant), or with the mixed inoculum MICOSAT F® Vite (CCS-Aosta) containing a microbial 120 consortium including AM fungi (MICO; 30 g for each plant). In detail, FMOS inoculum consisted of spores, extraradical mycelium, sorghum mycorrhizal roots and sorghum growth substrate and 121 122 each plant has been inoculated with about 1000 propagules. MICO inoculum, as stated in the

website (http://www.micosat.it/portfolio/fertilizzante-micosat-f-vite/), contains: Trichoderma viride, 123 124 T. harzianum, Pochonia chlamidosporia, Streptomyces spp. ST60, Streptomyces spp. SB14, 125 Streptomyces spp. SA51, Bacillus subtilis BA41, Pseudomonas fluorescens PN53, Pseudomonas 126 spp. PT65, Glomus spp. GB67, Glomus mosseae GP11, Glomus viscosum GC41 in the percentage of 40% crude inoculum (AM fungi) and 21.6% bacteria and saprotrophic fungi. In parallel, control 127 128 plants (CTRL) have been prepared using only the sterilized soil. Plants were grown in greenhouse conditions from July to October 2013 under natural light and temperature, with drip irrigation for 129 one hour every 15 days with slight intensification in days of high heat peaks. At least 33 plants for 130 each of the three considered conditions (CTRL, FMOS, MICO) were used. At the end of the 131 132 experiment, thirty randomly chosen 1-cm-long root segments per plant were treated in 10% KOH for four hours at room temperature, stained with 0.1% cotton blue in lactic acid and then fungal 133 colonization was quantified according to the Trouvelot system (Trouvelot et al. 1986) using the 134 135 MYCOCALC software. Root segments, obtained from at least fifteen plants, were analysed. The remaining roots were stored at -80°C until molecular analysis. 136

To evaluate fresh and dry root weight under the different conditions, five entire root systems for each treatment have been weighted (fresh weight, FW), dried at 60°C for three days and weighted again (dry weight, DW).

140

141 RNA extraction and Illumina sequencing

142 For the RNASeq experiment, roots were harvested from the CTRL, FMOS and MICO plants after three months from the inoculation. Total root systems were chilled in liquid N₂ and RNA was 143 extracted using the 'pine tree-method' (Chang et al. 1993) with the addition of 2% PVPP to the 144 extraction buffer. For each growth condition, we used two biological replicates, each containing the 145 146 pooled RNA from roots sampled from three plants. RNA quality and quantity controls have been performed using the Agilent 2100 Bioanalyzer. Ten micrograms of each RNA sample (RIN >8) 147 were sent to HUGEF (Torino, Italy) where the libraries were produced and sequenced using an 148 Illumina Genome Analyzer (Solexa). The six libraries were indexed, and single-end multiplexed 149 150 sequencing was performed using 100 bp length reads. The reads obtained from Illumina HiSeq were 151 processed using CASAVA pipeline version 1.8.2. (Illumina Inc, San Diego, CA, USA) and further 152 checked for sequence quality with the fastQC application (ver. 0.10.1).

154 **Bioinformatic methods**

155 Expression profiling and differential expression analysis

To determine gene expression levels, reads were mapped against *Vitis vinifera* 12x genome using TopHat version 2.0.12 with default parameters, and alignments were processed with Cufflinks version 2.2.0 (Trapnell et al. 2013). Cuffdiff was used to detect differentially expressed genes, and a false discovery rate (FDR) of 0.05 (Benjamini and Hochberg 1995) was set as a significance threshold. Sample clustering based on Jensen–Shannon distances between conditions and replicates was performed using the R package CummeRbund (Goff et al. 2013).

162 Functional annotation and GO-enrichment analysis

Vitis 163 vinifera annotations were retrieved from the VitisNet gene portal (https://www.sdstate.edu/ps/research/vitis/pathways.cfm; Grimplet et al. 2012). To identify protein 164 domains and CAZyme domains, V. vinifera predicted proteins were annotated with PFAM release 165 166 27 and dbCAN v.3 databases using Hmmer version 3.1b1 with default parameters. GO-terms and KEGG pathways annotation were performed with Blast2GO version 2.8 using default parameters 167 (Conesa et al. 2005). To detect significantly enriched GO-terms in differentially expressed genes 168 (DEGs), two-tailed Fisher Exact Tests were performed and an FDR of 0.05 was set as a significance 169 170 threshold.

171 **Quantitative RT-qPCR validation**

All the RNA samples were treated with the Turbo DNA-free[™] kit (Ambion, Austin, TX, USA) for 172 RT-qPCR analyses according to the manufacturer's instructions. The RNA samples were submitted 173 174 to a control reverse-transcription PCR to check for the absence of DNA contamination using the One Step RT-PCR kit (Qiagen) and primers specific for the grapevine elongation factor 1- α gene 175 176 (*VvEF1-a*, Reid et al. 2006; Table S1). First strand cDNA was synthesized from 500 ng of total RNA with the Superscript II reverse transcriptase kit (Invitrogen) following the manufacturer's 177 178 instructions. At the end of the reaction, the cDNA was diluted to 1:3 for the gene expression 179 analysis. Gene specific primers (Table S1) were designed using Primer 3 180 (http://bioinfo.ut.ee/primer3-0.4.0/primer3/). Quantitative RT-PCR (RT-qPCR) reactions were carried out in a 48-well StepOne[™] Real time PCR system instrument (Applied Biosystems), in a 181 182 final volume of 20 μ l, containing 10 μ l of 23 iQ SYBR Green Supermix, 4 μ l of primers 3 μ M, 5 μ l of water and 1 µl of cDNA template. The PCR program consisted of a holding stage (95°C for 10 183 min) and 40 cycles of 95°C for 15 sec. and 60°C for 1 min. A melting curve (55-95°C with a 184 heating rate of 0.5°C per 10 sec. and a continuous fluorescence measurement) was recorded at the 185

end of each run to assess amplification product specificity. All the reactions were performed with three technical replicates and three biological replicates. A portion of the grapevine VvEF1- α gene was used as the housekeeping gene for normalization (Chitarra et al. 2014), by subtracting the CT

value of $VvEF1-\alpha$ from the CT value of the candidate gene resulting from the Δ CT. The expression

ratios were calculated without the PCR efficiency correction from equation $2^{\Delta\Delta CT}$; where $\Delta\Delta CT$

191 represents the ΔC_T sample – ΔC_T control. Before calculating the ΔC_T , the technical replicates were

192 checked for their C_T value uniformity and for outliers, which led to the exclusion of any standard

- deviations above 0.2. The primer names and corresponding sequences are listed in Table S1.
- 194

195 Phosphorous and potassium determination in roots

To determine P and K, about 2 mg of root system (4 plants for each condition) were dried for two days at 60°C, ground and digested at 95°C for 1 hour in 1 ml HNO₃ 6M, filtered using a glass filter and diluted with distilled water 1:6. The cations were determined with a Perkin Elmer Optima 7000 (Perkin Elmer, Norwalk, Connecticut, USA) inductively coupled plasma-optical emission spectrometer (ICP-OES). Standard solutions were prepared from concentrated stock solutions (Merck Titrisol or Sigma Aldrich). High purity water (HPW) produced with a Millipore Milli-Q system was used throughout. The reagents used were of analytical grade.

203

204 Statistical analyses

All the data were subjected to statistical analysis using SYSTAT 10 software, applying the nonparametric Kruskal-Wallis test adopting a probability level of P < 0.05.

207

208 **Results**

209 Plant development and root features

After 3 months of growth under greenhouse conditions, all the plants presented a similar vegetative development (not shown). Fresh and dry root weight were evaluated (Figure S1): no statistically significant differences in root biomass were found among the three treatments (FMOS, MICO, CTRL) nor in root P and K concentration, where slightly higher values were recorded in FMOS plants (Figure S1).

The roots of the plants inoculated with the mixed inoculum (MICO) did not present a significant AMF colonization, with only three plants presenting traces of AM fungal colonization (Table S2): a

morphological quantification revealed a mean number of colonized fragments (F) of 2.6% and a mean intensity of mycorrhization (M) of 0.88 % (Figure 1). AM fungal colonization was observed in the FMOS roots, with a mean number of colonized fragments of 80.66 % (F) and a mean intensity of mycorrhization (M) of 48.93% (Figure 1). Arbuscules (a%) and vesicles were highly variable; their percentage ranged from 2.66 to 41.13 and from 4.86 to 21.45, respectively. No colonization structures were detected in the CTRL roots (not shown).

223

224 Analyses of RNA-Seq data: read number, transcriptome coverage and total expressed genes

Sequencing of RNA samples produced on average 20,147,903 of reads per sample (Table 1). Sequencing reads ranging from 12 to 32 million for each sample (Table 1) were mapped on the *Vitis vinifera* genome obtaining on average ~93.2% overall alignment rate. Based on the sum of transcript lengths, as reported in the current V1 annotation (39,893,396 bp that does not currently account for transcript isoforms), we reached an average transcriptome coverage of 24x for each replicate.

A look at the whole root transcriptome and at the differentially expressed genes

232 A total of 9,593 genes were expressed in all samples when using a cut-off value of RPKM > 0 to 233 declare a gene as expressed. Based on expression values, the samples clustered by condition (Table 234 S3, Figure S2). In order to identify genes involved in root development we mined the keyword 235 "root" from the gene descriptions of *Vitis vinifera* V1 annotation thus obtaining 57 candidate genes. 236 Among them, a total of 33 genes were expressed in all our samples setting a cut-off value of RPKM > 0 to declare a gene as expressed. This data set comprises genes that seem to be specific to the 237 root, as they are reported to be involved in different root developmental processes looking at GO 238 239 descriptions and Blast2GO (ver. 3.3) results (not shown). As a second step, transcriptional changes 240 were determined by comparing F. mosseae-colonized roots (FMOS) and those inoculated with the mixed inoculum (MICO) with un-inoculated control (CTRL) samples. This analysis revealed 539 241 and 737 differentially expressed genes (DEGs) in FMOS and MICO plants, respectively (Figure 242 243 2a). The mixed inoculum led to the regulation of a higher number of genes compared to the AMF 244 inoculum, and a large proportion of them (85%, on average) were down-regulated in both 245 treatments versus the control condition. Only about 30% of the FMOS DEGs were found in common with those of MICO roots, suggesting that the AM fungus and the mixed inoculum have 246 led to different transcriptome profiles. Most of them presented a common trend in the two 247 248 conditions (Table S4), but some instances were observed in which regulation in the two treatments

was in different directions. Some DE transcripts were specific to a single condition (Table S5). In
addition, for three DEGs, FPKM have been found only in MICO roots and not in the control, i.e.
genes annotated as coding for a putative arachidonic acid-induced protein DEA1
(VIT_12s0035g02000), an unknown protein (VIT_03s0132g00060) and a no hit protein
(VIT 13s0047g00580), although with low FPKM values (1,145; 0,697; 5,710 respectively).

254 Among the 10 most up-regulated genes, seven transcripts were common between to the two conditions (Table 2). The first, with a fold change of 2.85 and 3.64 in FMOS and MICO 255 256 respectively, was a gene coding for a putative uroporphyrin III methylase (VIT 13s0064g01470). A 257 corresponding (homolog) gene has been reported to be up-regulated in luxuriant (N+) treated 258 Eucalyptus plants versus limited (N-) plants (Camargo et al. 2014), while in Arabidopsis roots a low 259 expression has been found under Cd treatment (van de Mortel et al. 2008). VIT 03s0063g00370 260 and VIT 18s0001g03910, which code for a putative ferredoxin nitrite and a putative nitrate reductase, respectively, have a role in nitrate/nitrite assimilation, and might be regulated by nitrate, 261 262 as previously demonstrated in Arabidopsis (Wang et al. 2003). In the same list, we also detected a gene encoding a putative nitric oxide reductase (VIT 06s0004g04400) as well as a gene coding for 263 264 a nodulin belonging to the MtN21 family (VIT 01s0026g00550).

The expression of 14 genes randomly selected from those identified in the RNA-seq experiment was successfully validated by RT-qPCR (Figure S3 and Figure 3).

To have an overview of the regulation of the main metabolic processes and signalling pathways 267 involved in the different comparisons, we conducted GO enrichment analysis. Figure 4 shows the 268 269 enriched GO terms specific for FMOS and for MICO, while the GO terms over-represented in both 270 growth conditions in response to AM fungus and the mixed inoculum are represented in Figure 5. Differentially expressed transcripts were grouped in functional classes, on the basis of the specific 271 biological process in which they were involved (Table S4; Table S5; Figure 4a, b). Several GO 272 273 terms were over-represented and among them transport and transporter activity categories were 274 over-represented in MICO roots in addition to cell wall, membrane, cell component organization 275 (Figure 4a). Among the over-represented genes in FMOS roots, response to endogenous stimulus, response to abiotic stimulus, nucleus, RNA biosynthetic process, and cell cycle were annotated 276 277 (Figure 4b). Six over-represented functional GO classes were found in common between the two different treatments: cellular components organization, cell cycle, nucleus, extracellular region, 278 279 carbohydrate metabolic process, cell-wall (Figure 5). CAZymes domains analysis showed that 280 MICO sample expressed genes contains several annotated CAZymes domains which are represented also in the whole V. vinifera dataset, such as glycosiltransferases (GT), glycoside 281

hydrolases domains (GH), carbohydrate-binding modules (CBM) and carbohydrate esterases (CE)

- 283 (Figure 6).
- 284

285 Grapevine transcripts: specific responses to the AM fungus *versus* the mixed inoculum

To better explore the novel transcriptomic data set, and to further understand the grapevine response 286 287 to AM fungi, we studied in greater detail the expression profiles of genes described in the literature as specifically involved during AM symbiosis in legumes, rice and tomato (Fiorilli et al. 2009; 288 289 Guether et al. 2009; Hogekamp et al. 2011; Handa et al. 2015; Fiorilli et al. 2015). AM symbiosis is 290 mostly acknowledged for the improved nutrient exchange established between the two symbionts, 291 and regulated by the activities of fine-tuned plant and fungal transporter genes (Casieri et al. 2013; 292 Berruti et al. 2016a). In accord with this claim, a consistent group of plant transporters were identified as differentially expressed between treated (FMOS and/or MICO) and control plants 293 294 (Table S6). One of them is an inorganic phosphate transporter gene (VIT 16s0050g02370) that 295 shows homology with the mycorrhiza-inducible inorganic phosphate transporters such as LePT4 296 and OsPT11 (Table 2; Figure S3), and an oligopeptide transporter 4 (VIT 18s0001g07940). Several genes encoding for protein involved in the transport of molecules across cell membranes were also 297 298 up-regulated in the same FMOS roots. Among them, genes coding for a putative potassium (K^{+}) 299 transporter KUP1 (VIT 19s0027g01820), a sulphate transporter (VIT 05s0020g03970), a lysine 300 histidine transporter (LHT, VIT 01s0011g03180), and an organic cation transport protein OCT1 301 (VIT 17s0119g00080) were exclusively up-regulated in FMOS roots. By contrast, other 302 transporters were up-regulated in both the conditions: among them, two genes coding for putative nitrate transporters (VIT 17s0000g09470 and VIT 01s0127g00070) and a Zinc transporter 10 303 precursor (VIT 10s0042g01100) as well as three protease inhibitor/seed storage/lipid transfer 304 305 protein (LTP) genes (Table 2). An additional VIT 09s0002g05660 sulphate transporter Sulp family 306 was up-regulated under both treatments. In agreement with previous work (Zhang et al. 2010; 307 Hogekamp et al. 2011), four ABC-transporters were identified as co-induced in FMOS roots (VIT 07s0031g02550; VIT 13s0074g00690; VIT 16s0098g00570; VIT 04s0008g04790), and 308 309 only the last one was also up-regulated in MICO roots. Putative ammonium transporter genes 310 (VIT 04s0008g05080, VIT 00s1818g00010, VIT 00s0179g00310, VIT 07s0031g02950) were significantly down-regulated in MICO roots as well as a gene corresponding to a putative NIP 1;2 311 312 (VIT 10s0003g01830).

The presence of AM fungi leads to relevant changes in the hormonal plant profile (Gutjahr 2014), including gibberellins, which are predicted to modulate their concentrations during the symbiosis. A gene coding for a DELLA protein GAI1 (VIT 17s0000g10300) was found as up-regulated in FMOS and significantly down-regulated in MICO samples. Strigolactones not only represent a new class of plant hormones, but also may stimulate fungal branching while acting as pre-symbiotic molecules (Bonfante and Genre 2015). A gene involved in strigolactone (SL) biosynthesis, i.e. a carotenoid cleavage dioxygenase (CCD) 7 (VIT_15s0021g02190; *VvCCD7*) was found to be upregulated in FMOS samples. Interestingly, *VvCCD7* transcripts were not detected in MICO roots, consistent with the almost total absence of AM fungal colonization.

Homologs of nodulin genes (Denancè et al. 2014) were found to be differentially regulated in our 322 datasets. In FMOS, genes encoding two MtN3 nodulins (VIT 16s0050g02540 323 and VIT 17s0000g00820) were up- and down-regulated respectively, while two MtN21 genes were up-324 325 regulated (VIT 03s0017g02310 and VIT 01s0026g00550). One of these last (VIT 01s0026g00550) was up-regulated also in MICO roots together with another MtN21 gene 326 (VIT 04s0023g02730), while two others were down-regulated (VIT 13s0084g00090 and 327 328 VIT 08s0040g02500).

Among genes potentially involved in arbuscule formation and fungal accomodation, two genes encoding CESA cellulose synthase (VIT_10s0003g01560 and VIT_07s0005g04110), a gene coding for a cellulose synthase IRX3 (VIT_11s0037g00530), a chitinase class III (VIT_16s0050g02210), and a subtilisin serine endopeptidase gene (VIT_15s0048g01200) were exclusively up-regulated in FMOS roots. A gene encoding a laccase (VIT_08s0007g01910) was also up-regulated only in FMOS, in addition to a metallothionein (VIT_08s0007g00330).

335 Both FMOS and MICO conditions elicited the expression of several TF genes belonging to different groups, while other members inside these families were down-regulated (Table S3; Table 336 337 S4). In detail, genes belonging to GRAS, DOF, Zinc-finger, MYB and DREB transcription factor groups were found to be up-regulated in FMOS roots. Members inside these groups were also up-338 339 regulated in MICO samples. Additionally, it is worthwhile to note the regulation in AM-colonized 340 roots of several circadian-regulated genes as well as genes potentially involved in the response to 341 environmental stimuli (Table S3). Among them, the grapevine homolog to arachidonic acid-induced 342 protein DEA1 (VIT 12s0035g02000) and a gene coding for a protein belonging to the RD22-like 343 subfamily (VIT 04s0008g04150). By contrast, genes potentially involved in response to pathogens, 344 such as stilbene synthase genes, were down-regulated in FMOS roots as well as four genes coding 345 for Avr9/Cf-9 rapidly elicited protein 20. In contrast, no stilbene synthase genes were regulated in 346 MICO roots.

Among differentially expressed genes in MICO roots, we again found a consistent core of genes involved in nutrient transport, but interestingly several of them had a different ID than those for FMOS, revealing a specific response to the inoculum (Table S3; Table S4). Among them, two

high affinity nitrate transporter (VIT 06s0061g00310 and VIT 06s0061g00320), two additional 350 351 nitrate transporters (VIT 11s0016g05170 and VIT 18s0001g11280), a sulphate transporter 1.3 352 (VIT 18s0001g04910), a ZIP family transporter (VIT 19s0085g00740), an arsenite transport 353 protein (VIT 02s0025g03310), a Mg-chelatase subunit XANTHA-F (VIT 08s0007g08540) were up-regulated. Differently from the expression profile described in FMOS (Table S3; Table S4), 354 genes coding for putative ammonium transporters, and a gene coding for a cationic amino acid 355 transporter 2 (VIT 10s0003g04540) were down-regulated. Among genes involved in other 356 357 pathways that potentially could be affected by the presence of bacteria (i.e. hormonal balance and defense; Vacheron et al. 2013; Drogue et al. 2014), an auxin response factor 3 358 359 (VIT 10s0003g04100) is specifically up-regulated in addition to a gene coding for the ABA 8'hydroxylase CYP707A1 (VIT 02s0087g00710), which is a key catabolic enzyme and could be 360 involved in the regulation of ABA level (Okamoto et al. 2006). Ethylene responsive factors genes 361 362 (ERF), all were down-regulated in the MICO-treated roots, confirming previous results obtained in different beneficial plant-bacterium interactions (Verhagen et al. 2004; Drogue et al. 2014). Two 363 genes encoding pore-forming toxins (VIT 07s0005g06090 and VIT 07s0005g06110) were 364 365 specifically up-regulated in the MICO treatment.

366

367 Discussion

In this work we have developed new transcriptomic data sets that illustrate the main pathways activated in grapevine roots as well as those elicited by beneficial microbes. Although grapevine is a woody plant with economic relevance for berry production, limited attention has been given so far to its root system and how its transcriptome responds to AM fungi and Plant-Growth Promoting Bacteria (PGPBs). In addition, grapevine is characterized by a secondary growth pattern, but, with a few exceptions (Shu et al. 2016), all the transcriptional profiles following microbial colonization so far available refer to herbaceous crop plants.

In our experimental set up we investigated the impact of both a single AM fungus (*Funneliformis mosseae*), which is considered a symbiotic fungus for many crops, and a microbial consortium, which is commercialized as suitable for grapevine.

The two data sets derived from plants inoculated with microbes reveal some interesting similarities: they both present a limited number of differential expressed genes and a higher number of downregulated genes in respect to previous papers on the transcriptome profiles in AM- and/or PGPBcolonized roots. Many previous experiments demonstrated that up to 60-70% of genes were up regulated during AM symbiosis (Guether et al. 2009; Fiorilli 2009; Handa et al. 2015; Hogekamp et

al. 2011; Fiorilli et al. 2015). Nevertheless, examples of transcriptomes with prevalent down-383 384 regulated genes already have been reported. Drogue et al. (2014) have analyzed four different plant-385 microbe combinations using two Azospirillum strains and two rice cultivars, and only in Nip B510 386 combination the regulated genes were mainly down-regulated (203 up-regulated vs 2336 downregulated). Interestingly, in a recent study focused on the transcriptome of mycorrhizal litchi roots, 387 388 Shu, et al. (2016) found a number of down- (156) and up- (286) regulated genes, with a ratio between up- and down-regulated genes lower than other previous works. We cannot exclude that 389 woody plants differently modulate their root transcriptome in the presence of beneficial microbes, 390 or they may require a different and/or longer timing than herbaceous plants. 391

392

393 A single microbe *versus* a consortium

394 Overall, the analysis of the generated data sets revealed that the impact on the gene expression of a 395 single AM fungal species and of a complex microbial inoculum on the grapevine transcriptome was 396 diverse. The differences in transcriptome profiles mirrored morphological observations (Figure 1) showing a good AM colonization in FMOS and only traces of AM fungal hyphae in MICO roots. 397 398 Despite the complex microbial inoculum label indicated the presence of AM fungal propagules, the 399 AM colonization was found in low amount. This is in agreement with previous data obtained using similar microbial formulations produced by the same company: a very low AM fungal colonization 400 intensity was detected in *Camellia japonica* rooted cuttings (Berruti et al. 2013), and the AM fungal 401 402 taxa inoculated failed to colonize maize roots and lacked soil persistence (Berruti et al. 2016b). 403 However, the mixed inoculum provided us the opportunity to test a complex condition, because - as 404 in the soil – grapevine was in contact with multiple microbes. Looking at the genes significantly regulated in FMOS and MICO roots, genes belonging to the same categories (e.g. nutrient transport, 405 TF, cell wall metabolism) have been found to be up-regulated in both conditions, but several of 406 them had different IDs, suggesting a specific response to the specific inoculum. 407

408 The AM fungus activates many of the AM-symbiosis markers that are at the moment considered the 409 functional signatures of the symbiosis (Guether et al. 2009a; Gomez et al. 2009; Hogekamp et al. 2011). Among them major attention can be directed to a gene coding for a protein that shows a high 410 411 identity with LePT4, a phosphate transporter that probably also acts as a sensor of phosphate 412 availability in the soil and inside the root environment (Volpe et al. 2015). However, several up-413 regulated nutrient transporter genes (12) were identified, in agreement with that reported in Lotus japonicus where 43 nutrient transporters were identified as up-regulated in the transcriptome of 414 415 mycorrhizal roots (Guether et al. 2009a). In our work, several of the common up-regulated genes

between the two treatments (FMOS and MICO) are involved in nitrogen metabolism, mainly in 416 417 relation to nitrate, suggesting that in grapevine these beneficial root-associated microbes are 418 particularly efficient in stimulating plant responses to nitrogen, which is an essential element for all 419 grapevine processes (http://www.awri.com.au/wp-420 content/uploads/1 nutrition nitrogen fertilisation.pdf). In contrast, transcripts corresponding to 421 other genes which are considered functional marker genes of the AM symbiosis and expected to be 422 up-regulated in the presence of the AM fungus, such as for example ammonium transporter and NIP aquaporin genes (Guether et al. 2009b; Giovannetti et al. 2012), were found in AM-colonized 423 424 grapevine roots although not significantly up- or down-regulated in the comparison with CTRL 425 plants.

426 Nodulin genes, first described as legume genes involved in root nodule symbiosis development and also reported as up-regulated in AM symbiosis, showed a differential regulation between the two 427 428 datasets. Twelve MtN3/saliva/SWEET genes (Sugars Will Eventually be Exported Transporters) 429 have been reported in Vitis vinifera, while 23 MtN21/EamA-like/UMAMIT genes (Usually 430 Multiple Acids Move In and out Transporters) have been found (Denancè et al. 2014). Recently, plant SWEETs have been shown to be involved in the feeding of pathogenic microbes, and an 431 432 important role for SWEET transporters during the mycorrhizal symbiosis has been suggested, 433 although further analyses are needed to clarify their role during symbiosis (Manck-Götzenberger 434 and Requena 2016).

435

As expected, a core of differentially regulated genes involved cell wall-related genes. The induction 436 437 in mycorrhizal roots of a large number of genes related to membrane dynamics and cell wall metabolism is well documented, consistent with the concept that plant cells have an active role in 438 439 fungus accommodation via membrane proliferation and cell wall construction (Balestrini and 440 Bonfante, 2014). Here, three genes encoding three putative cellulose synthase, two CESA and one 441 IXR3 respectively, have been found to be up-regulated only in the presence of the AM fungus. 442 While CesA proteins are part of the cellulose synthase complex in higher plants (Taylor 2008), and 443 in L. japonicus, transcripts of a putative cellulose synthase, LjCesA, have been demonstrated to 444 accumulate in arbusculated cells, IXR3 (AtCesA7) has been reported as required for secondary wall 445 cellulose synthesis in Arabidopsis (Richmond and Sommerville 2000). Genes putatively involved in 446 lignin biosynthesis (Barros et al. 2015), such as a putative cinnamoyl-CoA reductase and a 447 cinnamyl alcohol dehydrogenasegene gene, also were found to be up-regulated in the presence of 448 the AM fungus. Because these genes already have been described during plant-PGPR (Plant Growth

Promoting Rhizobacteria) interactions (Vacheron et al. 2013), but never as AM-inducible, wesuggest that they may take part in a response that is specific for woody hosts.

Another novel set of genes which has never been reported as AM-inducible concerns circadianrelated genes (Carbonell-Bejerano et al. 2014). They have been deeply investigated in model plants like *Arabidopsis*, but also have been reported as crucial in ecologically relevant symbioses such as corals living with their photosynthtic algae (Sorek et al. 2014). Circadian clock-related genes have been identified in the grapevine genome and oscillation in their expression has been correlated with the daily oscillatory changes in the berry transcriptome at late ripening stages (Carbonell-Bejerano et al. 2014).

458 It already was reported that PGPBs can affect plant physiology and growth, including root system architecture, and that these modifications are accompanied by changes in plant transcriptome 459 profiles (Vacheron et al. 2013). An impact on the root transcriptome has been reported in PGPB-460 461 treated roots with several bacterial models, and the differences in the regulated transcripts were 462 mainly related to the used species/strain (Verhagen et al. 2004; Weston et al. 2012; Vacheron et al. 2013; Drogue et al. 2014; Spaepen et al. 2014). Interestingly, an Azospirillum strain (B510) that can 463 colonize the outer layer of rice root tissue leads to a repression of a wider set of genes involved in 464 465 stress response and defence than a strain that was shown to colonize only the rice-root surface 466 (Drogue et al. 2014). Indeed, some plant-associated bacteria are known as ISR (Induced Systemic 467 Resistance)- bacteria, while others directly promote plant growth, thus leading to different plant gene expression profiles. In our study, a mixed inoculum containing different bacteria 468 469 species/strains has mirrored a natural environment (where plants encounter different bacterial 470 species). Genes belonging to different categories (e.g. transcriptional regulation, nutrient transport, 471 hormonal balance, cell wall metabolism) have been identified as regulated in PGPB-treated roots, 472 showing an impact on different root processes, in agreement with previous transcriptome studies. In 473 addition, genes encoding pore-toxin proteins were found to be up-regulated in the MICO condition. 474 These proteins are the most common bacterial cytotoxins and are required for virulence in a large 475 number of important pathogens. Interestingly, pore-forming proteins with remarkably similar 476 structures to pore-forming toxins (PFTs) are found in vertebrates and constitute part of their 477 immune system (Dal Peraro and van der Goot 2016). The functions of these proteins has remained 478 unclear in higher plants, but their transcription levels were greatly increased under biotic stress 479 (Shao et al. 2015).

In conclusion, with an experimental set up which has allowed the plants to grow in a comparable way irrespective of the microbial inoculum, we found that AM fungi may elicit in grapevine most of the responses which have already been characterized in crop and herbaceous plants. This is a

- further confirmation that the symbiotic pathway operating in the plants as a consequence of the AM
 presence is very ancient and conserved irrespective of a plant's phylogenetic position.
- The mixed inoculum led to a very low colonization by AMF, but elicited an important transcriptional regulation, which, as a consequence, probably can be assigned predominantly to the presence of the PGPBs.
- Because grapevines live in association with multiple bacterial and fungal communities (Trouvelot et al. 2015; Pinto and Gomes 2016), our data offer a starting point to dissect the grapevine response both to a single microbe and to a mixed inoculum, offering a basis for the development of novel
- 491 approaches in vineyard practices.
- 492
- 493

494 **Conflict of interest**

- 495 The authors declare that they have no conflict of interest.
- 496

497 **Reference list**

498 Anesi A, Stocchero M, Dal Santo S, Commisso M, Zenoni S, Ceoldo S, Tornielli GB, Siebert TE,

Herderich M, Pezzotti M, Guzzo F (2015) Towards a scientific interpretation of the terroir concept:

- plasticity of the grape berry metabolome. BMC Plant Biology 15:191. doi: 10.1186/s12870-015-
- 501 0584-4

502 Balestrini R, Magurno F, Walker C, Lumini E, Bianciotto V (2010) Cohorts of arbuscular

- mycorrhizal fungi (AMF) in *Vitis vinifera*, a typical Mediterranean fruit crop. Environ Microbiol
 Rep 2:594-604.
- 505 Balestrini R, Bonfante P (2014) Cell wall remodeling in mycorrhizal symbiosis: a way towards
- 506 biotrophism. Front Plant Sci 5:237. doi: 10.3389/fpls.2014.00237
- Barros J, Serk H, Granlund I, Pesquet E (2015) The cell biology of lignification in higher plants.
 Ann Bot 115:1053-1074.
- 509 Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful 510 approach to multiple testing. J Royal Stat Soc 57:289-300. Retrieved from
- 511 <u>http://www.jstor.org/stable/2346101</u>
- 512 Berruti A, Borriello R, Della Beffa MT, Scariot V, Bianciotto V (2013) Application of nonspecific
- commercial AMF inocula results in poor mycorrhization in *Camellia japonica* L. Symbiosis 61:6376.

- 515 Berruti A, Lumini E, Balestrini R, Bianciotto V (2016a) Arbuscular mycorrhizal fungi as natural
- 516 biofertilizers: Let's benefit from past successes. Front Microbiol 6:1559. doi:
 517 10.3389/fmicb.2015.01559
- 518 Berruti A, Lumini E, Bianciotto V (2016b) AMF components from a microbial inoculum fail to
- colonize roots and lack soil persistence in an arable maize field. Symbiosis doi:10.1007/s13199-
- 520 016-0442-7
- 521 Bonfante P, Genre A (2010) Mechanisms underlying beneficial plant–fungus interactions in 522 mycorrhizal symbiosis. Nature Commun 1:48. doi:10.1038/ncomms1046
- 523 Bonfante P, Genre A (2015) Arbuscular mycorrhizal dialogues: do you speak 'plantish' or 524 'fungish'? Trends Plant Sci 20:150-154. doi: 10.1016/j.tplants.2014.12.002
- 525 Camargo EL, Nascimento LC, Soler M, Salazar MM, Lepikson-Neto J, Marques WL, Alves A,
- 526 Teixeira PJ, Mieczkowski P, Carazzolle MF, Martinez Y, Deckmann AC, Rodrigues JC, Grima-
- 527 Pettenati J, Pereira GA (2014) Contrasting nitrogen fertilization treatments impact xylem gene
- expression and secondary cell wall lignification in *Eucalyptus*. BMC Plant Biol 14:256. doi:
 10.1186/s12870-014-0256-9.
- 530 Carbonell-Bejerano P, Rodríguez V, Royo C, Hernáiz S, Moro-González LC, Torres-Viñals M,
- Martínez-Zapater JM (2014) Circadian oscillatory transcriptional programs in grapevine ripening
 fruits. BMC Plant Biology 14:78. doi: 10.1186/1471-2229-14-78
- Casieri L, Ait Lahmidi N, Doidy J, Fourrey C, Migeon A, Bonneau L et al (2013) Biotrophic
 transportome in mutualistic plant–fungal interactions. Mycorrhiza 23:597-625.
- Chang S, Pur Year J, Carney J (1993) A simple and efficient method for isolating RNA from pine
 trees. Plant Mol Biol Rep 11:113-116.
- 537 Chitarra W, Balestrini R, Vitali M, Pagliarani C, Perrone I, Schubert A, Lovisolo C (2014) Gene
- 538 expression in vessel-associated cells upon xylem embolism repair in *Vitis vinifera* L. petioles.
- 539 Planta 239:887-899. doi: 10.1007/s00425-013-2017-7
- 540 Conesa A, Gotz S, Garcia-Gomez JM, Terol J, Talòn M, Robles M. (2005) Blast2GO: a universal
- tool for annotation, visualization and analysis in functional genomics research. Bioinformatics
- 542 21:3674-3676. doi: 10.1093/bioinformatics/bti610
- 543 Corso M, Bonghi C (2014) Grapevine rootstock effects on abiotic stress tolerance. Plant Science
- 544 Today 1:108-113. doi.org/10.14719/pst.2014.1.3.64
- 545 Corso M, Vannozzi A, Maza E, Vitulo N, Meggio F, Pitacco A, Telatin A, D'Angelo M, Feltrin E,
- 546 Negri AS, Prinsi B, Valle G, Ramina A, Bouzayen M, Bonghi C, Lucchin M (2015) Comprehensive
- transcript profiling of two grapevine rootstock genotypes contrasting in drought susceptibility links

- the phenylpropanoid pathway to enhanced tolerance. J Exp Bot 66:5739-52. doi:
 10.1093/jxb/erv274
- 550 Dal Peraro M, van der Goot FG (2016) Pore-forming toxins: ancient, but never really out of 551 fashion. Nature Rev Microbiol 14:77-92. doi: 10.1038/nrmicro.2015.3
- 552 Dal Santo S, Tornielli GB, Zenoni S, Fasoli M, Farina L, Anesi A, Guzzo F, Delledonne M,
- 553 Pezzotti M (2013) The plasticity of the grapevine berry transcriptome. Genome Biology 14:R54.
- 554 Dal Santo S, Fasoli M, Negri S, D'Incà E, Vicenzi N, Guzzo F, Tornielli GB, Pezzotti M, Zenoni S
- (2016) Plasticity of the berry ripening program in a white grape variety. Front Plant Sci 7:970. doi:
 10.3389/fpls.2016.00970
- 557 de Andrés-de Prado R, Yuste-Rojas M, Sort X, Andrés-Lacueva C, Torres M, Lamuela-Raventós
- 558 RM (2007) Effect of soil type on wines produced from Vitis vinifera L. Cv. Grenache in
- commercial vineyards. J Agric Food Chem 55:779-786. doi: 10.1021/jf062446q
- 560 Denancé N, Szurek B, Noël (2014) Emerging functions of nodulin-Like proteins in non-nodulating
- plant species. Plant Cell Physiol 55:469-74. doi:10.1093/pcp/pct198
- 562 Drogue B, Sanguin H, Chamam A, Mozar M, Llauro C, Panaud O, Prigent-Combaret C, Picault N,
- 563 Wisniewski-Dyé F (2014) Plant root transcriptome profiling reveals a strain-dependent response
- during *Azospirillum*-rice cooperation. Front Plant Sci 5:607. doi: 10.3389/fpls.2014.00607
- 565 Du Y-P, Jiang E-S, Wang F-P, Zhang S-Z, Zhai H (2014) Gene expression profiling of rootstock
- 566 '140Ru' and Vitis vinifera L. cv. 'Crimson Seedless' grape roots infected with grape phylloxera.
- 567 Plant Growth Regulation 73:1-8.
- FAO (2013) FAOSTAT database collections. Food and Agriculture Organization of the United
 Nations. Available via FAOSTAT: http://faostat3.fao.org
- 570 Fasoli M, Dell'Anna R, Dal Santo S, Balestrini R, Sanson A, Pezzotti M, Monti F, Zenoni S (2016)
- 571 Pectins, hemicelluloses and celluloses show specific dynamics in the internal and external surfaces
- of grape berry skin during ripening. Plant Cell Physiol 57:1332-1349. doi: 10.1093/pcp/pcw080
- 573 Fiorilli V, Catoni M, Miozzi L, Novero M, Accotto GP, Lanfranco L (2009) Global and cell-type
- 574 gene expression profiles in tomato plants colonized by an arbuscular mycorrhizal fungus. New
- 575 Phytol 184:975-987. doi: 10.1111/j.1469-8137.2009.03031.x
- 576 Fiorilli V, Vallino M, Biselli C, Faccio A, Bagnaresi P, Bonfante P (2015) Host and non-host roots
- 577 in rice: cellular and molecular approaches reveal differential responses to arbuscular mycorrhizal
- 578 fungi. Front Plant Sci 6:636. doi: 10.3389/fpls.2015.00636
- 579 Flexas J, Barón M, Bota J, Ducruet J-M, Gallé A, Galmés J, Jiménez M, Pou A, Ribas-Carbó M,
- 580 Sajnani C, Tomàs M, Medrano H (2009) Photosynthesis limitations during water stress acclimation

- and recovery in the drought-adapted *Vitis* hybrid Richter-110 (*V. berlandieri×V. rupestris*). J Exp
- 582 Bot 60:2361-2377. doi: 10.1093/jxb/erp069
- Handa Y, Nishide H, Takeda N, Suzuki Y, Kawaguchi M, Saito K (2015) RNA-seq Transcriptional
- 584 Profiling of an Arbuscular Mycorrhiza Provides Insights into Regulated and Coordinated Gene
- 585 Expression in Lotus japonicus and Rhizophagus irregularis. Plant Cell Physiol 8:1490-511. doi:
- 586 10.1093/pcp/pcv071
- 587 Hogekamp C, Arndt D, Pereira PA, Becker JD, Hohnjec N, Kuster H (2011) Laser microdissection
- 588 unravels cell-type-specific transcription in arbuscular mycorrhizal roots, including CAAT-box
- transcription factor gene expression correlating with fungal contact and spread. Plant Physiol 157:2023-2043.
- Holland TC, Bowen P, Bogdanoff C, Hart MM (2014) How distinct are arbuscular mycorrhizal
 fungal communities associating with grapevines? Biol Fertil Soils 50:667-674.
- 593 Giovannetti M, Balestrini R, Volpe V, Guether M, Straub D, Costa A et al (2012) Two putative-
- aquaporin genes are differentially expressed during arbuscular mycorrhizal symbiosis in *Lotus japonicus*. BMC Plant Biol 12:186
- 596 Giovannetti M, Tolosano M, Volpe V, Kopriva S, Bonfante P (2014) Identification and functional
- 597 characterization of a sulfate transporter induced by both sulfur starvation and mycorrhiza formation
- in *Lotus japonicus*. New Phytol 204:609-619.
- Goff L, Trapnell C and Kelley D (2013) cummeRbund: Analysis, exploration, manipulation, and
 visualization of Cufflinks high-throughput sequencing data.
- Gomez SK, Javot H, Deewatthanawong P, Torres-Jerez I, Tang Y, Blancaflor EB et al (2009)
 Medicago truncatula and *Glomus intraradices* gene expression in cortical cells harboring
 arbuscules in the arbuscular mycorrhizal symbiosis. BMC Plant Biol 9:10.
- 604 Grimplet J, J Van Hemert, P Carbonell-Bejerano, J Diaz-Riquelme, J Dickerson, A Fennell, M
- 605 Pezzotti, JM Martinez-Zapater (2012) Comparative analysis of grapevine whole-genome gene
- 606 predictions, functional annotation, categorization and integration of the predicted gene sequences.
- 607 BMC Res Notes 5:213. doi: 10.1186/1756-0500-5-213
- 608 Guether M, Balestrini R, Hannah MA, Udvardi MK, Bonfante P (2009a) Genome-wide 609 reprogramming of regulatory networks, transport, cell wall and membrane biogenesis during
- 610 arbuscular mycorrhizal symbiosis in Lotus japonicus. New Phytol 182:200-212. doi:
- 611 10.1111/j.1469-8137.2008.02725.x
- 612 Guether M, Neuhäuser B, Balestrini R, Dynowski M, Ludewig U, Bonfante P (2009b) A 613 mycorrhizal-specific ammonium transporter from *Lotus japonicus* acquires nitrogen released by
- arbuscular mycorrhizal fungi. Plant Physiol 150:73-83.

- 615 Gutjahr C (2014) Phytohormone signaling in arbuscular mycorhiza development. Curr Opin Plant
- 616 Biol 20: 26-34. doi: 10.1016/j.pbi.2014.04.003
- Jaillon O, Aury J, Noel B, Policriti A, Clepet C, Casagrande A et al (2007) The grapevine genome
- sequence suggests ancestral hexaploidization in major angiosperm phyla. Nature 449: 463-467.
- 619 Koundouras S, Marinos V, Gkoulioti A, Kotseridis Y, van Leeuwen C (2006) Influence of vineyard
- 620 location and vine water status on fruit maturation of nonirrigated cv. Agiorgitiko (*Vitis vinifera* L.).
- Effects on wine phenolic and aroma components. J Agric Food Chem 54: 5077-5086.
- 622 Lovisolo C, Lavoie-Lamoureux A, Tramontini S, Ferrandino A (2016) Grapevine adaptations to
- water stress: new perspectives about soil/plant interactions. Theor Exp Plant Physiol 28: 53-66. doi:
- 624 10.1007/s40626-016-0057-7
- 625 Lumini E, Orgiazzi A, Borriello R, Bonfante P, Bianciotto V (2010) Disclosing arbuscular
- 626 mycorrhizal fungal biodiversity in soil through a land-use gradient using a pyrosequencing627 approach. Environ Microbiol 12: 2165-2179.
- 628 Manck-Götzenberger J, Requena N (2016) Arbuscular mycorrhiza symbiosis induces a major
- transcriptional reprogramming of the potato SWEET sugar transporter family. Front Plant Sci
 7:487. doi: 10.3389/fpls.2016.00487
- 631 Marè C, Aprile A, Roncaglia E, Tocci E, Corino LG, De Bellis L, Cattivelli L (2013) Rootstock and
- soil induce transcriptome modulation of phenylpropanoid pathway in grape leaves. J Plant Interact
- 633 8:334-349. doi: 10.1080/17429145.2012.754958
- Milli A, Cecconi D, Bortesi L, Persi A, Rinalducci S, Zamboni A, Zoccatelli G, Lovato A, Zolla L,
- Polverari A (2012) Proteomic analysis of the compatible interaction between *Vitis vinifera* and
- 636 *Plasmopara viticola*. J Proteomics 75:1284-302. doi: 10.1016/j.jprot.2011.11.006
- 637 Pantaleo V, Vitali M, Boccacci P, Miozzi L, Cuozzo D, Chitarra W, Mannini F, Lovisolo C,
- 638 Gambino G (2016) Novel functional microRNAs from virus-free and infected *Vitis vinifera* plants
- under water stress. Scientific Reports 6:20167. doi: 10.1038/srep20167
- 640 Perrone I, Pagliarani C, Lovisolo C, Chitarra W, Roman F, Schubert A (2012) Recovery from water
- stress affects grape leaf petiole transcriptome. Planta 235:1383-1396. doi: 10.1007/s00425-0111581-y
- Pinto C, Gomes AC (2016) *Vitis vinifera* microbiome: from basic research to technological
 development. BioControl 61:243. doi:10.1007/s10526-016-9725-4
- 645 Reid KE, Olsson N, Schlosser J, Peng F, Lund ST (2006) An optimized grapevine RNA isolation
- 646 procedure and statistical determination of reference genes for real-time RT-PCR during berry
- 647 development. BMC Plant Biol 6:27. doi:10.1186/1471-2229-6-27

- Richmond TA, Somerville CR (2000) The cellulose synthase superfamily. Plant Physiol 124:495-498.
- 650 Ruzicka D, Chamala S, Barrios-Masias FH, Martin F, Smith S, Jackson LE, Barbazuk WB,
- 651 Schachtman DP (2013) Inside Arbuscular mycorrhizal roots molecular probes to understand the
- 652 symbiosis. Plant Genome 6, No 2.
- 653 Schreiner RP, Mihara KL (2009) The diversity of arbuscular mycorrhizal fungi amplified from
- 654 grapevine roots (*Vitis vinifera* L.) in Oregon vineyards is seasonally stable and influenced by soil 655 and vine age. Mycologia 101:599-611. doi:10.3852/08-169
- 656 Shao R, Xin L, Mao J, Li L, Kang G, Yang Q (2015) Physiological, ultrastructural and proteomic
- responses in the leaf of maize seedlings to polyethylene glycol-stimulated severe water deficiency.
- 658 Int J Mol Sci 16:21606-21625. doi: 10.3390/ijms160921606
- 659 Shu B, Li W, Liu L, Wei Y, Shi S (2016) Transcriptomes of arbuscular mycorrhizal fungi and
- Litchi host interaction after tree girdling. Front Microbiol 7:408. doi: 10.3389/fmicb.2016.00408
- 661 Sorek M, Díaz-Almeyda EM, Medina M, Levy O (2015) Circadian clocks in symbiotic corals: The
- duet between *Symbiodinium* algae and their coral host. Marine Genomics 14:47-57.
- Spaepen S, Bossuyt S, Engelen K, Marchal K, Vanderleyden J (2014) Phenotypical and molecular
 responses of Arabidopsis thaliana roots as a result of inoculation with the auxin-producing
 bacterium *Azospirillum brasilense*. New Phytol 201:850-861. doi:10.1111/nph.12590
- Tombesi S, Nardini A, Frioni T, Soccolini M, Zadra C, Farinelli D, Poni S, Palliotti A (2015)
 Stomatal closure is induced by hydraulic signals and maintained by ABA in drought-stressed
 grapevine. Sci Rep 5, Article number: 12449. doi:10.1038/srep12449
- 669 Trapnell C, Hendrickson DG, Sauvageau M, Goff L, Rinn JL, Pachter L (2013) Differential
- analysis of gene regulation at transcript resolution with RNA-seq. Nature Biotech 31:46-53. doi:
 10.1038/nbt.2450
- Trouvelot A, Kough JL, Gianinazzi-Pearson V (1986) Estimation of VA mycorrhizal infection levels. Research for methods having a functional significance. In Proceedings 1st European symposium, Physiological and genetical aspects of mycorrhizae, Dijon. Centre National de la Recherche Scientifique, Dijon; Institut National de la Recherche Agronomique, Dijon; Station d'Amelioration des Plantes, Paris, France, pp 217-221
- 677 Trouvelot S, Bonneau L, Redecker D, van Tuinen D, Adrian M, Wipf D (2015) Arbuscular
- 678 mycorrhiza symbiosis in viticulture: a review. Agron Sustain Dev 35:1449-1467.
- 679 doi:10.1007/s13593-015-0329-7

- 680 Vacheron J, Desbrosses G, Bouffaud M-L, Touraine B, Moënne-Loccoz Y, Muller D, Legendre L,
- Wisniewski-Dyé F and Prigent-Combaret C (2013) Plant growth-promoting rhizobacteria and root
 system functioning. Front Plant Sci 4:356. doi: 10.3389/fpls.2013.00356
- van De Mortel JE, Schat H, Moerland PD, Van Themaat EVL, Van Der Ent S, Blankestijn H,
- 684 Ghandilyan A, Tsiatsiani S, Aarts MGM (2008) Expression differences for genes involved in lignin,
- glutathione and sulphate metabolism in response to cadmium in Arabidopsis thaliana and the
- 686 related Zn/Cd-hyperaccumulator Thlaspi caerulescens. Plant Cell Environ 31:301-324.

687 doi:10.1111/j.1365

- 688 Velasco R, Zharkikh A, Troggio M, Cartwright DA, Cestaro A, Pruss D et al. (2007) A high quality
- draft consensus sequence of the genome of a heterozygous grapevine variety. PLoS ONE 2:1-18.
- 690 Verhagen, B. W., Glazebrook, J., Zhu, T., Chang, H. S., van Loon, L. C., and Pieterse, C. M.
- 691 (2004). The transcriptome of rhizobacteria-induced systemic resistance in Arabidopsis. Mol Plant
- 692 Microbe Interact 17:895-908. doi: 10.1094/MPMI.2004.17.8.895
- 693 Vitali M, Chitarra W, Galetto L, Bosco D, Marzachi C, Gullino ML, Spanna F, Lovisolo C (2013)
- Flavescence dorée phytoplasma deregulates stomatal control of photosynthesis in *Vitis vinifera*.
 Ann Appl Biol 162:335-346. doi:10.1111/aab.12025
- 696 Vivier MA, Pretorius JS (2002) Genetically tailored grapevines for the wine industry. Trends
 697 Biotech 20:472-478. doi: 10.1016/S0167-7799(02)02058-9
- Volpe V, Giovannetti M, Sun X-G, Fiorilli V, Bonfante P (2015) The phosphate transporters LjPT4
- and MtPT4 mediate early root responses to phosphate status in non mycorrhizal roots. Plant Cell
- 700 Environ doi: 10.1111/pce.12659
- 701 Zhang Q, Blaylock LA, Harrison MJ (2010) Two Medicago truncatula half-ABC transporters are
- essential for arbuscule development in arbuscular mycorrhizal symbiosis. Plant Cell 22:1483-1497.
- 703 doi: 10.1105/tpc.110.074955
- Zenoni S, Ferrarini A, Giacomelli E, Xumerle L, Fasoli M, Malerba G, Bellin D, Pezzotti M,
- 705 Delledonne M (2010) Characterization of transcriptional complexity during berry development in
- 706 Vitis vinifera using RNA-Seq. Plant Physiol 152:1787-95. doi: 10.1104/pp.109.149716
- 707 Wang R, Okamoto M, Xing X, Crawford NM (2003) Microarray analysis of the nitrate response in
- *Arabidopsis* roots and shoots reveals over one thousand rapidly responding genes and new linkages
 to glucose, trehalose-6-P, iron and sulfate metabolism. Plant Physiol 132:556-567.
- 710 Weston DJ, Pelletier DA, Morrell-Falvey JL, Tschaplinski TJ, Jawdy SS, Lu TY, Allen SM, Melton
- 711 SJ, Martin MZ, Schadt CW, et al (2012) Pseudomonas fluorescens induces strain-dependent and
- strain-independent host plant responses in defense networks, primary metabolism, photosynthesis,
- and fitness. Mol Plant Microbe Interact 25:765-778.

Description	Total number of reads	Total alignments	Total unique aligned reads	Total aligned bases	Uniquely aligned mapping reads
CTRL 1	19,445,924	17,649,261	16,238,647	882,463,050	15,358,770
CTRL 2	32,353,647	28,847,668	26,381,472	1,442,383,400	24,908,376
FMOS 1	26,154,442	23,614,153	21,657,536	1,180,707,650	20,483,703
FMOS 2	12,134,327	11,470,488	10,513,263	573,524,400	9,949,366
MICO 1	15,730,900	15,332,942	14,077,415	766,647,100	13,344,301
MICO 2	19,006,336	18,441,962	16,874,712	922,098,100	15,954,651

8 ,

718 Table 2. List of the 10 genes most up- or down-regulated in FMOS and MICO conditions, in

719 comparison with control plants.

V.vinifera_Gene_id	Sample	UP/DOWN	Gene description
VIT_01s0026g00550	FMOS	up	nodulin MtN21 family
VIT_03s0063g00370	FMOS	up	Nitrite reductase
VIT_13s0064g01290	FMOS	up	basic helix-loop-helix (bHLH) family
VIT_13s0064g01470	FMOS	up	urophorphyrin III methylase
VIT_16s0050g02540	FMOS	up	nodulin MtN3 family
VIT_05s0062g01160	FMOS	up	pectinesterase family
VIT_06s0004g04400	FMOS	up	nitric-oxide reductase, cytochrome b-
			containing subunit I
VIT_18s0001g03910	FMOS	up	nitrate reductase 2 (NR2)
VIT_14s0068g01580	FMOS	up	basic helix-loop-helix (bHLH) family
VIT_08s0007g01910	FMOS	up	Laccase
VIT_01s0026g00550	MICO	up	nodulin MtN21 family
VIT_03s0063g00370	MICO	up	Nitrite reductase
VIT_13s0064g01290	MICO	up	basic helix-loop-helix (bHLH) family
VIT_13s0064g01470	MICO	up	urophorphyrin III methylase
VIT_18s0001g03910	MICO	up	nitrate reductase 2 (NR2)
VIT_14s0068g01580	MICO	up	basic helix-loop-helix (bHLH) family
VIT_17s0000g05620	MICO	up	integral membrane HPP family protein
VIT_04s0023g03540	MICO	up	Thaumatin SCUTL1
VIT_06s0004g04400	MICO	up	nitric-oxide reductase, cytochrome b-
			containing subunit I
VIT_10s0003g04880	MICO	up	Ferredoxin:nadp+ Oxidoreductase
	FMOG		PETH OFFICE
VII_02s0025g02650	FMOS	down	Cellulase CEL2
VII_05s0020g021/0	FMOS	down	Sugar transporter ERD6-like 16
VII_05s0020g03740	FMOS	down	lipid transfer protein
VII_12s0028g02800	FMOS	down	isoflavone methyltransferase/ Orcinol
VIT 15c0046c01600	FMOS	down	o-methylitansierase 1 00mt1 acidic endochitinase (CHIB1)
VIT_14s0060g00760	FMOS	down	galactinal synthese
$VIT_02_0028_002800$	FMOS	down	galactinol synthase
VIT_17s0053g02800	FMOS	down	alpha expansin 1 precursor
VIT_01s0011g06180	FMOS	down	blight associated protein p12 precursor
$VIT_02_{0}012_{0}0020$	FMOS	down	Expansion like D1
$VIT_02s0012g00830$	FMOS MICO	down	Callulasa CEL 2
$VIT_02_{0}023g_{0}2030$	MICO	down	gibborallin 20 ovidesa
$VIT_0280234g00010$ $VIT_07c0104c01220$	MICO	down	Auvin response factor 2
$VII_0/S0104g01230$ $VIT_10c0116c01620$	MICO	down	
$VIT_1/_{0}068 - 01610$	MICO	down	DELLA protain DGL 1 (DGA lika
v11_145000801010	WIICO	uowii	protein 1)
VIT_07s0104g00360	MICO	down	early-responsive to dehydration
VIT_13s0019g02560	MICO	down	subtilisin protease C1
VIT 00s0665g00020	MICO	down	Carrier protein, Mitochondrial

VIT_12s0055g00950	MICO	down	receptor-like kinase 902
VIT_10s0003g05390	MICO	down	FAD-linked oxidoreductase 1

721 Legen	ds
-----------	----

Fig. 1. Colonization rate in grapevine roots after treatment with MICO and FMOS. F%, Frequency of mycorrhiza in the root system; M%, Intensity of the mycorrhizal colonization in the root system; a%, Arbuscule abundance in mycorrhizal parts of root fragments; A%, Arbuscule abundance in the root system; v%, vescicle abundance in mycorrhizal parts of root fragments.
Fig. 2. Venn diagrams of differentially-expressed genes (DEGs). a) Venn diagram of all DEGs. b) Venn diagram of up-regulated DEGs. c) Venn diagram of down-regulated DEGs.

Fig. 3. Correlation between log2 relative expression values measured by RT-qPCR and RNAsequencing analyses.

Fig. 4. GO enrichment in differentially-expressed genes (DEGs). a) Biological process enriched
 GO-terms for MICO samples. b) Biological process enriched GO-terms for FMOS samples.

Fig. 5. GO enrichment in differentially-expressed genes (DEGs). Biological process enriched GOterms for DEGs in common between FMOS and MICO samples against the whole reference *V*. *vinifera* gene space.

Fig. 6. The numbers of CAZymes functional domains identified in FMOS and MICO samples. On
 the rows, PL=polysaccharide lyases, GT=glycosyltransferases, GH=glycoside hydrolases,
 CE=carbohydrate esterases, CBM=carbohydrate-binding modules and AA= auxiliary activities.

740 Supplementary materials

Fig. S1. Fresh and dry root weight and P and K determination. Data are expressed as a mean ± SD.

Fig. S2. Dendrogram showing the clustering of RNA-seq samples in two major sub-groups basedon their expression signatures.

Fig. S3. RT-qPCR validation of the relative expression for a subset of genes randomly selected from the RNAseq experiment in the FMOS versus CTRL (a) and MICO versus CTRL (b) comparison. Blue and red bars represent the relative expression (log2fold change) recorded in the RT-qPCR and RNAseq experiments, respectively. Asterisks indicate statistically significant data (p<0.05).

Table S1. List of the oligonucleotides used for RT-qPCR experiments.

751 752 753	Table S2. Colonization rate, in grapevine roots after treatment with MICO and FMOS, for each plant.
754	Table S3. Gene expression data.
755 756	Table S4 . Differential expression analysis results and functional annotation of common DEGs inFMOS and MICO samples.
757 758	Table S5 . Differential expression analysis results and functional annotation of DEGs specific ofFMOS or MICO samples.
759 760	Table S6 . Transporters related DEGs differentially expressed in MICO and FMOS (above) and specific for one of the two samples (below).
761	
762	
763	
764	
765	
766	
767	
768	
769	
770	
771	
772	
773	
774	
775	





