

Proteome adaptations under contrasting soil phosphate regimes of Rhizophagus irregularis engaged in a common mycorrhizal network

Ghislaine Recorbet, Silvia Calabrese, Thierry Balliau, Michel Zivy, Daniel Wipf, Thomas Boller, Pierre-Emmanuel Courty

▶ To cite this version:

Ghislaine Recorbet, Silvia Calabrese, Thierry Balliau, Michel Zivy, Daniel Wipf, et al.. Proteome adaptations under contrasting soil phosphate regimes of Rhizophagus irregularis engaged in a common mycorrhizal network. Fungal Genetics and Biology, 2021, 147, pp.103517. 10.1016/j.fgb.2021.103517 . hal-03313640

HAL Id: hal-03313640 https://hal.inrae.fr/hal-03313640

Submitted on 3 Feb 2023

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License

Version of Record: https://www.sciencedirect.com/science/article/pii/S1087184521000013 Manuscript_83e563f4bc4ed08e44342b932f0100e4

1	Proteome adaptations under contrasting soil phosphate regimes of Rhizophagus irregularis engaged in a
2	common mycorrhizal network
3	Ghislaine Recorbet ¹ , Silvia Calabrese ² , Thierry Balliau ³ , Michel Zivy ³ , Daniel Wipf ¹ , Thomas Boller ² , Pierre-
4	Emmanuel Courty ¹
5	
6	¹ Agrécolologie, AgroSup Dijon, CNRS, Université de Bourgogne, INRAE, Université Bourgogne Franche-
7	Comté, Dijon, France
8	² Department of Environmental Sciences, Botany, Zurich-Basel Plant Science Center, University of Basel, Basel,
9	Switzerland
10	³ PAPPSO, GQE - Le Moulon, INRAE, Univ. Paris-Sud, CNRS, AgroParisTech, Univ. Paris-Saclay, 91190 Gif-
11	sur-Yvette, France.
12	
13	Corresponding author: Ghislaine Recorbet, email:ghislaine.recorbet@inrae.fr
14	
15	
16	Highlights:
17	• The ERM of <i>R. irregularis</i> was grown between poplar and sorghum under Pi limitation or not
18	• Shotgun proteomic and qRT-PCR approaches shed light on the ERM adaptive mechanisms to Pi
19	availability
20	• Lipids are used as the main C source for fungal development in low-Pi condition
21	• Pi mobilization and N catabolism are stimulated in low-Pi condition
22	
23	Keywords: mycorrhizal symbiosis, shotgun proteomic, Rhizophagus irregularis, extra-radical mycelium,
24	common mycelial network, phosphate nutrition.
25	

26 Abstract

27 For many plants, their symbiosis with arbuscular mycorrhizal fungi plays a key role in the acquisition of 28 mineral nutrients such as inorganic phosphate (Pi), in exchange for assimilated carbon. To study gene regulation 29 and function in the symbiotic partners, we and others have used compartmented microcosms in which the extra-30 radical mycelium (ERM), responsible for mineral nutrient supply for the plants, was separated by fine nylon nets 31 from the associated host roots and could be harvested and analysed in isolation. Here, we used such a model 32 system to perform a quantitative comparative protein profiling of the ERM of *Rhizophagus irregularis* BEG75, 33 forming a common mycorrhizal network (CMN) between poplar and sorghum roots under a long-term high- or 34 low-Pi fertilization regime. Proteins were extracted from the ERM and analysed by liquid chromatography-35 tandem mass spectrometry.

36 This workflow identified a total of 1301 proteins, among which 162 displayed a differential amount 37 during Pi limitation, as monitored by spectral counting. Higher abundances were recorded for proteins involved 38 in the mobilization of external Pi, such as secreted acid phosphatase, 3',5'-bisphosphate nucleotidase, and 39 calcium-dependent phosphotriesterase. This was also the case for intracellular phospholipase and 40 lysophospholipases that are involved in the initial degradation of phospholipids from membrane lipids to 41 mobilize internal Pi. In Pi-deficient conditions. The CMN proteome was especially enriched in proteins assigned 42 to beta-oxidation, glyoxylate shunt and gluconeogenesis, indicating that storage lipids rather than carbohydrates 43 are fuelled in ERM as the carbon source to support hyphal growth and energy requirements. The contrasting 44 pattern of expression of AM-specific fatty acid biosynthetic genes between the two plants suggests that in low Pi 45 conditions, fatty acid provision to the fungal network is mediated by sorghum roots but not by poplar. Loss of 46 enzymes involved in arginine synthesis coupled to the mobilization of proteins involved in the breakdown of 47 nitrogen sources such as intercellular purines and amino acids, support the view that ammonium acquisition by 48 host plants through the mycorrhizal pathway may be reduced under low-Pi conditions. This proteomic study 49 highlights the functioning of a CMN in Pi limiting conditions, and provides new perspectives to study plant 50 nutrient acquisition as mediated by arbuscular mycorrhizal fungi.

51

52 **1. Introduction**

53 Phosphorus (P), as a component of nucleic acids, chemical energy (ATP), cell membrane phospholipids, 54 signal transduction processes, and regulation of enzyme activities, is an essential element for all living 55 organisms. Because P also serves critical roles in photosynthesis, low P availability is a major factor constraining 56 plant growth and metabolism in many soils worldwide. Thus, application of large amounts of Pi fertilizers are 57 used ensure plant productivity in most current agricultural systems (Smith et al. 2011). Crop fertilization has 58 notably tripled the rate of consumption of P, increasing crop production, while in the same time accelerating soil 59 degradation and water eutrophication (Conley et al. 2009). In addition, P fertilizers are mainly derived from 60 mined rock phosphate deposits, and these are predicted to become a limiting factor for food production within 61 the next century (van Vuuren et al. 2010).

62 An alternative to the input of Pi fertilizers is to exploit the mechanisms developed by many P-limited 63 plants to improve P use efficiency in agro-ecosystems, in particular the mutualistic symbiosis with arbuscular 64 mycorrhizal fungi (AMF) of the phylum Glomeromycota (Tedersoo et al. 2018). Approximately 80% of 65 terrestrial plant species engage in this symbiosis, including the majority of agricultural crops (Smith and Read, 66 2008). AMF benefit crop productivity because of their ecosystem services (Gianinazzi et al. 2010). One 67 prominent function of AMF consists in their contribution to plant Pi acquisition, absorbed from the soil solution 68 through hyphal scavenging of soil volumes that are not accessible by plant roots (Smith et al. 2011). It has been 69 estimated that inoculation with AMF might result in a reduction of approximately 80% of the recommended 70 fertilizer P rates under certain conditions (Jakobsen 1995). In return, AMF are supplied with host organic carbon, 71 in the form of sugar and lipid (Gutjahr and Parniske 2013; Roth and Paszkowski 2017).

72 In arbuscular mycorrhizal (AM) plants, the extra-radical mycelium (ERM) that develops outside of the 73 host root, is the component of mycorrhiza that mines rhizospheric and bulk soil to acquire scarce nutrients and 74 translocate them to the fungus-root interface where transfer to the host plant is performed (Olsson et al. 2014). 75 Building on the ERM, the AM phosphate pathway involves the uptake of soil Pi at the soil-fungus interface 76 through high-affinity fungal phosphate transporters (PTs) (Garcia et al. 2016; Wang et al. 2017). Pi accumulates 77 in the vacuoles of extraradical hyphae as polyphosphate (polyPi) (Ezawa et al. 2004). PolyPi chains are thought 78 to be transferred by a tubular vacuolar network into the intraradical hyphae and arbuscules (Uetake et al. 2002). 79 Pi is hydrolyzed from polyPi by a fungal phosphatase and exopolyphosphatase, after which the Pi is released into 80 the plant interfacial apoplast of arbuscules (Javot et al. 2007). In this manner, AM fungi give plants access to Pi 81 beyond the depletion zone that develops around the roots.

82 Despite the key role of the ERM in mineral plant nutrition, current knowledge regarding the global 83 metabolic mechanisms by which the ERM responds to Pi availability remains scarce. Using *Rhizophagus* grown 84 with Lotus japonicus under phosphorus-deficient conditions, Kikuchi et al. (2014) reported the induced 85 expression of the genes encoding Pi symporters, P-type ATPases and polyP polymerase four hours after Pi 86 application to Pi-starved hyphae. Howerver, because of the time-dependent outcome of symbiotic relationships 87 in response to nutrient availability (Olsson et al. 2006), it is likely that the metabolism of external hyphae may 88 differ between short- and long-term Pi shortage. In addition, as AM symbionts exhibit a broad host spectrum, 89 ERM can connect the roots of the same or different plant species through the formation of a common mycelial 90 network (CMN), which can transfer nutrients to several plants simultaneously (Walder et al. 2012; Felbaum et al. 91 2014; Bücking et al. 2016; Calabrese et al. 2019). The ERM provides extensive pathways for nutrient fluxes 92 through the soil and among plants, and its functioning presumably relies on the existence of a complex 93 regulation of fungal metabolism with regard to nutrient sensing, production of specific enzymes, and resource 94 partitioning between host roots and their fungal symbionts (Leake et al. 2004; Walder et al. 2012). Although 95 CMN are known to influence plant establishment/survival, physiology and defence (Gorzelak et al. 2015), 96 information is scarce regarding the characterization of the CMN metabolic adaptations to contrasting Pi 97 fertilization regimes.

98 In a previous study, when performing metabolome analysis on the ERM of the AMF Rhizophagus 99 irregularis associated to Populus trichocarpa and Sorghum bicolor, we observed that the metabolite profile of 100 the ERM was not significantly affected by Pi availability (Calabrese et al. 2019). However, not only in mammals 101 and plants, but also in fungi, studies over the years have revealed that metabolic pathways are strongly regulated 102 by post-transcriptional control of the proteome, including protein degradation, allosteric changes, and 103 posttranslational modifications, which expand the organism's capacity to respond to nutrient availability 104 (Gutterridge et al. 2010; Plaxton and Tran, 2011; Lan et al. 2012). This is especially true for Pi that acts as an 105 allosteric activator or inhibitor of many enzymes (Gregory et al. 2009). In this regard, proteomics may likely be 106 able to decipher key mycelial metabolic pathways involved in fungal adaptation to differential Pi levels by 107 providing a valuable overview of expression changes of the most abundant proteins in the cell (Alexova and 108 Millar, 2013). Of note, although mass spectrometry (MS) does not provide information about enzyme activity, 109 protein amounts coupled to functional grouping correlate with microbial activity (Wilmes and Bond, 2006; 110 Siggins et al. 2012). With the aim to enhance our understanding of the CMN fungal metabolic pathways as 111 affected by Pi availability, we performed a quantitative comparative protein profiling of the ERM grown

between poplar and sorghum roots under a long-term, high- or low-, Pi fertilization regime. To decipher the fungal proteome, proteins were extracted from the ERM and analysed by liquid chromatography-tandem mass spectrometry (LC-MS/MS). Changes in protein amount between the ERM proteomes obtained under high and low-Pi conditions were further monitored by spectral counting. The biological significance of the Pi-responsive proteins is discussed with special regard to fungal energetic resources and Pi scavenging/recycling strategies.

117

118 **2.** Materials and Methods

119 2.1. Biological material, microcosms and phosphate fertilization

120 Experiments were performed as previously described (Calabrese et al. 2019), using Populus trichocarpa 121 cuttings (clone 10174, Orléans, France) and Sorghum bicolor (L.) Moench, cv Pant-5. After sterilization in 2.5% 122 KClO for 10 min, seeds were germinated in the dark at 25 °C for three days. Microcosms were set-up in tripartite 123 compartments consisting of two root hyphal compartments (1020 ml each) attached to one central hyphal 124 compartment (1100 ml) (Supplementary Figure 1). Root hyphal and central compartments were filled with 925 125 and 1000 g of an autoclaved (120°C, 20 min) quartz sand (Alsace, Kaltenhouse, Trafor AG, Basel): zeolithe 126 (Symbion, Czech Republic) substrate (1:1, w:w), respectively. Poplar cuttings were planted into one of the 127 outside compartments and sorghum seedlings into the other. Both plants were inoculated with Rhizophagus 128 irregularis, isolate BEG75 (Inoculum Plus, Dijon, France). For inoculation, plants were supplemented with 1 ml 129 of a spore suspension (110 spores in 1 ml of 0.01 M citrate buffer, pH 6). Compartments were separated by two 130 21 µM meshes and one 3 mm mesh, to allow the AMF to grow from one compartment to the other but to avoid 131 plant roots protruding into the hyphal compartment. Plants were fertilized once a week with 10 ml Hoagland 132 solution without Pi, until all plants showed signs of phosphate depletion. From the 22nd week, either high-Pi (560 133 µM) or low-Pi (28 µM) containing Hoagland solution was applied to the middle hyphal compartment for 9 134 weeks. For each P treatment, the experimental setup was replicated three times. Plants were grown under 135 controlled condition 16 h of light [220 µE m⁻² s⁻¹] at 25 °C and 8 h of dark at 20 °C, constant relative aerial 136 humidity of 65 %.

137

138 2.2. Harvest, mycorrhizal colonization measurements, and P extraction

139 The ERM was extracted by suspending the substrate of the hyphal compartment with tap water and 140 fishing the hyphae from the surface using a 32 μ M mesh. This step was repeated several times. Afterwards, the 141 cleaned ERM samples were snap frozen in liquid nitrogen and stored at -80 °C. Roots from the plant 142 compartments were removed from substrate under tap water and cut into 1 cm pieces. Samples (about 100mg) 143 were either snap frozen and stored at -80 °C for further gene expression analysis or stored for root colonization 144 measurements. For that purpose, roots were immersed in 10% KOH and stored at 4 °C overnight. The next day, 145 the roots were rinsed and immersed in lactic-acid glycerol water (1:1:1, v:v:v) for destaining. Total colonization 146 count was performed using the magnified intersection method (McGonigle et al. 1990). Mycorrhizal parameters 147 were subjected to arcsine square root transformation before comparison of means using Student's *t*-test with a 148 value of p < 0.05 considered to be statistically significant.

To determine the total P concentration in poplar and sorghum, root and shoot dried samples were ground using a ball mill. Up to 500mg were used for the modified P extraction method by Murphy and Riley (1962).

152

153 **2.3.** Protein extraction, prefractionation, and digestion

154 For the two Pi fertilization regimes, protein extraction from frozen ERM (1 to 2g equivalent fresh 155 weight) was performed on three biological replicates using the phenol extraction protocol described by Dumas-156 Gaudot et al. (2004). Briefly, ERM was ground into liquid nitrogen and homogenised in 10 ml of 0.5 M Tris-157 HCl, pH 7.5, lysis buffer that contained 0.7 M sucrose, 50 mM EDTA, 0.1 M KCl, 10 mM thiourea, 2 mM 158 PMSF and 2% (v/v) β -mercaptoethanol. One volume of Tris-buffered phenol was added and, after mixing for 30 159 min, the phenolic phase was separated by centrifugation and rinsed with another 10 ml of lysis buffer. Proteins 160 were precipitated overnight at -20 °C after adding 5 volumes of methanol containing 0.1 M ammonium acetate. 161 The pellet, recovered by centrifugation, was rinsed with cold methanol and acetone and dried under nitrogen gas. 162 Proteins were solubilized in 200 µl of Laemmli buffer (Laemmli et al. 1970) before ultracentrifugation during 163 30 min (Beckman Airfuge, 30 psi). Protein amount in the supernatant was measured according to the Bradford 164 method (Bradford, 1976).

165

166 2.4. Sample pre-fractionation and protein digestion

167 For each biological replicate (n = 3) of the two treatments, fungal proteins (20 µg) were pre-fractionated 168 by a 0.7 cm migration on 12% SDS-PAGE. After Coomassie Brillant Blue staining, each lane was cut into seven 169 slices of equal size (1 mm), washed in distilled water and destained using 100 mM NH₄CO₃ in 50% acetonitrile. 170 A reduction step was performed by addition of 40 µl of 10mM dithiotreitol in 50 mM NH₄HCO₃ for 30 min at 56 171 °C. The proteins were alkylated by adding 30 µl of 55 mM iodoacetamide in 50 mM NH₄HCO₃ and allowed to

172 react in the dark at room temperature for 45 min. Gel sections were washed in 50 mM NH₄HCO₃, then ACN, and 173 finally dried for 30 min. In-gel digestion was subsequently performed for 7 h at 37 °C with 125 ng of modified 174 trypsin (Promega) dissolved in 20% (v/v) methanol and 20 mM NH₄CO₃. Peptides were extracted successively 175 with 0.5% (v/v) TFA and 50% (v/v) ACN and then with pure ACN. Peptide extracts were dried and suspended in 176 25 µl of 0.05% (v/v) TFA, 0.05% (v/v) HCOOH, and 2% (v/v) ACN.

- 177
- 178

2.5. Liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis

179 Peptide separation was performed using an Ultimate 3000 RSLCnano (Thermo fisher Scientific, 180 Waltham, Massachusetts, USA). Peptides were first desalted using a PepMap100 C18 trap column (5 µm, 100Å, 181 Dionex) with 2% ACN (v/v) and 0.1% HCOOH (v/v) in water for 3min at 15 µl/min. Peptides were further 182 separated on a pepmap100 C18 column (5 µm, 15cm x 75 µm, Dionex). The mobile phase consisted of a 183 gradient of solvents A: 0.1% HCOOH (v/v), 2% ACN (v/v) in water and B: 80% ACN (v/v), 0.1% HCOOH 184 (v/v) in water. Separation was set at a flow rate of 0.3 µl/min using a linear gradient of solvent B from 1 to 32% 185 in 32 min, followed by an increase to 35 % in 2 min and finally to 98% for 3 min. Eluted peptides were analysed 186 with a LTQ-Orbitrap Discovery (Thermo Electron) using a nanoelectrospray interface. Ionization (1.3 kV 187 ionization potential) was performed with a liquid junction and a non-coated capillary probe (10 µm i.d.; New 188 Objective). Peptide ions were analysed using Xcalibur 2.0.7 with the following data-dependent acquisition steps 189 (1) full MS scan (mass to charge ratio (m/z) 300-1400, centroid mode in orbitrap), (2) MS/MS (qz = 0.22, 190 activation time = 50 ms, and collision energy = 35%, centroid mode in ion trap). Step 2 was repeated for the 191 three major ions detected in step 1 with a minimal intensity of 500 relative abundance. Dynamic exclusion was 192 set to 30 s.

193

194

2.6. Protein identification and quantification

195 Spectra search was performed against the protein sequences from the R. irregularis 196 Gloin1_GeneCatalog_proteins (version 20120510, 30282 entries) downloaded from the server 197 http://genome.jgi.doe.gov/Gloin1/Gloin1 using the X!Tandem software (version 2015.04.01.1) (Langella et al. 198 2017). Enzymatic cleavage was declared as a trypsin digestion with one possible missed cleavage in first pass. 199 Cysteine carbamidomethylation were set to static, while methionine oxidation, protein n-terminal acetylation 200 with or without excision of methionine, dehydration of N-terminal glutamic acid, deamination of N-terminal 201 glutamine and N-terminal carbamidomethyl-cysteine as possible modifications. Precursor mass precision was set to 10 ppm with a fragment mass tolerance of 0.5 Da. Identified proteins were filtered and grouped using X!Tandem according to (i) the tolerated presence of at least two peptides with an *E-value* smaller than 0.01 and (ii) a protein *E-value* (calculated as the product of unique peptide *E-values*) smaller than 10^{-5} .

Quantification of proteomic data was achieved by spectral counting using normalized spectral abundance factor (NSAF) analysis (Zybailov et al. 2006). A NSAF value was calculated for each protein in the six replicates (three biological replicates x two conditions). As NSAF represent percentages, all data were arcsine square root-transformed to obtain a distribution of values that could be checked for normality using the Kolmogorov-Smirnov test. For each protein identified, significant differences (p < 0.05) between transformed NSAF values originating from low- and high-Pi data sets were analysed using the Welch-test (degrees of freedom = n-1), which is compatible with unequal variances between groups (Staher et al. 2014).

212

213 2.7. In silico analysis

214 The number of trans-membrane (TM) domains and KOG (euKaryotic Orthologous Groups) functional 215 classification were inferred from the JGI genomic resource for *R*. irregularis 216 (https://mycocosm.jgi.doe.gov/Rhiir2_1/Rhiir2_1.home.html). Prediction of protein subcellular localization was 217 WoLF PSORT (https://www.genscript.com/wolf-psort.html), performed using CELLO 218 (http://cello.life.nctu.edu.tw) and DeepLoc 1.0 (http://www.cbs.dtu.dk/services/DeepLoc/). Secreted proteins 219 were predicted according to the Fungal Secretome and Subcellular Proteome KnowledgeBase 2.1 (FunSecKB2; 220 http://proteomics.ysu.edu/secretomes/fungi2/index.php) after Blastp search. Orthologous sequence search was 221 performed with EnsemblPlants (https://plants.ensembl.org/index.html).

222

223 **2.8. RNA extraction**

Total RNA was extracted from lyophilized extra-radical mycelia (Sánchez-Rodríguez et al. 2008; Calabrese et al. 2017; Calabrese et al. 2019), and frozen plant samples using the RNeasy Plant Mini Kit (Qiagen, Courtaboeuf, France). RNA extracts were DNase treated with the DNA-freeTM Kit, DNase Treatment and Removal Reagents (AMBION® by life technologies). Total RNA was quantified with the Qbit RNA BR Assay kit and purity was estimated using the Nanodrop (ND-1000, Witec, Switzerland).

229

230 **2.9.** Reverse transcription and qRT-PCR

231 Complementary DNAs (cDNAs) from three biological replicates were obtained using the 232 iScriptTMcDNA Synthesis Kit (BIO RAD Laboratories, Paolo Alto, CA, United States), using 200 ng of total 233 RNA per reaction. For quantification a two-step quantitative RT-PCR approach was used. Gene specific primers 234 were designed with Primer-BLAST (https://www.ncbi.nlm.nih.gov/tools/primer-blast) and tested as well in 235 amplify 3.1 (http://engels.genetics.wisc.edu/amplify). Target gene expressions were normalized to the expression 236 of the reference gene translation elongation factor in *R. irregularis*. Quantitative RT-PCRs were run in a 7500 237 real-time PCR system (Roche) using the following settings: 95 °C for 3 min and then 40 cycles of 95 °C for 30 s, 238 60 °C for 1 min and 72 °C for 30 s. The number of replicates comprised three biological and three technical 239 replicates per treatment. Differences in gene expression between applied conditions were tested by either 240 Student's t-test or one-way ANOVA using SPSS Statistics, version 22 (IBM, Chicago, USA).

241

242 **3. Results and discussion**

243 **3.1.** Mycorrhizal colonization and plant P content

244 In agreement with previous data (Calabrese et al. 2019), the hyphal colonization and the percentage of 245 vesicles at harvest were not significantly different between low and high P treatments in sorghum and poplar 246 (Supplementary Figure 2). However, the relative abundance of arbuscules was significantly enhanced in 247 sorghum roots in the low-P conditions relative to the high-P treatment, thereby supporting the view that 248 mycorrhization sustains P acquisition in sorghum during phosphate limitation (Calabrese et al. 2019). 249 Consistently, the total P content in AM sorghum, which reached 350 µg on average, was similar between the two 250 fertilization regimes, while it was significantly (p < 0.05) lower (250 µg) in P-limited AM poplar. To monitor Pi 251 acquisition through the fungal side, we measured in the ERM the expression of the high affinity transporter 252 *RiPT1* that was previously found to be regulated in ERM by external P concentration (Maldonado-Mendoza et 253 al. 2001; Calabrese et al. 2019). As shown in Supplementary Figure 3, RiPT1 was up-regulated by external 254 low-P concentrations in the ERM indicating that *RiPT1* participates in soil Pi uptake in our system.

255

256 **3.2.** Comparative proteomics

Fungal proteins were phenol-extracted from three biological replicates for each condition, and analysed using LC-MS/MS after pre-fractionation by SDS-PAGE. Using a an *E-value* smaller than 0.01 per peptide and at least two peptides for correct protein assignment, a total of 1301 non-redundant proteins were identified in the six fractions analysed (two treatments x three biological replicates). These 1301 fungal proteins were sorted in Supplementary Table 1 with respect to their predicted number of *trans*-membrane (TM) domains and KOG functional classification. As anticipated from the lower abundance of proteins integral to membranes relative to soluble proteins (Vit and Petrak, 2017), proteins with a predicted TM domain (11.4%) were underrepresented in the pool of identified proteins (**Table S1**).

265 To identify the fungal proteins responsive to Pi availability, we compared the means of NSAF values (n266 = 3) between low- and high-Pi treatments for each of the 1301 proteins identified in Table S1. A total of 162 267 proteins (12.4%) showed a significant (p < 0.05) differential amount between the two conditions. As listed in 268 Supplementary Table 2, this repertoire encompassed 108 and 54 fungal proteins, the abundance of which 269 increased and decreased under Pi limitation, respectively. According to the MetaCyc database and the MIPS 270 Functional Catalogue (Ruepp et al. 2004; Caspi et al. 2014), these 162 differentially-accumulated proteins 271 belong to 19 biochemical pathways highlighted as framed bold letters in Figure 1, and ten biological processes 272 listed in Table S2, respectively. Figure 2 that displays the quantitative distribution of Pi-responsive protein 273 within MIPS functional categories further indicates that low-Pi fertilization led to an increase in processes 274 related to energetics, protein degradation, metabolism and cell growth, concomitantly to a depletion of proteins 275 involved in protein folding.

276 One of the prominent fungal metabolic responses observed in the ERM under Pi limitation was the 277 broad activation of energetic pathways, including β -oxidation of fatty acids (FAs) (13 proteins), the 278 mitochondrial electron transport chain (ETC)/ membrane-associated energy conservation mechanisms (13 279 proteins), the glyoxylate/tricarboxylic acid (TCA) cycle and glycolysis/gluconeogenesis (five proteins each) 280 (Table S2, Figure 1). Another obvious feature of low-Pi-responsive fungal proteins was the decreased 281 abundance of 13 proteins mediating protein folding such as chaperones and disulfide isomerases, the depletion of 282 the 26S ubiquitin-proteasome system (six proteins) together with the activation of peptide catabolism (six 283 peptidases) (Table S2, Figure 2). As regards amino acid biosynthesis upon low-Pi fertilization, Figure 1 shows a 284 noticeable increased amount of enzymes belonging to the methyl cycle and transsulphuration pathway that 285 contribute to cysteine synthesis from methionine, while there was a concomitant depletion of enzymes involved 286 in arginine biosynthesis (ornithine carbamoyltransferase, arginosuccinate lyase). Pi limitation also led to 287 enhanced amounts of enzymes involved in purine catabolism (five proteins) and phospholipid hydrolysis, 288 including phospholipases and phosphatidyl decarboxylase (Figure 1). Finally, a similar trend was observed for 289 metabolic pathways related to the synthesis of cell wall precursors, including chitin and galactomannan, a 290 polymer of mannose and galactose (Figure 1).

291 To further underpin this metabolic reprogramming, we searched the literature for the main AM fungal 292 pathways that have been reported to be transcriptionally regulated with special regard to the data published 293 relative to R. irregularis genes responsive to a differential Pi supply (Kikuchi et al. 2014; Vijayakumar et al. 294 2016; Sugimura and Saito, 2017; Xu et al. 2017). Using the primers listed in Supplementary Table 3, the 295 relative transcript levels in hyphae under low- and high-Pi conditions were monitored by quantitative RT-PCR. 296 Results displayed in Table S3 support higher transcript abundance in low-Pi conditions for fungal genes related 297 to β-oxidation (mitochondrial carnitine-acylcarnitine carrier protein), gluconeogenesis (fructose-1,6-298 bisphosphatase, phosphoenolpyruvate carboxykinase), glyoxylate shunt (malate synthase, isocitrate lyase), TCA 299 cycle (aconitase), phosphate sensing (phosphoserine phosphatase, SPX domain), and decreased transcription of 300 fungal genes involved in arginine biosynthesis (ornithine carbamoyltransferase OTC/ARG3), and protein folding 301 (chaperone HSP104).

302

303 **3.3.** Hyphal development and mobilization of Pi pools are stimulated by Pi limitation

304 Under Pi limitation, AM fungal colonization is expected to assist the plant mineral element uptake 305 through the development of extra-radical hyphae that access poorly mobile soil nutrients beyond the depletion 306 zone formed around absorbing roots (Olsson et al. 2002; Olsson et al. 2006). In this study, we identified several 307 low-Pi-responsive proteins that are required for the delivery at of secretory vesicles involved in plasma 308 membrane increase at the growing apex (Wessels, 1993; Levin, 2005; Steinberg, 2007). Data in Table S2 show 309 that Pi limitation enhanced the abundance of Cdc42 and Sec4, which are involved in the formation of lateral 310 hyphal branches, and membrane traffic coupled to polarized vesicle exocytosis, respectively (Virag et al. 2007; 311 Donovan and Bretscher, 2015). This was also the case for enzymes involved in fungal cell wall construction, 312 including the biosynthesis of chitin and galactomannan (Figure 1). In addition to a greater effective surface area 313 for absorbing nutrients, another microbial strategy to improve the acquisition of external P consists of the 314 secretion of proteins that increase the release of Pi from the soil to solution (Richardson and Simpson, 2011; Rai 315 et al. 2013). When searching for secreted proteins as predicted by their extracellular localization according to 316 WoLF PSORT, CELLO, or DeepLoc together with exploring the fungal secretome knowledgebase FunSecKB2, 317 we identified in Table S2 enhanced amounts under Pi limitation for a tartrate-resistant acid phosphatase type 5, a 318 3',5'-bisphosphate nucleotidase, and a calcium-dependent phosphotriesterase. These enzymes hydrolyse 319 phosphate esters to mobilize Pi from the soil, the latter being made available for fungal uptake by high-affinity Pi 320 transporters (Hinsinger, 2001; Plaxton and Tran, 2011). Consistently, the expression in the fungal ERM of the high affinity Pi:H⁺ transporter *RiPT1* was enhanced under low-Pi supply relative to high-Pi conditions (**Figure S3**), concomitantly to the induction of the vacuolar transporter chaperone VTC4 (**Table S2**). VTC couples synthesis of polyP to its translocation across the tonoplast, thereby avoiding its toxic accumulation in the cytosol (Gerasimaite et al. 2014). Of note, both VTC4 and phosphoserine phosphatase, which were induced in the fungal proteome in low-Pi conditions (**Table S2**), belong to the ten proteins identified in *R. irregularis* that contain a SPX domain (Ezawa and Saito, 2018), which acts as a sensor for inositol pyrophosphate signalling molecules (Wild et al. 2016).

328 The low-Pi-responsive proteins identified in **Table S2** pointed to lipid remodelling events in the fungal 329 ERM. Among them was the increased abundance of lysophospholipases that contribute to the initial degradation 330 of phospholipids from membrane lipids. A similar trend was observed for a member of phospholipase D family, 331 which has been implicated in glycerophospholipid catabolism during Pi starvation (Li et al. 2006; Jeong et al. 332 2017). As previously observed in Zea mays roots facing P starvation (Calderon-Vazquez et al. 2008), the protein 333 phosphatidylinositol transfer protein SEC14 shows a decreased abundance in low-Pi conditions (Table S2). In 334 yeast, inactivation of SEC14 increases the turnover of the phospholipid phosphatidylcholine (Patton-Vogt et al. 335 1997). Overall, these results suggest a breakdown of fungal membrane phospholipids upon Pi limitation, which 336 allows internal Pi mobilization, and makes the lipid moiety diacylglycerol available for non-phosphorus-lipid 337 biosynthesis (Figure 1). In this context, it has been shown that R. irregularis ERM contains glycosylated sterols 338 and glycosylated sphingolipids. As these glycolipids are phosphate-free, they can replace phospholipids in the 339 membranes during P deprivation (Wewer et al. 2014). In this line of reasoning, gene expression profiling of R. 340 irregularis in roots of Lotus japonicus has also highlighted the repression of fungal glycerophospholipid 341 metabolism when plants were shifted to high-Pi conditions (Vijayakumar et al. 2016).

342 Collectively, these results indicate that in low-Pi conditions not only external Pi is taken up by an 343 extending ERM, but Pi may be also mobilized from internal pools through phospholipid catabolism

344

345 3.4. Lipids are used as the main C source for fungal development under Pi limitation

AMF are obligate biotrophs that depend on the supply of carbon (C) from the host in the form of sugar and lipid (Luginbuehl et al. 2017; McLean et al. 2017; Choi et al. 2018). Extra-radical hyphae are unable to synthesize storage lipids, which are transported from the intra-radical mycelium to the ERM (Pfeffer et al. 1999). Currently, it is believed that the transfer of triacylglycerols (TAG) and glycogen to the ERM would provide the energy required both for active P uptake processes from the soil and the synthesis of new C skeletons needed to extend the ERM in search of new P resource (Bago et al. 2003; Rich et al. 2017; Roth and Paszkowski 2017). As
anticipated by Bago and collaborators (2000) according to ¹³C labelling experiments and low glycolytic enzyme
activities in the ERM of AMF, there was a large body of evidence at the proteome level that storage lipids rather
than sugars are fuelled in the ERM as the C source to support hyphal extension under low-Pi conditions.

355 This conclusion was first drawn from the activation under Pi limitation of the β -oxidation of FAs, as 356 inferred from the increased amount of proteins related to FA activation (long chain acyl-CoA synthetase and 357 ligase, acyl-CoA synthetase), carnitine shuttle (mitochondrial carnitine-acylcarnitine carrier protein), and 358 Lynen's helix (acyl-CoA dehydrogenase, enoyl-CoA hydratase, hydroxyacyl-CoA dehydrogenase, ketoacyl-CoA 359 thiolase) (**Table S2**). Second, after β -oxidation, the final product acetyl-CoA can enter the glyoxylate cycle that, 360 convert lipids into carbohydrates through the enzymes isocitrate lyase and malate synthase in fungal 361 glyoxysomes (Lammers et al. 2001). Consistently, the latter enzyme displayed an enhanced abundance in the 362 fungal ERM upon Pi limitation (Table S2). In support of the recruitment of the glyoxylate cycle, there was a 363 depleted amount of isocitrate dehydrogenase (Table S2), the inactivation of which is believed to force the C flux 364 toward the glyoxylate shunt (Holms, 1996). Third, after the glyoxylate cycle, its net product, succinate, can be 365 imported into mitochondria by dicarboxylate carriers before being oxidized to fumarate by mitochondrial 366 succinate dehydrogenase that couples the TCA cycle with the ETC (Kunze et al. 2006). The abundance of these 367 proteins was notably higher in low-Pi conditions (Table S2). Fourth, the glyoxylate cycle enables cells to utilise 368 FAs to fuel gluconeogenesis, as mirrored in the current study by the increased amount upon Pi limitation of 369 enzymes specific for gluconeogenesis, including phosphoenolpyruvate carboxykinase and fructose-1,6-370 bisphosphatase (Table S2). In agreement with the higher abundance of gluconeogenic enzymes, there was in 371 low-Pi conditions a depleted amount of fructose-2,6-bisphosphatase that generates fructose-2,6-bisphosphate, an 372 allosteric inhibitor of fructose-1,6-bisphosphatase (Benkovic and de Maine, 1982; Figure 1). Finally, parallel to 373 gluconeogenesis, there was an increased amount in low-Pi conditions of the trehalose-6-phosphate synthase 374 complex that is connected to glycogen biosynthesis through Glc6P, a potent activator of glycogen synthase 375 (François and Parrou, 2001). Within this line, two glycogenesis enzymes, namely phosphoglucomutase and 376 UDP-glucose pyrophosphorylase, displayed a higher abundance under Pi limitation (Figure 1). In support of an 377 enhanced glycogen biosynthesis in response to low-Pi conditions, data listed in Table S3 include a decreased 378 transcription of the fungal gene encoding glycogen phosphorylase, which catalyses the hydrolysis of glycogen.

379 One rationale behind the oxidation of FAs as the C source to support hyphal growth in Pi-limited 380 condition is that TAG provide energy twice as much as carbohydrates (Berg et al. 2002). In addition, unlike 381 glycogen catabolism via glycogenolysis and glycolysis, TAG metabolism through beta-oxidation and 382 gluconeogenesis is coupled to the release of Pi (Figure 1). This metabolic route implies that energy provision to 383 the fungal ERM during Pi limitation would depend on TAG import from the intra-radical mycelium and plant 384 monoacylglycerol provision to the fungus (Luginbuehl et al. 2017). To test this scenario in our experimental 385 conditions, we compared between Pi-replete and Pi-starved conditions expression changes in the plant genes 386 KASI (β-keto-acyl ACP synthase I), FatM (ACP-thioesterase), RAM2 (glycerol-3-phosphate acyl transferase 387 REDUCED ARBUSCULAR MYCORRHIZA2), and STR2 (half-size ABC transporter STUNTED 388 ARBUSCULE 2), which form an AM-specific operational unit for lipid biosynthesis and transport in 389 arbuscocytes (Keymer et al. 2017). According to the list of AM-specific lipid biosynthetic/transport genes 390 identified in Medicago truncatula (Bravo et al. 2016), we used EnsemblPlants to download the corresponding 391 poplar and sorghum orthologs, a step that retrieved 14 and 15 sequences, respectively (Table S4). After profiling 392 their expression in the roots of poplar and sorghum in the two Pi conditions, we observed a contrasting pattern of 393 transcript abundance between the two plants in response to Pi deficiency (Table S4). While the mRNA level of 394 genes coding KASI, FATM, and STR2 obviously increased in sorghum roots upon low Pi supply, the transcript 395 abundance of genes coding KASI, FATM and RAM2 conversely decreased in poplar roots in the same 396 condition. This pattern of gene expression strongly suggests that in low Pi conditions, FA provision to the ERM 397 is mediated by sorghum roots but not by poplar. Noteworthy, these results are reminiscent of the asymmetry in 398 the terms of trade observed by Walder et al. (2012) between flax and sorghum connected by a CMN, where 399 contrary to flax, sorghum invested massive amounts of C.

400

401 **3.5.** N catabolism is stimulated by Pi limitation

402 The current model of N assimilation in AM symbiosis includes the synthesis of arginine in ERM, its 403 transfer to the IRM, where it is broken down to release N for transfer to the host plant (Cruz et al. 2007; Tian et 404 al. 2010; Felbaum et al. 2012). In accordance with the mechanisms of mutual exclusion of anabolism and 405 catabolism of arginine, data in Table S2 indicate a decreased amount of arginosuccinate lyase and ornithine 406 carbamoyltransferase that are involved in arginine biosynthesis. This occured concomitantly to an increased 407 abundance for the enzymes urease and ornithine aminotransferase (OAT), which use the products of arginine 408 catabolism (Govindarajulu et al. 2005; Funck et al. 2008). Given that ammonium in the ERM inhibits the activity 409 of arginase and urease (Cruz et al. 2007; Bücking and Kafle, 2015), these results not only indicate that arginine 410 is catabolized in the ERM, but also that ammonium acquisition by host plants through the mycorrhizal pathway

411 may be reduced in low-Pi conditions. To investigate this possibility, we monitored in low- and high-Pi 412 conditions using qRT-PCR the transcriptional regulation of the ammonium transporters PtAMT3.1 and 413 SbAMT3.1 known to display an AM-specific induction in poplar and sorghum roots, respectively (Koegel et al. 414 2013; Calabrese et al. 2019). Results in Supplementary Figure 4 show that the induction of *PtAMT3.1* and 415 SbAMT3.1 was only observed in poplar and sorghum AM roots, respectively. In addition, the lower expression 416 of these transporters in AM roots under low-Pi conditions pointed to a decreased ammonium transfer through the 417 AM pathway in both host plants when facing a long-term Pi shortage. These findings agree with the study of 418 Johnson (2010), who reported that AM symbiosis provides the nutrient (P or N) that is most limiting for the host. 419 They are also reminiscent of what observed during *R. irregularis* spore germination in response to root exudates, 420 during which the recruitment of OAT has been proposed to reflect the mobilization of internal nitrogen stores to 421 meet a higher demand for ammonium due to increased fungal growth (Gachomo et al. 2009).

422 There were additional clues within the low-Pi responsive proteins for the mobilization of intracellular N 423 catabolic enzymes that are required for the utilization of secondary N sources when preferred nitrogen sources 424 such as ammonium, are limiting in filamentous fungi (Marzluf 1993). As regards the utilization of purines, five 425 enzymes involved in purine catabolism, namely AICAR transformylase, 5' nucleotidase, purine nucleoside 426 phosphorylase, urate oxidase and urease, displayed an increased abundance in low-Pi conditions (Table S2, 427 Figure 1). Of note, similarly to OAT, purine catabolic enzymes require nitrogen derepression. The fungal 428 proteins enriched in low-Pi conditions also included intra-cellular components of the proteasome system, the 429 centre for protein degradation, while there was a depletion of several chaperones involved in protein folding 430 (Table 1). This indicates that in low-Pi conditions, proteins enter the degradation machinery rather than the 431 chaperone/repair pathway. We also observed an increase in 20S proteasome levels, in concert with an reduction 432 in 26S proteasome subunits (Table S2). As the 26S proteasome degrades proteins in an ATP-dependent reaction, 433 while the 20S proteasome does not so, it has been proposed that increased ration of 20S to 26S proteasomes 434 would allow preservation of protein degradation capacity, while saving energy during nutrient deprivation 435 (Bajorek et al. 2003). The mobilization of internal nitrogen catabolic enzymes that are required for the utilization 436 of secondary nitrogen sources overall supports a higher demand for ammonium due to increased fungal growth. 437 Because finely branched hyphae degenerate within a few days (Bago et al. 1998; de Vries et al. 2009), it can be 438 proposed that during long-term Pi deficiency, cell lysis may be a major reservoir for nitrogen in the fungal CMN. 439 AMF can also release peptidases and proteases into the soil that enable the cleaving of organically bound 440 nitrogen (Bonfante and Genre, 2010; Behie and Bidochka, 2014; Bücking and Kafle, 2015). Data in Table S2 highlight two fungal proteins accumulating in low-Pi conditions, a subtilisin-related protease and a peptidase
 M28, which may fulfil this function according to their predicted extracellular localization.

Taken together, arginine and purine catabolism together with extra- and intra-cellular proteolysis indicate that during long-term Pi deficiency, the fungal ERM uses available nitrogen sources from breakdown of external and internal resources for its own metabolism.

446

447 Conclusions

448 As summarized in Figure 3, shotgun proteomic and qRT-PCR approaches add novel aspects to the Pi 449 deficiency responses of an AMF engaged in a common mycorrhizal network. Among them is the mobilization of 450 nitrogen catabolic enzymes that are required for the utilization of secondary nitrogen sources. As regards 451 phosphate, they include the identification of secreted 3',5'-bisphosphate nucleotidase and phosphotriesterase that 452 can hydrolyse phosphate esters to mobilize P from the soil. Likewise, the low-Pi-responsive fungal proteome 453 also indicates that Pi may be mobilized from internal Pi pools through fatty acid breakdown coupled to 454 gluconeogenesis, and phospholipid catabolism. The contrasting pattern of expression of AM-specific fatty acid 455 biosynthetic genes between the two plants suggests that in low Pi conditions, fatty acid provision to the fungal 456 network is mediated by sorghum roots but not by poplar. A relevant future challenge will be to assess the 457 contribution of these pathways to plant Pi acquisition.

458

459 Acknowledgments

This project was supported by the Swiss National Science Foundation (Grant n° PZ00P3_136651 to P-EC and n°
127563 to TB). The authors also acknowledge the financial support from the Burgundy Franche-Comté Regional
Council and the division of Plant Health and Environment of the French National Institute for Agricultural, Food
and Environment (INRAE).

464

465 Authors' contributions

P-E. C., S.C., and G.R. carried out the experiments. T.B. and M.Z. performed the mass spectrometry analysis of
the samples. G.R. analysed the results and wrote the manuscript with support from all authors. P-E. C., T.B., and
D.W. supervised the project.

469

470 Declaration of Competing Interest

- 471 The authors declare no conflicts of interest.
- 472

473 **References**

- 474 Alexova, R., Millar, A.H., 2013. Proteomics of phosphate use and deprivation in plants. Proteomics 13, 609-623.
- 475 Bago, B., Azcon-Aguilar, C., Goulet, A., Piché, Y., 1998. Branched absorbing structures (BAS): a feature of the
- 476 extraradical mycelium of symbiotic arbuscular mycorrhizal fungi. New Phytol. 139, 375-388.
- 477 Bago, B., Pfeffer, P.E., Abubaker, J., Jun, J., Allen, J.W., Brouillette, J., Douds, D.D., Lammers, P.J., Shachar-
- 478 Hill, Y., 2003. Carbon export from arbuscular mycorrhizal roots involves the translocation of carbohydrate as
- 479 well as lipid. Plant Physiol. 131, 1496-507.
- Bago, B., Pfeffer, P.E., Shachar-Hill, Y., 2000. Carbon metabolism and transport in arbuscular mycorrhizas.
 Plant Physiol. 124, 948-958.
- Bajorek, M., Finley, D., Glikman, M.H., 2003. Proteasome disassembly and downregulation is correlated with
 viability during stationary phase. Curr. Biol. 13, 1140-1144.
- 484 Behie, S.W., Bidochka, M.J., 2014. Nutrient transfer in plant-fungal symbioses. Trends Plant Sci. 19, 734-740.
- Benedetto, A., Magurno, F., Bonfante, P., Lanfranco, L., 2005. Expression profiles of a phosphate transporter
 gene (*GmosPT*) from the endomycorrhizal fungus *Glomus mosseae*. Mycorrhiza 15, 620-627.
- 487 Benkovic, S.J., de Maine, M.M., 1982. Mechanism of action of fructose 1,6-bisphosphatase. Adv. Enzymol.
 488 Related Areas Mol. Biol. 53, 45-82.
- Berg, J.M., Tymoczko, J.L., Stryer, L., 2002. Triacylglycerols are highly concentrated energy stores, in:
 Freeman, W.H. (Ed.), Biochemistry. 5th edition. New York, Section 22.1.
 https://www.ncbi.nlm.nih.gov/books/NBK22369/.
- Bonfante, P, Genre, A., 2010. Mechanisms underlying beneficial plant-fungus interactions in mycorrhizal
 symbiosis. Nat. Comun. 1, 48.
- Bradford, M.M., 1976. A rapid and sensitive method for the quantification of microgram quantities of protein
 utilizing the principle of protein-dye binding. Anal. Biochem. 72, 248-254.
- Bravo A., York, T., Pumplin, N., Mueller, L.A., Harrison, M.J., 2016. Genes conserved for arbuscular
 mycorrhizal symbiosis identified through phylogenomics.Nat. Plant. 2, 15208.
- 498 Bücking, H., Kafle, A., 2015. Role of arbuscular mycorrhizal fungi in the nitrogen uptake of plants: current
- 499 knowledge and research gaps. Agronomy 5, 587-612.

- Bücking, H., Mensah, J.A., Fellbaum, C.R., 2016. Common mycorrhizal networks and their effect on the
 bargaining power of the fungal partner in the arbuscular mycorrhizal symbiosis. Commun. Integr. Biol. 9,
 e1107684.
- 503 Calabrese, S., Cuisant, L., Sarazin, A., Niehl, A., Erban, A., Brulé, D., Recorbet, G., Wipf, D., Roux, C., Kopla,
- 504 J., Boller, T, Courty, P.E., 2019. Imbalanced regulation of fungal nutrient transports according to phosphate
- 505 availability in a symbiocosm formed by poplar, sorghum, and *Rhizophagus irregularis*. Front. Plant Sci. 10,
- 506 1617.
- 507 Calabrese, S., Kohler, A., Niehl, A., Veneault-Fourrey, C., Boller, T., Courty, P. E. ,2017. Transcriptome
 508 analysis of the *Populus trichocarpa-Rhizophagus irregularis* mycorrhizal symbiosis: Regulation of plant and
 509 fungal transportomes under nitrogen starvation. Plant Cell Physiol. 58, 1003-1017.
- 510 Calderon-Vazquez, C., Ibarra-Laclette, E., Caballero-Perez, J., Herrera-Estrella, L., 2008. Transcript profiling of
- *Zea mays* roots reveals gene responses to phosphate deficiency at the plant- and species-specific levels. J.
 Exp. Bot. 59, 2479-2497.
- Caspi, R., Altman T., Billington, R., Dreher, K., Foerster, H., Fulcher, C.A., Holland, T.A., Keseler, I.M.,
 Kothari, A., Kubo, A., Krummenacker, M., Latendresse, M., Mueller, L.A., Ong, Q., Paley, S., Subhraveti,
- P., Weaver, D.S., Weerasinghe, D., Zhang, P., Karp, P.D., 2014. The MetaCyc database of metabolic
 pathways and enzymes and the BioCyc collection of pathway/genome databases. Nucleic Acids Res. 42,
 D459–D471.
- 518 Choi, J., Summers, W., Paszkoski, U., 2018. Mechanisms underlying establishment of arbuscular mycorrhizal
 519 symbioses. Annu. Rev. Phytopathol. 56, 135-60.
- 520 Conley, D.J., Pearl, H.W., Howarth, R.W., Boesch, D.F., Sitzinger, S.P., Havens, K.E., Lancelot, C., Linkens,
- 521 G.E., 2009. Controlling eutrophication: nitrogen and phosphorus. Science 323, 1014-1015.
- 522 Cruz, C., Egsgaard, H., Trujillo, C., Ambus, P., Requena, N., Martins-Loução, M.A., Jakobsen, I., 2007.
 523 Enzymatic evidence for the key role of arginine in nitrogen translocation by arbuscular mycorrhizal fungi.
- 524 Plant Physiol. 144, 782-792.
- de Vries, F.T., Bååth, E., Kuyper, T.W., Bloem, J., 2009. High turnover of fungal hyphae in incubation
 experiments. FEMS Microbiol. Ecol. 67, 389–396.
- 527 Donovan, K.W., Bretscher, A., 2015. Tracking individual secretory vesicles during exocytosis reveals an ordered
 528 and regulated process. J. Cell Biol. 210, 181-189.

- 529 Dumas-Gaudot, E., Valot, B., Bestel-Corre, G., Recorbet, G., St-Arnaud, M., Dieu, M., Raes, M., Saravanan,
- 530 R.S., Gianinazzi, S., 2004. Proteomics as a way to identify extra-radicular fungal proteins from *Glomus*
- 531 *intraradices*-RiT-DNA carrot root mycorrhizas. FEMS Microbiol. Ecol. 48, 401-411.
- Ezawa, T., Cavagnaro, T.R., Smith, S.E., Smith, F.A., Ohtomo, R., 2004. Rapid accumulation of polyphosphate
 in extra-radical hyphae of an arbuscular mycorrhizal fungus as revealed by histochemistry and a
- 534 polyphosphate kinase/luciferase system. New Phytol. 161, 387–392.
- 535 Ezawa, T., Saito, K., 2018. How do arbuscular mycorrhizal fungi handle phosphate? New insight into fine536 tuning of phosphate metabolism. New Phytol. 220, 1116-1121.
- 537 Fellbaum, C.R., Gachomo, E.W., Beesetty, Y., Choudary, S., Strahan, G.D., Pfeffer, P.E., Kiers, E.T., Bücking,
- H., 2012. Carbon availability triggers fungal nitrogen uptake and transport in arbuscular mycorrhizal
 symbiosis. P. Natl. Acad. Sci. USA. 109, 2666-2671.
- 540 Fellbaum, C.R., Mensah, J.A., Cloos, A.J., Strahan, G.E., Pfeffer, P.E., Kiers, E.T., Bücking, H., 2014. Fungal
- nutrient allocation in common mycorrhizal networks is regulated by the carbon source strength of individualhost plants. New Phytol. 203, 646–656.
- François, J., Parrou, J.L., 2001. Reserve carbohydrates metabolism in the yeast *Saccharomyces cerevisiae*.
 FEMS Microbiol. Rev. 25, 125-45.
- 545 Funck, D., Stadelhofer, B., Koch, W., 2008. Ornithine-δ-aminotransferase is essential for arginine catabolism but
 546 not for proline biosynthesis. BMC Plant Biology 8, 40.
- 547 Gachomo, E., Allen, J.W., Pfeffer, P.E., Govindarajulu, M., Douds, D.D., Jin, H., Nagahashi, G., Lammers, P.J.,
 548 Shachar-Hill Y., Bücking, H., 2009. Germinating spores of *Glomus intraradices* can use internal and
 549 exogenous nitrogen sources for de novo biosynthesis of amino acids. New Phytol. 184, 399-411.
- Garcia, K., Doidy, J., Zimmermann, S., Wipf, D., Courty, P.-E., 2016. Take a trip through the plant and fungal
 transportome of mycorrhiza. Trends Plant Sci. 21, 937-950.
- 552 Gerasimaite, R., Sharma, S., Desfougères, Y., Schmidt, A., Mayer, A., 2014. Coupled synthesis and 553 translocation restrains polyphosphate to acidocalcisome-like vacuoles and prevents its toxicity. J. Cell Sci.
- 554 127, 5093-5104.
- Gianinazzi, S., Gollotte, A., Binet, M.N., van Tuinen, D., Redecker, D., Wipf, D. 2010. Agroecology: the key
 role of arbuscular mycorrhizas in ecosystem services. Mycorrhiza 20, 519–530.
- 557 Gorzelak, M.A., Asay, A.K., Pickles, B.J., Simard, S.W., 2015. Inter-plant communication through mycorrhizal
- networks mediates complex adaptive behaviour in plant communities. AoB Plants 7, plv050.

- Govindarajulu, M., Pfeffer, P.E., Abubaker, J., Douds, D.D., Allen, J.W., Bücking, H., Lammers, P.J., Shachar
 Hill, Y., 2005. Nitrogen transfer in the arbuscular mycorrhizal symbiosis. Nature 435, 819-823.
- Gregory, A.L., Hurley, B.A., Tran, H.T., Valentine, A.J., She, Y.M., Knowles, V.L., Plaxton, W.C., 2009. *In vivo* regulatory phosphorylation of the phosphoenolpyruvate carboxylase AtPPC1 in phosphate-starved
 Arabidopsis thaliana. Biochem. J. 420, 57–65.
- 564 Gutjahr, C., Parniske, M., 2013. Cell and developmental biology of arbuscular mycorrhiza symbiosis. Ann. Rev.
 565 Cell Dev. Biol. 29, 593–617.
- 566 Gutteridge, A., Pir, P., Castrillo, J.I., Charles, P.D., Lilley, K.S., Oliver, S.G., 2010. Nutrient control of 567 eukaryote cell growth: a systems biology study in yeast. BMC Biol. 8, 68.
- Hinsinger, P., 2001. Bioavailability of soil inorganic P in the rhizosphere as affected by root-induced chemical
 changes: a review. Plant Soil 237, 173-195.
- Holms, H., 1996. Flux analysis and control of the central metabolic pathways in *Escherichia coli*. FEMS
 Microbiol Rev. 19, 85-116.
- Jakobsen, I., 1995. Transport of phosphorus and carbon in VA mycorrhizas, in Varma, A., Hock, B. (Eds.),
 Mycorrhiza. Springer, Berlin, pp. 297-324.
- Javot, H., Pumplin, N., Harrison, M.J., 2007. Phosphate in the arbuscular mycorrhizal symbiosis: transport
 properties and regulatory roles. Plant Cell Environ. 30, 310-322.
- 576 Jeong, K., Baten, A., Waters, D.L., Pantoja, O., Julia, C.C., Wissuwa, M., Heuer, S., Kretzschmar, T., Rose, T.J.,
- 577 2017. Phosphorus remobilization from rice flag leaves during grain filling: an RNA-seq study. Plant
 578 Biotechnol. J. 15, 15-26.
- Johnson, N.C., 2010. Resource stoichiometry elucidates the structure and function of arbuscular mycorrhizas
 across scales. New Phytol. 185, 631–647.
- Keymer, A., Pimprikar, P., Wewer, V., Huber, C., Brands, M., Bucerius, S.L., Delaux, P.M., Klingl, V.,
 Röpenack-Lahaye, E.V., Wang, T.L., Eisenreich, W., Dörmann, P., Parniske, M., Gutjahr, C., 2017. Lipid
- transfer from plants to arbuscular mycorrhiza fungi. Elife. 6, e29107.
- 584 Kikuchi, Y., Hijikata, N., Yokoyama, K., Ohtomo, R., Handa, Y., Kawaguchi, M., Saito, K., Ezawa, T., 2014.
 585 Polyphosphate accumulation is driven by transcriptome alterations that lead to near-synchronous and near-
- equivalent uptake of inorganic cations in an arbuscular mycorrhizal fungus. New Phytol. 204, 638-649.
- 587 Koegel, S., Boller, T., Lehmann, M.F., Wiemken, A., Courty, P.E., 2013. Rapid nitrogen transfer in the Sorghum
- 588 *bicolor-Glomus mosseae* arbuscular mycorrhizal symbiosis. Plant Signal. Behav. 8, 8.

- 589 Kunze, M., Pracharoenwattana, I., Smith, S.M., Hartig, A., 2006. A central role for the peroxisomal membrane
 590 in glyoxylate cycle function. Biochim. Biophys. Acta 1763, 1441-1452.
- Laemmli, U.K., Amos, L.A., Klug, A., 1970. Correlation between structural transformation and cleavage of the
 major head protein of T4 bacteriophage. Cell 7, 191-203.
- 593 Lammers, P., Jun, J., Abubaker, J., Arreola, R., Gopalan, A., Bago, B., Hernández-Sebastiá, C., Allen, J.W.,
- 594 Douds, D.D., Pfeffer, P.E., Shachar-Hill, Y., 2001. The glyoxylate cycle in an arbuscular mycorrhizal fungus.
 595 Carbon flux and gene expression. Plant Physiol. 127, 1287-1298.
- Lan, P., Li, W., Schmidt, W., 2012. Complementary proteome and transcriptome profiling in phosphate-deficient
 Arabidopsis roots reveals multiple levels of gene regulation. Mol. Cell Proteomics 11, 1156-1166.
- 598 Langella, O., Valot, B., Balliau, T., Blein-Nicolas, M., Bonhomme, L., Zivy, M., 2017. X!TandemPipeline: A
- tool to manage sequence redundancy for protein inference and phosphosite identification. J. Proteome Res. 2,494-503.
- Leake, J., Johnson, D., Donnelly, D., Muckle, G., Boddy, L., Read, D., 2004. Networks of power and influence:
 the role of mycorrhizal mycelium in controlling plant communities and agroecosystem functioning. Can. J.
 Bot. 82, 1016-1045.
- Levin, DE., 2005. Cell wall integrity signaling in *Saccharomyces cerevisiae*. Microbiol. Mol. Biol. Rev. 69, 262291.
- Li, M., Welti, R., Wang, X., 2006. Quantitative profiling of arabidopsis polar glycerolipids in response to
 phosphorus starvation. Roles of phospholipases Dζ1 and Dζ2 in phosphatidylcholine hydrolysis and
 digalactosyldiacylglycerol accumulation in phosphorus-starved plants. Plant Physiol. 142, 750-761.
- 609 Luginbuehl, L.H., Menard, G.N., Kurup, S., van Erp, H., Radhakrishnan, G.V., Breakspear, A., Oldroyd, G.E.,
- Eastmond, P.J., 2017. Fatty acids in arbuscular mycorrhizal fungi are synthesized by the host plant. Science.
 356, 1175-1178.
- 612 Maldonado-Mendoza, I.E., Dewbre, G.R., Harrison, M.J., 2001. A phosphate transporter gene from the extra-
- radical mycelium of an arbuscular mycorrhizal fungus *Glomus intraradices* is regulated in response to
 phosphate in the environment. Mol. Plant Microbe Interact. 14, 1140–1148.
- McLean, A.M., Bravo, A., Harrison, M.J., 2017. Plant signaling and metabolic pathways enabling arbuscular
 mycorrhizal symbiosis. Plant Cell 29, 2319-2335.

- 617 McGonigle T. P., Miller M. H., Evans D. G., Fairchild G. L., Swan J. A. (1990). A new method which gives an
- 618 objective measure of colonization of roots by vesicular—arbuscular mycorrhizal fungi. New Phytol. 115,
- 619 495-501.
- Marzluf, G.A., 1993. Regulation of sulfur and nitrogen metabolism in filamentous fungi. Ann Rev. Microbiol.
 47, 31-55.
- Murphy, J., Riley, J.P., 1962. A modified single solution method for the determination of phosphate in natural
 waters. Anal. Chim. Acta 27, 31-36.
- 624 Olsson, O., Olsson, P.A., Hammer, E.C., 2014. Phosphorus and carbon availability regulate structural
 625 composition and complexity of AM fungal mycelium. Mycorrhiza 24, 443-451.
- 626 Olsson, P.A., Hansson, M.C., Burleigh, S.H., 2006. Effect of P availability on temporal dynamics of carbon
- allocation and *Glomus intraradices* high-affinity P transporter gene induction in arbuscular mycorrhiza.
 Appl. Environ. Microbiol. 72, 4115-4120.
- 629 Olsson, P.A., van Aarle, I.M., Allaway, W.G., Ashford, A.E., Rouhier, H., 2002. Phosphorus effects on
 630 metabolic processes in monoxenic arbuscular mycorrhiza cultures. Plant Physiol. 130, 1162-1171.
- Patton-Vogt, J..L, Griac, P., Sreenivas, A., Bruno, V., Dowd, S., Swede, M.J., Henry, S.A., 1997. Role of the
 yeast phosphatidylinositol/phosphatidylcholine transfer protein (Sec14p) in phosphatidylcholine turnover and
 INO1 regulation. J. Biol. Chem. 272, 20873-20883.
- 634 Pfeffer, P.E., Douds, D.D., Bécard, G., Shachar-Hill, Y., 1999. Carbon uptake and the metabolism and transport
 635 of lipids in an arbuscular mycorrhiza. Plant Physiol. 120, 587-598.
- Plaxton, W.C., Tran, H.T., 2011. Metabolic adaptations of phosphate-starved plants. Plant Physiol. 156, 10061015.
- Rai, A, Rai, S, Rakshit, A., 2013. Mycorrhiza-mediated phosphorus use efficiency in plants. Environ. Exp. Biol.
 11, 107-117.
- Rich, M.K., Nouri, E., Courty, P.E., Reinhardt, D., 2017. Diet of Arbuscular Mycorrhizal Fungi: Bread and
 Butter? Trends Plant Sci. 22, 652-660.
- Richardson, A.E., Simpson, R.J., 2011. Soil microorganisms mediating phosphorus availability update on
 microbial phosphorus. Plant Physiol. 156, 997-1005.
- 644 Roth, R., Paszkowski, U., 2017. Plant carbon nourishment of arbuscular mycorrhizal fungi. Curr. Opin. Plant
- 645 Biol. 39, 50-56.

- Ruepp, A., Zollner, A., Maier, D., Albermann, K., Hani, J., Mokrejs, M., Tetko, I., Güldener, U., Mannhaupt, G.,
 Münsterkötter, M., Mewes, H.W., 2004. The FunCat, a functional annotation scheme for systematic
 classification of proteins from whole genomes. Nucleic Acids Res. 14, 5539-5545.
- 649 Sánchez-Rodríguez, A., Portal, O., Rojas, L. E., Ocaña, B., Mendoza, M., Acosta, M., Jiménez, E., Höfte, M.,
- 650 2008. An efficient method for the extraction of high-quality fungal total RNA to study the *Mycosphaerella*651 *fijiensis-Musa* spp. Interaction.Mol Biotechnol. 40, 299-305.
- 652 Siggins, A., Gunnigle, E., Smith, F.A., 2012. Exploring mixed microbial community functioning: recent
 653 advances in metaproteomics. FEMS Microbiol. Ecol. 80, 265-280.
- Smith, S.E., Jakobsen, I., Grønlund, M., Smith, F.A., 2011. Roles of arbuscular mycorrhizas in plant phosphorus
 nutrition: interactions between pathways of phosphorus uptake in arbuscular mycorrhizal roots have
 important implications for understanding and manipulating plant phosphorus acquisition. Plant Physiol. 156,
 1050-1057.
- 658 Smith, S.E., Read, D.J., 2008. Mycorrhizal symbiosis, third ed. Academic Press and Elsevier, London.
- Staehr, P., Löttgert, T., Christmann, A., Krueger, S., Rosar, C., Rolčík, J., Novák, O., Strnad, M., Bell, K.,
 Weber, A.P., Flügge, U.I., Häusler, R.E., 2014. Reticulate leaves and stunted roots are independent
 phenotypes pointing at opposite roles of the phosphoenolpyruvate/phosphate translocator defective in *cue1* in
 the plastids of both organs. Front. Plant Sci. 5, 126.
- 663 Steinberg, G., 2007. Hyphal growth: a tale of motors, lipids, and the Spitzenkörper. Eukaryot. Cell 6, 351-360.
- 664 Sugimura, Y., Saito, K., 2017. Transcriptional profiling of arbuscular mycorrhizal roots exposed to high levels of
- phosphate reveals the repression of cell cycle-related genes and secreted protein genes in *Rhizophagus irregularis*. Mycorrhiza 27, 139-146.
- Tedersoo, L., Sánchez-Ramírez, S., Kõljalg, U., Bahram, M., Döring, M., Schigel, D., May, T., Ryberg, M.,
 Abarenkov, K., 2018. High-level classification of the Fungi and a tool for evolutionary ecological analyzes.
 Fungal Divers. 90, 135-159.
- Tian, C., Kasiborski, B., Koul, R., Lammers, P.J., Bücking, H., Shachar-Hill, Y., 2010. Regulation of the
 nitrogen transfer pathway in the arbuscular mycorrhizal symbiosis: Gene characterization and the
 coordination of expression with nitrogen flux. Plant Physiol. 153, 1175-1187.
- 673 Uetake, Y., Kojima, T., Ezawa, T. and Saito, M. 2002, Extensive tubular vacuole system in an arbuscular
 674 mycorrhizal fungus, *Gigaspora margarita*. New Phytologist 154: 761-768.

- van Vuuren, D.P., Bouwman, A.F., Beusen, A.H.W., 2010. Phosphorus demand for the 1970–2100 period: A
 scenario analysis of resource depletion. Global Environ. Change 20, 428-439.
- Vijayakumar, V., Liebisch, G., Buer, B., Xue, L., Gerlach, N., Blau, S., Schmitz, J., Bucher, M., 2016. Integrated
 multi-omics analysis supports role of lysophosphatidylcholine and related glycerophospholipids in the *Lotus*
- *japonicus–Glomus intraradices* mycorrhizal symbiosis. Plant Cell Environ. 39, 393–415.
- Virag, A., Lee, M.P., Si, H., Harris, S.D., 2007. Regulation of hyphal morphogenesis by *cdc42* and
 *rac1*homologues in *Aspergillus nidulans*. Mol. Microbiol. 66, 1579-1596.
- Vit, O., Petrak, J., 2017. Integral membrane proteins in proteomics. How to break open the black box? J.
 Proteomics 153, 8-20.
- Walder, F., Niemann, H., Natarajan, M., Lehmann, M.F., Boller, T., Wiemken, A., 2012. Mycorrhizal networks:
 common goods of plants shared under unequal terms of trade. Plant Physiol. 159, 789-797.
- Wang, W., Shi, J., Xie, Q., Jiang, Y., Yu, N., Wang, E., 2017. Nutrient exchange and regulation in arbuscular
 mycorrhizal symbiosis. Mol. Plant 10, 1147-1158.
- Wewer, V., Brands, M., Dörmann, P., 2014. Fatty acid synthesis and lipid metabolism in the obligate biotrophic
 fungus *Rhizophagus irregularis* during mycorrhization of *Lotus japonicus*. Plant J. 79, 398-412.
- 690 Wild, R., Gerasimaite., R, Jung, J.Y., Truffault, V., Pavlovic, I., Schmidt, A., Saiardi, A., Jessen, H.J., Poirier,
- K., Hothorn, M., Mayer, A., 2016. Control of eukaryotic phosphate homeostasis by inositol polyphosphate
 sensor domains. Science 352, 986-990.
- 693 Wessels, J.G.H., 1993. Wall growth, protein excretion and morphogenesis in fungi. New Phytol. 132, 397-413.
- Wilmes, P., Bond, P.L., 2006. Metaproteomics: studying functional gene expression in microbial ecosystems.
 Trends Microbiol. 14, 92-97.
- Ku, L.J., Jiang, X.L., Hao, Z.P., Li, T., Wu, Z.X., Chen, B.D., 2017. Arbuscular mycorrhiza improves plant
 adaptation to phosphorus deficiency through regulating the expression of genes relevant to carbon and
 phosphorus metabolism. Chinese J. Plant Ecol. 41, 815-825.
- 699 Zybailov, B., Mosley, A.L., Sardiu, M., Coleman, M.K., Florens, L., Washburn, M.P., 2006. Statistical analysis
- 700 of membrane proteome expression changes in *Saccharomyces cerevisiae*. J. Proteome Res. 5, 2339-2347.
- 701

702 Figure legends

Figure 1. Schematic overview of the fungal metabolic pathways responsive to low-Pi fertilization, as predicted
by spectral counting. Metabolic pathways, as inferred from MetaCyc, are indicated in framed bold letters. Red

705 and green colours refer to increased and decreased protein amounts, respectively. Yellow arrow indicates 706 allosteric inhibition, while blue arrow indicates allosteric activation. ACOX, acyl-coenzyme A oxidase; AICAR, 707 aminoimidazole carboxamide ribotide; CoA, coenzyme A; DAH7P, 3-deoxy-D-arabino-heptulosonate 7-708 phosphate; DHAP, dihydroxyacetone phosphate; ERM, extra-radical mycelium; ETC, electron transport chain; 709 FA, fatty acid; FOX2, bifunctional hydroxyacyl-CoA dehydrogenase/enoyl-CoA hydratase; GPX, glutathione 710 peroxidase; GSH, reduced glutathione; GST, glutathione S-transferase; H₂O₂, hydrogen peroxide; H₂S, hydrogen 711 sulfide; IMP, inosine monophosphate; IRM, intra-radical mycelium; NADPH, nicotinamide adenine dinucleotide 712 phosphate; P, phosphate; PA, phosphatidic acid; PCM1, phosphoacetylglucosamine mutase; Pi, inorganic 713 phosphate; PPi, inorganic pyrophosphate; PPP, pentose phosphate pathway; PSD, phosphatidylserine 714 decarboxylase; SAM, S-adonesylmethionine; SH, sulfhydryl; SSG, glutathione disulfide; TAG, triacylglycerol; 715 TCA, tricarboxylic acid; UDP, uridine diphosphate; UPS, ubiquitin-proteasome system; UQ, ubiquinone; UQH₂, 716 ubiquinol

717

Figure 2. Functional distribution of the fungal proteins that display significantly different NSAF values when comparing low- and high-Pi fertilization regimes, as listed in Table2. Bars refer to the number of proteins per functional category, which show either an increased (dark colour) or a decreased (grey colour) NSAF value in low-Pi conditions. Only biological processes accounting for more than 5% of the 162 Pi-responsive proteins identified in Table S2 are presented.

723

724 Figure 3. Overview of the metabolic pathways responsive to low-Pi fertilization, as inferred from shotgun 725 proteomic and qRT-PCR approaches. Red and green colours refer to increased and decreased protein and 726 transcript (*) abundance, respectively. AAs, amino acids; Acetyl-ACP, acetyl-acyl carrier protein; ERM, extra-727 radical mycelium; ETC, electron transport chain; FA, fatty acid; FatM, ACP-thioesterase; IRM, intra-radical 728 mycelium; KASI, β-keto-acyl ACP synthase I; MAG, monoacylglycerol; OAA, oxaloacetate; RAM2, glycerol-729 3-phosphate acyl transferase REDUCED ARBUSCULAR MYCORRHIZA 2; STR2, half-size ABC transporter 730 STUNTED ARBUSCULE 2; TAG, triacylglycerol; TCA, tricarboxylic acid ;VTC, vacuolar transporter 731 chaperone.

732

733





