- 1 Laccaria bicolor MiSSP8 is a small-secreted protein decisive for the
- 2 establishment of the ectomycorrhizal symbiosis
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- 26 protein.

Abstract

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29 The ectomycorrhizal symbiosis is a predominant tree-microbe interaction in forest ecosystems 30 sustaining tree growth and health. Its establishment and functioning implies a long-term and 31 intimate relationship between the soil-borne fungi and the roots of trees. Mycorrhiza-induced 32 Small Secreted Proteins (MiSSPs) are hypothesized as keystone symbiotic proteins, required 33 to set up the symbiosis by modifying the host metabolism and/or building the symbiotic 34 interfaces. 35 L. bicolor MiSSP8 is the third most highly induced MiSSPs in symbiotic tissues and it is also 36 expressed in fruiting bodies. The MiSSP8-RNAi knockdown mutants are strongly impaired in 37 their mycorrhization ability with *Populus*, with the lack of fungal mantle and Hartig net 38 development due to a lack of hyphal aggregation. MiSSP8 C-terminus displays a repetitive 39 motif containing a kexin cleavage site, recognized by KEX2 in vitro. This suggests MiSSP8 40 protein might be cleaved into small peptides. Moreover, the MiSSP8 repetitive motif is found 41 in other proteins predicted secreted by both saprotrophic and ectomycorrhizal fungi. Thus, our 42 data indicate that MiSSP8 is a small-secreted protein involved at early stages of 43 ectomycorrhizal symbiosis, likely by regulating hyphal aggregation and pseudoparenchyma 44 formation.

## Introduction

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Forest soils contain a wide diversity of microorganisms displaying multiple nutrition modes from saprotrophy to pathogenicity, through mutualism (Buée *et al.*, 2009; Uroz *et al.*, 2010; Fierer *et al.*, 2007). Tree-associated microbes are considered as key drivers of tree health, productivity, and ecosystem functionality (Berg *et al.*, 2014; 2015). In particular, fungal communities are primary contributors to carbon and nitrogen cycling in forest ecosystems (van der Hejden, 1998; Lindahl and Tunlid, 2015). Trees forming ectomycorrhizae (ECM) with soil-borne fungi dominate temperate and boreal forest ecosystems (Brundrett, 2009). These mutualistic interactions rely on bidirectional exchanges of nutrients, which happen in mycorrhized roots. The host tree provides carbon derived from its photosynthesis to the

56 fungus, whereas in return ECM fungi provide nitrogen, phosphorus and water. 57 Ectomycorrhizal fungi thus inhabit a dual ecological niche, forest soils and tree root cells, 58 requiring two contrasting ways of life: saprotrophic in soil (for nitrogen acquisition) and 59 biotrophic within plant living tissues. Therefore, most ECM fungi develop (i) extramatrical 60 mycelium exploring the rhizospheric surrounding soil, (ii) aggregated fungal hyphae 61 ensheathing fine lateral roots named mantle, and finally (iii) a highly branched network of 62 fungal hyphae (called the Hartig net) within the apoplastic space of epidermal and cortical 63 root cells (Martin et al., 2016). The Hartig net constitutes the biotrophic interface required for 64 efficient nutrient exchanges. In particular environmental conditions (humidity, temperature), 65 host-derived carbon can be used to build the fruiting body, a spore-releasing structure made of hyphal aggregation (Kües and Navarro-González, 2015; Genre and Bonfante, 2012; 66 67 Lakkireddy et al., 2011). 68 Despite their critical ecological roles, only a few ECM interactions have been studied at the 69 molecular level. Notably, mechanisms mediating the early steps of ECM symbiosis 70 development remain mostly uncharacterized (Daguerre et al., 2017; Martin et al., 2016). First, 71 a pre-contact phase during which plant and fungi communicate is a prerequisite for successful 72 root colonization. Diffusible molecules such as fungal auxins and sesquiterpenes likely 73 mediate this communication and trigger an increase in the lateral roots formation (Felten et 74 al., 2009; Ditengou et al., 2015; Krause et al., 2015; Vayssières et al., 2015). Then, fungal 75 accommodation within the apoplast of root cells requires controlling both hyphal and host 76 root development (Martin et al., 2016). For example, in the poplar-L. bicolor model, both 77 ethylene and jasmonic acid treatments restrain in planta fungal colonization i.e. restrict the 78 intradical hyphal network (Plett et al., 2014). The symbiotic interface at the Hartig net derives 79 from remodeling of both fungal and plant cell walls (Balestrini and Kottke, 2016). Cell wall 80 carbohydrates and proteins (e.g. hydrophobins, mannoproteins) are thus likely to actively 81 contribute to in planta fungal colonization (for review, Balestrini and Kottke, 2016) and to 82 efficient nutrient exchanges. Formation of the Hartig net also leads to a massive fungal 83 colonization within the apoplast of colonized roots, without eliciting strong defence responses 84 (Martin et al., 2016). Considering ECM symbiosis as a biotrophic plant-fungal interaction,

85 secreted fungal molecules likely govern plant colonization by subverting host immunity and 86 manipulating its metabolism to promote the symbiosis establishment and/or functioning (Plett 87 and Martin, 2015; Lo Presti *et al.*, 2015). 88 Genome-wide analysis of Laccaria bicolor has led to the identification of 98 proteins, named 89 MiSSPs (Mycorrhiza-induced Small Secreted Proteins), up-regulated in symbiotic tissues 90 (Martin et al., 2008). Among them, only MiSSP7 has been described as a symbiosis effector 91 so far. MiSSP7 is secreted by the fungus and it enters the host cells in which it localizes 92 within the nucleus. Moreover, MiSSP7-RNAi mutants are impaired in ectomycorrhiza 93 formation (Plett et al., 2011). Inside the nucleus, MiSSP7 interacts with the Populus 94 trichocarpa PtJAZ6 (JAsmonate Zim domain 6), a co-receptor of jasmonic acid (Plett et al., 95 2014). MiSSP7 stabilizes PtJAZ6, avoiding its degradation in the presence of jasmonic acid, 96 leading to the repression of target genes' transcription (Plett et al., 2014). Preliminary results 97 have shown that genes involved in plant cell wall remodelling and plant defence responses 98 might be these target genes (Plett et al., 2014). Furthermore, effector proteins, such as SP7 99 from the arbuscular mycorrhizae fungus Rhizophagus irregularis (Kloppholz et al., 2011), as 100 well as PIIN\_08944 and FGB1 (Fungal Glucan-Binding 1, PIIN\_03211) from the root 101 endophyte Piriformospora indica (Akum et al., 2015; Wawra et al., 2016) also suppress host 102 immunity promoting root colonization and symbiosis. These data support the concept 103 whereby a mutualistic symbiont uses its repertoire of secreted proteins to set up the symbiosis 104 by suppressing host immunity and/or targeting cell-wall remodeling. 105 Considering the diversity of secreted proteins used by pathogens to alter the host metabolism, 106 mycorrhizal fungi might use an equivalent strategy to setup their interaction. However, 107 available literature on such proteins required for the symbiosis establishment is still very poor 108 and functional analyses of MiSSPs are required to clarify and detail how mycorrhizal fungi 109 communicate with their host plant to establish interaction (Plett and Martin, 2015). The 110 Mycorrhiza-induced Small Secreted Protein of 8 kDa, i.e MiSSP8 (JGIv2 ID #388224), 111 displays the third highest induction in mature ectomycorrhizal root tips and is also up-112 regulated in fruiting-bodies (Martin et al., 2008). MiSSP8 is, with other MiSSPs, part of the 113 "core" regulon expressed during the colonization of two hosts, Populus trichocarpa and

114 Pseudotsuga menziesii (Plett et al., 2015). The proteins associated to this core regulon have 115 hence been hypothesized to be the key genetic determinants required for the symbiosis 116 development in L. bicolor (Plett et al., 2015). In this study, we report the functional analysis 117 of L. bicolor MiSSP8 by using a combination of experimental and in silico approaches. Here, 118 we identify MiSSP8 as a key symbiosis factor required for L. bicolor mycorrhization ability, 119 mantle formation and subsequent Hartig net development. This symbiosis factor also has a 120 repetitive motif at its C-terminus containing a kexin-like cleavage site, which is recognized by 121 KEX2 in vitro and liberate four short peptides. The repetitive motif of MiSSP8 is found in 122 other fungal proteins, mostly from ECM and saprotrophic fungi. All together our data indicate 123 that MiSSP8 is involved in fungal mantle and Hartig net formation, potentially by regulating 124 hyphal aggregation, and suggest a part of the symbiotic toolbox used by the ECM fungi to 125 initiate symbiosis is already present in the genome of their saprotrophic ancestors.

#### Material and methods

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## 128 Microorganism and plant material

- 129 Saccharomyces cerevisiae strains YTK12 (Jacobs et al., 1997), MaV103 and MaV203
- (Invitrogen) were propagated in YAPD medium (1% yeast extract, 2% peptone, 2% glucose,
- and 40 mg/L adenine) and cultured at 30°C. The ectomycorrhizal fungus L. bicolor Maire P.
- D. Orton strains (S238N and RNAi-lines) were maintained at 25°C on modified Pachlewski
- medium P5 +/- 150µg.ml<sup>-1</sup> of hygromycin B (Pachlewski and Pachlewska, 1974, Di Battista
- 134 et al., 1996). The hybrid Populus tremula x Populus alba (INRA clone 717-1-B4) cuttings
- were micropropagated *in vitro* and grown on half MS medium (Murashige and Skoog, 1962)
- in glass culture tubes under a 16 h photoperiod at 24°C in a growth chamber. L. bicolor
- basidiocarps were harvested beneath inoculated Douglas fir in a nursery. Three fruiting body
- samples from two developmental stages were harvested; the "early" stage corresponding to
- just emerging fruiting body and "late" stage corresponds to ones with "open" cap. Samples
- 140 from both developmental stages were divided into stipe and cape prior to RNAs extraction
- 141 (Fig. S1).
- To test for altered cell wall susceptibility, Congo red (150 µg/mL; Sigma-Aldrich) was added

to the medium (Fig. S2).

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#### Yeast secretion trap assay

Functional validation of the predicted signal peptide of MiSSP8 was done using the yeast signal-sequence trap assay (Plett *et al.*, 2011). Briefly, full-length sequences of MiSSP8 with or without its signal peptide were cloned into pSUC2-GW, a plasmid carrying the invertase SUC2 lacking both its initiation methionine and signal peptide. Yeast strain YTK12 was transformed with 200 ng of the plasmid using the lithium acetate method (Gietz and Schiestl, 2008). All transformants were confirmed by PCR with vector-specific primers and grown on yeast minimal medium with glucose (SD-W medium: 0.67% Yeast Nitrogen Base without amino acids, 0.075% tryptophan dropout supplement, 2% glucose and 2% agar). To assess invertase secretion, overnight yeast cultures were diluted to an  $O.D_{600} = 1$  and  $20 \mu l$  of dilution were plated onto YPSA medium (1% yeast extract, 2% peptone, 2% sucrose, and 1  $\mu g.mL^{-1}$  antimycin A, inhibitor of cytochrome c oxidase). The YTK12 strains transformed with either the pSUC2-GW empty vector or containing SUC2SP (yeast invertase with signal peptide) were used as negative and positive controls, respectively.

#### Genetic transformation of L. bicolor

- 161 The ihpRNA expression cassette/transformation vector was constructed using the
- 162 pHg/SILBAy vector system (Kemppainen et al, 2005; Kemppainen and Pardo, 2009). The
- full-lenght MiSSP8 cDNA was amplified from oligo(dT)18 synthesized S238N cDNA (First
- Strand cDNA Kit, (Fermentas) using the following gene specific primers:
- 165 MISSP8-SnaBiFor: CTTCTACGTAATGTATTTCCACACTCTTTTCG
- 166 MISSP8-HindIIIRev: TGTCAAGCTTTCAATCACTATCGCGCCTC.
- 167 The cDNA was TA-cloned into pCR®2.1-TOPO® (Invitrogen) for sequencing and the
- 168 corresponding plasmid was used as PCR template to obtain the amplicons needed for ihpRNA
- expression cassette construction. The cloning into the pSILBAy vector was carried out using
- 170 the SnaBI, HindIII, BgIII and StuI restriction sites in pSILBAy. Primers used for
- amplification of the MiSSP8 sequence arms were:

172 MISSP8-SnaBIFor: CTTCTACGTAATGTATTTCCACACTCTTTTCG 173 MISSP8-HindIIIRev: TGTCAAGCTTTCAATCACTATCGCGCCTC 174 MISSP8-BglIIRev: TGTCAGATCTTCAATCACTATCGCGCCTC 175 Completed ihpRNA expression cassette was further cloned as a full length SacI linearized 176 pSILBAy plasmid into the T-DNA of the binary vector pHg to create pHg/pSγMiSSP8. The pHg/pSyMiSSP8 was used for transforming L. bicolor dikaryotic strain S238N with 177 178 Agrobacterium tumefaciens strain AGL1 (Kemppainen and Pardo, 2005). The transformed fungal strains were selected with 300 µg.mL<sup>-1</sup> hygromycin B (Invitrogen) and were later 179 maintained under 150 µg.mL<sup>-1</sup> hygromycin B selection pressure on modified P5 medium. 180 181 Four pHg/pSyMiSSP8 L. bicolor transformant strains were used for further molecular and 182 physiological analyses. 183 184 Molecular analyses of Laccaria bicolor transformants 185 Plasmid rescue of the right border (RB) - linked gDNA was carried out with BamHI cut and 186 self-ligated L. bicolor gDNA according to Kemppainen et al. (2008). Sequencing of the 187 rescued plasmids was done using M13/pUC-reverse primer (-26)17 mer. Left border (LB) 188 TAIL-PCR was done according to Kemppainen et al., 2009, using three T-DNA specific 189 nested primers LB1.3 (Mullins et al., 2001) and the arbitrary primer AD2 (Liu et al., 1995). 190 L3/AD2 amplified TAIL-PCR products were TA-subcloned into pCR®2.1-TOPO® 191 (Invitrogen) and sequenced with the L3 primer. All the PCR reactions were carried out using 192 Tpersonal thermocycler (Biometra ®) and PCR chemicals from Fermentas. Sequencing 193 reactions were purchased from Macrogen Sequencing Service (Seoul, SK). 194 195 In vitro mycorrhization experiments 196 For in vitro mycorrhization tests between P. tremula x alba INRA717-1B4 and L. bicolor, we 197 used a "sandwich" co-culture system described in Felten et al. (2009). After three weeks of 198 co-incubation of poplar cuttings with L. bicolor, at least 10 to 20 biological replicates were 199 analysed for the percentage of colonized roots with the L. bicolor wild-type strain S238N or 200 with the four L. bicolor MiSSP8-RNAi lines. Two independent empty vector transformants of

- 201 L. bicolor (ev7 and ev9 lines, Plett et al, 2011) were also tested for their ability to colonize
- 202 roots.

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- 203 Motif analysis
- For the identification of proteins sharing the DWRR motif found in MiSSP8 sequence, the
- 205 following regular expression
- DW[K/R]-x(2,20)-DW[K/R]-x(2,20)-DW[K/R]-x(2,20) was used on a set
- 207 (Table S1) of fungal and oomycete proteomes available at the MycoCosm database (March
- 208 2019) using the PS-Scan software (de Castro *et al*, 2006) with the options -g (greedyness off)
- 209 -v (overlaps off) -p (pattern search). Only protein sequences from published genomes starting
- 210 with a methionine were kept for further analysis (Table S1). In order to assess conservation of
- the DWRR motif, protein sequences retrieved by PS-Scan were further analyzed by GLAM2
- 212 (Gapped Local Alignments Motifs) software v 4.11.0 (Frith et al., 2008) with default
- 213 parameters. Similar GLAM2 analysis was performed on shuffled sequences as control. The
- 214 presence of a signal peptide in the protein sequence was assessed with SignalP v 4.1 (Petersen
- 215 et al, 2011) with default parameters.

## 217 Microscopy analysis of poplar-ECM roots.

- 218 ECM root tips of poplar-L. bicolor (WT or MiSSP8-RNAi lines) were fixed 24 h in 4%
- 219 paraformaldehyde in PBS buffer (100 mM phosphate buffer, 2.7 mM KCl and 137 mM NaCl
- 220 pH 7.4) at 4°C. The root segments were embedded in agarose 5% and cut into 25 μm radial
- sections with a Leica VT1200S Leica vibratome (Leica Microsystems). Sections were
- categorized according to their distance (100, 200 and 600 µm) from the root apex. 25 µm-
- width sections were stained with 10µg.mL<sup>-1</sup> wheat germ agglutinin (WGA)–Alexa Fluor®
- 488 Conjugate (W21404, ThermoFisher, France) and 1µg.mL<sup>-1</sup> propidium iodide (Invitrogen,
- France). To compare the development of the Hartig net between samples, sections between
- 226 200 and 600 µm distance from the root apex were analyzed.

#### Cell imaging by confocal laser-scanning microscopy

- 229 Transversal sections of ECM root tips were viewed by a Zeiss LSM780 (Carl Zeiss AG,
- 230 Germany) confocal laser scanning microscope system. Images were obtained with objective
- 231 CAPO-40x/1.2 water-immersion objective, acquired sequentially to exclude excitation and
- emission crosstalk (when required). Spectral deconvolution was used to assess specificity of
- the emission signal. Images were processed with ZEN (Carl Zeiss AG) software.

## Quantitative RT-PCR

- For the expression of *MiSSP8* in colonized root tips, extramatrical and free-living mycelium,
- 237 total RNA was extracted from 100 mg biological material using the RNeasy Plant Mini Kit
- (Qiagen), extraction buffer RLC was supplemented with 2% PEG8000. An on column DNase
- I treatment was included in the protocol. 500 ng of total RNA was converted into cDNA in a
- 240 20 µl reaction using the High-Capacity cDNA Reverse Transcription Kit (Applied
- 241 Biosystems, Life technologies, Thermo Fisher Scientific) according to manufacturer's
- 242 instructions and subsequently 1/5 diluted in sterile RNAse and DNAse free water. Real-time
- 243 qPCR was performed in an optical 96-wells plate with a StepOne sequence detection system
- 244 (Applied Biosystems, Life Technologies) and fast cycling conditions (20 s at 95°C, 40 cycles
- of 3 s at 95°C and 30s at 60°C). Each 10 µl reaction contained 2X Fast SYBR green Master
- 246 Mix (Applied Biosystems, Life Technologies), 300 nM gene-specific forward and reverse
- primers, water and 10 ng of cDNA. Data were expressed relatively to the sample with the
- 248 highest expression level (2 -(Ct-Ctmin)) and normalized against six reference genes (JGIv2 IDs
- 249 #293350, #611151, #313997, #446085, #246915 and #319764). Stability of the reference
- 250 genes was confirmed by geNorm analysis (Vandesompele et al., 2002). The normalization
- 251 factor NF for each sample was calculated as the geometric mean of the relative expression
- 252 level of the six reference genes. Consequently, the expression level of MiSSP8 for an
- 253 individual sample was obtained by the formula 2 -(Ct-Ctmin)/NF according to Vandesompele et
- al. (2002). Finally, data were rescaled relative to the FLM.
- 255 For the fruiting body expression, cDNA was obtained from 500 ng of total RNA using the i-
- 256 Script cDNA reverse transcription kit (Biorad) in a final volume of 20 µL. RT-qPCR
- 257 reactions were performed on 10 ng cDNA and 300 nM forward and reverse primers in each

reaction, using the RotorGene (Qiagen) with the standard cycle conditions: 95 °C for 3 min; 40 cycles at 95 °C for 15 s and 65 °C for 30 s, followed by a melting curve analysis (temperature range from 65 °C to 95 °C with 0.5 °C increase every 10 s). In this particular case, transcript abundance was normalized using *L. bicolor* histone H4 (JGIv2 ID# 319764) and ubiquitin (JGIv2 ID #446085) encoding genes. Amplification efficiency (E) was experimentally measured for each primer pair and was taken in account for calculation of normalized expression (Pfaffl *et al.*, 2001).

## Production of recombinant protein and biochemical analysis

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MiSSP8\(Delta I-20\) (i.e. devoid of its first twenty amino-acid residues) was synthesized by Genecust (Luxembourg) and subcloned into the pET-28a-CPDSalI vector (Shen et al., 2009) between NcoI and SalI restriction sites. The resulting plasmid was subsequently used for the transformation of the Rosetta2 (DE3) pLysS strain of E. coli (Novagen). The expression of the recombinant protein (ending with DSDVD in C-ter) was performed at 37°C in Lysogeny broth (LB) medium supplemented with 30 µg.mL<sup>-1</sup> of kanamycin and 34 µg.mL<sup>-1</sup> of chloramphenicol. When the cell culture reached an O.D<sub>600</sub> of 0.7, recombinant protein expression was induced by the addition of 0.1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG), and the cells were grown for further 4 hours at 37°C. Cells were then harvested by centrifugation (13000rpm, 5min), resuspended in a 30 mM Tris/HCl pH 8.0, 200 mM NaCl lysis buffer and lysed by sonication. The cell extract was centrifuged at 35000 g for 25 min at 4°C to remove cellular debris and aggregated proteins. C-terminal His-tagged proteins were purified by gravity-flow chromatography on a nickel nitrilotriacetate (Ni-NTA) agarose resin (Qiagen) according to the manufacturer's recommendations followed by an exclusion chromatography on a Superdex75 column connected to an ÄKTA Purifier<sup>TM</sup> (GE Healthcare). CPD-tag was cleaved using 200 µM inositol-6-phosphate as described by Schen et al. (2009) and removed by size exclusion chromatography. Circular dichroism experiments were carried out with 75 µM of recombinant MiSSP8 in 10 mM sodium phosphate buffer pH 7 using a Chirascan Plus (Applied Photophysics).

In vitro digest assay of recombinant MiSSP8 by KEX2.

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Recombinant MiSSP8 protein (75µg) was incubated with 0.02 units of recombinant KEX2

protease (MoBiTec) in 5 mM CaCl<sub>2</sub>, 50 mM Tris pH7 at room temperature during 1, 2, 4 or 6

hours then was stopped by heating at 95°C for 10 minutes. 50% of each samples was analysed

on Tris-Tricine-precast 12-15% polyacrylamide gels (Biorad), stained with Coomassie blue.

## Mass spectrometry analysis using LC Q-TRAP of KEX2-digested MiSSP8.

294 Synthesized peptide standard (DSDW) obtained from Genecust was solubilized and diluted to

10<sup>-8</sup>M in 2% acetonitrile in water and stored at -20°C. *In vitro* KEX2/MiSSP8 digest assay

was diluted in 100 µl of 2% acetonitrile in water and stored at -20°C. The U-HPLC 3000

(Dionex) was equipped with a reverse-phase column Acquity UPLC BEH-C18 (2.1 x 150

mm, 1.7 µm, Waters). Separation was done with a gradient of eluent A (water/0.1% formic

acid) and eluent B (acetonitrile), started at 5% B for 1 min, followed by a 8 min gradient to

300 100% B, followed by an isocratic step at 100% B for 2 min, a 2 min gradient back to 5% B,

equilibration step 2 min at 5% B, before starting another analysis, at a constant flow rate of

302 300µl/min. 10µl of each samples were injected.

303 The mass spectrometer used was a 4500 Q-Trap mass spectrometer (Applied Biosystems,

304 Foster City, USA) with an electro-spray ionization source in the positive ion mode. The

capillary voltage was fixed at 4500 V and the source temperature at 400°C. Optimizations of

the source parameters were done using the peptide standards DSDW at 10<sup>-5</sup> M in 2 %

acetonitrile by infusion at 7µl. min-1, using a syringe pump. For MS/MS analysis, enhanced

product ions (EPI) mode was used. Declustering potential was fixed at 110 V. Fragmentations

were induced by collision induced dissociation (CID) with nitrogen at a collision energy of 40

V. For the multiple reaction monitoring (MRM) mode, each precursor ions and b/y ions were

calculated for the predicted peptide using Protein Prospector. For relative quantification, the

transitions selected for each peptide were: 522.5>185 (DSDW, RT: 5.0 min), 550.5>175

313 (DSDVD, RT: 4.5 min), 678.5>254 (RDSDW, RT: 5.0 min), 678.5>361 (DSDWR, RT: 4.4

314 min), 834.5>517 (DSDWRR, RT: 4.0 min), 834.5>361 (RDSDWR, RT: 4.0 min), 834.5>313

(RRDSDW, RT: 4.0 min). The intensity of peak height was measured in counts per second

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Results

## MISSP8 is up-regulated both in ECM root tips and fruiting body.

Transcript profiling of L. bicolor free-living mycelium and P. trichocarpa colonized root tips have highlighted the presence of >50 Mycorrhiza-induced Small Secreted Proteins (MiSSPs) in this ECM fungus. Of these, MiSSP8 displayed the third highest induction in mature ectomycorrhizal root tips (Martin et al., 2008). To determine the regulation of MiSSP8 throughout ECM development, we investigated MiSSP8 expression during in vitro ECM time course with P. tremula x alba using real time-qPCR. At 7 days post-contact, fungal hyphae started colonization of fine roots and form the mantle. At 14 days, the Hartig net is present and at 21 days, fully mature ECM tissues have developed. The very low and constitutive-level of MiSSP8 expression in free-living mycelium (FLM) was set as a reference. MiSSP8 was upregulated in ECM root tips from day 7 to day 14, reaching its maximum induction at this latter time point (Fig. 1A). The expression decreased to reach the same level as in FLM at 21 days (mature ECM). Expression of MiSSP8 in the extraradical mycelium (i.e. the part of the rhizospheric mycelium not in contact with the roots) was the same as FLM all along the time course (Fig. 1A). In order to investigate the expression level of MiSSP8 in a non-symbiotic tissue, we performed qRT-PCR on L. bicolor fruiting body tissues (stipe and cap) at two different developmental stages (early and late) (Fig. 1B, Fig. S1). The expression of MiSSP8 in L. bicolor sporocarps is higher than its expression in FLM and ECM root tips. Overall, MiSSP8 was strongly induced during fruiting body-development and P. tremula x alba root colonization (mantle and Hartig net formation).

#### MiSSP8 is secreted as indicated by the yeast invertase secretion assay

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MiSSP8 is a protein containing 70 amino acids, the first twenty residues of which encode a predicted signal peptide as predicted by SignalP v4.1 (Fig. 1C). According to the hydrophobicity plot, the mature MiSSP8 is a hydrophilic protein with charged amino acids exposed to solvent (Fig. 1C) but there is no predicted secondary structure (Fig. S2A). A circular dichroism experiment performed on the recombinant protein produced in E. coli further confirms the lack of secondary structure of MiSSP8 (Fig. S2B). In order to confirm that the predicted signal peptide is functional and properly processed, the full length MiSSP8 (including its signal peptide) was fused to the yeast invertase SUC2, which catalyzes the hydrolysis of sucrose. The transformed yeasts were able to grow on a minimal medium supplemented with sucrose and antimycin (Fig. 1D, bottom right) like the yeast transformed with a full-length invertase (Fig. 1D, bottom left). Yeasts transformed with an empty vector control (Fig. 1D, top left) or an invertase fused to MiSSP8 lacking its signal peptide (Fig. 1D, top right) did not grow on the same medium. This demonstrates that the signal peptide of MiSSP8 is properly recognized and processed in yeast and triggered the secretion of the invertase. Therefore, it is likely that L. bicolor secretes MiSSP8 into the extracellular space during root colonization and fruiting body development.

# MiSSP8-repetitive motif is shared with proteins containing repeats from saprotrophic and ectomycorrhizal fungi

The mature MiSSP8 is composed of 50 amino acids, with a predicted molecular weight of 6105.32 Da and a theoretical pI of 5.12. At its C-terminus, MiSSP8 contains a 25 amino acids long sequence carrying a repetitive motif (i.e. DWRR), repeated four times consecutively. This protein displayed sequence similarities with only one protein of *Laccaria amethystina* (JGIv2 #676588), a close relative of *L. bicolor*, according to BLASTP search on NCBI and JGI MycoCosm databases. Altogether, these data suggest that MiSSP8 is a natively unstructured protein without sequence similarities with previously characterized proteins. To identify additional proteins with a similar motif at their C-termini, we used a pattern search algorithm and identified in total 38 proteins from 23 published fungal proteomes (Fig. 2,

372 Table S1). The DW[K/R]R containing proteins identified were mostly associated to 373 saprotrophs (32/38) and ectomycorrhizal (5/38) fungi including also one arbuscular 374 mycorrhizal fungus, but none pathogenic fungi (Fig. 2, Table S1). Most of the proteins 375 identified above using pattern search algorithm (34/38) had a predicted signal peptide (Fig. 2). 376 No protein domains were detected by PROSITE database, except for two proteins from the 377 white rot fungus Schizophyllum commune, which possess a N-terminal aspartic peptidase A1 378 domain (Fig. 2). GLAM2 motif analysis found an enrichment of a DWR/KR motif, named 379 DW[K/R]R thereafter. 380 The median size of the proteins was 138.5 amino acids, ranging from 70 to 738 amino acids 381 (Fig. 3A). The number of repetitions of the conserved peptide varied from three to seventeen, 382 with no correlation between the size of the protein and the number of motifs (Fig. 3B, Fig. 383 S3). RNA-Seq expression data from *Pinus pinaster* root tips colonized by the ectomycorrhizal 384 fungus Hebeloma cylindrosporum showed that the DWRR containing protein from H. 385 cylindrosporum (JGI IDv2: 440029) was upregulated during root colonization (Doré et al., 386 2015; GEO accession number GSE63868/GSE66156). GLAM2 motif analysis found an 387 enrichment of a DWR/KR motif, named DW[K/R]R thereafter (Fig. 3C, Fig. S3). The 388 DW[K/R]R repetitive motif is therefore shared between proteins predicted as secreted for the 389 most part and associated to saprotrophs and mycorrhizal fungi. 390 391 MiSSP8 possess a repetitive motif containing a kexin cleavage site, recognized *in vitro* by 392 the yeast KEX2 protease. 393 The presence of a repetitive motif in the protein sequence suggests that MiSSP8 might be 394 processed post-translationally in order to become active. Interestingly, the motif DW[K/R]R 395 contains [K/R]R residues, which are known recognition sites for KEX2 proteases in fungi 396 (Mizuno et al., 1988; Mizuno et al., 1989). In fungi, an additional proteolytic cleavage of the 397 KEX2-peptides occurs by KEX1 to withdraw the two amino acids KR or RR. The processing 398 of MiSSP8 at the identified cleavage sites (i.e. RR at each DWRR repeats) by KEX1 and

KEX2 would release one peptide of 27 amino acids, three short peptides of four amino acids

(DSDW) and a final three amino acids long peptide DSD. The sole action of the KEX2

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protease will lead to the release of the DSDWRR peptide. In order to assess whether this motif is recognized by KEX2, we performed in vitro digest assay of recombinant mature MiSSP8 with recombinant yeast KEX2 protein. MiSSP7, a L. bicolor small-secreted protein of 7kDa, which does not contain the (K/R)R motif, was used as negative control. After 2h of incubation, the amount of undigested MiSSP8 decreased and was not detectable after 4h of incubation. For the same incubation period, MiSSP7 was not degraded (Fig. 4). Using mass spectrometry, we assessed whether the predicted DSDWRR or additional peptides were produced to confirm the specificity of the *in vitro* enzyme test. The separation and analytical detection parameters by mass spectrometry were fixed using the synthetic standard DSDW. We obtained a limit of detection of the standard DSDW at 10<sup>-8</sup> M for 10 µl injected, at a retention time of 5.08 min. In the products of digestion, we searched for C-terminal DSDVD, the peptide DSDW and other predicted peptides with one to two additional arginine (R) before/after DSDW (Figure 1C). Among the major detected peptides, we observed the C terminal peptide DSDVD, DSDWR and DSDWRR, meaning that as predicted, KEX2 can hydrolyze MiSSP8 after two arginines (formation of DSDVD and DSDWRR), and between the two remaining arginines, to form DSDWR (Fig. S4).

#### RNAi-mediated knockdown of MiSSP8 encoding gene impairs mycorrhization rate

Since *MiSSP8* expression is induced during the early steps of ectomycorrhizal symbiosis development, we assessed whether MiSSP8 is required for establishment of symbiotic interaction. Generation of knockout mutants through homologous recombination has not been accomplished so far in *L. bicolor*. However, genetic tools to obtain *L. bicolor* strains with significantly reduced target gene expression levels through RNA interference (RNAi) are available (Kemppainen *et al.*, 2009; Kemppainen and Pardo 2010). RNAi hairpin targeting the MiSSP8 transcripts was expressed in *L. bicolor* through *A. tumefaciens* mediated transformation (ATMT). A transgenic ATMT *Laccaria* library was generated and 24 randomly selected independent RNAi lines were passed through consecutive hygromycin B selection steps. Among these, four *Laccaria* RNAi lines were analyzed at molecular level (Plett *et al.*, 2011; Table S2). Real-time qPCR analysis confirmed down-regulation of *MiSSP8* 

in the four transgenic lines, with a reduction from 82% to 95% compared to the empty-vector control lines (Fig. 5A). We quantified the ability of these transgenic RNAi lines to colonize *P. tremula x alba* roots *in vitro*. The mycorrhization rate, defined as percentage of ECM root tips formed over the total number of lateral roots, was found significantly reduced with the mutants, being 45% for the wild-type and empty-vector controls and only 10 to 15% for the four *L. bicolor MiSSP8*-RNAi lines (Fig. 5B). The mycelial growth of each of the *MiSSP8*-RNAi line was tested on rich agar medium or on Congo Red-containing medium and it was similar to the wild-type strain S238N (Fig. S5). Neither did the RNAi mutant show any growth defect nor altered fungal cell wall susceptibility compared to the wild-type fungus. Altogether, it demonstrates MiSSP8 is required for the establishment of the symbiosis with poplar.

# RNAi-mediated knockdown of *MiSSP8* impairs formation of fungal mantle and subsequent Hartig net development

To estimate whether MiSSP8 is necessary for fungal mantle and Hartig net development (two hallmarks of the mature ectomycorrhizal root tips), we performed microscopy analysis on ECM poplar root tips formed. After one week of contact with poplar roots, the fungal mantle formed by *MiSSP8*-RNAi lines was strongly unstructured and thinner than the one obtained with the reference strain (empty vector (ev) control). In addition, whereas ev control already formed first steps of Hartig net, the *MiSSP8*-RNAi lines did not display any kind of fungal colonization. After two weeks, *MiSSP8*-RNAi lines displayed one or two layers of fungal hyphae forming the mantle (Fig. 5C). This observation contrasts with the fungal mantle formed with the control transformant strain, which displayed a thick and well-organized fungal sheath with several layers of fungal hyphae stacked on each other. More strikingly, *MiSSP8*-RNAi lines were impaired in their ability to form the Hartig net even after two weeks of contact (Fig. 5C). Altogether, these results highlight the involvement of MiSSP8 in the differentiation of the fungal mantle precluding hyphal expansion to form the Hartig net.

#### Discussion

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MiSSP8 is required for symbiosis development, likely through its role for hyphal aggregation. Several MiSSPs from L. bicolor, including MISSPs, are known to be part of the "core" regulon expressed during the colonization of two hosts, *Populus trichocarpa* and *Pseudotsuga* menziesii (Plett et al., 2015). These proteins have hence been hypothesized to be key genes required for the symbiosis development in L. bicolor (Plett et al., 2015). The strong decrease in the ability to form mycorrhizae by the MiSSP8-targeted RNAi lines is consistent with this hypothesis. In addition, our study clearly shows that MiSSP8 is directly involved in the symbiosis establishment, through a role played in mantle formation and the subsequent Hartig net development. Real-time qPCR performed on free-living mycelium, symbiotic tissues and fruiting bodies showed a high level of MiSSP8 expression both in mycorrhizal root tips and fruiting bodies but not the free-living mycelium. This expression profile suggests an involvement of MiSSP8 in both symbiosis-related (i.e. formation of ectomycorrhiza) and non-symbiosis-related processes (i.e. fruiting body formation). Both the ectomycorrhizal mantle sheath and the fruiting body tissues are composed of pseudoparenchyma, a pseudo-tissue made of aggregated hyphae that looks like the plant parenchyma (Brunner and Scheidegger, 1992; Peterson and Farquhar, 1994). Ectomycorrhizae developed by MiSSP8-RNAi lines display a disorganized fungal mantle and no Hartig net formation. We speculate that this phenotype results of the lack of fungal aggregation. Hyphae from mantle are indeed glued together and embedded into an extracellular material composed of glycoproteins and fungal polysaccharides (e.g. chitin, β 1-3 glucans) (Massicotte et al., 1990; Dexheimer et al., 1994). Previous studies propose a sequential role of hydrophobins, repellent and polypeptides with a RGD motif (e.g. SRAP32), in the aggregation of hyphae during mantle development in the ectomycorrhizal fungus Pisolithus tinctorius (reviewed by Martin et al., 1999). MiSSP8 is repetitive protein with a kexin cleavage site sharing similarities with other

repetitive proteins from saprotrophs

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Since MiSSP8 displays high sequence similarity with only one L. amethystina gene, we conclude that MiSSP8 is a Laccaria-specific gene, upregulated both in symbiosis and in fruiting bodies. These lineage-specific genes may have been formed de novo or may derived from neofunctionalization of duplicated genes or from ancestral genes that have strongly diverged due to selection pressure (Kohler et al., 2015; Pellegrin et al., 2015). However, despite the absence of sequence similarities, a MCL (Markov Cluster Algorithm) analysis performed on a set of 49 fungal genomes had previously retrieved 33 proteins containing a similar motif as MiSSP8 (Kohler et al, 2015). With the current analysis, and using a different search approach, we also identify additional fungal proteins harboring the same repetitive motif than MiSSP8 as well as a kexin endoproteinase cleavage site (K/R)R (Mizuno et al., 1988; Mizuno et al., 1989). In addition, we showed this cleavage site is recognized in vitro by yeast KEX2. Despite several trials, we were not able to detect the predicted released peptides in ectomycorrhiza root tips or L. bicolor fruiting bodies (data not shown). This could be due to post-translational modifications of the peptides or a fixation to extracellular components such as fungal cell wall carbohydrates or glycoproteins. We can therefore only suggest that the proteins containing the (DWRR)<sub>n</sub> motif could be processed by kexin prior to their secretion or go through post-translational modifications in order to become active and release such peptides. Several fungal peptides are produced from KEX2-processed precursor proteins (Le Marquer et al, 2019) e.g., cyclic peptides with mycotoxic activity in Ascomycota such as phomopsins (Ding et al., 2016) or ustiloxins (Umemura et al., 2014). On the other hand, Ustilago maydis Rep1 protein is also cleaved by kexin-protease and has a structural role in the fungal cell wall (Wösten et al., 1996; Teertstra et al., 2006). A 11 amino acid long peptide processed by KEX2 in *Cryptococcus neoformans* is required for virulence and to activate the sexual program (Homer et al., 2016; Tian et al., 2018). Since MiSSP8-RNAi lines are not impaired in hyphal growth or fungal cell-wall sensitivity, it is unlikely that MiSSP8 or its derived peptides are involved in fungal cell wall structure. In addition, MiSSP8 does not bind to fungal cell wall sugars (data not shown). The precise role of MiSSP8 and its derived peptides in the development of the fungal mantle will require further research.

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The fungal proteins identified in our study exhibit the (DW[K/R]R)<sub>n</sub> motif at their C-termini with a variable number of repetitions but they do not share sequence similarities at their Ntermini. Variations in number of tandem repeats is thought to provide functional diversity (Verstrepen et al, 2005), suggesting that the identified DW[K/R]R-containing proteins might be involved in various cellular processes and carry out different functions. If KEX2processed, the variable number of repeats would lead to different number of released peptides. A comparative phylogenomics analysis of 49 fungal genomes revealed that ECM fungi have evolved several times from saprotrophic ancestors, these being either white rot, brown rot or soil decayers, by developing a set of symbiotic genes with rapid turnover, and in particular gain of genes such as MiSSPs (Kohler et al., 2015). Moreover, a recent comparative analysis of five Amanita genomes, two ECM symbionts and three asymbiotic species, concluded that several genetic components of the toolkit used by ECM symbionts is already encoded in the genome of their saprotrophic relatives, explaining the recurrent emergence of the ECM symbiosis over time (Hess et al., 2018). Consistent with this finding, MiSSP8 shares similarities with other saprotrophic proteins through its fungal-specific repetitive motif. Since MiSSP8 is likely playing a role in the formation of the pseudoparenchyma of both nonsymbiotic (basidiocarp) and symbiotic (ECM) structures, we propose that MiSSP8 function, initially required for L. bicolor fruiting body formation, has been recruited for the establishment of the symbiosis. This highlights the dual use of the given secreted protein into symbiotic and non-symbiotic processes. Our analysis also suggest that there are two categories of MiSSPs: the orphan ones, such as MiSSP7 and the ones preexisting in saprotrophic ancestors, such as MiSSP8.

In conclusion, we have characterized the Mycorrhiza induced Small Secreted Protein of 8 kDa (MiSSP8) by combining functional and *in silico* approaches. We demonstrate that MiSSP8 has a functional secretion signal peptide and it contains a repetitive motif containing kexin cleavage sites recognized *in vitro* suggesting the protein might be cleaved in order to become functional. Our data show that MiSSP8 or its derived peptides are decisive factors for symbiosis establishment. The DWRR repetitive motif being also found in SSPs from

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saprotrophic fungi, we propose that MiSSP8 or its derived peptides could have been initially linked to fruiting body development by participating in pseudoparenchyma formation through fungal hyphae aggregation before being recruited for symbiosis establishment. **Acknowledgments** This research was sponsored by the Genomic Science Program, US Department of Energy, Office of Science, Biological and Environmental Research as part of the Plant-Microbe Interfaces Scientific Focus Area (http://pmi.ornl.gov) and the Laboratory of Excellence ARBRE (grant no. ANR-11-LABX-0002 ARBRE). The research was supported by the Institut National de la Recherche Agronomique and the University of Lorraine (Ph.D. scholarship to CP and YD). Both Région Lorraine Research council and the European Fund for Regional Development give funding for the Functional Genomics Facilities at Institut National de la Recherche Agronomique-GrandEst. Part of the work was supported by Laboratoire Recherche Sciences Végétales - UPS CNRS, MetaToul (Metabolomics and Fluxomics Facitilies, Toulouse, France, www.metatoul.fr) and the French National infrastructure for metabolomics and fluxomics, www.metabohub.fr, MetaboHUB-ANR-11-INBS-0010. JR was an AgreenSkills Marie Sklodowska Curie postdoctoral fellow co-funded by the European Commission (FP7-267196). We thank Barbara Montanini and Simone Ottonello for sharing the pSUC-modified vectors. We thank Alexandre Kriznik from the Platform of Biophysics and Structural Biology-UMS 2008 IBSLor (CNRS-INSERM-UL) for his technical assistance in circular dichroism and isothermal calorimetry experiments. Funding to A.P. and M.K. was provided by grants from Universidad Nacional de Quilmes, Consejo Nacional de Investigaciones Científicas y Técnicas, and Agencia Nacional de Promoción Científica y Tecnológica, Argentina. **Authors contributions** 

- 569 CVF and FM designed and managed the project; CP, YD, FG, MK, JR, VP, NFdF, AH, CVF
- 570 performed the experiments; CP, CVF, MK, AP, VP, NFdF analyzed the data; CP performed
- 571 the bioinformatic analysis; CP and CVF wrote the manuscript and all authors revised it.

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- 779 **Legends of Figures**
- 780 Figure 1: MiSSP8 is a secreted repeat-containing protein highly expressed in
- ectomycorrhizal root tips and fruiting body tissues.
- 782 (A) Real-time qPCR time course performed on Laccaria bicolor extraradical mycelium
- (ExM), free-living mycelium (FLM) and during development of *Populus tremula* x *alba* ECM
- 784 root tips. Expression of *MiSSP8* in FLM was set as a reference. Error bars represent standard
- deviation from three biological replicates.
- (B) Real-time qPCR analysis on the expression of *MiSSP8* versus two reference genes in the
- fruiting body of *L. bicolor*, the cap and stipe, at both early and late stages of development.
- 788 Stars indicate significant differences (p-value cut-off < 0.05) using Welch t-test (Welch,
- 789 1947). Error bars represent standard deviation from three biological replicates.
- 790 (C) MiSSP8 sequence, domain organization and hydrophobicity score as obtained by
- Protscale using the amino-acid scale from Kyle and Doolittle (1982) with a window size of 5.
- 792 (**D**) Yeast signal trap assay shows that the signal peptide predicted in MiSSP8 sequence is
- 793 functional in yeast. Saccharomyces cerevisiae was transformed either with empty vector
- 794 control (top left), mature MiSSP8 (lacking its signal peptide) fused to mature invertase
- 795 (MiSSP8Δ1-20::Invertase, top right), full-length invertase (with its signal peptide, bottom
- 796 left) or full length MiSSP8 fused to mature invertase (MiSSP8::Invertase, bottom right).
- 797 Transformed yeasts were plated on growth medium containing sucrose as the sole source of
- 798 carbon.

- 800 Figure 2: The DW[K/R]R proteins possess a repetitive structure and are likely secreted.
- 801 The modular structure of the proteins is shown. Protein domains have been identified by
- PROSITE and signal peptides are predicted by SignalP v4.1.
- 804 Figure 3: MiSSP8-repetitive motif is found in proteins from saprotrophic and
- 805 ectomycorrhizal fungi.
- 806 (A) Size distribution (in amino acids) of the DW[K/R]R-containing proteins within the
- 807 different fungal lifestyles identified.

- 808 **(B)** Number of repetitions identified in the DW[K/R]R-containing proteins.
- 809 (C) Identification of the DWRR motif by GLAM2 software. The set of 38 sequences retrieved
- with PS-Scan has been submitted to GLAM2 run with default parameters.

## Figure 4. Recombinant MiSSP8 is cleaved by KEX2 protease in vitro.

- 814 (A) Visualization of proteolytic digests of recombinant MiSSP8 and MiSSP7 proteins by
- recombinant KEX2 protease, separated on Coomassie-stained-polyacrylamide gels. A time
- course over 1, 2, 4 and 6h of incubation was performed. Sizes are indicated in kDa. MiSSP7
- 817 is used as negative control since it is not a putative substrate for KEX2 protease.+ and -
- indicate the presence or absence of KEX2, respectively.
- 819 (B) Detected peptides by LC-MS/MS after 4h of digestion by the recombinant KEX2 protease
- 820 of the MiSSP8 protein. Theoretical formed peptides are DSDWRR and DSDVD but
- 821 additional combinations of cleavages were hypothesized. The transitions selected for each
- 822 peptide were: m/z 522.5>185 (DSDW, RT: 5.0 min), 550.5>175 (DSDVD, RT: 4.5 min),
- 823 678.5>254 (RDSDW, RT: 5.0 min), 678.5>361 (DSDWR, RT: 4.4 min), 834.5>517
- 824 (DSDWRR, RT: 4.0 min), 834.5>361 (RDSDWR, RT: 4.0 min), 834.5>313 (RRDSDW, RT:
- 4.0 min). Peak height intensity is measured in counts per second (cps). Mean of 3 independent
- replicates, standard error of the mean is reported.

## 828 Figure 5: Knockdown of MiSSP8 impairs the establishment of ectomycorrhizal

829 symbiosis.

827

- 830 (A) Expression was measured by qRT-PCR. All values are shown as mean  $\pm$  standard
- deviation; n = 3. Stars indicate that expression is statistically different than control line based
- on a non-parametric Kruskal-Wallis test followed by a Games-Howell post-hoc test (Ruxton
- and Beauchamp, 2008). Adjusted p-value (FDR) cut-off = 0.05. ev indicates fungal lines
- transformed with an empty vector (i.e. negative control).
- 835 (B) Knockdown of MiSSP8 strongly decreases the number of ectomycorrhizal root tips
- compared to the wild-type strain or the empty vector controls. Percentage of poplar (*Populus*

861

865

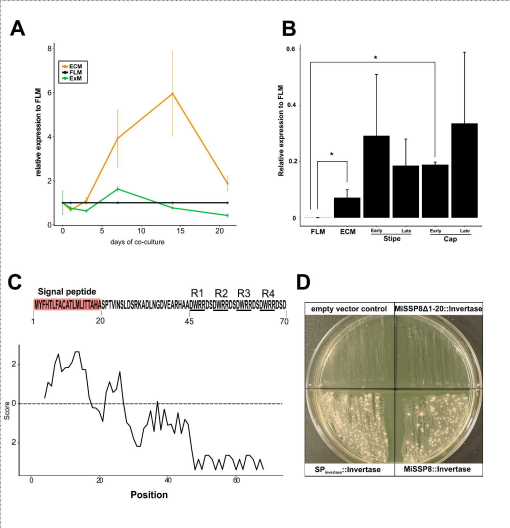
837 tremula x alba) mycorrhizal root tips formed by wild-type L. bicolor S238N and two empty 838 vector transformation controls (ev7 and 9) versus four independent L. bicolor MiSSP8-RNAi 839 lines. All values are shown as mean ± standard deviation; n=25; Stars indicate significant 840 difference from wild-type, ev7 and ev9 using t-test and a p-value cut-off < 0.05. (C) Transversal section of ectomycorrhizal root tips formed by ev7 control transformant line 842 and MiSSP8-RNAi line 3. The MiSSP8-RNAi line 3 displays a loose mantle and no Hartig net 843 compared to the empty vector control strain (ev7). Scale bars: 10 µm. Propidium iodide stains 844 plant cell walls and nuclei (red) and WGA-Alexa 488 stains fungal cell walls (green). Green 845 signal in root cortical cells correspond to aspecific WGA-Alexa 488 fixation to plant cell 846 walls and not to fungal cells. 847 848 Figure S1: Fruiting body of L. bicolor at two different developmental stages. The cap and 849 stipe of fruiting bodies from L. bicolor were harvested at early and late developmental stages 850 in order to study the expression of MiSSP8 during the development of the fruiting body. 851 852 Figure S2: In silico prediction and experimental validation of MiSSP8 secondary 853 structure. (a) Secondary structure of MiSSP8 as predicted by the JPred 4 server (Drozdetskiy 854 et al, 2015). (b) Analysis of recombinant MiSSP8 secondary structure by circular dichroism. 855 856 Figure S3: Correlation between protein size sequences and occurrence of the 857 **DWRR/DWRR-like motif.** No correlation found between the size of the DW[K/R]R protein 858 sequences identified and the number of motifs they contain (A), even after removing the 859 outliers (B). 860 Figure S4: Spectrum obtained for the peptide DSDWR in LC-MS/MS. The selected 862 precursor ion (m/z) 678.6) is corresponding to the peptide DSDWR  $(M+H)^+$ ; after 863 fragmentation (CE 40V), y ions are observed as predicted in Protein Prospector (m/z 175.1; 864 361.1; 476.2; 563.2; 678.2).

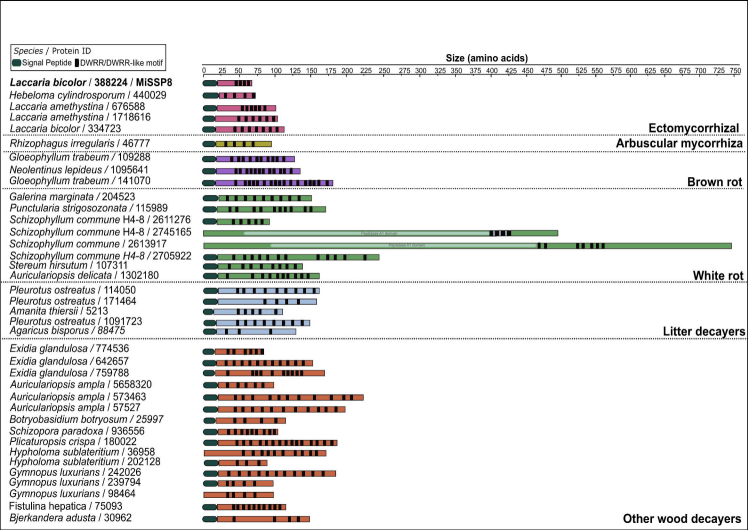
Figure S5. *L. bicolor missp8* RNAi mutants are not impaired in their saprotrophic growth in both control medium (P5) or under cell-wall stress conditions.

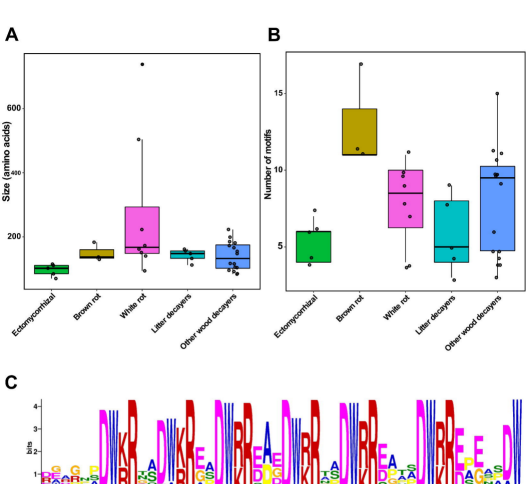
Viability of *Missp8* RNAi lines was assessed by growing the different strains used (Wildtype (WT), empty vector control (ev7) and the four RNAi lines generated on either control medium (P5) or under cell-wall stress condition (Congo Red).

Supplementary Table S1: List of genomes scanned and proteins sequences containing from two to twenty repetitions of the DW[K/R]R motif.

Supplementary Table S2: Molecular analysis of *Laccaria bicolor Missp8*-RNAi lines. Molecular analysis of the four RNAi lines generated expressing RNAi hairpin targeting the Missp8 encoding transcript.







Α

Intensity (cps)

