The transcriptome of the arbuscular mycorrhizal fungus Glomus intraradices (DAOM 197198) reveals functional tradeoffs in an obligate symbiont

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**Summary**

The arbuscular mycorrhizal symbiosis is arguably the most ecologically important eukaryotic symbiosis, yet it is poorly understood at the molecular level. To provide novel insights into the molecular basis of symbiosis-associated traits, we report the first genome-wide analysis of the transcriptome from *Glomus intraradices* DAOM 197198.

We generated a set of 25 906 nonredundant virtual transcripts (NRVTs) transcribed in germinated spores, extraradical mycelium and symbiotic roots using Sanger and 454 sequencing. NRVTs were used to construct an oligoarray for investigating gene expression.

We identified transcripts coding for the meiotic recombination machinery, as well as meiosis-specific proteins, suggesting that the lack of a known sexual cycle in *G. intraradices* is not a result of major deletions of genes essential for sexual reproduction and meiosis. Induced expression of genes encoding membrane transporters and small secreted proteins in intraradical mycelium, together with the lack of expression of hydrolytic enzymes acting on plant cell wall polysaccharides, are all features of *G. intraradices* that are shared with ectomycorrhizal symbionts and obligate biotrophic pathogens.

Our results illuminate the genetic basis of symbiosis-related traits of the most ancient lineage of plant biotrophs, advancing future research on these agriculturally and ecologically important symbionts.

**Introduction**

The arbuscular mycorrhizal (AM) symbiosis between fungi in the Glomeromycota (Schüssler et al., 2001) and plants involves over two-thirds of all known plant species, including important crop species, such as wheat (*Triticum aestivum*), rice (*Oryza sativa*), maize (*Zea mays*), soybean (*Glycine max*) and poplar (*Populus* spp.). This mutualistic symbiosis, involving one of the oldest fungal lineages, is arguably the most ecologically and agriculturally important symbiosis in terrestrial ecosystems (Fitter et al., 2011). The extraradical mycelium (ERM) of the symbiont acts as an extension of the root system and increases the uptake of key nutrients, particularly phosphorus (P) and zinc (Zn) and possibly also nitrogen (N) (Smith & Smith, 2011). These fungi are therefore crucial to plant growth (Smith & Read, 2008) and also define the diversity of plants in ecosystems (van der Heijden et al., 1998). Furthermore, because the colonization of plants by AM fungi can also result in a 20% net increase in photosynthesis (Smith & Read, 2008), these universal mycosymbionts make a very large, poorly understood contribution to the global carbon cycling budget of ecosystems.

The Glomeromycota are unique in that their spores and coenocytic hyphae contain multiple nuclei in a common cytoplasm. No sexual cycle is known, although anastomosis and nuclear movement between
hyphae of the same species have been described (Giovannetti et al., 2001), as well as genetic exchange between AM fungal individuals (Croll et al., 2008, 2009). Furthermore, the concept of an individual is unclear as nuclei within a single AM fungus appear to be genetically different in some species, which raises substantial questions about the natural selection, population genetics and gene expression of these highly unusual organisms (Jany & Pawlowska, 2010; Sanders & Croll, 2010). AM fungi are thought to grow clonally, but controversy clouds their ploidy, their genome size, and whether or not they can reproduce sexually (for a review, see Sanders & Croll, 2010). Evidence for recombination has been provided, but whether mating and meiosis are involved is unknown (Croll & Sanders, 2009). There is therefore a need to identify genes whose products are required for proper completion of meiotic recombination.

Root colonization by AM fungi follows a series of distinct steps (for a review, see Bonfante & Genre, 2010) starting with a presymbiotic molecular dialogue that involves root-released strigolactones (Akiyama et al., 2005) and AM fungal signaling molecules, such as lipochitooligosaccharides (Maillet et al., 2011). AM fungi are obligate mutualistic symbionts that can only grow for a limited time without colonizing a susceptible host root (Bécard et al., 2004). This has been suggested to be a consequence of some nutritional deficiencies and loss of metabolic pathways, such as an absence of de novo fatty acid synthesis (Bago et al., 2000), during the asymbiotic and presymbiotic phases related to putative genome erosion (Ercolin & Reinhardt, 2011). After spore germination, the first contact at the root surface is marked by the differentiation of the fungal hyphae into an appressorium from which a penetration hypha invades the root epidermis (Genreet al., 2005). After the fungus has traversed the outer cell layers it spreads in the inner cortex, that is, intraradical mycelium (IRM), and forms highly branched structures inside cortical cells, so-called arbuscules. Arbuscules are the main site of nutrient transfer from the fungus to the plant (Javot et al., 2007; Bonfante & Genre, 2010; Smith & Smith, 2011). Arising from the colonized roots, the ERM proliferates in the growth medium, where it absorbs and assimilates the available nutrients before their transfer to the host plant. Major transcriptome shifts are thus expected during these different key developmental stages, but little is known of the repertoire of effector-like proteins, membrane transporters and assimilative enzymes involved in these different steps of symbiosis development and functioning.

The ability to establish a sophisticated zone of interaction, such as the haustorium in pathogenic oomycetes and fungi interacting with plants, requires sophisticated host defense suppression (Dodds & Rathjen, 2010), which is predominantly achieved via secreted proteins delivered into the host cell (Kamoun, 2007). The protein SP7 of Glomus intraradices is secreted and transferred to the plant cell nucleus in colonized roots of Medicago truncatula, where it binds to the host ethylene-responsive transcriptional factor regulating the expression of several defense-related genes (Kloppholz et al., 2011). It remains to be determined whether G. intraradices expresses genes coding for additional secreted effector-like proteins during its interaction with the host plant.

The mycorrhizal colonization leads to quantitative and qualitative changes in the host transcriptome. Plant genes that are specifically regulated during the establishment and/or functioning of the AM symbiosis have been identified by both targeted and high-throughput expression profiling in several sequenced model and crop plants (Güimil et al., 2005; Balestrini & Lanfranco, 2006; Gomez et al., 2009; Grunwald et al., 2009; Guether et al., 2009a). In contrast, regulation of gene expression in the AM fungal symbionts caused by interactions with roots has so far been poorly explored, and studies have mainly been based on targeted approaches providing only narrow insights into fungal adaptation to the symbiotic mode (Gomez et al., 2009; Seddas et al., 2009; Tani et al., 2009; Kuznetsova et al., 2010). Most studies have focused on P transport and metabolism (Viereck et al., 2004; Benedetto et al., 2005; Balestrini et al., 2007; Javot et al., 2007; Gómez-Ariza et al., 2009; Grunwald et al., 2009) and primary carbon metabolism (Bago et al., 2000,
Further studies adopting a microarray or high-throughput sequencing approach are definitely warranted, because they could provide a wider understanding of how genetic changes in the plant affect overall patterns of fungal gene expression, as well as the impact of the fungal genotypes on symbiosis fitness and plant growth (Angelard et al., 2010).

Biotrophs are widely accepted to intimately interact and co-evolve with their hosts. The genetic changes that brought about the evolution of obligate biotrophy in the Glomeromycota are, however, unknown. Recent research on obligate biotrophic pathogens, such as the powdery mildew fungus *Blumeria graminis* (Spanu et al., 2010), the downy mildew oomycete *Hyaloperonospora arabidopsis* (Baxter et al., 2010), the oomycetous white rust *Albugo laibachii* (Kemen et al., 2011), and the poplar and stem rusts *Melampsora larici-populina* and *Puccinia graminis*, respectively (Duplessis et al., 2011), reveals a close correlation between the biotrophic life style and gene losses in primary and secondary metabolism. In the oomycetes, all haustorium-forming species have lost the thiamine biosynthetic pathway, suggesting that haustorial oomycetes obtain thiamine from the host (Kemen et al., 2011). It appears that biotrophy in these obligate plant pathogens is also associated with a convergent loss of secondary metabolic enzymes and an extremely reduced set of carbohydrate-active enzymes devoted to plant cell wall depolymerization. Genes for nitrate and nitrite reductases, a nitrate transporter, and sulfite reductase are also often missing. This streamlining of metabolism presumably reflects adaptation to life within host cells, as it is not observed in nonobligate biotrophic pathogens. Whether similar events have also characterized AM fungal evolution remains to be determined.

As mentioned above, transcriptome studies have been pursued on AM roots in multiple plant species under a variety of experimental conditions, but they have mainly focused on the host plant transcriptome. No single study has yet brought together genome-wide transcriptomic data for the fungal component of this important symbiosis. The aim of this study was to establish a comprehensive, genome-wide inventory of gene expression in *G. intraradices* DAOM197198 by sequencing cDNA libraries from different fungal structures (germinated spores, extra- and intraradical mycelium, and arbuscules). Through transcriptomic analyses, we also wished to identify which of these genes were induced upon symbiosis and, further, to assign the differentially expressed transcripts to key symbiotic mechanisms, such as nutrient transport and assimilation, host colonization, and signaling pathways. Several factors have led to the choice of *G. intraradices* for the first large-scale transcriptome sequencing of an AM fungus. As a symbiont, *G. intraradices* is highly effective in mobilizing, taking up and transferring mineral nutrients, such as inorganic orthophosphate ions (Pi), N and sulfur (S), from soils to plants (Govindarajulu et al., 2005; Allen & Shachar-Hill, 2009; Tian et al., 2010; Smith & Smith, 2011) and it readily colonizes many plants, including agriculturally important crop species (e.g., alfalfa (*Medicago truncatula*), poplar, rice and wheat) as well as model plants such as *M. truncatula* and *Lotus japonicus*. *Glomus intraradices* is one of the most studied AM fungi as it rapidly colonizes its host plants, and it is a model species for dissecting the molecular and cellular biology of the Glomeromycota (Seddas et al., 2009; Sanders & Croll, 2010). It is readily amenable to *in vitro* culture on transformed host roots (Chabot et al., 1992) and is the only species for which spores are available commercially in pure form in large quantities.

The comparative analyses of gene repertoires in *G. intraradices*, pathogenic obligate biotrophs and ectomycorrhizal fungi offer insights into genes that may be involved in obligate biotrophy and mycorrhizal symbioses. In the absence of a whole-genome sequence for any member of the phylum Glomeromycota (Martin et al., 2008a), the availability of large-scale expressed sequenced tag (EST) collections represents the core foundation for understanding genome functionality in the Glomeromycota.
Materials and Methods Biological material

*Glomus intraradices* Schenck & Smith DAOM 197198 (recently reassigned to *G. irregularare* and then *Rhizophagus irregularis* (Blaszk., Wubet, Renker & Buscot) C. Walker & A. Schüßler comb. nov.; see Stockinger et al., 2009) was produced in monoxenic cultures maintained on *Agrobacterium rhizogenes*-transformed carrot (*Daucus carota*; clone DC2) roots (Bécard & Fortin, 1988). Germinated spores were produced as described by Chabot et al. (1992). Biological materials used for cDNA library construction and microarray transcript profiling are summarized in Table 1. Protocols for producing the biological materials, RNA purification and cDNAs are described in the online Supporting Information Methods S1. For array profiling, *G. intraradices* ERM was grown on liquid M medium without sucrose for 3 wk before harvesting. *Glomus intraradices*- and mock-inoculated roots of *M. truncatula* were prepared as described for the cDNA library AKNA (Methods S1). Germinated spores were sampled as described for the cDNA library AKND (Methods S1) (Balestrini et al., 2007). Samples were snap-frozen in liquid N₂.

| Table 1. *Glomus intraradices* tissues used in the extraction of transcripts for cDNA library construction and oligoarray profiling
<table>
<thead>
<tr>
<th>Samples</th>
<th>Sanger sequencing</th>
<th>454 sequencing</th>
<th>Array profiling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germinated spores</td>
<td>CCHU</td>
<td></td>
<td>Three replicates</td>
</tr>
<tr>
<td>Germinated spores mix = spores + spores in GR24 + spores in API</td>
<td></td>
<td>EXTA</td>
<td></td>
</tr>
<tr>
<td>Germinated spores in GR24</td>
<td></td>
<td></td>
<td>AKNC</td>
</tr>
<tr>
<td>Germinated spores in API</td>
<td></td>
<td></td>
<td>AKNB</td>
</tr>
<tr>
<td><em>Daucus carota</em> ERM</td>
<td>CACE</td>
<td></td>
<td>EXTB</td>
</tr>
<tr>
<td><em>Medicago truncatula</em> ERM</td>
<td></td>
<td></td>
<td>Three replicates</td>
</tr>
<tr>
<td><em>M. truncatula</em> IRM</td>
<td></td>
<td>AKNA*</td>
<td>Three replicates</td>
</tr>
<tr>
<td>LMD-microdissected arbuscule-containing cells</td>
<td></td>
<td></td>
<td>AKND</td>
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</table>

GR24, striogalactone GR24; API, apigenin; ERM, extraradical mycelium; IRM, intraradical mycelium; AKNA, AKNB, AKNC, AKND; CACE, CCHU, EXTA and EXTB stand for cDNA library IDs. AKNA, *M. truncatula* IRM; AKNB, germinated spores incubated in apigenin for 24 h; AKNC, germinated spores incubated in the striogalactone GR24 for 24 h; AKND, laser-microdissected *M. truncatula* arbuscule-containing cells; CACE, *M. truncatula* ERM; CCHU, germinated spores; EXTA, germinated spore mix; EXTB, *M. truncatula* ERM; LMD, laser microdissection system.

*The cDNA library AKNA has been constructed using mRNA from a mix of the three replicates also used for microarray transcript profiling.*

EST sequencing, filtering and assembly

Sanger sequencing was performed on ABI3730xl analyzers (Applied Biosystems, Nutley, NJ, USA) and pyrosequencing on the Genome Sequencer FLX System (454 Life Sciences/Roche, Nutley, NJ, USA). Reads (797 394) were filtered and trimmed for low quality, low complexity, and adaptor sequences using SeqClean (TIGR (The Institute for Genomic Research); http://sourceforge.net/projects/seqclean/) (Supporting Information Fig. S1). Sequences ≤ 100 bp were discarded. The resulting high-quality sequences were screened to detect potential bacterial and fungal contaminant sequences, and plant sequences present in cDNAs from symbiotic tissues. Sequences having a GC% higher than that of *G. intraradices* (i.e. ≥ 45%) were removed (Fig. S2). Sequences with a high nucleotide sequence similarity (BLASTN e-value ≤ 1e-5, score ≥ 150, % identity ≥ 95) with identified contaminants, *Aspergillus* spp. and *Chromobacterium violaceum*, as well as the *M. truncatula* genome and transcripts, and other plant DNA (e.g. carrot) were also removed (Fig. S1). Finally, to avoid the loss of *G. intraradices* outlier sequences, removed sequences were aligned to the *G. intraradices* draft genome using BLASTN (e-value ≤ 1e-5, score ≥ 150) at the INRA
GlomusDB website (http://mycor.nancy.inra.fr/IMGC/GlomusGenome/index3.html) and retained when sequence similarity was ≥ 95%. Filtered reads were then assembled using the MIRA assembler (Chevreux et al., 2004). To choose the best assembly, several assemblies were constructed using different parameters and several indexes were measured, including the number of nonredundant virtual transcripts (NRVTs), the percentage of assembled reads and the percentage of NRVTs with hits against gene models from the taxonomically related, although distant, Mucoromycotina *Rhizopus oryzae* (http://genome.jgi-psf.org/Rhior3/Rhior3.home.html) (Supporting Information Table S1). The assembly generated by MiraSearchESTSNPs was selected. The number of detected NRVTs increased with the number of reads and the rarefaction curve did not reach a plateau (Fig. S3a). However, many of these sequences may be the result of DNA pyrosequencing errors creating false transcripts. This is an acute problem for the AT-rich sequences of *G. intraradices*. In contrast, the number of protein sequences documented in the Swiss-Prot DNA database (http://www.uniprot.org/) and *R. oryzae* gene repertoire showing a similarity to *G. intraradices* NRVTs plateaued at approx. 5500 homologs (Fig. S3b), suggesting that most of the transcriptome of *G. intraradices* was covered. This was confirmed by a TBLASTN search (cut-off e-value of 1e-5) using the protein sequences of the core eukaryotic genes (CEGs) (Parra et al., 2009); 245 (98.7%) of the 248 CEG proteins were found in the current *G. intraradices* transcriptome. The missing protein sequences were a metal-binding protein, a spindle assembly checkpoint protein and a monooxygenase involved in coenzyme Q (ubiquinone) biosynthesis. Eighty-six per cent of NRVTs matched the current *G. intraradices* S2.5 Mb-genome assembly (Martin et al., 2008a), confirming that most filtered, high-quality ESTs were from *G. intraradices*.

Single-nucleotide polymorphisms (SNPs) in NRVTs were identified with MiraSearchESTSNPs using the following parameters: minimum reads, 2 (Sanger) or 4 (454); minimum quality value, 30 (Sanger) or 25 (454); minimum neighbor quality value, 20.

For the digital northern gene expression, the number of reads for each NRVT in each library was counted and the relative frequency (reads of a given NRVT divided by the total number of reads) was obtained. Significant differences in gene expression between libraries for each NRVT were calculated (Audic & Claverie, 1997).

EST sequences are available at the National Center for Biotechnology Information (NCBI) (accession numbers GW091323–GW125581and GW086621–GW090678). A MySQL/PHP database compiling NRVT sequences and annotations is available at the INRA GlomusDB website. This server can also be used to query the ESTs and the draft genome assembly using BLAST programs.

**Functional annotation**

NRVTs were compared against Swiss-Prot using BLASTX with a significance threshold (e-value < 1e-5). Gene annotations were assigned to each NRVT based on the best BLAST hits. Sequences were searched against the *G. intraradices* draft genome contigs (version ‘test14’) at the INRA GlomusDB using BLASTN (e-value ≤ 1e-5). NRVTs were also compared with proteins of Basidiomycota (*Cryptococcus neoformans*, *Laccaria bicolor*, *M. larici-populina*, *Phanerochaete chrysosporium* and *Ustilago maydis*), Ascomycota (*Aspergillus nidulans*, *B. graminis*, *Botrytis cinerea*, *Magnaporthe grisea*, *Neurospora crassa* and *Tuber melanosporum*) and Mucoromycotina (*R. oryzae* and *Mucor circinelloides*) using BLASTX (e-value ≤ 1e-5). To identify conserved protein domains in NRVTs, predicted protein sequences were compared with the Eukaryotic Orthologous Groups (KOG) database (Tatusov et al., 2003). The counts of each KOG domain by species were transformed into a z-scores matrix to center the data, and visualization
was performed using MeV (MultiExperiment Viewer) (Saeed et al., 2006). The gene ontology (GO) terms (Ashburner et al., 2000) were assigned to each sequence using BLAST2GO (Conesa et al., 2005). Enrichment analysis of GO annotations was carried out with BINGO (http://www.psb.ugent.be/cbd/papers/BiNGO/Home.html) using the hypergeometric test and Benjamini and Hochberg’s false discovery rate (FDR) multiple testing correction \( P \leq 0.05 \) (Maere et al., 2005).

To predict *G. intraradices* metabolic pathways, sequences were queried for Kyoto Encyclopedia of Genes and Genomes (KEGG) Orthology (KO) assignments (Kanehisa & Goto, 2000) using the KEGG Automatic Annotation Server (KAAS) (Moriya et al., 2007) and results were then plotted into the KEGG Global metabolic atlas (map01100) using iPATH (Letunic et al., 2008). A high-resolution version of this metabolic pathway map can be downloaded at: http://mycor.nancy.inra.fr/IMGC/GlomusGenome/download/Gi_metabolic_map.pdf.

To identify sequences encoding secreted proteins, NRVTs were translated into protein using FrameFinder (http://www.ebi.ac.uk/~guy/estate/). *In silico* predictions of secreted proteins were then carried out using SignalP 3.0 (Bendtsen et al., 2004), TargetP 1.1 (Emanuelsson et al., 2000) and tmhmm 2.0 (prediction of transmembrane helices in proteins) (Krogh et al., 2001) as previously reported in Duplessis et al. (2011). Small secreted proteins (SSPs) were selected based on an arbitrary cut-off of 300 amino acids (Duplessis et al., 2011).

**Oligoarray design**

For the design of custom oligoarrays, a MIRA assembly (v. 1.0) comprising 22,254 NRVTs was constructed using cDNA libraries CACE, CCHU, EXT and EXTB. At the time, sequences from the AKNA, AKNB, AKNC and AKND libraries were not available. To ensure that all sequences were sufficiently long to allow the design of high-quality probes, we selected the subset of sequences \( \geq 200 \) bp in length \((n = 19,465)\). To remove redundancy among sequences that would produce cross-hybridization, the assembled NRVTs were clustered based on nucleotide sequence similarity using BLASTClust (http://www.ncbi.nlm.nih.gov/Web/Newsltr/Spring04/blastlab.html). Within each cluster, sequences were ranked by length, and the longer member of each cluster was selected \((n = 15,932)\). We also selected singleton sequences with matches in NCBI databases and with length \( \geq 200 \) bp \((n = 2046)\). We then added 2327 NRVTs generated by the Paracel Transcript Assembler (Striking Development; http://www.paracel.com/) (S. Rensing, unpublished results) which are not found in the MIRA NRVT set. We also included 12 *G. intraradices* reference sequences downloaded from the NCBI, 23 *G. intraradices* mitochondrial sequences (C. Roux, unpublished results) and 70 *G. intraradices* sequences generated from the manual annotation of the *G. intraradices* genome and not found in the NRVTs. Finally, 4471 remaining singleton Sanger sequences without matches in public databases were added. For 2477 sequences of this set, no probe could be designed, probably because of their relative high A + T content, which resulted in a higher-than-usual prevalence of homopolymer runs and other forms of low-complexity sequences. This procedure produced a total set of 22,404 sequences containing 14,828 MIRA NRVTs, 1976 PARACEL NRVTs, 5496 singletons and 104 additional sequences. This set did not include NRVTs from symbiotic tissues not available when the custom array was constructed. However, c. 80% of the NRVTs used to construct the oligoarray were found by BLASTN in the 25,906 MiraSearchESTSNPs NRVTs.

The *G. intraradices* expression array \((4 \times 72K)\) manufactured by Roche NimbleGen Systems Limited (Madison, WI, USA) (http://www.nimblegen.com/products/exp/index.html) contained three independent,
nonidentical, 60-mer probes per sequence. Included in the oligoarray were \textit{G. intraradices} sequences, and 5785 random 60-mer control probes.

\textbf{Transcript profiling}

Whereas biological samples used for cDNA sequencing were produced by several different collaborators, biological materials used for microarray transcript profiling were generated in triplicate in a small number of laboratories: spores at the Laboratoire de Recherche en Sciences Végétales (Université de Toulouse, France), \textit{M. truncatula} ERM at the Departamento de Microbiología del Suelo y Sistemas Simbióticos (Estación Experimental del Zaidín, CSIC, Granada, Spain) and \textit{M. truncatula} IRM at the Boyce Thompson Institute for Plant Research (Tower Road, Ithaca, NY, USA). cDNA synthesis was carried out at INRA-Nancy.

RNA was extracted from mycorrhizal roots with Trizol reagent (Invitrogen Corporation, Carlsbad, CA, USA) with additional phenol : chloroform (1 : 1, v/v) purification steps, from ERM and spores using the RNeasy Plant Mini Kit (Qiagen) with RLT lysis buffer. RNA from mycorrhizal roots was treated with Turbo DNase I (Ambion Inc., Austin, TX, USA) and purified with the RNeasy MinElute CleanUp kit (Qiagen). DNase treatment of RNA from ERM was performed using RNase-free DNase (Qiagen). RNA quality was checked before cDNA synthesis using the Bio-Rad Experion analyzer. RNA preparations (three biological replicates) were amplified using the SMART PCR cDNA Synthesis Kit (Takara Bio Europe/Clontech, Saint-Germain-en-Laye, France) according to the manufacturer’s instructions. Single dye labeling of cDNA samples, hybridization procedures and data acquisition were performed at the NimbleGen facilities (NimbleGen Systems, Reykjavik, Iceland) following their standard protocol. As ERM and IRM materials were not produced in strictly similar conditions, transcript profiles of ERM and IRM should be compared with caution.

Average expression levels were calculated for each gene from the independent probes on the array and were used for further analysis. Raw array data were normalized by the robust multiarray average (RMA) routine using the ARRAYSTAR software (Dnastar Inc., Madison, WI, USA). A transcript was deemed expressed when its signal intensity was three-fold higher than the mean signal-to-noise threshold (cut-off value) of the random oligonucleotide probes present on the array (50–100 arbitrary units). The maximum signal intensity values for the most abundant transcripts were c. 65 000 arbitrary units. A control hybridization was performed with cDNA from uninfected \textit{M. truncatula} roots to evaluate any plant RNA cross-hybridization with \textit{G. intraradices} sequences and to detect any plant-derived probes on the array. The later probe signals were discarded. A Student’s \( t \)-test with FDR correction (Benjamini–Hochberg) was applied to the data using ARRAYSTAR. Transcripts with a significant \( P \)-value (< 0.05) and \( \geq \) five-fold change in transcript level were considered to be differentially expressed. The complete expression data set is available as series (accession number GSE29866) at the Gene Expression Omnibus at NCBI (http://www.ncbi.nlm.nih.gov/geo/).

\textbf{Validation of oligoarray data}

To obtain the biological material for the PCR validation of oligoarray data, 6000 arbuscule-containing cells from \textit{M. truncatula} roots (2000 cells for each of three biological replicates) were microdissected using a laser microdissection system (LMD) according to Balestrini \textit{et al.} (2007). ERM from monoxenic \textit{M. truncatula} cultures and c. 6000 germinated spores were sampled, snap-frozen in liquid N\(_2\) and stored at –80°C until used. Total RNA was extracted using the RNeasy Microarray Tissue Mini Kit (Qiagen, Courtaboeuf, France) following the manufacturer’s instructions. All reverse transcriptase (RT)-PCR assays were carried out using the One Step RT-PCR kit (Qiagen). DNA contamination in RNA samples was
evaluated using the *G. intraradices* elongation factor *GintEF1α* gene specific primers for *G. intraradices*. To determine the amount of extracted RNA, a semiquantitative RT-PCR using *GintEF1α* specific primers was performed. Reactions were carried out in a final volume of 25 μl. Amplification reactions were run for 40 cycles of 94°C for 30 s, 58°C for 30 s, and 72°C for 30 s, and an aliquot of the PCR reaction was taken after 35, 37 and 40 cycles.

Results and Discussion

An obligate biotroph with a large gene repertoire

The cDNA libraries of *G. intraradices* listed in Table 1 were sequenced to assess the size and diversity of the fungus gene repertoire and explore the responses of the AM fungus at key developmental stages, germinated spores, ERM and IRM. After sequencing, the *de novo* hybrid clustering of the 437 411 filtered reads produced 25 906 NRVTs (Table S1); 87% of the reads were in the assembly corresponding to 20.3 Mbp of nonredundant sequences. Only 10 823 NRVTs (41.8%) matched database sequences (cut-off *e*-value of 1e-5) in BLAST searches against genes from fungi other than Glomeromycota (Fig. S4). Homologous NRVTs corresponded to 8773 unique known genes in the DNA databases. Among the predicted peptide sequences, 9035 and 9265 showed significant sequence similarity to proteins from *R. oryzae* and *M. circinelloides*, respectively, distantly related fungi in the order Mucorales (Fig. S4). NRVTs that did not match any of the known genes still gave highly relevant hits against the *G. intraradices* draft genome, suggesting that these represent novel orphan genes. As many *G. intraradices* transcripts are represented by two NRVTs (i.e. aligning to the 5′ or 3′ end of the transcript) with an NRVT:gene ratio of 1.4, the number of nonredundant expressed transcripts in *G. intraradices* was c. 18 500 (25 906/1.4).

Based on BLASTClust analysis, the number of NRVTs found in multigene families was low, with 120 (90% sequence identity; 80% sequence coverage) to 500 (90% sequence identity; 50% sequence coverage) NRVTs, most of the gene families having only two members.

Although the vast majority of genes encoding enzymes of primary metabolism are retained in obligate biotrophic pathogens (downy and powdery mildews, white rust, and poplar and stem rusts), notable exceptions include anaerobic fermentation, biosynthesis of glycerol from glycolytic intermediates, biosynthesis of thiamine, and nitrate and sulfate assimilation (Baxter et al., 2010; Spanu et al., 2010; Duplessiset al., 2011; Kemen et al., 2011). Out of the 25 906 *G. intraradices* NRVTs, 5296 (20.4%) had significant matches in the KEGG pathway database and were assigned to 175 KEGG pathways. Most KEGG enzymes were mapped to NRVTs (see the metabolic pathway map online at http://mycor.nancy.inra.fr/IMGC/GlomusGenome/download3.php?select=anno), indicating the occurrence of these active metabolic processes in *G. intraradices* mycelium. In contrast to oomycete and fungal biotrophic pathogens, transcripts coding for nitrate and nitrite reductases, nitrate transporter, and sulfite reductase were highly expressed in *G. intraradices*. However, *G. intraradices* has lost the thiamine biosynthetic pathway (Table S2), as have most haustorium-forming species (Kemen et al., 2011). We infer that *G. intraradices* obtains thiamine from the host. There is no invertase transcript in the *G. intraradices* transcriptome (Table S2), implying that this fungus depends on the plant to both provide and hydrolyse sucrose, with the glucose moiety then preferentially transferred to the mycobiont. This is
consistent with earlier observations (Schaarschmidt et al., 2006). Additional genes missing from biotrophic pathogens (Spanu et al., 2010), such as those coding for the allantoin permease DAL4p, the uracil permease FUR4p, and several enzymes involved in aromatic amino acid metabolism and detoxification, were also missing from *G. intraradices* (Table S2). Although some metabolic pathways are missing in *G. intraradices*, we hypothesize that evolution to biotrophy in *G. intraradices* was not initiated by massive loss of metabolic complexity as observed in obligate biotrophic pathogens (Spanu et al., 2010). The mycobiont retains the ability to take up and assimilate nutrients from its soil environment.

**Protein domain distributions**

About half (5056) of the 10,823 peptide sequences from *G. intraradices* with sequence similarity to documented proteins in databases were shared with all other fungal species and can thus be considered ubiquitous among fungi. The comparison of protein sets of *G. intraradices* with those of *R. oryzae* and *M. circinelloides* showed that 1076 proteins were unique to these basal lineages of aspetate, coenocytic fungi. Interestingly, a small set of *G. intraradices* protein sequences were only shared with one or other of the sequenced mycorrhizal symbionts, the ascomycete *T. melanosporum* (49 NRVTs) or the basidiomycete *L. bicolor* (107 NRVTs). They mainly code for hypothetical proteins with no known function, although a few predicted proteins belong to signaling pathways. A total of 9321 NRVTs contained at least a part of a conserved protein domain in the KOG database. Compared with other fungal gene repertoires, *G. intraradices* showed an overrepresentation of proteins involved in signaling pathways and ubiquitin-related metabolism (Fig. 1). The expansion of the tyrosine kinase-encoding gene family involved in signaling pathways is also a feature of the ectomycorrhizal *L. bicolor* genome (Fig. 1; Martin et al., 2008b).
Sequence polymorphism of transcripts

There were a total of 43,872 SNPs in 3963 NRVTs (15.3% of the total number of NRVTs; 2.1 SNP/kb); 1102 NRVTs contained < 5 SNPs (Fig. 2a). When this SNP analysis was conducted on NRVT regions having a similarity with known protein coding sequences, 846 polymorphic NRVTs (8%; 1.3 SNP/kb) were identified among 10,823 homologs. Most polymorphic NRVTs contained < 5 SNPs (Fig. 2b). The NRVTs with known function having the highest SNPs were a 60S ribosomal protein L17 (15 SNPs) and a Ras-related Rab (11 SNPs). The presence of multiple SNPs in hundreds of NRVTs confirmed the within-isolate DNA sequence polymorphism that has repeatedly been reported for a number of genes in *G. intraradices* (Croll et al., 2008; Sanders & Croll, 2010). The transcriptome data show that this polymorphism is widespread in the genome, and that > 1 variant of each polymorphic gene is transcriptionally active. This implies that the sequence variation may be of functional importance.
Symbiosis induces alterations in the *G. intraradices* transcriptome

Little is known about the impact of symbiosis on the mycobiont transcriptome and molecular factors driving developmental pathways (Güimil *et al.*, 2005; Gomez *et al.*, 2009; Seddas *et al.*, 2009; Kuznetsova *et al.*, 2010; Sanders & Croll, 2010). Our transcript profiling thus provides the first large-scale discovery of fungal symbiosis-related genes. Of the 18751 coding sequences detected by oligoarrays, 395 (2.1%) and 569 (3.0%) were up- and down-regulated (≥ five-fold; *P*-value ≤ 0.05), respectively, in IRM in comparison to germinated spores, whereas 202 (1.1%) and 74 (0.4%) were up- and down-regulated, respectively, in ERM in comparison to germinated spores. Highly up-regulated genes in IRM and ERM are shown in Tables 2, 3. In this study, mRNA concentrations were used as a proxy of protein concentrations. Gene expression is, however, a multistep process that involves the transcription, translation and turnover of messenger RNAs and proteins, and transcript levels are not always related to protein levels and enzyme activities (Schwanhäusser *et al.*, 2011). It remains to be determined whether the observed changes in transcripts lead to alteration of the proteome, enzyme activities and/or metabolic fluxes. In addition, regulatory enzymes exhibiting allosteric properties are likely to be effective agents for fine tuning of metabolic fluxes in symbiotic tissues.

The majority of IRM- and ERM-induced genes were lineage-specific genes, as 80% of the most highly up-regulated transcripts coded for orphan proteins (Tables 2). Differentially expressed genes coding for proteins of known function were categorized into functional classes (Tables 3). Genes overrepresented in IRM vs germinated spores showed a GO enrichment (FDR-corrected *P* ≤ 0.05) for several categories of biological processes associated with ion transport, lipid/steroid metabolism and DNA replication (data not shown). Out of 25 transcripts that appeared to be highly up-regulated in IRM, 17 were validated by RT-PCR (data not shown). While 13 out of 17 transcripts were detected in at least one other stage of the life cycle.
besides arbuscules, four were arbuscule-specific transcripts (Table S5). Three were orphan genes, while the fourth showed similarity to a multidrug resistance protein (ATP Binding Cassette (ABC) transporter).

As the oligoarray was constructed before the 454 ESTs from IRM and LMD arbuscule-containing cells were obtained, we also carried out a digital northern analysis based on the differential frequency of 454 reads (Audic & Claverie, 1997). This transcript profiling confirmed the above oligoarray expression patterns (data not shown). However, we detected a significant set of IRM transcripts coding for SSPs not detected in ERM and spores. These SSPs showed a striking up-regulation in IRM and arbuscule-containing cells (Table 4). The most highly up-regulated IRM orphan transcript ‘step3_c3163’ coded for an SSP of 280 amino acids of unknown function with similarity to proteins of several biotrophic basidiomycetes (e.g. *L. bicolor* and *M. larici-populina*). This transcript was only detected in mycorrhizal roots and arbuscules.
Table 3: All transcripts (nonredundant virtual transcripts (NRVTs)) with functional annotation that were up-regulated by at least 10-fold in *G. intraradices*.

<table>
<thead>
<tr>
<th>NRVT ID</th>
<th>Ratio</th>
<th>IRM expression level</th>
<th>Putative function</th>
<th>Functional categories</th>
</tr>
</thead>
<tbody>
<tr>
<td>EXT825980_b0</td>
<td>17.6</td>
<td>8653</td>
<td>MFS monocarboxylic acid transporter</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c20826</em></td>
<td>13.7</td>
<td>25 099</td>
<td>Amino acid transporter</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c3620</em></td>
<td>27.4</td>
<td>21 270</td>
<td>MFS nitrate transporter</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c10810</em></td>
<td>34.5</td>
<td>22 106</td>
<td>MFS nitrate transporter</td>
<td>Transport</td>
</tr>
<tr>
<td>EXT118370_b0</td>
<td>333.4</td>
<td>1353</td>
<td>Iron permease</td>
<td>Transport</td>
</tr>
<tr>
<td>GI_8222_c1</td>
<td>236.2</td>
<td>18 627</td>
<td>ZIP (Zrt- and Irt-like) Zn transporter</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c13887</em></td>
<td>15.8</td>
<td>8222</td>
<td>P-type ATPase</td>
<td>Transport</td>
</tr>
<tr>
<td>GI_5720_c1</td>
<td>12.5</td>
<td>3337</td>
<td>Cation transport-related protein</td>
<td>Transport</td>
</tr>
<tr>
<td>EXT11701_b0</td>
<td>24.7</td>
<td>7920</td>
<td>ABC transporter</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c14346</em></td>
<td>16.1</td>
<td>1921</td>
<td>P-type ATPase</td>
<td>Transport</td>
</tr>
<tr>
<td><em>G. c9272</em></td>
<td>77.1</td>
<td>3766</td>
<td>Fatty acyl-CoA-kinase</td>
<td>Lipid metabolism</td>
</tr>
<tr>
<td>EXT126208_b0</td>
<td>10.6</td>
<td>9156</td>
<td>Peroxisomal 3-ketoacyl-CoA-thiolase</td>
<td>Lipid metabolism</td>
</tr>
<tr>
<td><em>G. c8771</em></td>
<td>40.2</td>
<td>9435</td>
<td>Phosphatidylglycerol/phosphatidylinositol transfer</td>
<td>Lipid metabolism</td>
</tr>
<tr>
<td><em>G. c605_c1</em></td>
<td>48.2</td>
<td>23 206</td>
<td>Lipase</td>
<td>Lipid metabolism</td>
</tr>
<tr>
<td><em>G. c11625</em></td>
<td>15.4</td>
<td>26 531</td>
<td>Acetyl-CoA acetyltransferase</td>
<td>Lipid metabolism</td>
</tr>
<tr>
<td><em>G. c15563</em></td>
<td>10.0</td>
<td>10 357</td>
<td>Adenine phosphoribosyl transferase</td>
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<tr>
<td>GI_1390_c2</td>
<td>32.1</td>
<td>15 107</td>
<td>Ribonucleoside-diphosphate reductase</td>
<td>Nucleotide metabolism</td>
</tr>
<tr>
<td><em>G. c2075</em></td>
<td>23.9</td>
<td>12 730</td>
<td>Thymidylate synthase</td>
<td>Nucleotide metabolism</td>
</tr>
<tr>
<td><em>G. c12729</em></td>
<td>26.4</td>
<td>10 113</td>
<td>Endoporphathase</td>
<td>Phosphate metabolism</td>
</tr>
<tr>
<td>EXT108707_b0</td>
<td>18.8</td>
<td>17 134</td>
<td>Endoporphathase</td>
<td>Phosphate metabolism</td>
</tr>
<tr>
<td><em>G. c4991</em></td>
<td>27.0</td>
<td>26 571</td>
<td>Carbonic anhydrase</td>
<td>Carbon metabolism</td>
</tr>
<tr>
<td>GI_8293_c1</td>
<td>43.4</td>
<td>7083</td>
<td>Cytochrome P450</td>
<td>Secondary metabolism</td>
</tr>
<tr>
<td><em>G. c7119</em></td>
<td>10426.8</td>
<td>10 427</td>
<td>Cytochrome P450</td>
<td>Secondary metabolism</td>
</tr>
</tbody>
</table>

| EXT818295_b0 | 27.7  | 20 682               | Multikopper oxidase                                   | Secondary metabolism         |
| *G. c1906* | 10.0  | 11 236               | Glutathione S-transferase                             | Secondary metabolism         |
| *G. c1140* | 10.9  | 24 021               | Iron sulfur cluster assembly protein                  | Protein modification         |
| *G. c14063* | 21.0  | 17 704               | Ubiquitin-conjugating enzyme                          | Protein modification         |
| *G. c23386* | 56.8  | 13 312               | Protease, Ulp1 family                                 | Protein modification         |
| *G. c2553* | 13.6  | 3141                 | Ubiquitin/ribosomal protein 527a                     | Protein modification         |
| *G. c8660* | 10.0  | 8892                 | Menaquinone biosynthesis                              | Coenzyme metabolism         |
| *G. c1425* | 13.6  | 22 927               | Steroidogenic acute regulatory protein                | Signal transduction          |
| CAC68046_g1 | 11.3  | 2292                 | Mitogen-activated protein kinase                      | Signal transduction          |
| *G. c7236b* | 13.5  | 2738                 | Tyrosine-like kinase                                  | Signal transduction          |
| *G. c6409* | 50.5  | 37 996               | Calmodulin-binding protein                            | Signal transduction          |
| *G. c7412* | 33.1  | 50 044               | Calmodulin-binding protein                            | Signal transduction          |
| *G. c3994* | 14.4  | 5290                 | Chitin synthase                                       | Cell wall biosynthesis       |
| *G. c1601* | 40.8  | 50 329               | Cell wall anchor domain protein                       | Cell wall biosynthesis       |
| CHU1751_g1 | 1480.7 | 8199                 | Centromere protein-related protein                    | Cell cycle control           |
| *G. c3811* | 27.2  | 9906                 | Kinetochore protein                                   | Cell cycle control           |
| *G. c7300* | 17.9  | 20 774               | Targeting protein for Xilip2                         | Cell cycle control           |
| *G. c2880* | 50.3  | 16 495               | Kinesin-like protein                                  | Cytoskeleton                 |
| *G. c2425* | 39.1  | 4799                 | DNA topoisomerase type II                             | Chromatin structure          |
| *G. c1747* | 1248.8 | 22 989               | Chromatin assembly factor 1                          | Chromatin structure          |
| *G. c006* | 26.3  | 37 147               | Histone H3.1                                         | Chromatin structure          |
| CHU9624_1b | 12.5  | 450                  | DNA repair protein RAD51                               | Replication, repair          |
| *G. c9197* | 12.7  | 7425                 | DNA replication licensing factor                     | Replication, repair          |
| *G. c1484* | 73.5  | 2903                 | Uracil-DNA glycosylase                                | Replication, repair          |
| *G. c13628* | 31.5  | 3433                 | BTB/POZ domain-containing protein                     | Transcription                |
| *G. c18585* | 13.2  | 4791                 | Zn finger-containing protein                          | Transcription                |
| *G. c10966* | 80.6  | 9500                 | Zn finger-containing protein                          | Transcription                |
| *G. c1198* | 11.2  | 25 711               | Yabby-like transcription factor                       | Translation initiation factor |
| *G. c739* | 11.3  | 11 896               | Translation initiation factor 3                       | Translation                   |

IRM compared with germinated spores based on oligonucleotide profiling (ratio ≥ 10; P-value < 0.05).

ABC, ATP-binding cassette; BTB (for BR-C, Ttk and bab) or POZ (for Fox virus and Zinc finger); IRM, intrasradical mycelium; MFS, major facilitator family.
The majority of IRM- and ERM-induced genes were lineage-specific genes, as 80% of the most highly up-regulated transcripts coded for orphan proteins (Tables 2). Differentially expressed genes coding for proteins of known function were categorized into functional classes (Tables 3). Genes overrepresented in IRM vs germinated spores showed a GO enrichment (FDR-corrected \( P \leq 0.05 \)) for several categories of biological processes associated with ion transport, lipid/steroid metabolism and DNA replication (data not shown). Out of 25 transcripts that appeared to be highly up-regulated in IRM, 17 were validated by RT-PCR (data not shown). While 13 out of 17 transcripts were detected in at least one other stage of the life cycle besides arbuscules, four were arbuscule-specific transcripts (Table S5). Three were orphan genes, while the fourth showed similarity to a multidrug resistance protein (ATP Binding Cassette (ABC) transporter).

As the oligoarray was constructed before the 454 ESTs from IRM and LMD arbuscule-containing cells were obtained, we also carried out a digital northern analysis based on the differential frequency of 454 reads (Audic & Claverie, 1997). This transcript profiling confirmed the above oligoarray expression patterns (data not shown). However, we detected a significant set of IRM transcripts coding for SSPs not detected in ERM and spores. These SSPs showed a striking up-regulation in IRM and arbuscule-containing cells (Table 4). The most highly up-regulated IRM orphan transcript ‘step3_c3163’ coded for an SSP of 280 amino acids of unknown function with similarity to proteins of several biotrophic basidiomycetes (e.g. L. bicolor and M. larici-populina). This transcript was only detected in mycorrhizal roots and arbuscules.

<table>
<thead>
<tr>
<th>NRVT ID</th>
<th>IRM (reads)</th>
<th>Arbuscules (reads)</th>
<th>Spores (reads)</th>
<th>Putative function</th>
<th>Size (aa)</th>
<th>Cys</th>
</tr>
</thead>
<tbody>
<tr>
<td>step3_c3163</td>
<td>62</td>
<td>4</td>
<td>0</td>
<td>Hypothetical protein</td>
<td>280</td>
<td>12</td>
</tr>
<tr>
<td>remain_c2375</td>
<td>24</td>
<td>18</td>
<td>0</td>
<td>No hit</td>
<td>156</td>
<td>3</td>
</tr>
<tr>
<td>step3_c3357</td>
<td>15</td>
<td>2</td>
<td>0</td>
<td>No hit</td>
<td>149</td>
<td>0</td>
</tr>
<tr>
<td>remain_c4548</td>
<td>12</td>
<td>4</td>
<td>0</td>
<td>M2D2-related lipid recognition domain</td>
<td>173</td>
<td>4</td>
</tr>
<tr>
<td>step3_c2834</td>
<td>11</td>
<td>19</td>
<td>0</td>
<td>No hit</td>
<td>61</td>
<td>9</td>
</tr>
<tr>
<td>remain_c2379</td>
<td>11</td>
<td>7</td>
<td>0</td>
<td>No hit</td>
<td>120</td>
<td>0</td>
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<tr>
<td>step3_c3282</td>
<td>10</td>
<td>1</td>
<td>0</td>
<td>No hit</td>
<td>86</td>
<td>2</td>
</tr>
<tr>
<td>remain_c8131</td>
<td>9</td>
<td>0</td>
<td>0</td>
<td>Hypothetical protein</td>
<td>150</td>
<td>3</td>
</tr>
<tr>
<td>remain_c2779</td>
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<td>20</td>
<td>0</td>
<td>No hit</td>
<td>67</td>
<td>0</td>
</tr>
<tr>
<td>remain_c8897</td>
<td>7</td>
<td>1</td>
<td>0</td>
<td>No hit</td>
<td>71</td>
<td>7</td>
</tr>
<tr>
<td>remain_c12045</td>
<td>5</td>
<td>3</td>
<td>0</td>
<td>No hit</td>
<td>122</td>
<td>4</td>
</tr>
</tbody>
</table>

These sequences have no probe on oligoarrays.

aa, amino acids; # Cys, number of cysteine residues; NVRT, nonredundant virtual transcript.

Recent studies of plant–microbe interactions involving pathogenic and ectomycorrhizal fungi have shown that many lineage-specific orphan genes code for effector proteins playing key roles in host colonization and in planta accommodation by controlling the plant immune system (De Wit et al., 2009; Plett et al.,
Several obligate biotrophic pathogens and ectomycorrhizal symbionts have a decreased repertoire of carbohydrate-acting enzymes involved in the degradation of plant cell wall (PCW) polysaccharides (Martin et al., 2008b, 2010; Baxter et al., 2010; Spanu et al., 2010; Duplessis et al., 2011). We identified 139 NRVTs encoding carbohydrate-active enzymes in the G. intraradices transcriptome (Table S6). They included enzymes that are involved in acquiring and converting carbohydrates and in maintaining fungal cell wall plasticity. Genes encoding glycosyl hydrolases (GHs) involved in degrading PCW lignocellulosic polymers were not found in the IRM predicted proteome. No genes encoding GH from families GH6 and GH7 (which include cellobiohydrolases and are involved in the attack of crystalline cellulose), polysaccharide lyases or proteins with cellulose-binding motif 1 (CBM1) were identified (Table S6). Enzymes that attack amorphous cellulose, hemicellulose, and pectin (e.g. β-glucosidases and cellobiose dehydrogenases of family GH3, GH10 xylanases, and GH28 pectinases) were absent. Genes encoding carbohydrate esterases and secreted feruloyl esterase which could hydrolyze cross-links in PCW were also not expressed. However, genes encoding GH5 endoglucanases (which act on mannann and/or cellulose) and a GH9 cellobiohydrolase were identified (Table S6). As observed in ectomycorrhizal symbionts (Martin et al., 2008b, 2010; Nagendran et al., 2009), the minimal set of PCW-degrading enzymes in the transcriptome of G. intraradices might be an evolutionary adaptation to avoid the release of polysaccharide fragments and their detection by the host immune system during the obligate biotrophic phase of the fungus. By contrast, a large set of genes encoding enzymes involved in chitin metabolism were expressed, including chitin synthases, chitin deacetylases, and chitinases (Table S6). They probably play a role in remodeling the fungal cell wall during growth and symbiosis. The abundance of transcripts coding for dolichyl-related mannosylation enzymes suggests an intense biosynthesis of cell wall mannans and mannoproteins.

Nutrient assimilation

We identified numerous transcripts coding for different predicted proteins associated with the uptake and assimilation of major soil nutrients, such as nitrate and Pi, and conversion of metabolites shuttled between partners (amino acids and carbohydrates) (Table S7). Although a few genes involved in nutrient transport and assimilation have previously been identified in G. intraradices and Glomus mosseae (Benedetto et al., 2005; Govindarajulu et al., 2005; Javot et al., 2007; Tian et al., 2010), there is no comprehensive molecular description of N and P assimilation pathways, despite the crucial importance of metabolite exchange in this symbiosis (Kiers et al., 2011; Smith & Smith, 2011). The striking induction of several genes coding for metal transporters suggests that metals, such as Zn, are important for plant colonization, and the expression pattern of genes coding for assimilative enzymes fully supports rapid uptake, translocation and transfer of metabolites, as discussed in the following sections.

Nitrogen metabolism In the G. intraradices-colonized M. truncatula roots, transcripts for transporters and enzymes involved in N uptake and assimilation showed high constitutive expression, supporting the contention that a high N turnover is taking place in the symbiosis (Gomez et al., 2009; Guether et al.,

2011). The set of differentially expressed G. intraradices orphan genes included the effector protein SP7, which interacts with the pathogenesis-related transcription factor Ethylene Response Factor (ERF19) in the host nucleus (Klopholz et al., 2011) (Table S4). Given the key results obtained for SP7, the role played by other mycorrhiza-up-regulated SSPs of G. intraradices should be elucidated, as well as the identity of plant-based signals that may control their expression within the root space.
Following reciprocal by Kiers and Phosphate metabolism *Glomus intraradices* expressed a wide spectrum of secreted phosphatase transcripts, including those coding for the *p*-nitrophenylphosphatase Pho13p, acid phosphatase Pho3p, repressible alkaline phosphatase Pho8p, and magnesium-dependent phosphatase, able to act on various phosphate esters (Table S9). No phytase transcript was detected. *Glomus intraradices* has a combination of low-affinity and high-affinity transporter genes, including the high-affinity Pi transporter gene *GintPT* (equivalent to *Pho84p* from *Saccharomyces cerevisiae*), and genes encoding the low-affinity Pi transporter Pho91p and the Na+/Pi symporter Pho89p (Table S9). The transcripts coding for the interactor protein Pho88p which promotes maturation and trafficking of Pho84p in yeast have also been identified. Pi transporter genes are expressed in spores, ERM and IRM. The expression pattern of *GintPT* is in agreement with the results obtained on *GmosPT* in *G. mosseae* (Benedetto et al., 2005) and the observed expression of GmosPT in arbuscules (Balestrini et al., 2007; Gómez-Ariza et al., 2009). Although high net transfer of P from soil to the plant takes place via the mycobiont (Kiers et al., 2011; Smith & Smith, 2011), our findings suggest that the IRM may re-absorb some of the Pi released in the symbiotic apoplastic space. As shown by Kiers et al. (2011), the fungus might exert some control on the delivery of nutrients to its host plant, and reciprocal rewards stabilize cooperation in the mycorrhizal symbiosis.

Following Pi uptake, PolyP accumulates in *G. intraradices* hyphae and is stored in the vacuolar compartments, where it buffers cytoplasmic Pi concentrations. PolyP is translocated along hyphae toward the interfacial apoplast (Viereck et al., 2004; Javot et al., 2007; Hijiikata et al., 2010). The rapid synthesis of PolyP is thus crucial for the maintenance of effective hyphal Pi uptake (Tani et al., 2009). Transcripts coding for the PolyP polymerase/Vacuolar Transporter Chaperone Complex Vtc4p protein involved in the synthesis and transfer of PolyP to the vacuole (Howorth et al., 2009), and for its accessory proteins, such as Vtc1p, were expressed in *G. intraradices* (Table S9). Transcripts for vacuolar endopolyphosphatase Ppn1p and the exopolyphosphatase Ppx1p were also identified. The endopolyphosphatase transcript was strikingly up-regulated in IRM. Several components of the complex phosphate regulatory system (PHO regulon) found in *S. cerevisiae* and *N. crassa* were expressed in *G. intraradices*, including the ankyrin-repeat-containing NUC2-related protein (Pho81p), the cyclin-dependent protein kinase regulator Pho80p and the mitogen-activated serine/threonine Cdc2 (Cell Cycle Control 2) cyclin-dependent protein kinase Pho85p (Table S9), suggesting putative regulatory similarities.

**Lipid metabolism** The profuse development of fungal membranes associated with arbuscule development in host cells dramatically increases the need for plasma membrane fatty acids and sterols. The oligoarray analysis identified transcripts in IRM coding for the subunits α and β of the fatty acid synthase complex,
together with the acetyl-CoA carboxylase transcript (Table S10), confirming that the mycobiont does possess fatty acid synthetic capacities (Table S10). Similarly, mycorrhiza-induced genes predicted to be involved in fatty acid metabolism are up-regulated in *M. truncatula* (Gomez et al., 2009), suggesting a complementary regulation of fungal and plant lipid metabolism. In this host plant, the gene *MtMSBP1*, encoding a membrane steroid (progesterone)-binding protein, is also induced early by a diffusible AM fungal signal produced by *G. intraradices* branched hyphae (Kuhn et al., 2010). The high activity of enzymes involved in sterol and steroid metabolism in IRM, together with the AM fungal induction of *MtMSBP1*, might be related to the need to alter sterol metabolism (e.g. lanosterol) to allow plasma membrane invagination and intracellular accommodation of the fungal symbiont in the cortical cells.

### An expanded inventory of conserved meiotic genes provides evidence for cryptic sex

The lack of an observed sexual stage in any member of the Glomeromycota led to the suggestion, which has been debated, that AM fungi were ancient aseous. Results from Croll & Sanders (2009) strongly suggest that recombination occurred among some *G. intraradices* genotypes in the field, although the majority of populations examined so far have been clonal (Rosendahl, 2008). As Glomeromycota could represent one of the earliest diverging fungal lineages, their meiotic processes could represent an ancestral state. We surveyed the transcriptome of *G. intraradices* for a set of sex and meiotic genes conserved among eukaryotes (Malik et al., 2008) (Table S11) and identified several ‘meiosis-specific’ genes (*HOP2* (Homologous-pairing protein 2) and *MND1* (Meiotic nuclear division protein 1)) which are only known to function in meiosis in other eukaryotes (Malik et al., 2008). These genes are hypothesized to be present in organisms with sexual ancestry (Table S11). The homolog of a transcript coding for the key meiotic recombinase *SPO11* (REC12) was not found, but has been identified in the genomic sequences of *Glomus diaphanum* (MUCL 43196), *G. intraradices* (DAOM 197198), *Glomus clarum* (DAOM 234281) and *Glomus cerebriforme* (DAOM 227022) (Halary et al., 2011). Transcripts coding for high-mobility group (HMG) domain-containing transcriptional factors with a significant similarity (55%) to the *sexP* and *sexM* genes from *Phycomyces blakesleeanus* (Idnurmet et al., 2008) were also retrieved. These genes are master switches controlling mating type in fungi. Mechanisms controlling the *G. intraradices* sexual cycle need to be further examined within AM fungal populations as they are likely to allow the mixing of nuclei and subsequent recombination among different individuals of this fungus interacting in soil.

In summary, induced expression of genes coding for membrane transporters and SSPs during the symbiotic interaction and the lack of expression of hydrolytic enzymes acting on PCW polysaccharides are hallmarks of *G. intraradices*. These results extend conserved patterns of gene expression profiles observed in obligate biotrophic pathogens (Spanu et al., 2010; Kemen et al., 2011) and ectomycorrhizal symbionts (Plett & Martin, 2011) to the Glomeromycota lineage. By contrast, obligate biotrophy in *G. intraradices* is not associated with a striking reduction of metabolic complexity (e.g. lack of N and S assimilation pathways), as observed in many obligate biotrophic pathogens, so that the ability to interact with the soil environment with respect to nutrient uptake is maintained in the symbiotic fungus. Finally, we can hypothesize that biotrophy in AM fungi has evolved through a series of steps requiring effectors, such as the secreted SP7 (Kloppholz et al., 2011), coupled with a reduced inventory of PCW-hydrolyzing enzymes to suppress or attenuate host defense reactions, and weak selection forces to maintain certain biosynthetic pathways if products (e.g. thiamine and sucrose) can be directly obtained from the host. The present comprehensive repertoire of *G. intraradices* genes, the first for Glomeromycota, provides a basis for future research in environmental genomics and for accessing symbiosis-related functional features in other members of this unique phylum.
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References


