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Symbiosis with an endobacterium increases the fitness of a mycorrhizal fungus, raising its bioenergetic potential

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Running title: Endobacterial impact on the AM fungus fitness
Abstract

Arbuscular Mycorrhizal Fungi (AMF) occur in the rhizosphere and in plant tissues as obligate symbionts, playing key roles in plant evolution and nutrition. AMF possess endobacteria, and genome sequencing of the endobacterium *Candidatus Glomeribacter gigasporarum* revealed a reduced genome and a dependence on the fungal host. To understand the effect of bacteria on fungal fitness, we used next-generation sequencing to analyse the transcriptional profile of *Gigaspora margarita* in the presence and in the absence of its endobacterium. Genomic data on AMF are limited; therefore, we first generated a gene catalogue for *G. margarita*. Transcriptome analysis revealed that the endobacterium has a stronger effect on the pre-symbiotic phase of the fungus. Coupling transcriptomics with cell biology and physiological approaches, we demonstrate that the bacterium increases the fungal sporification success, raises the fungal bioenergetic capacity, increasing ATP production, and eliciting mechanisms to detoxify reactive oxygen species. By using TAT peptide to translocate the bioluminescent calcium reporter aequorin, we demonstrated that the line with endobacteria had a lower basal intracellular calcium concentration than the cured line. Lastly, the bacteria seem to enhance the fungal responsiveness to strigolactones, the plant molecules that AMF perceive as branching factors. Although the endobacterium exacts a nutritional cost on the AMF, endobacterial symbiosis improves the fungal ecological fitness by priming mitochondrial metabolic pathways and giving the AMF more tools to face environmental stresses. Thus, we hypothesize that, as described for the human microbiota, endobacteria may increase AMF innate immunity.

Keywords: Arbuscular Mycorrhizal Fungi/ Endosymbiotic bacteria/ Mitochondria/ Transcriptome profiling
Introduction

Arbuscular mycorrhizal fungi (AMF) are crucial drivers of plant evolution: as the most widespread component of the plant microbiota, they occupy a double niche, thriving in the soil (pre-symbiotic phase) and inside root tissues (symbiotic phase). In both niches, they play a key role in nutrient cycling and plant health by taking up minerals such as phosphorous, and delivering them to their host plants (Bonfante & Genre, 2010). Notwithstanding their massive presence in the soil, AMF live as obligate biotrophs, which require organic carbon from their host plant. The spores and hyphae of AMF contain thousands of nuclei, making classical genetic approaches unsuitable. Also, many AMF contain endobacteria in their cytoplasm, leading to a further, unexpected increase in their genetic complexity (Bonfante & Anca, 2009). One type of AMF endosymbiont is the rod-shaped, Gram-negative beta-proteobacterium (Bonfante et al., 1994) Candidatus Glomeribacter gigasporarum (CaGg), which symbioses only with members of the Gigasporaceae family (Bianciotto et al., 2003; Mondo et al., 2012). The CaGg genome sequence (Ghignone et al., 2012) revealed that Glomeribacter endobacteria are nutritionally dependent on the fungal host (Ghignone et al., 2012); however, their contribution to host fitness remains unclear. Removal of CaGg from the host AMF causes limited changes in spore morphology and no evident impact on mycorrhization (Lumini et al., 2007). The CaGg endobacteria have small genomes, physiological dependence on their hosts, and vertical transmission, suggesting that these endobacteria live as mutualistic associates of the AMF. These features convincingly reveal that the endobacterium depends on the fungus, but do not explain why the hosts fungus maintains the endobacterium, despite its energetic cost (Ghignone et al., 2012).

Here, we used next-generation sequencing to produce a genome-wide transcriptional profile of Gigaspora margarita in the presence and in the absence of its endobacterium. This analysis, based on our de novo assembly of the transcriptome of G. margarita, showed large-scale changes in gene
expression related to the presence of the endobacterium and revealed effects mostly targeting the mitochadrion. Our results, confirmed by cell biology approaches and physiological measurements, indicate that the endobacterium increases the environmental fitness of the fungus, raising its bioenergetic capacity and potentially acting as a driver for priming the fungal innate immune response.

Materials and methods

Biological material

Fungal isolate and spore production

Spores of Gigaspora margarita Becker and Hall (BEG 34, deposited at the European Bank of Glomeromycota) containing (B+) or not (B-) the CaGg endobacteria were used in this study. B- spores were obtained from B+ spores as described in Lumini et al., 2007. The absence of endobacteria in the B- line was routinely checked following the protocol described in Salvioli et al., 2008. All the spores were maintained and propagated by using white clover (Trifolium repens) as trap plant. Briefly, clover plants were inoculated with 100-150 spores and after 3 months new spores were generated and collected by the wet sieving technique (Gerdemann and Nicolson, 1963). To monitor spore production, soil samples from selected pots were sampled 3-4 times each year, and the spores were collected and counted.

Spore germination and mycorrhization

Spores were divided in batches of 100, surface sterilized with Chloramine T (3% P/V) and streptomycin sulphate (0.03% P/V), some batches were germinated in 1 ml of sterile distilled water for 10 days in the dark at 30°C (germinating spores), while others were germinated in 1 ml of sterile distilled water for 3 days in the dark at 30°C and in a solution 10^{-7} M of the synthetic strigolactone
analogue GR24 (Chiralix, The Netherlands) for 7 more days (SL-treated). After 10 days, the
germinated spores and their germinating mycelium were collected, immediately frozen in liquid
nitrogen, and crushed with a pestle and mortar for further RNA extraction.

600 B+ and 600 B- spores were used to produce mycorrhizal seedlings of *Lotus japonicus* (Regel)
K. Larsen by using the “Millipore sandwich” method (Novero *et al.*, 2002). After 4 weeks, *L.
*japonicus* roots were observed under a stereomicroscope and extraradical mycelium and
mycorrhizal roots were sampled and frozen in liquid nitrogen for RNA extraction (symbiotic stage).

*Treatment with oxidant agent and strigolactones*

Sterilized spores (B+ and B-) were placed in a multi-well plate (30 spores in each well) and treated
with different concentrations of H$_2$O$_2$ (100 mM, 10 mM, 2 mM, 1 mM 0.75 mM, 0.5 mM, 0.3 mM
and 0.25 mM), GR24 (10$^{-7}$ M) or sterile distilled water. Spores were observed under a
stereomicroscope after 3 days of treatment at 30°C to check the germination rate. For each
treatment, at least 90 spores belonging to 3 different wells were observed. To understand whether
the oxidant agent can also lead to early transcriptional changes, a new set of sterilized spores were
treated with H$_2$O$_2$ 0.3 mM, GR24 (10$^{-7}$M), or sterile distilled water for 3 days at 30°C, frozen in
liquid nitrogen, and used for RNA extraction. The extracted material was processed as described
above.

*Molecular analyses*

*RNA extraction and sample preparation for sequencing*

Total RNA was extracted using the RNeasy Microarray Tissue Mini Kit (Qiagen, Germany). The
concentration and quality of the nucleic acids were assessed with a Nanodrop1000 (Thermo
Scientific, Wilmington, NC, USA), and the integrity was checked with the Bioanalyzer instrument
(Agilent Technologies, Santa Clara, CA, USA). For details, see Supplemental Text.
Real time q-PCR assays

For RT-qPCR validation, RNA was extracted as previously described and treated with the TURBO DNA-free kit (Life Technologies, Carlsbad, CA, USA). The samples were then reverse transcribed using Superscript II Reverse Transcriptase (Life Technologies, Carlsbad, CA, USA). Quantitative real time PCR experiments and data analysis were carried out as described in Salvioli et al, 2012, using as a reference gene for transcript normalization the G. margarita elongation factor (Tef). The primer names and corresponding sequences are listed in Table S10.

Generation of data, bioinformatics and phylogenetic analyses

Generation of Data Sets 1 and 2 and de novo transcriptome assembly

In the absence of a reference genome, a de novo assembly was generated using reads from four in vitro normalized paired-end libraries (Data Set 1, see below) obtained from the B+ line of G. margarita containing the endobacterium and sampled at four stages of the fungal life cycle (quiescent spores, germinating spores, spores treated with strigolactone, and extraradical mycelium), without replicates, and 14 single-end libraries (Data Set 2, see below) obtained from both the B+ strain and the cured line (B- line) sampled at three stages of the fungal life cycle (germinating spores, spores treated with strigolactone, and symbiotic mycelium thriving inside the roots. In total, 18 libraries were produced (Table S11). Data Set pre-process is described in Supplementary Materials and methods. The de novo assembly of Data Set 1 and 2 libraries was performed on a 60 core and 256 GB RAM machine, running Ubuntu server 12.04 LTS, using Trinity v.Trinityrnaseq_r20131110 (Grabherr et al., 2011). Detailed description of the assembly process is provided in Supplementary Materials and methods.
The *G. margarita* BEG34 Transcriptome Shotgun Assembly project (Bioproject PRJNA267628; Biosamples SAMN03216569-SAMN03216586) has been deposited at DDBJ/EMBL/GenBank under the accession GBYF00000000. The version described in this paper is the first version, GBYF01000000. Downstream analyses performed on the assembled transcripts are detailed in Supplementary Materials and methods.

**Calling differentially expressed genes**

For DEG identification, DESeq2 1.2.8 Bioconductor package was run with local fit nd betaPrior parameter set to TRUE. Independent filtering was enabled (Anders & Huber, 2010; Love et al., 2013). A False Discovery Rate (FDR) of 0.05 was set as threshold for DEG calling. The number of DEGs for each contrast is reported in Figure S4 According to current standards for RNA-seq (ref: Encode project standards), at least two biological replicates for each condition were used. Sample clustering (Figure S11;) was performed on rlog-transformed data (Love et al., 2013).

**GO enrichment analyses and KEGG maps**

GO enrichment analyses were conducted with the goseq bioconductor package version 1.14.0, while KEGG pathway pictures, KO (KEGG Orthology) mappings were first obtained from KAAS (KEGG Automatic Annotation Server; [http://www.genome.jp/kegg/kaas/](http://www.genome.jp/kegg/kaas/)) using as query trinity-assembled Gigaspora transcripts against the KEGG GENES database. Details are provided in the Supplemental material.

**Phylogenetic analysis**
A phylogenetic tree based on the comparison of whole proteomes of *G. margarita* and various fungal genomes was constructed using CVTree v3 using the default parameters (Xu & Hao, 2009). Fungal proteomes were those available as built-in fungal database proteomes in CVTree. Additional proteomes not available in CVtree databases were added for the analysis, namely *Tuber melanosporum* version ASM15164v1.21 (retrieved from ftp://ftp.ensemblgenomes.org/), *Rhizophagus irregularis*, and *Mucor circinelloides* (retrieved from NCBI queries).

Other bioinformatic techniques

Unless otherwise stated, further graphical outputs were generated with scripts as available in DESEQ2 package vignette (Love *et al.*, 2013).

Cellular and physiological analyses

Confocal and ultrastructural analysis

Single *G. margarita* spores from the B+ and B- lines were placed on microscope slides in 20 μl of Bacteria Counting Kit component A (B-7277; Molecular Probes) diluted 1:1,000 according to the manufacturer’s directions. The Bacteria Counting Kit contains the SYTO BC bacterial stain which is a high-affinity nucleic acid stain that easily penetrates both gram-positive and gram-negative bacteria. The spores were then crushed with a coverslip, incubated in the dark for at least 5 min, and observed under a Leica TCSSP2 confocal microscope (excitation 488 nm; emission 520 nm) to detect the endobacteria. To perform ultrastructural observations, parallel sets of single spores were processed by high-pressure freezing followed by freeze substitution, as described in Desirò *et al.*, 2014. After cutting and counterstaining, thin sections were observed under a transmission electron microscope Philips CM10.
Mitochondria staining with MitoTracker Green FM.

Mitochondria were stained with the fluorescent probe MitoTracker Green (Life Technologies). Five sterilized spores were placed on a microscope slide along with 50 μl of MitoTracker Green 1 μM and propidium iodide (50 μg/ml), spores were crushed with a coverslip and observed with a Leica TCSSP2 confocal microscope (excitation 490 nm; emission 515 nm) after 10 min of incubation in the staining solution. At least 20 spores were observed for the B+ and B- lines. Images taken at the same magnification were used to measure the mitochondrial diameter: 89 mitochondria belonging to 8 B+ spores and 76 mitochondria belonging to 7 B- spores were evaluated. Data were subjected to statistical analysis using the Kruskal-Wallis test for nonparametric data.

TAT-aequorin-based Ca²⁺ measurements

Ca²⁺ measurements were carried out in germinated spores of G. margarita (samples of 200 spores, germinated for 10 d) of both B+ and B- lines as described (Moscatiello et al., 2014).

ATP detection assay

Total levels of ATP were determined by using a Luminescent ATP Detection Assay kit (Abcam) according to the manufacturer’s instructions. The ATP assay is based on the production of light caused by the reaction of ATP with added luciferase and D-luciferin. The emitted light is proportional to the cellular ATP concentration. Luminescence was determined in 3 d germinating spores of G. margarita (B+ and B- lines) and total amounts of cellular ATP were calculated by using an ATP standard dilution series.

Phosphate measurements in roots and shoots from mycorrhizal clover plants

The P contents were determined as described in Supplemental materials.
Results and Discussion

To perform a comprehensive analysis of the effect of CaGg on the transcriptome of Gigaspora margarita, in the absence of a reference genome for G. margarita, we first created a de novo assembly of the G. margarita transcriptome. To this end, we developed a two-step strategy, where high-throughput sequencing of cDNA (RNA-seq) from the strain of G. margarita containing the endobacterium (B+ line) was performed, creating Data Set 1. We used these data to generate a preliminary, de novo assembly of the G. margarita transcriptome. We then conducted a second RNA-seq experiment to compare the B+ line and the cured line (B- line) at three stages of the fungal life cycle (germinating spores, spores treated with strigolactone, and symbiotic mycelium thriving inside roots, Figure S1), creating Data Set 2. These two sets of data were used to generate the final de novo assembly.

The Gigaspora transcriptome

Analysis of Data Sets 1 and 2 with the Trinity program created a combined de novo assembly comprising 86,183 transcripts (isoforms), with length greater than 350 bp, corresponding to 35,029 total potential genes. This gene count is in line with the about 28,000 genes reported for R. irregularis, the closest relative of G. margarita so far sequenced (Lin et al., 2014; Tisserant et al., 2013). The G. margarita transcripts had an average GC content of 31.96%, a median contig length of 781 bp, and an average contig length of 1185.77 bp; the contig N50 based on all transcripts was 1,683 bp.

Only 10,936 contigs (12.7%) were successfully annotated with Blast2GO (E-value filter= 1.0E-6; Annot_cutoff=55; GO_weight=5; HSP-Hit_Cov_cutoff=0), and BLAST searches identified 4,904 contigs (5.7%) that matched only sequences in the refseq_protein database (cut-off e-value of 1e-5), but were not further annotated.
The top BLAST hits included genes from species sharing similarities with *G. margarita*, such as *Laccaria bicolor* (Basidiomycota) and *Tuber melanosporum* (Ascomycota), two fungi with symbiotic lifestyles (Martin & Selosse, 2008; Martin et al., 2010), along with saprotrophic or pathogenic fungi such as *Schizophyllum commune*, *Coprinopsis cinerea*, *Postia placenta*, *Cryptococcus neoformans*, and *Ustilago maydis* (Figure S2). The complete collection of *G. margarita* and *Rhizophagus irregularis* transcripts was used to search for their Best Reciprocal Hits (BRH). We found that the two organisms share 6,276 transcripts (5,901 with annotation; Table S1) and *G. margarita* expressed most of the transcripts described as key features of the *R. irregularis* genome. These transcripts include phosphate transporters, saccharide transporters, chitinases, chitin synthases, and HMG-box transcription factors putatively involved in mating-type recognition (see Table S2 for a comprehensive list). AM fungi are considered to be asexual, but the huge number of mating type genes in *R. irregularis* (Tisserant et al., 2013) and in *G. margarita* opens the question of their biological meaning. Consistent with the analysis of the *R. irregularis* genome, the *G. margarita* transcriptome has cobalamin-dependent and cobalamin-independent methionine synthases, both sharing very high identity with the homologous genes in the *R. irregularis* genome. Both types of methionine synthases are expressed at comparably high levels during the presymbiotic stage of fungal development. Since *CaGg* possesses the operon for synthesis of vitamin B12, the finding that *G. margarita* produces a transcript for a putative lysosomal cobalamin transporter potentially involved in vitamin B12 import, is of particular interest. Also in agreement with one of the key features of *R. irregularis* genome, we found no plant-cell wall degrading enzymes among the *G. margarita* transcripts. Pfam analysis identified 3,838 domains (Table S3). As observed in *Rhizophagus* (Tisserant et al., 2013), *G. margarita* showed an overrepresentation of proteins involved in signaling pathways and ubiquitin-related metabolism. Interestingly, for both fungi, the most abundant domains occurred in genes encoding tyrosine kinases and Sel1 (see
Supplementary data). In addition, we also identified 177 transcripts that belong to CaGg, on the basis of their sequences (Ghignone et al., 2012).

Thus, this first glimpse at the G. margarita transcript-repertoire reveals that Rhizophagus irregularis and G. margarita (notwithstanding their deep differences in phylogeny, life cycle, and ecological strategies) have a strict genetic relatedness, as shown by a cladogram based on the comparison of the whole proteome of G. margarita with other fungal genomes (Figure S3).

Candidatus Glomeribacter gigasporarum affects the G. margarita transcriptomic profile most during the fungal presymbiotic phase

Due to the complexity of the experimental design (two fungal lines investigated at three stages, Figure S1), we first used DESeq2 to identify differentially expressed genes (DEGs). As a validation test, we randomly selected some DEGs to check by RT-qPCR (Table S4) and found that the RT-qPCR confirmed the differential expression detected by RNA-seq. We then compared the numbers of DEGs in different conditions to understand whether the main driver of gene expression in G. margarita depends on the life cycle stage or on the presence/absence of the endobacterium (see also supplemental data). We identified the most DEGs (9,609) when comparing the germinating spores containing the bacterium (B+) with the germinating spores in the cured B- line (G condition). Comparison of the germinating B+ spores with B- spores treated with the synthetic strigolactone GR24 (SL condition) identified 3,427 differentially expressed transcripts. We found very few transcripts that were differentially expressed during the symbiotic phase (Figure S4); this could be due to the fungal dilution in the root tissues and/or to a limited effect of the bacteria on the fungus during the symbiosis. This second option is consistent with our quantification of mycorrhizal colonization in clover, which found no difference between the B+ and B- lines (Figure S5), as previously shown for sorghum and carrot (Lumini et al., 2007).
The presence of \textit{CaGg} inside the fungal cytoplasm seems to be crucial for AMF function during the presymbiotic phase, when the fungus develops in the rhizosphere, outside the protection offered by the plant cell. Confocal and electron microscopy suggest that bacteria are diluted along the intraradical symbiotic mycelium (Bianciotto \textit{et al.}, 1996), while spores (Figure 1) act as a reservoir of these microbial communities (Desirò \textit{et al.}, 2014). This result opens the question of what advantages the endobacterium provides to its fungal host during this phase.

As a second step, we used the DEGs identified by DESeq to estimate Gene Ontology (GO) term enrichment, with GOseq (Young \textit{et al.}, 2010), using a threshold FDR=0.1. Consistent with the presence of more DEGs in the G condition, the GOseq analysis yielded 48 and 5 enriched GO terms in the G and in the SL conditions, respectively. For the G condition, the most relevant enriched GO terms deal with membrane processes and transport, regulation by phosphorylation, signal transduction, and oxidoreductase activity (Figure 2). By contrast, in the SL condition, only the ATP binding and protein phosphorylation terms were particularly relevant. These data further suggest that the endobacterial presence exerts a more relevant influence on the fungus at the germinating spore stage. However, specific changes also occur in the SL condition, indicating that the bacterial presence and the strigolactone treatment could interact in influencing the \textit{G. margarita} transcript profile.

\textit{Candidatus Gromeriibacter gigasporarum affects the expression of genes involved in growth, development, and transport in its fungal host}

Among the enriched GO categories in the G condition, the presence of the endobacterium affected membrane processes, and in particular chitin metabolism (Table S5). Several differentially expressed chitin synthase transcripts were identified, and five of them were upregulated in the B+ fungus in the G condition, while only one putative chitinase was downregulated. Also, several
transcripts related to putative chitin deacetylases, i.e. the enzymes that catalyze the deacetylation of chitin in chitosan, were downregulated.

In addition to the remodelling of the fungal wall, the endobacterium affects the expression of genes containing the mating type domains (Table S1 and S2), raising the question whether these genes could be involved in the production of the asexual conidia, as reported for *Penicillium chrysogenum* (Bohm *et al.*, 2013). To test whether the presence of the bacteria affected spore formation, we monitored the spore production of *G. margarita* over three years (Table S6) and found that the cured line produced only the 50% of the spores produced by the *G. margarita* line containing the bacteria.

The presence of the endobacterium also affected genes in the transport category (Figure 3a). For example, a plasma membrane iron permease containing an FTR1 domain was among the most-upregulated genes in the B+ line, in both the G and SL conditions, as also confirmed by RT-qPCR validation assays (Table S4). By contrast, genes related to nitrogen transport were downregulated in the B+ line; in particular, eight putative ammonium transporters were downregulated (Figure 3a), Phosphate uptake is a crucial trait of AM fungi and, accordingly, many transcripts identified in the *G. margarita* transcriptome encode different phosphate transporters (Table S7). Among them, we identified a full-length sequence coding for a phosphate:H symporter belonging to the major facilitator superfamily. This sequence shares the highest similarity with the phosphate transporter described in the extraradical mycelium of *Glomus intraradices* (Maldonado-Mendoza *et al.*, 2001). Another major facilitator superfamily high affinity P transporter (comp36913_c0), previously described in *G. margarita* and present in GenBank (GI:591140015), showed the highest expression and upregulation. The closest characterized relative of such sequence is Pho84, which has been demonstrated to function as a transceptor in yeast (Popova *et al.*, 2010).
These results open new questions on the potential effect of \textit{CaGg} on phosphorous transport from the fungus to the host plant. The endobacterium receives phosphate from its fungal host (Ghignone \textit{et al.}, 2012); the resulting stronger Pi-gradient probably causes more-sustained Pi uptake by the fungus from the soil, and more phosphate flow towards both the bacterium and the plant. To understand whether this had a positive or negative impact on the plant, we measured the phosphate in roots and shoots of clover plants colonized by \textit{G. margarita} and found that the plants colonized by the B+ line contained significantly more Pi than the plants colonized by the B- cured line (Figure 3b).

Taken as a whole, the impact of the endobacterium on the fungal transcriptome during the presymbiotic phase leads to functional changes that increase the fungal ecological fitness and may also have deep consequences for the third partner in the interaction, i.e. the plant.

\textit{Candidatus} Glomeribacter gigasporarum affects transcription of the fungal mitochondrial genes and interferes with the strigolactone response

To better describe the effect of the endobacterial presence on the fungal presymbiotic phase, we assigned KO (KEGG Orthology) terms to the transcript datasets through the KAAS server and then used the Pathview tool to map the pathways (Luo & Brouwer, 2013). This allowed us to illustrate the pathways represented in the fungal transcriptome, and to examine the effects of the endobacterium.

The most impressive changes in pathways in the fungal host involved oxidative phosphorylation, which was strongly induced by the presence of endobacteria (Figure 4). In particular, mitochondrial genes (Pelin \textit{et al.}, 2012) such as those encoding NADH dehydrogenase and Cytochrome oxidase I, were upregulated (Table 1). The transcriptome data therefore point to increased ATP production in the presence of the endobacteria. To validate this observation, we used a luminescence assay to quantify cellular ATP concentrations. Indeed, the cured line of \textit{G. margarita} produced significantly
less ATP (71.18 ± 10.38%) when compared to the line containing the endobacterium (Student’s *t*-test, *p*<0.05). By contrast, the cured line showed enhanced pentose phosphate metabolism, suggesting that an alternative pathway produces reducing power in the absence of the endobacterium (Figure S6a).

Since complex I is responsible for mitochondrial proliferation, mitochondria were monitored by Mitotracker staining in the spores of *G. margarita* containing, or not containing, the endobacteria (Figure 5). The two conditions showed roundish mitochondria in similar numbers; however, mitochondria from the cured line, at 2.4 μm in diameter, were significantly larger than those from the B+ line, at 1.7 μm in diameter (Figure S7). Transmission electron microscopy showed that the mitochondrial cristae were flattened in both lines, but the matrix of mitochondria in the cured line was more electron-transparent. Interestingly, in Alzheimer’s disease patients, where mitochondrial dysfunction has an early and preponderant role in the disease, oxidative injury may disturb the structure and fission of mitochondria, resulting in enlarged mitochondria (Moreira *et al.*, 2010). The results suggest that the absence of the endobacterium leads to changes that affect mitochondrial morphology.

The impact of the endobacterium on fungal mitochondrial genes was further enhanced in the SL condition (Table 1). Strigolactones stimulate mitochondrial activity in the germinating spores of the AMF *Gigaspora rosea*, which does not contain endobacteria (Besserer *et al.*, 2006). Our analysis surprisingly reveals that the endobacterium and the strigolactone treatment lead to a similar transcriptomic response under the tested conditions (seven days after germination), i.e. upregulation of the genes encoding NADH dehydrogenase and Cytochrome oxidase I (COX 1) in the fungus.

According to the concept of the dual effects of mitochondria (Moreira *et al.*, 2010), we wondered whether the higher mitochondrial activity might, on the one hand, lead to increased production of
ATP and reactive oxygen species (ROS) or, on the other hand, might give the fungus the capacity to better face oxidative stresses.

**Gigaspora margarita and its obligate endobacterium both express ROS-scavenger genes**

To test the hypothesis of crosstalk between the fungal oxidative response, the presence of the bacterium, and the treatment with the strigolactone analogue GR24, the transcriptome data were screened to identify genes involved in the detoxification of ROS. Some of these ROS-related genes were differentially expressed (Figure 6 and Table S8). In detail, thioredoxin reductase, peroxiredoxins, and glutathione peroxidase, which are ubiquitous molecules involved in ROS detoxification, were upregulated by the presence of the endobacterium. Selenocompound metabolism, which is known to be involved in antioxidant functions, was also upregulated in both the G and SL conditions (Figure S6b). Another class of ROS-detoxifying molecules, the superoxide dismutases (SOD), was also affected by the endobacterial presence: two copper-zinc SODs were upregulated in the G condition, and a copper-zinc SOD already characterized in *G. margarita*, and reported as upregulated during spore germination (Lanfranco *et al.*, 2005) was upregulated by the presence of the endobacterium in the G condition. Interestingly, the copper-zinc SOD was also detected as one of the most expressed proteins in a preliminary analysis comparing the fungal proteome in the presence and absence of the endobacterium (Salvioli *et al.*, 2010). Also, RT-qPCR validated the constitutive expression of some of these respiratory and ROS-related genes (Table S4), confirming their higher basal expression in the presence of the endobacterium. Interestingly a screen of the bacterial genome (Ghignone *et al.*, 2012) also revealed ROS-related genes, including a copper-Zn SOD and a thioredoxin peroxidase, in addition to the genes involved in oxidative phosphorylation (Table S9).

In conclusion, the availability of the *G. margarita* transcriptome, its mitochondrial genome, and the *CaGg* genome allowed us to show that the AM fungus with its endobacterium has more constitutive
tools to face oxidative stress, expressing a double set of ROS-scavenger genes. Irrespective of the presence of the endobacterium, GR24 treatment enhanced both the respiratory activities and the ROS-related responses, suggesting that the fungus perceives strigolactone as a xenobiotic, and that its effects mostly target the mitochondrion.

The fungal-bacterial association enables a rapid transcriptional response to oxidative stress and strigolactone treatment

To examine the hypothesis that fungi with the endobacteria react to oxidative stress more actively than the fungi without the endobacteria, we treated the B+ and B- fungal lines with different concentrations of the ROS hydrogen peroxide (H₂O₂). To understand whether this ROS causes rapid transcriptional changes, and whether strigolactone can mimic these stress-induced changes, we used RT-qPCR to measure the transcript levels of the mitochondrial and ROS-related genes that we identified as DEGs in the RNAseq experiment (Figure 7). Assays were performed on RNA extracted from B+ and B- spores treated with 0.3 mM H₂O₂ or GR24, after 3 hours of incubation. A set of control spores, treated only with water and collected at the same timepoint, was also included. The control experiment confirmed the trend detected in the RNA-seq data, i.e. that fungal genes encoding NADH dehydrogenase and COX 1 were expressed to higher levels in the B+ line than in the cured line, as well as glutathione peroxidase, which detoxifies H₂O₂, and Cu/Zn SOD, which transforms reactive oxygen species into H₂O₂ (blue columns, Figure 7). Hydrogen peroxide treatment stimulated the expression of some of the ROS-related transcripts in both lines (Figure S8). However, the fold ratio between the two lines points to a higher expression in the B+ spores, with a significant value for thioredoxin reductase (orange columns, Figure 7). These transcriptomic results suggest that the B+ line is constitutively more equipped to face oxidative stress conditions.

In agreement with previous observations (Besserer et al., 2006), the 3-hr treatment with GR24 increased the transcription of genes encoding NADH Dehydrogenase and COX 1 in both the lines
(Figure S8), with a significantly higher fold-change ratio in the B+ line for COX1 (yellow column, Figure 7). By contrast, the results were less clear-cut when ROS-related genes were considered: only the transcript of glutathione peroxidase, which uses H₂O₂ as a substrate, was strongly upregulated in the B+ spore, when compared to the cured line. On the other hand, the Cu/Zn SOD transcripts were higher after the GR24 treatment in the cured line (Figure S8), suggesting that, irrespective of the bacterium, strigolactone affects fungal ROS homeostasis.

To check whether the endobacterium directly perceives oxidative stress and strigolactone, we measured the expression of some genes involved in ROS-detoxification (Figure S9), under the same conditions of brief H₂O₂/GR24 treatment. While no significant changes were detected following GR24 treatment, an expression decrease was observed after the H₂O₂ ROS treatment, in comparison with the germinating spores, probably due to the antimicrobial effects of the H₂O₂. This result demonstrates that the bacterium itself does not cooperate with the fungus in ROS detoxification after H₂O₂/GR24 treatment, but its presence and its constitutively expressed ROS-related genes contribute to the ROS homeostasis of *G. margarita*. A regulated synthesis of ROS plays a role not only in plant communication (Gilroy S, 2014), but also in fungal morphogenesis, growth, and development, as suggested for fungi interacting with plants or animals (Abbà *et al.*, 2009; Mu *et al.*, 2014; Ryder *et al.*, 2013).

**The presence of the endobacterium affects the basal intracellular calcium concentration**

One of the most investigated aspects of the AM-plant interaction is the calcium-mediated signaling pathway, which is elicited in the plant host by molecules from the AMF, by the symbiotic pathway (Genre *et al.*, 2013; Oldroyd, 2013). To examine intracellular calcium, we used the TAT peptide, which acts as a potent nanocarrier to translocate macromolecules into living cells, to deliver the bioluminescent calcium reporter aequorin inside the AMF. Previous work with this system demonstrated that *G. margarita* responds to environmental stresses, as well as symbiotic signals,
with transient changes in the intracellular calcium concentration ([Ca\(^{2+}\)](i)) (Moscatiello et al., 2014).

We hypothesized that the B+ and B- lines of *G. margarita* perceive environmental and/or symbiotic signals differently. Using TAT-aequorin, we monitored [Ca\(^{2+}\)](i) in germinating spores under different conditions (basal status, cold shock, and GR24 treatment), using the [Ca\(^{2+}\)](i) as a fast readout of the fungal response. The cured line in the three conditions showed an unexpected behaviour: the basal [Ca\(^{2+}\)](i) was significantly higher than that in the B+ fungus (Figure 8), suggesting that calcium constitutively accumulates in the cured line. Cold shock thus elicited a comparatively weaker response in the cured line: the calcium concentration started from a higher level, caused a lower peak, and then remained higher than in the B+ line throughout the experiment.

Interestingly, GR24 is considered a symbiotic molecule, and GR24 treatment elicited a calcium signature comparable to that induced by an environmental stress condition, causing a fast and transient [Ca\(^{2+}\)](i) increase in the B+ line (Moscatiello et al., 2014). However, as observed for the cold shock, the cured line showed a weaker response, since the amplitude of the change in [Ca\(^{2+}\)](i), calculated as the difference between the peak and the resting level (Δ[Ca\(^{2+}\)](i)), appeared reduced.

Calcium has a crucial role in the establishment of symbiotic events: plants perceive microbial signals from rhizobia and AMF, activating oscillations in calcium concentration, which persist from minutes to hours (Oldroyd, 2013). Rhizobia and AMF also perceive plant signals, eliciting a transient calcium elevation (Moscatiello et al., 2010; Moscatiello et al., 2014). The relevance of calcium signalling as a mediator of extra- and intracellular stimuli is well described for many filamentous fungi (Bencina et al., 2005), and for *Rhizophagus* (Liu et al., 2013).

A permanent increase in intracellular calcium has generally been considered as harmful for the cell (Zhivotovsky & Orrenius, 2011). We hypothesize that the absence of the bacterium affects calcium signaling, leading to an increase in the basal [Ca\(^{2+}\)](i), which negatively affects fungal cellular metabolism, possibly by interfering with ATP synthesis. Indeed, the low solubility of Ca\(^{2+}\) with phosphates may interfere with ATP-based metabolism, which is an essential feature of all living
Our experiments further indicate that the bacterium may interfere with the regulatory signaling network, which also includes ROS (Gilroy S, 2014); the endobacteria influence the intracellular calcium level in a still-unknown way, and potentially alter fungal ATP synthesis. It could be speculated that the endobacterium functions as an additional Ca\(^{2+}\) store in both hyphae and spores, thereby cooperating in fine-tuning the [Ca\(^{2+}\)] in the AMF. Irrespective of this, the TAT experiment also shows that GR24 causes a reaction in the fungus similar to the reaction to environmental stress.

**Conclusion**

In summary, here we used a combination of molecular, cellular, and physiological approaches to reveal that the obligate endobacterium CaGg affects the biology of the AMF *Gigaspora margarita*, and that its effect is most relevant during the presymbiotic phase, when the fungus, unprotected by its plant host, encounters the biotic and abiotic stimuli in the soil (Figure 9).

The changes we detected in the fungal transcriptional landscape suggest that the presence of the endobacterium tunes a huge number of metabolic pathways, including spore production, fungal wall remodeling, and mineral nutrient uptake and transport, also leading to an unexpected positive impact on the phosphate content of plant roots and shoots. However, among the many differentially expressed genes, the most interesting indicated that the mitochondrion is a primary target of the effects of the presence of the endobacteria. These effects enhance the positive and negative effects of the mitochondria: on the one hand, in the fungal line with its ancient endobacterium (Mondo *et al.*, 2012), activation of respiration leads to a higher ATP production. On the other hand, this line also shows stronger activation of genes involved in ROS generation and detoxification than the cured line. Since the endobacterium also constitutively activates respiratory and ROS-scavenger genes, we suggest that the AMF hosting the endobacterium has twice the tools to face environmental stresses. Thus, we hypothesize that, as described for the human microbiota (Chu &
Mazmanian, 2013), the intracellular endobacteria may prime the level of innate immunity in the AMF. However, differently from other eukaryotes, experimental data exploring the existence of innate immunity in fungi do not seem to be available. The control of the intracellular concentration of calcium could be one of the key processes, first regulating the bioenergetic status of the fungus. Lastly, the comparison of the fungal transcriptome after treatment with a synthetic strigolactone allowed us to reveal that both the endobacterium and strigolactone produced a comparable response in the fungal mitochondria, possibly because the prokaryote and GR24 (Besserer et al., 2006) have a similar effect on the expression of respiratory genes. We suggest that strigolactones, which are usually considered as plant symbiotic signals, are first perceived by the AMF as foreign molecules (xenobiotics), activating a fast and transient \([\text{Ca}^{2+}]_i\) increase, as well as elevated respiration, eventually leading to ROS production and detoxification. In the absence of the bacterium, and in line with the overall slower metabolism shown by the cured line, all these processes are attenuated.

Taken as a whole, our data indicate that the endobacterium improves the fitness of its fungal host during the pre-symbiotic rhizospheric phases. Unlike *Rhizophagus* (Hempel et al., 2007; Varela-Cervero et al., 2015), *Gigaspora* mostly produces spores, as pre-symbiotic propagules, which represent the exclusive structures for the success of colonization, and *Gigaspora* has taken advantage of the endobacterium. This symbiosis has proven beneficial to the AMF, from an evolutionary point of view, thus potentially offsetting its nutritional cost for the fungus.

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Conflict of Interest

The authors declare no conflict of interest

Supplementary information is available at the ISME Journal web site.


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Figure 1: A composite picture illustrating the location of Candidatus Glomeribacter Gigasporarum. a: G. margarita spores (S) observed under a stereomicroscope produce a network of germinating hyphae (Gh); b: a squashed spore reveals the fungal nuclei (N) and a multitude of endobacteria (arrows) after staining with Bacteria Counting Kit component A and observation by confocal microscopy. The red line is drawn to suggest the fungal wall; C: a bacterium observed by electron microscopy reveals the multilayered Gram negative wall; it is located inside the fungal cytoplasm (FC), limited by a membrane of fungal origin. Bars correspond to 230 μm in a, 13 μm in b and 0.35 μm in c.

Figure 2: Gene Ontology (GO) term enrichment as estimated for the comparison B+ versus B- cured germinating spores. The analysis was performed with the Goseq package (Bioconductor), using a threshold FDR= 0.1.

Figure 3: a: Graphical representation of the regulated transcripts referring to the transporter category, as identified in the comparison B+ vs B- under germinating (G) and strigolactone-treated (SL) conditions. Each square corresponds to a regulated transcript. b: Phosphorous mineral content in clover plants (roots and shoots) colonized by B+ and B- G. margarita lines. Data are mean ± SD of three biological replicates. Statistically supported differences are indicated with different letters according to a Kruskal-Wallis non parametric test at p<0.05.

Figure 4: Top: KEGG-based depiction of mitochondrial oxidative phosphorylation in G. margarita. Functions supported by upregulated or non-regulated transcripts are shown in red or grey, respectively. Bottom: list of the transcripts related to mitochondrial oxidative phosphorylation as upregulated by the endobacterial presence under G and SL conditions.

Figure 5: Confocal (left) and electron microscopy (right) images illustrating the shape and morphology of mitochondria from B+ and cured (B-) G. margarita spores. Mitochondria
(arrow) were detected by staining with MitoTracker Green, and fungal nuclei (N) with propidium iodide. The inset on the right column reveals the cristae organization and the differences in the matrix. Bars correspond to 5 μm in a, c and their insets, to 0.7 μm in b and to 0.16 μm in the inset, and to 0.7 μm in d and to 0.4 μm in the inset.

**Figure 6: Clustering and heatmap analysis of ROS-related genes.** DESeq2-normalized and rlog-transformed expression data for differentially expressed ROS-related genes in germinating spores and strigolactone-treated spores (both B+ and B-): samples were clustered with the heatmap.2 function in the gplots 2.14.2 Bioconductor package. Pink and yellow bars at the top of the heatmap were added to mark, respectively, the B+ and B- major tree branches as generated by clustering algorithm.

**Figure 7: Relative quantification of gene expression as obtained for a set of mitochondrial and ROS-related genes.** Fold change is calculated for each gene in the B+ versus B- line in germinating, H2O2- and strigolactone-treated spores with the basal expression recorded in the B- condition (threshold line at Fold change=1). Statistically significant data (Kruskal-Wallis non parametric test, p<0.05) are marked with an asterisk.

**Figure 8: TAT-aequorin-based Ca2+ measurements in Gigaspora margarita germinated spores.** Ca2+ assays were performed in germinated fungal spores (10 d) after 1 h incubation with 30 μM TAT-aequorin. Intracellular free Ca2+ concentration was monitored in resting conditions (A) and after treatment (arrow) with a cold shock (B) or strigolactone GR24 (10^-6 M) (C). The reported traces represent typical observed responses (n=3). Black trace, B+ line. Grey trace, cured line.

**Figure 9: Schematic summary of the mechanisms by which the endobacterium affects Gigaspora margarita metabolism.** Starting from the left, the drawing illustrates how in the presence of the endobacterium, the fungus Gigaspora margarita upregulates genes involved in respiration (1), leading to higher ATP production (2). This status is mirrored by morphological
changes in the mitochondrial ultrastructure (3). Thanks to the upregulation of SOD, H$_2$O$_2$ is
produced and further detoxified to water by ROS-scavenger enzymes, like glutathione peroxidase (4). Intracellular calcium homeostasis is efficiently maintained (5). Treatment with GR24, a synthetic strigolactone induces some transcriptional responses that mimic the endobacterium presence, suggesting that there is interference between the microbe presence and the plant hormone treatment (6). The oxidative burst is correlated with higher spore production in many fungi: this feature is indeed present in _G. margarita_, which contains endobacteria (7). Since _G. margarita_ exclusively colonizes plants starting from spores (8), this feature represents an evolutionary benefit (9). Red arrows, transcriptome results; green arrows, data from biochemical, cellular and physiological experiments; interrupted arrow, hypotheses based on the literature.
Figure 1

A composite picture illustrating the location of G. gilvum. (a) G. gilvum spores (S) observed under a stereomicroscope produce a network of germ tubes (T); (b) a squashed spore reveals the fungal nature (C) and a multitude of endobacteria (Bacteria) after staining with basic fuchsin. The arrow is drawn to suggest the fungal wall; (c) a bacterium observed by electron microscopy reveals the multilayered Gram-negative wall. It is located inside the fungal cytoplasma (FC), limited by a membrane of fungal origin, that correspond to 0.5 μm in a, 0.3 μm in b and 0.35 μm in c.

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Gene Ontology (GO) term enrichment as estimated for the comparison S+ versus S− cured germinating spores. The analysis was performed with the Scoreq package (Bioconductor), using a threshold false discovery rate=0.1.
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(a) Graphical representation of the regulated transcripts, referring to the transporter category, as described in the comparison G vs SL under experiment (G) and SL-treated (SL) conditions. Each square corresponds to a regulated transcript. (b) Phosphorus mineral content in dollar plants (roots and shoots) normalized to the end of -c (marginal) lines. Data are projections of three biological replicates. Statistically supported differences are indicated with the same letters according to a Kruskal-Wallis non-parametric test at P<0.05.

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Figure

TAT-sequeirin-based Ca$^{2+}$ measurements in $O$. manganea germinated spores. Ca$^{2+}$ assays were performed in germinated fungal spores (10 days) after 1h incubation with 30μM TAT-sequeirin. Intracellular free Ca$^{2+}$ concentration was monitored in resting conditions (a) and after treatment (arrow) with a cold shock (b) or SL GR24 (10$^{-6}$ M) (c). The reported traces represent typical observed responses (n=3). Black trace, B+ line; Grey trace, cured line.
Schematic summary of the mechanisms by which the endobacterium affects *G. marginata* metabolism. Starting from the left on the top, the drawing illustrates how in the presence of the endobacterium, the fungus *G. marginata* upregulates genes involved in respiration, leading to higher ATP production. This status is mirrored by morphological changes in the mitochondrial ultrastructure. Thanks to the upregulation of SOD, H$_2$O$_2$ is produced and further detoxified to water by ROS-scavenging enzymes, like glutathione peroxidase. Intracellular calcium homeostasis is efficiently maintained. Treatment with G20A, a synthetic SL, induces some transcriptional responses that mimic the endobacterium presence, suggesting that there is interference between the microbe presence and the plant hormone treatment. The oxidative burst is correlated with higher spore production in many fungi; this feature is indeed present in *G. marginata*, which contains endobacteria. As *G. marginata* exclusively colonizes plants starting from spores, this feature represents an evolutionary benefit. Red font, transcriptome results; green font, data from biochemical, cellular and physiological experiments; interrupted line, hypotheses based on the literature. A full colour version of this figure is available at The ISME Journal Online.