

1 **The genome of *Geosiphon pyriformis* reveals ancestral traits linked to the**
2 **emergence of the arbuscular mycorrhizal symbiosis**

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17 **Abstract**

18 Arbuscular mycorrhizal fungi (AMF, sub-phylum Glomeromycotina) are prominent root
19 symbionts of land plants. This symbiotic association, named arbuscular mycorrhizal symbiosis
20 (AMS), allows the fungus to receive photosynthetically fixed carbohydrates from the plant in
21 exchange of improved nutrients and water uptake by the roots. AMS is considered to have played
22 a key role in the colonization of the land by plants, but how this symbiotic ability emerged is not
23 entirely understood due to lack of genomes from AMF representatives of basal phylogenetic
24 nodes. To address this, we sequenced the genome of the AMF *Geosiphon pyriformis*, a species
25 involved in the only known fungal endosymbiosis with a cyanobacterium. This species is
26 unknown to undergo AMS, yet we find that its genome carries all hallmarks of AMF obligate
27 plant biotrophy, including a reduced set of plant cell wall degrading enzymes and an absence of

1 genes for the production of sugars and fatty acids. Using comparative genomics and available
2 RNA-seq data from a model species, we identify a set of genes lost in the MRCA of
3 Glomeromycotina, and a putative core gene set differentially regulated during AMS. Overall, our
4 findings indicate that the mechanisms involved in AMS appeared prior to the emergence of
5 Glomeromycotina. These also provide a basis for future research on symbiotic mechanisms in
6 prominent plant mutualists and primary insights into a fungus-cyanobacteria endosymbiosis.

7

1 **Introduction**

2 Within the continuum of interactions between organisms, from parasitism to mutualism, the
3 symbiosis formed by plants and arbuscular mycorrhizal fungi (AMF) (subphylum
4 Glomeromycotina) [1] is amongst the most prominent [2]. Over 70% of known land plant
5 species have the potential to form this symbiosis (AMS) [3]. AMS plays a major role in extant
6 terrestrial ecosystems as it allows plants to efficiently acquire poorly soluble soil nutrients [4].
7 During AMS, the fungal partner colonizes the plant roots (or the thalli of root-less plants), and
8 forms small tree-like structures called arbuscules within the plant cell that promote bidirectional
9 nutrient transfers between partners. Thereby, AMF receives photosynthetically fixed
10 carbohydrates from the plant and, in return, the fungus delivers phosphate, nitrogen and other
11 trace elements as well as water to its host. This plant-fungus symbiosis dates back more than 400
12 million years [5], and is considered as one of the key innovations that allowed the colonization of
13 lands by plants [6] .

14 Genomic and genetic analyses conducted in diverse plant species have started to shed light on
15 the molecular mechanisms that allowed the evolution of this symbiosis on the host side [7,8]. By
16 contrast, how and when AMS abilities emerged in AMF remains elusive. Comparative
17 phylogenomics could be used to understand the evolution AMS [7,8], however, such approaches
18 require the availability of genome data covering the diversity of AMF lineages. Based on
19 ribosomal RNA (rRNA) phylogenies, AMF are separated into four orders, namely
20 Diversisporales, Glomerales, Archaeosporales and Paraglomerales. To date, only genomes from
21 Diversisporales and Glomerales have been sequenced, while only fragmentary protein coding
22 datasets are available for basal phylogenetic nodes that include the Archaeosporales (*Ambispora*
23 *leptoticha*) and Paraglomerales (*Paraglomus occultum*) [9].

24 Among the Archaeosporales, *Geosiphon pyriformis* stands out. This species is the only fungus
25 known to produce endosymbiosis with nitrogen-fixing cyanobacteria (*Nostoc punctiforme*), and
26 it was proposed that this association represent the AMF ancestral state [10–12]. As opposed to
27 other AMF, which penetrate the plant cells to form arbuscules, *G. pyriformis* forms long fungal
28 cells (‘bladders’) that enclose cyanobacteria. Once in the bladder, the cyanobacterium is
29 photosynthetically active and fixes nitrogen, and in exchange the fungus delivers inorganic
30 nutrients and provides water. The bladder membrane, which encloses the cyanobacteria cells, is

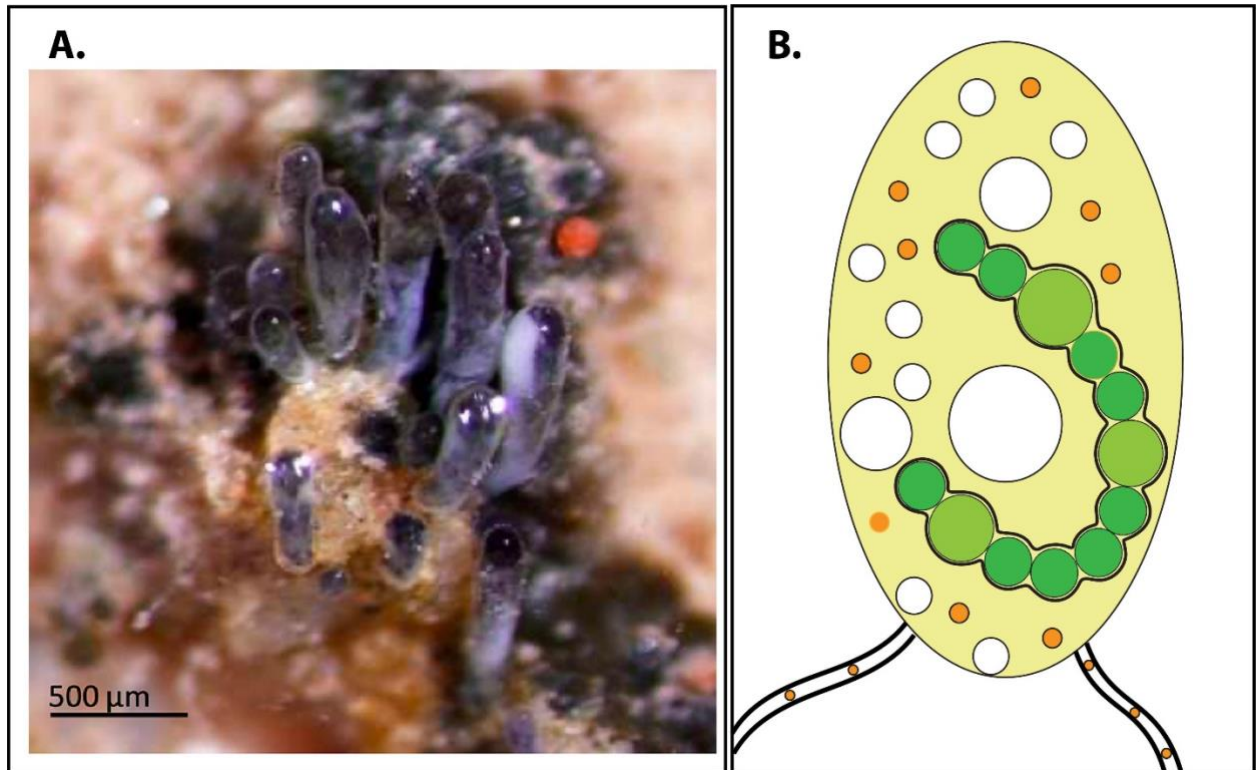
1 an extension of the fungal plasma membrane [14]. The first AMF monosaccharide transporter
2 (GpMST1) was found using the *G. pyriformis*-*N. punctiforme* endosymbiosis system [15,16],
3 and despite almost four decades of research on this species *G. pyriformis* was never found to
4 form intracellular symbiotic structures such as arbuscules, [15,17,18].

5 Arguably, *G. pyriformis* represents an ideal candidate to investigate the origin of AMS and the
6 emergence of a unique endosymbiosis. Here, we aimed to advance knowledge in these questions
7 by sequencing the genome of *G. pyriformis* using a re-discovered isolate.

8 **Results**

9 *General genome characteristics of Geosiphon pyriformis*

10 The only known culture of *G. pyriformis* was lost over a decade ago (A. Schüßler, pers. comm).
11 In an attempt to rediscover *G. pyriformis*, we searched and identified symbiotic bladders of the
12 *G. pyriformis*-*N. punctiforme* symbiosis (Figure 1) at the only known stable habitat of this
13 species in the Spessart Mountains near the village of Bieber in Germany [19].



14

15

1 **Figure 1: a.** Image of *G. pyriformis* bladders in soil from its natural habitat in the Spessart
2 mountains. **b.** Schematic representation of *G. pyriformis* bladders containing *Nostoc* cells (based
3 on M. Kluge 2002). *Nostoc* cells are shown in dark green, and heterocysts (differentiated cell that
4 carries out nitrogen fixation) are shown in light green. Bladders contain *G. pyriformis* nuclei
5 (orange), and several vacuoles (white). Aseptate hyphae reach out and extend from the bladders.

6 Upon cultivation, we extracted DNA and RNA from active bladders. Total DNA was subjected to
7 5Kb-mate-pair and 125 bp paired-end Illumina sequencing, producing respectively 47 and 81
8 million 125 bp paired ends and 5kb mate-pairs reads. *G. pyriformis* reads were identified using a
9 read binning approach recently implemented to assemble the genome of *Diversispora epigaea*
10 *epigaea*[16], and upon identification these were assembled into a 129 MB assembly and 795
11 scaffolds with an average read coverage of 118X. In parallel, total RNA was subjected to 150 bp
12 paired-end Illumina sequencing. The resulting RNA-seq reads were mapped onto the *G. pyriformis*
13 genome assembly using STAR [20] and used for genome annotation after implementing
14 RepeatMasker[18]. This procedure identified of 24195 genes in *G. pyriformis*, resulting in a
15 BUSCO gene repertoire completeness of 96.2% (3.1% complete duplicated). The gene counts,
16 estimated genome size, and genome statistics are all similar to those of model AMF species [13,21]
17 and are indicative of high genome completeness (Table 1, Supplementary Table 1 and
18 Supplementary Figure 1). SignalP showed that, among all the genes identified in *G. pyriformis*,
19 365 represent putative secreted proteins (Supplementary Table 2) and 27% of these are candidate
20 effectors (Supplementary Table 3) [10,11]. We also identified 19 putative secreted CAZymes in
21 *G. pyriformis*; in line with numbers found in other AMF species (Supplementary Table 4 and
22 Supplementary Table 5).

23 AMF genomes carry a substantial fraction of transposable elements (TE) [10,11,25,26], and we
24 found that *G. pyriformis* has undergone similar TE expansions. The expansion of Gypsy
25 transposable elements in *G. pyriformis* is evident in comparison to all other AMF genomes (Figure
26 2). With regards to TE, we find no evidence that *G. pyriformis* carries a two-speed genome
27 (Supplementary Figure 2) [29]. Two-speed genomes are characterized by the presence of TE-poor
28 and gene dense regions that a clearly separate from others that contain rapidly evolving genomic
29 regions that usually carry less genes, abundant TE, and other repeat elements [30,31].

30



4 **Figure 2** Bubble plot containing all transposable elements in the genomes of glomeromycotina
 5 and mucoromycotina genomes used in this study. The figure shows the expansion of Gypsy
 6 elements in *G. pyriformis*.

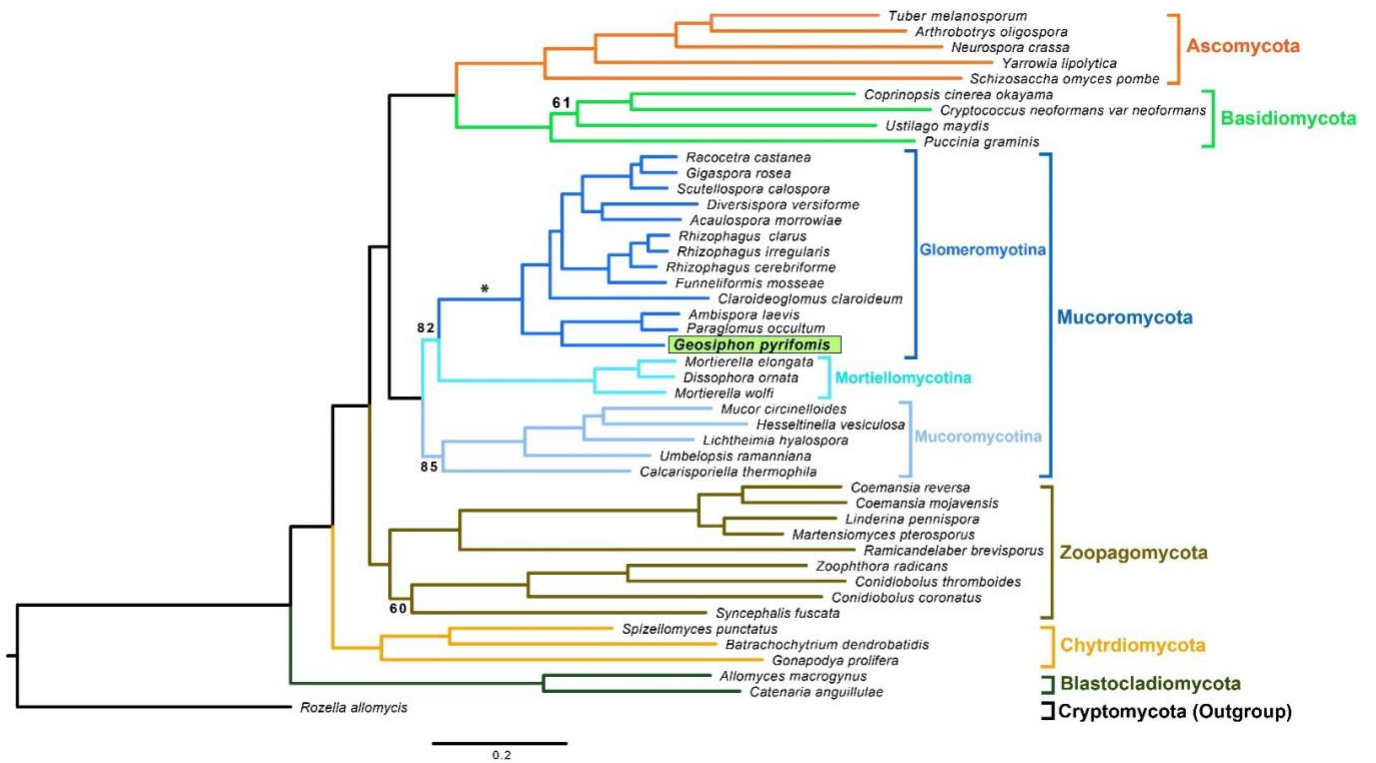
7 Using genome data and single nucleus data it was recently shown that AMF carry two genome
 8 organizations – i.e. homokaryotic (co-existing nuclei carry one parental haploid genotype) or
 9 dikaryotic (two parental genotypes co-exist in the mycelium) [26–29]. Mapping reads onto the *G.*
 10 *pyriformis* genome revealed reduced levels of polymorphism (0.5 SNP/Kb) and allelic frequencies
 11 suggesting that this species carries low nuclear diversity and likely homokaryotic
 12 (Supplementary Figure 3).

13 *Placement of G. pyriformis based on phylogenomics*

14 The *G. pyriformis* genome annotation was used to identify the phylogenetic placement of this
 15 species using amino-acid sequences. In this case, we used a set of 434 conserved fungal single
 16 copy genes (data available at DOI: 10.5281/zenodo.1413687) to construct a phylogenetic tree of
 17 the fungal kingdom. Phylogenomics supports the monophyly of Glomeromycotina and its close

1 relationship with Mortierellomycotina within the phylum Mucoromycota [23] (Figure 3). Within
 2 Glomeromycotina, *G. pyriformis* groups within a monophyletic clade with *Ambispora leptoticha*
 3 and *Paraglomus occultum*, which diverged around 287 MYA (Supplementary Figure 4). This
 4 clade is distinct from more diverged nodes that contain sequenced representatives from
 5 Glomerales and Diversisporales [22]. The current placement of *G. pyriformis* as a sister lineage
 6 to *A. leptoticha* and *P. occultum* has full statistical support, and is favored by 73% of the gene
 7 sequences we used. Alternative topologies that place, for example, *G. pyriformis* at a basal node
 8 to all AMF (Alt-T1; Supplemental Figure 5, Supplemental Table 6) or as being associated with
 9 Glomerales or Diversisporales (Alt-T2 and Alt-T3; Supplemental Figure 5, Supplemental Table
 10 6) were all rejected significantly using statistical tests implemented in IQ-TREE, including the
 11 KH, SH, ELW, and AU tests (Supplemental Table 6).

12 **Figure 3:** Phylogenetic tree representing the evolutionary relationships of fungi and placement



13 of *G. pyriformis* in Glomeromycotina clade. The tree was resolved using maximum likelihood
 14 phylogenetic reconstruction with IQ-TREE on a concatenated alignment of 434 protein coding
 15 genes. Numbers indicate nodes with less than 100% bootstrap support. Branches are coloured
 16 according to their phylum. Phylogenetic tests for all alternative topological placements of *G.*

1 *pyriformis* were rejected. The asterisk denotes the location where inferred gene losses and gains
2 occurred in the MRCA of Glomeromycotina.

3 ***The genome of Geosiphon pyriformis uncovers shared gene features in the Glomeromycotina***

4 Phylogenomics revealed that *G. pyriformis* is a member of a clade that diverged early from the
5 lineage encompassing the already sampled Diversisporales and Glomerales. As such, the *G.*
6 *pyriformis* genome fills the gap in the genomic coverage of major AMF phylogenetic clades. With
7 this data in hand, we first searched for genetic features that arose in the most recent common
8 ancestors (MRCA) of all Glomeromycotina by comparing orthogroups from five available AMF
9 genomes and four other members of the Mucoromycota as outgroups using OrthoFinder [39]. This
10 analysis identified 661 gains and 344 losses that occurred before the divergence of the
11 Glomeromycotina (Supplemental Table 7 and 8).

12 Among the 344 orthogroups classified as lost in the MRCA of the Glomeromycotina, we note the
13 missing key enzymes involved in essential metabolic functions, such as sugar and thiamine
14 metabolisms, or in the biosynthesis of fatty acids. These genes are also referred to as “Missing
15 Glomeromycotina Core Genes” (MGCGs; Supplementary Table 8, Supplementary Table 9), and
16 our analysis reveals that these have also been lost by *G. pyriformis*. Other key losses that affect all
17 sequenced Glomeromycotina include enzymes that actively degrade plant cell wall
18 (Supplementary Table 10). Among the 661 orthogroups gained in the MRCA, most encode for
19 proteins involved in signaling pathways (e.g. protein kinases), protein–protein interactions (e.g.
20 the tetratricopeptide repeat, Sell1, the homodimerization BTB (Broad-Complex, Tramtrack and
21 Bric a brac), and WD-40 domain-containing proteins) and High Mobility Box (HMG)
22 (Supplementary Table 11).

23 Comparative genomics also showed that *G. pyriformis* carries the same signatures of sexual
24 reproduction found in AMF relatives. These include a complete set of meiosis-specific genes
25 (Supplementary Table 12) [32], and a highly conserved genomic locus with architecture and
26 sequence similarity to the mating-type (MAT) locus of basidiomycetes [33,34] (Supplementary
27 Figure 6).

28

29

1 ***Regulation of orthogroups gained in the MRCA of Glomeromycotina***

2 We investigated available gene expression data from the model AMF *Rhizophagus irregularis*,
3 and found that 7 and 39 orthogroups (with a total of 8 and 272 genes; Supplementary Table 13
4 and Supplementary Table 14) gained in the MRCA of *Glomeromycotina* are respectively
5 upregulated and downregulated across all four experimental conditions in the model AMF
6 *Rhizophagus irregularis*.

7 These conditions include symbiotic associations between *R. irregularis* and distinct plant hosts
8 (*Medicago truncatula*, *Brachypodium distachyon*) [7,43], and others based on laser-capture
9 microdissection arbuscule-specific gene expression[34,35,37,39]. We investigated the putative
10 function of these differentially regulated genes by identifying protein motifs along their coding
11 sequences, and found that these are involved in a myriad putative cellular function, though these
12 mostly include protein tyrosine kinases, cytochrome p450, as well as FAD binding
13 (Supplemental Table 13 and 14). Lastly, one differentially regulated OG (OG0001728) shows
14 evidence of originating from horizontal gene transfer from bacteria – i.e. this orthogroup is
15 shared between bacteria and fungi (Supplementary Figure 7).

16 ***Distinct genomic features of Geosiphon pyriformis***

17 Besides the evolution of AMS, the acquisition of the *G. pyriformis* genome offers an opportunity
18 to identify innovations linked to the emergence of the only known cyanobacteria – fungus
19 endosymbiosis. To identify such innovations, hierarchical clustering and abundance of Pfam
20 domains was performed using available genomes in the Glomeromycotina and representatives of
21 Mucoromycotina and Mortierellomycotina (Supplementary Figure 8). This analysis revealed a
22 significant overrepresentation of 16 protein domains in *G. pyriformis* compared to relatives in the
23 Mucoromycota – e.g. Lipase_3, RNase_H, Retrotrans gag domains, dUTPase, Spuma_A9PTase,
24 Myb_DNA-bind_6 (Supplementary Table 15; Supplementary Table 16).

25 We also sought evidence of horizontal gene transfers (HGT) between partners of the unique
26 *Geosiphon-Nostoc* endosymbiosis, and found 18 genes with potential bacterial within in the *G.*
27 *pyriformis* genome (Supplementary Table 17). Among putative HGT, two are protein encoding
28 genes with significant sequence conservation with *Nostoc* and Gamma proteobacteria
29 homologues (Supplementary Figure 9, Supplementary Figure 10). All putative HGT are located

1 within contigs with average coverage and surrounded by genes of AMF origin, suggesting these
2 do not represent contaminants.

3 **Discussion**

4 ***MRCA of all extant Glomeromycotina carried hallmarks of mutualism and obligate biotrophy***

5 Genome data from a representative of the basal node of the AMF phylogeny filled an important
6 gap in understanding the origin of AMS. Specifically, it allowed us to conclude that the MRCA of
7 all extant Glomeromycotina carried hallmarks of mutualism and obligate biotrophy – i.e. a lack of
8 genes for fatty acids and thiamine biosynthesis and nutrition, and a reduced number of genes that
9 actively degrade plant cells. As such, the mechanisms involved in AMS appeared prior to the
10 emergence of Glomeromycotina and a represent a synapomorphy of this sub-phylum. *G.*
11 *pyriformis* has also conserved genomic signatures of sexual reproduction, as well as an apparent
12 low nuclear polymorphism. Both traits are thus conserved across Glomeromycotina and are in
13 stark contrast with the notion that these organisms represent an ancient asexual lineage.

14 The retention of a conserved Glomeromycotina gene set in *G. pyriformis*, including a sub-set of
15 these involved in plant cell wall degradation, is surprising given that this species was never seen
16 producing mycorrhizae. As such, this retention could reflect an intrinsic capacity for *G. pyriformis*
17 to undergo classic (but rare) mycorrhizal associations with plants under the right conditions.
18 Although speculative at this point, this hypothesis is supported by the identification of rare
19 *Geosiphon*-like sequences in environmental samples [26–28,36].

20 ***Novel symbiotic abilities and horizontal gene transfers in G. pyriformis***

21 As a result of losses in fatty acid biosynthesis genes, AMF are entirely dependent on the host plant
22 they associate with to obtain lipids [45,47,49–51]. Within this context, our findings suggest that,
23 during the switch from regular AMS to a fungal- cyanobacteria symbiosis, *G. pyriformis* has
24 evolved novel strategies to obtain lipids from its new host through the expansion of specific gene
25 motifs. Specifically, the *G. pyriformis* genome carries a striking over-representation of Lipase 3
26 protein domains that hydrolyze ester linkages of fatty acids. As *Nostoc* spp is known to produce a
27 wide variety of extracellular lipids in high amounts [52,53], it is possible that these abundant lipids
28 are released in the environment (like many other cellular compounds released by cyanobacteria

1 [43,45,46,48]) and are then broken down by lipases to be used as an energy resource by *G.*
2 *pyriformis*.

3 As we find evidence of bacteria-like genes in the *G. pyriformis* genome, our work also suggests
4 that the co-existence of multiple endosymbionts and *G. pyriformis* nuclei within restricted bladders
5 offers some opportunities for horizontal gene exchange. Although none of the putative bacterial
6 genes we identified in *G. pyriformis* are functionally related, there is evidence that one is
7 differentially regulated during AMF symbiosis, supporting the notion that bacteria-like genes can
8 play a major role in fungal evolution [58].

9 ***Identification of a putative AMF core-symbiotic toolkit***

10 The *G. pyriformis* genome also enabled the identification of a putative core AMF symbiotic
11 toolkit conserved in all the sampled Glomeromycotina. This set of genes is differentially
12 regulated in model AMF during symbiotic interactions with different plant hosts, including
13 dicots, monocots and non-vascular plants, and thus provides a basis for future research on
14 symbiosis-related mechanisms in these plant symbionts. The identification of a core set of gene
15 gains specifically regulated during mycorrhizal symbiosis, and their conservation across the
16 Glomeromycotina phylogeny, also provides support for the early emergence of symbiosis-
17 specific gene functions in AMF over 400 million years ago, contemporaneously with the
18 evolution of the first land plants [6,55,56]. Lastly as genetic transformation is currently
19 unfeasible in Glomeromycotina, only assumptions can be proposed for the function of these
20 putative core genes. However, as some encode for chitin synthases, one attractive hypothesis
21 could be that some evolved for the production of short-chain chitooligosaccharides or lipo-
22 chitololigosaccharides that are known symbiotic signals triggering the activation of the symbiotic
23 program on the host plant [46,48].

24 **Materials and Methods**

25 **Cultivation of *G. pyriformis* samples from natural habitat**

26 *G. pyriformis* was sampled during autumn in the only known stable habitat near the village
27 Bieber in the Spessart (Germany). Active bladders of the *Geosiphon-Nostoc* endosymbiosis were
28 found in slightly acidic soil (pH 5). The bladders occurred close to the hornwort *Anthoceros* spp.
29 and the liverwort *Blasia pusilla* L., as these plants harbor the cyanobacteria needed to trigger the

1 *Geosiphon-Nostoc* endosymbiosis. After sampling spores and bladders were transferred to the
2 institute in Průhonice and cultured in beakers [57], which contain a small pot with a sterile
3 mixture of sand and soil (from the original habitat). The cultures were grown in a climate
4 chamber at 18°C with 14 h light and 10 h night. The substrate is kept wet by a filter paper, which
5 reaches from the substrate into a water reservoir in the beaker. To be maintained over time,
6 cyanobacteria be frequently added to the cultures. For our cultures, *Nostoc punctiformis* was
7 obtained from the Culture Collection of Algae (SAG) at the University of Göttingen (Germany)
8 as strain SAG69.79 [60] [59].

9 ***Genome and transcriptome sequencing and assembly***

10 High quality DNA was extracted from active bladders of *G. pyriformis* and *Nostoc punctiforme*
11 using the NucleoSpinII Plant kit (Machery-Nagel) and purified with the genomic DNA clean-up
12 kit (Machery-Nagel) using the manufactures recommendations. Total DNA was sent to FASTERIS
13 (Switzerland) for library Illumina library preparation and sequencing using on 150 paired end and
14 5kb mate pairs inserts (illumina Nextera mate pair kit). Sequencing was performed using the
15 Illumina HiSeq 4000 platform. Total RNA was extracted using the RNeasy-Mini Kit (Qiagen) as
16 per instructions of the manufacturer for library RNA-seq Illumina library preparation with
17 sequencing of 150 cycles and paired ends.

18 Poor quality and adapter sequences were trimmed using Trimmomatic [61] with parameters the
19 following parameters of ILLUMINACLIP:2:30:10 SLIDINGWINDOW:5:20 LEADING:5
20 TRAILING:5 MINLEN:50. The resulting 1 GB of non-redundant metagenome reads were
21 assembled using metaSPAdes V3.12.0 [62]. Assembled contigs were binned on the basis of tetra
22 nucleotide signature using CONCOCT [63], following part of the procedure used to assembly the
23 genome of *Diversispora epigaea* [64]. Binned clusters were annotated using BLAST v 2.6.0+ [65]
24 and clusters containing bacterial hits were removed. Using this approach, 21 bona-fide AMF
25 clusters were retained, and used as reference to filter original paired-end and mate pair reads with
26 BLAT v. 36x1 [66]. The reads which had mapped to the filtered contigs and matched by BLAT
27 were then extracted to build cleaned sequence libraries that were assembled with MaSuRCA 3.3.0
28 [67]. Additional round of nr BLAST searches on MaSuRCA assembled contigs were performed
29 to further remove contaminating bacteria. K-mer (k = 21) based methods were used on filtered

1 reads to estimate genome size of *G. pyriformis* using jellyfish 1.1.12 and plotted in GenomeScope
2 2.0 [68,69].

3 ***Genome annotation***

4 Protein coding genes were predicted using Funannotate V1.7.4
5 (<https://funannotate.readthedocs.io/>) [CITE 10.5281/zenodo.3679386], which automates gene
6 prediction. Assembled transcripts using Trinity [64] and Rnaseq reads mapped bam file were used
7 as transcript evidence for gene call.

8 Transposable elements were predicted using TransposonPSI [81]. Repeat sequences were first
9 identified using RepeatModeler with multiple numbers of iterations. The iteration with the most
10 number of repeats were then used for soft-masking the genome with REPEATMASKER (open
11 4.0.646) [82]. Output files generated from above procedures were used to identify repeat along the
12 assembly. The completeness of genome assembly was assessed with BUSCO version 2.0 [83] with
13 default parameters using the fungal gene dataset [fungi_odb9]
14 [<http://buscodev.ezlab.org/datasets/fungiodb9.tar.gz>] with HMMER version 3.2.1 searches [84].

15 Putative gene functions were identified using Diamond BLASTX [85]. Pfam domain analysis
16 were performed using hmmscan [76] and Carbohydrate-active enzymes (CAZYme) were
17 identified using the dbCAN CAZy database [88]. Putative CAZymes were further verified
18 through comparisons of data from Morin et al 2019 [21]. Secretory proteins were identified using
19 previously published pipelines [21,89], and effectors were identified using EffectorP 2.0 [90]. The
20 putative MAT loci of *Paraglomus* sp., and *R. irregularis* were identified by BLAST search
21 procedures. The tests for a two-speed genome analysis was performed in part by measuring
22 intergenic distance among genes in genome using a R script (code available in GitHub repository)
23 [30]. For this study, published genomes of additional Glomeromycotina, Mortierellomycotina, and
24 Mucoromycotina were downloaded from JGI portal MycoCosm database [DOI:
25 10.1093/nar/gkt1183].

26 ***SNP calling and allele frequency analysis***

27 Filtered Mate Pair and Paired End reads were mapped onto the assembled *G. pyriformis* genome
28 using the BWA-MEM v 0.7.17 algorithm [91] and sorted into a BAM file using samtools (v 1.9)
29 [92]. Variants were called using FREEBAYES v1.2.0 [93] and filtered using vcftools [94].

1 Filtering cutoffs and procedures were as described by in Ropars et al. 2016 and Morin et al. 2019
2 [21,33]. Quality filtered variants and SNPs which passed filtering were used for constructing allele
3 frequency plot using a custom R script (code available in GitHub repository).

4 ***Phylogenetic analysis and molecular dating***

5 The phylogenomic analyses employed a set of 434 generally conserved and single-copy proteins
6 in fungi (data available at DOI: 10.5281/zenodo.1413687), which were developed through efforts
7 of the 1000 fungal genomes project and provided in the Joint Genome Institute MycoCosm site
8 [1,95,96]. Profile-Hidden-Markov-Models of these markers were searched in the *Geosiphon*
9 predicted protein sequences using HMMER3 (v3.1b2) and recovered 393 homologs (out of the
10 434) in total. The 434 markers in 45 included fungal genomes were further collapsed into 57
11 partitions using a greedy search embedded in PartitionFinder v.2.1.1 for consistent phylogenetic
12 signals [97]. Phylogenetic trees were produced using the PHYling pipeline (data available DOI:
13 10.5281/zenodo.1257002) and with maximum likelihood method implemented in IQ-TREE
14 (v.1.7-beta9) [98]. Concordance factors across the tree were calculated using the package
15 implemented in IQ-TREE.

16 The divergence time of *Geosiphon* sp. from the clade of "*Ambispora leptoticha* and *Paraglomus*
17 *occultum*" was estimated using the R8S v1.81 [99] with the phylogenetic tree reconstructed from
18 the earlier step. We employed five calibration constraints to calibrate the tree, including the
19 crown groups of Fungi (1100 MYA) [84], Dikarya (772 MYA) [84], Chytridiomycota (>573
20 MYA) [85–87], the MRCA of Chytridiomycota and terrestrial fungi (>750 MYA) [85–87], and
21 Glomeromycotina (>460 MYA) [88]. The divergence time of each clade was inferred using the
22 Langley-Fitch method with Powell algorithm [89–91].

23 **Alternative topology test and dating analyses**

24 To test the likelihood of other possible phylogenetic placements of *G. pyriformis*, we first
25 reconstructed the associated phylogenetic trees using constraint tree topology as illustrated in
26 Supplementary Figure 5 via “-g” option of the IQTREE package (iqtree-1.7-beta9) [92]. We then
27 compared our best tree (shown as Figure 3) with alternative topologies to compute the log-
28 likelihoods of the trees using Kishino-Hasegawa test, Shimodaira-Hasegawa test, expected
29 likelihood weight, and approximately unbiased test via “-zb” and “-au” parameters in IQTREE
30 [93–97]. All tests were performed with 10,000 resampling estimated log-likelihood (RELL)

1 method for reliable results. The best-fit substitution models for the genome-scale data matrix
2 were estimated using ModelFinder implemented in IQTREE package [98].

3 ***Detection of putative horizontal gene transfers***

4 To identify genes in *Geosiphon* that have potential origin in cyanobacteria, we compared the
5 *Geosiphon* sp. genome to the available fungal and cyanobacterial genomes. To highlight
6 potential HGT genes, we used a Python script (available in github repository) [105] to filter out
7 genome component in *G. pyriformis* with higher similarity score to cyanobacteria than any fungi,
8 excluding the *G. pyriformis* itself.

9 ***Gene orthology and evolution of symbiotic specific genes***

10 Orthogroups resulting from the OrthoFinder run were parsed using a custom Python script. To be
11 retained, an orthogroup had to fill the following conditions: any sequence from the non-AMS
12 fungi, at least one sequence of *Geosiphon pyriformis*, one sequence of either *Rhizophagus*
13 *irregularis* or *Rhizophagus cerebriforme* and at least one sequence of *Gigaspora rosea* or
14 *Diversispora epigaea*. Reciprocally, orthogroups that could correspond to gene losses in the AMS
15 fungi were extracted by retaining orthogroups with no sequences of AMS fungi and at least one
16 sequence of each non-AMF fungi.

17 Orthogroups showing evidence of regulation in symbiotic conditions in *R. irregularis* were
18 subjected to Maximum Likelihood (ML) analysis to check for the absence of non-AMS species.
19 First, proteins contained in orthogroups were searched against the nine proteomes of 9 distinct
20 species using the BLASTp+ v2.9.0 [109] with default parameters and an e-value threshold fixed
21 at 1e-05 (threshold was set to 1e-03 when no non-AMS species sequences were identified). Then,
22 proteins were aligned using MUSCLE v3.8.31 [110] with default parameters and resulting
23 alignment trimmed to remove positions with more than 80% of gaps using trimAl v1.4rev22 [111].
24 Prior to ML reconstruction, best fitting evolution model was tested using ModelFinder⁹⁸ and then
25 ML analysis was performed using IQ-TREE v1.6.1 [112] with 10,000 replicates of SH-aLRT.
26 Trees were visualized and annotated with the iTOL platform v5.5 [113].

27 After first round of phylogeny, orthogroups showing an AMF specific pattern were blasted
28 against the full MycoCosm database (1565 proteomes, last accessed: 03/01/2020) to confirm the

1 AMF-specific pattern and a phylogenetic analysis was performed following the procedure
2 described above.

3 ***Differential expression analysis and combination of expression data to orthogroups***

4 Expression data of *Rhizophagus irregularis* in four conditions were used to select orthogroups
5 containing gene significantly deregulated in symbiosis for further analysis. Paired-end reads
6 were trimmed and fragments mapped onto Rhiir2_1 genome assembly of *R. irregularis*
7 (https://genome.jgi.doe.gov/Rhiir2_1/). Stringent settings of mapping were used (similarity and
8 length read mapping criteria at 98% and 95%, respectively). Genes differentially expressed
9 (DEG) in planta compared to extraradical mycelium were identified after EdgeR [108]
10 normalization with a false discovery rate (FDR) correction using CLC Genomic Workbench
11 (Qiagen). We retained genes showing an expression > 2- or < -2-fold times in planta compared to
12 extraradical hyphae (FDR \leq 0.05). Sets of 2683, 2518 and 2410 DEG were found in *M.*
13 *truncatula*, *B. distachyon* and *L. cruciata* respectively. Detailed information on the data are
14 available at the National Center for Biotechnology Information (NCBI) Gene Expression
15 Omnibus (GEO) portal (accession no GSE67926). The analysis performed on RNA-seq data
16 from arbuscocytes in *M. truncatula* [44] presented 6359 DEG.

17 **Code availability**

18 All scripts used to analyze are archived in <https://github.com/madhubioinfo/Geosiphon>.

19 **Data availability**

20 Genome assembly is available in NCBI with accession number of JAAOMT000000000. Genome
21 sequencing reads are submitted in SRA with accession number of SRR11466073, Bioproject
22 PRJNA610605, Biosample SAMN14307302. RNA-seq reads are available in SRA with
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8

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1 **Table 1: Summary statistics for genome assembly of sequenced *G. pyriformis* and other**
 2 **species from Glomeromycotina and selected Mucoromycota used in this study**

Genomes	Assembly size	No of scaffolds	Scaffold N50	Largest scaffold (Kb)	Total Gap%	Repeat %	Busco completeness %	GC %
<i>Geosiphon Pyriformis</i>	129	795	703	2733.91	0.023	64.35	96.2	29.25
<i>Gigaspora rosea VI.0</i>	597.95	7526	734	1204.75	7.92	63.44	97.9	28.81
<i>Rhizophagus ceribriforme DAOM227022 VI.0</i>	136.89	2592	266	709.02	17.60	24.77	98.3	26.55
<i>Rhizophagus irregularis DAOM 197198V2.0</i>	136.80	1123	129	1375.86	5.06	26.38	98	27.53
<i>Diversispora versiformis strain IT104</i>	147	731	434	2010.39	0.061	43.6	98.2	25.1
<i>Rhizopus microsporus ATCC11559 VI</i>	25.97	131	8	2782.17	2.41	4.68	98.6	37.48

<i>Mucor</i> <i>Circinelloides</i> CBS 277.49 V2.0	36.59	26	4	6050.25	0.00	20.38	97.2	42.17
<i>Phycomyces</i> <i>Blakesleeanus</i> NRRL1555 V2.0	53.94	80	11	4452.46	1.06	9.74	96.9	35.78
<i>Mortierella</i> <i>Elongata</i> AG-77	49.86	473	31	1526.29	0.30	4.63	99.7	48.05

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2 **Supplementary Figure legends**

3 **Supplementary Figure 1:** Plot showing genome size of assembled genome. The x-axis
4 represents the frequency of the number of times a given k-mer is observed in filtered illumina
5 data of *Geosiphon pyriformis*. The y-axis represents the total number of k-mers with a given
6 frequency.

7 **Supplementary Figure 2:** Heatmap showing evidence for a “one-speed genome” in *Geosiphon*
8 *Pyriformis*. Density plot of the intergenic distances on the 5' end (x axis) against the 3' end (y
9 axis) for each gene on the genome. Distances for all genes are binned and showed in color scale
10 using a blue to red. red indicating the largest number of genes in a bin. Putative effectors are in
11 black dots and found in gene dense region. In case of two speed genome, genome is separated in
12 two compartment gene dense (containing essential housekeeping genes) and gene sparse region
13 (containing secretory or effector/virulence related genes). Putative effector candidates are found
14 in gene sparse repeat rich region.

1 **Supplementary Figure 3:** Genome wide allele frequency of *G. Pyriiformis* shows it is a
2 homokaryon. Allele distribution is based on read counts of filtered bi-allelic SNPs, and
3 overlapped by density curves (black). Blue vertical lines represent the 0.5 allele frequency.

4 **Supplementary Figure 4:** Divergence time estimation of the *Geosiphon* sp. Branches lengths
5 are proportional to time with the scale shown at the bottom. Estimated ages are labeled at each
6 node. The ultrametric tree was produced using the r8s v1.81 program with the maximum
7 likelihood tree obtained from the IQTREE program as the input.

8 **Supplementary Figure 5:** Alternative phylogenetic placements of *Geosiphon* examined in the
9 alternative topology tests. Three alternative topologies with the *Geosiphon* at the earliest
10 diverged position within Glomeromycotina (Alt_T1), grouped with
11 the *Rhizophagus* and *Funneliformis* clade (Alt_T2), and joined with
12 the *Gigaspora* and *Acaulospora* clade (Alt_T3) were compared to the best tree produced in the
13 present study given the genome-wide data set. The log-likelihoods of each of the four topologies
14 were computed using the IQTREE package and the results including the Kishino-Hasegawa test,
15 Shimodaira-Hasegawa test, expected likelihood weight, and approximately unbiased test were
16 summarized in Supplementary Table 6.

17 **Supplementary Figure 6:** Transcriptional directions of the putative AMF mating-type locus –
18 i.e. HD1-like and HD2 - and their flanking regions in *G. pyriiformis* and *R. irregularis*. The
19 putative MAT locus and two surrounding genes are displayed in figure. Based on sequences
20 initially identified by Ropars *et al.* 2016.

21 **Supplementary Figure 7:** Maximum Likelihood phylogeny of the OG0001728 obtained using
22 the model: VT+R8. Bacterial proteins from nr database are indicated with red branches, fungal
23 proteins from JGI database are indicated in orange branches. Sequences in blue constitute the
24 original orthogroups.

25 **Supplementary Figure 8:** Presence and abundance of the different Pfam domain-containing
26 proteins in the five genomes of Glomeromycota including *G. pyriiformis* and Mucoromycota
27 species. The heat map represents Pfam domain counts in each of the genomes. Only the 100 most
28 represented domains are shown here. For visualization, the pfam abundance values are
29 transformed into z -scores, which measures the relative enrichment (red) and depletion (green) of

1 each motif. The data were visualized and clustered using MultiExperiment Viewer
2 (<http://www.tm4.org/mev.html>).

3 **Supplementary Figure 9:** Maximum likelihood phylogeny using WAG model of selenium
4 binding protein from *G. pyriformis* showing HGT with other prokaryotes and eukaryotic
5 relatives. Bootstrap values are provided for each node.

6 **Supplementary Figure 10:** Maximum likelihood phylogeny using WAG model of Molybdenum
7 cofactor carrier protein from *G. pyriformis* showing HGT with homologues from prokaryotic and
8 eukaryotic relatives. Bootstrap values are provided for each node.

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