













Desert truffle genomes reveal their reproductive modes and new insights into plant–fungal interaction and ectendomycorrhizal lifestyle

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Summary

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- Desert truffles are edible hypogeous fungi forming ectendomycorrhizal symbiosis with plants of Cistaceae family. Knowledge about the reproductive modes of these fungi and the molecular mechanisms driving the ectendomycorrhizal interaction is lacking.
- Genomes of the highly appreciated edible desert truffles *Terfezia claveryi* Chatin and *Tirmania nivea* Trappe have been sequenced and compared with other Pezizomycetes. Transcriptomes of *T. claveryi* × *Helianthemum almeriense* mycorrhiza from well-watered and drought-stressed plants, when intracellular colonizations is promoted, were investigated.
- We have identified the fungal genes related to sexual reproduction in desert truffles and desert-truffles-specific genomic and secretomic features with respect to other Pezizomycetes, such as the expansion of a large set of gene families with unknown Pfam domains and a number of species or desert-truffle-specific small secreted proteins differentially regulated in symbiosis. A core set of plant genes, including carbohydrate, lipid-metabolism, and defence-related genes, differentially expressed in mycorrhiza under both conditions was found.
- Our results highlight the singularities of desert truffles with respect to other mycorrhizal fungi while providing a first glimpse on plant and fungal determinants involved in ecto to endo symbiotic switch that occurs in desert truffle under dry conditions.

Introduction

The so-called desert truffles are a group of edible hypogeous fungi that establish mycorrhizal symbiosis with annual and perennial shrubs belonging to Cistaceae, a plant family adapted to arid and semiarid areas (Kovács & Trappe, 2014; Roth-Bejerano *et al.*, 2014). These areas are characterized by an aridity index AI < 0.5—that is, the ratio between the annual precipitation and potential evapotranspiration (United Nations Educational, Scientific and Cultural Organization, 1979)—and by poorly fertile soils with sandy texture and low inputs of organic matter (Bonifacio & Morte, 2014). Under these environmental conditions, mutualistic symbionts promote growth and survival of their host plants. For example, *Terfezia claveryi* mycorrhizas promote *Helianthemum almeriense* survival during drought periods by altering its physiological and

nutritional parameters (Morte *et al.*, 2010), such as increased phosphorus, nitrogen (N), and potassium uptake (Morte *et al.*, 2000). As a result of this interaction, a fine-tune regulation of both plant and fungal aquaporin expression (Navarro-Ródenas *et al.*, 2013; Marqués-Gálvez *et al.*, 2020) and hydrogen peroxide content in roots (Marqués-Gálvez *et al.*, 2019) also occur. The type of mycorrhiza established by these fungi is known as an ectendomycorrhiza (EEM). EEMs are characterized by the co-occurrence of an intercellular Hartig net, intracellular hyphae penetrating the cortex cells, where they form coil-like structures, and a thin and disordered fungal mantle surrounding the colonized roots (Morte *et al.*, 1994; Yu *et al.*, 2001). The occurrence of either intercellular or intracellular mycorrhizal structures depends on several factors. *In vitro* conditions, high auxin, high phosphate, and/or high water content favour the intercellular mycorrhizal type, whereas field conditions, low auxin, low phosphate, and/or low water availability favour the intracellular mycorrhizal type (Gutiérrez *et al.*, 2003; Zaretsky *et al.*, 2006a; Navarro-Ródenas *et al.*, 2012, 2013).

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Among desert truffles, *Terfezia* and *Tirmania* ascocarps (fruiting bodies) are well-known delicacies and largely marketed mainly because of their taste and nutritional values (Kagan-Zur *et al.*, 2014). *Terfezia claveryi* (Honrubia *et al.*, 2002; Morte *et al.*, 2008) and *Terfezia boudieri* (Slama *et al.*, 2010; Kagan-Zur *et al.*, 2014), both associating with *Helianthemum* spp. as host plants, are the only ones that have been successfully cultivated, the former in Spain and the latter in Tunisia and Israel. Since most climate models point to increased temperature and lower precipitation rates for the next decades, areas covered by arid and semiarid ecosystems are expected to increase (Schlesinger *et al.*, 1990; Lavee *et al.*, 1998; Huang *et al.*, 2016). Thus, desert truffles are becoming a promising, alternative crop in regions such as the Mediterranean basin and the Middle East, where desertification will cause an alarming increase in dry lands (Morte *et al.*, 2017). *Helianthemum almeriense* Pau is a drought deciduous shrub that is well represented in the Mediterranean basin. This species establishes EEM symbiosis with several desert truffles in the wild, including *T. claveryi*. The symbiotic relationship between *H. almeriense* and *T. claveryi* has been widely used as a model for the research of desert truffle symbiosis at a basic and applied level—see Morte *et al.* (2017) for review. However, at the present time, the information on the molecular mechanics involved in this particular symbiosis is scarce.

Several studies have revealed the specific genomic signature of ectomycorrhizal, arbuscular mycorrhizal, or ericoid mycorrhizal fungi (Martin *et al.*, 2008, 2010; Tisserant *et al.*, 2013; Kohler *et al.*, 2015; Chen *et al.*, 2018; Martino *et al.*, 2018; Murat *et al.*, 2018; Morin *et al.*, 2019; Sun *et al.*, 2019; Venice *et al.*, 2020), but similar studies have not yet been carried out on EEM fungi. These genome and transcriptome analyses have identified hallmarks defining the mycorrhizal lifestyle. Abundant and highly upregulated specific effector-like small secreted proteins (SSPs) together with convergent losses of secreted carbohydrate-active enzymes (CAZymes) are two of the most recurrent traits associated with the mycorrhizal lifestyle (Kohler *et al.*, 2015). Although complementary DNA–amplified fragment length polymorphism analyses have been carried out to identify genes related to *T. boudieri* × *Cistus incanus* symbiosis (Zaretsky *et al.*, 2006b), and the genome of *T. boudieri* has been recently published (Murat *et al.*, 2018), many key aspects of the biology of EEM desert truffles remain to be elucidated. In addition, the impossibility to mate these fungi under controlled conditions has prevented mycologists shedding light into their reproductive modes (Roth-Bejerano *et al.*, 2004). This problem could be overcome by studying the structure and gene content of the mating type (*MAT*) locus, the master regulator of sexual reproduction in fungi. In the *MAT* locus of *Ascomycetes*, the *MAT1-1-1* and *MAT1-2-1* genes, which regulate the expression of sex-specific pheromone biosynthetic and signalling genes, are present (Debuchy *et al.*, 2010). The two major mating systems in these fungi are heterothallism and homothallism. In heterothallic species, the isolates carry one of the two versions of the *MAT* locus, with either *MAT1-1-1* or *MAT1-2-1* genes, and these two versions are referred to as idiomorphs instead of alleles, because overall DNA sequence homology between idiomorphs is absent

(Turgeon & Debuchy, 2007). In these fungi, only isolates that differ at *MAT* are sexually compatible. Conversely, in homothallic species, the isolates carry both *MAT1-1* and *MAT1-2* genes in their genomes, and thus they are self-compatible (Bennett & Turgeon, 2017). So far, *MAT* loci have not been identified in any desert truffle, and therefore their mating mode (homothallism vs heterothallism) is unknown (Murat *et al.*, 2018). As shown for *Tuber melanosporum* (Rubini *et al.*, 2011, 2014; Linde & Selmes, 2012; Zampieri *et al.*, 2012; Murat *et al.*, 2013; Le Tacon *et al.*, 2014), *Tuber borchii* (Mello *et al.*, 2017; Leonardi *et al.*, 2019), and *Tuber aestivum* (Molinier *et al.*, 2016; Splivallo *et al.*, 2019), the characterization of the *MAT* locus has been critical for disclosing the reproduction mode of these fungi and, in turn, for designing new cultivation strategies.

Here, we sequenced the genomes of *T. claveryi* strain T7 and *Tirmania nivea* strain G3 and compared them with those of other Pezizomycetes. We also assessed both plant and fungal transcriptomes of the symbiosis between *T. claveryi* and *H. almeriense* under well-watered (favouring intercellular colonization) and drought-stress (favouring intracellular colonization) conditions. We aimed to decipher the specific genomic and transcriptomic features of desert truffles compared with other Pezizaceae species. Our findings pave the way for a better understanding of sexual and vegetative propagation modes of the desert truffles and highlight singularities in their genomes and transcriptomes that may shape the ectendomycorrhizal interaction between these fungi and their host plants.

Materials and methods

Strains and fungal material for genome sequencing

Terfezia claveryi strain T7 was isolated from an ascocarp collected in an *H. almeriense* × *T. claveryi* experimental field located in Zarzadilla de Totana (Murcia, Spain) in April 2007. *Tirmania nivea* strain G3 was isolated from an ascocarp collected close to *Helianthemum lippi* plants in an arid natural site, located in Abu Dhabi (United Arab Emirates), in February 2014. Both strains have been maintained *in vitro* by subculturing every 3–6 months, using modified Melin–Norkrans (MMN) and/or optimal MMN (MMNo; Arenas *et al.*, 2018) liquid shaking culture (75 rpm) or solid media at 23°C in the dark.

Genome sequencing, assembly, and annotation

Genomic DNA and total RNA from axenic liquid culture of *T. claveryi* and *T. nivea* mycelia were isolated by the cetyl trimethylammonium bromide method (Chang *et al.*, 1993). Genome sequencing details can be found in Supporting Information Methods S1. They were annotated with the Joint Genome Institute (JGI) Annotation Pipeline, which detects and masks repeats and transposable elements (TEs), predicts genes, characterizes each translated protein with sub-elements such as domains and signal peptides, chooses a best gene model at each locus to provide a filtered working set, clusters the filtered sets into draft gene families, ascribes functional descriptions, and creates a JGI

MycCosm genome portal with tools for public access and community-driven curation of the annotation (Grigoriev *et al.*, 2014; Kuo *et al.*, 2014).

Phylogenetic analyses and comparative genomics

For phylogenetic tree construction and genome comparison, 3156 single-copy orthologues were selected among 13 published fungal genomes belonging to the Pezizomycetes class (Table S1). Single-copy orthologues are well suited for phylogenetic analyses since they result from speciation events and go beyond the issues of conventional markers (Debray *et al.*, 2019). Orthologous genes among the selected fungi were identified using FASTORTHO program with the parameters set to 50% identity, 50% coverage, inflation 3.0 (Wattam *et al.*, 2014). Protein sequences were downloaded from the JGI MycoCosm database (<https://genome.jgi.doe.gov/mycosm/home>). Clusters with single-copy genes were identified and aligned with MAFFT 7.221 (Katoh & Standley, 2013), ambiguous regions eliminated, and single-gene alignments concatenated with GBLOCKS 0.91b (Cruickshank, 2000). A phylogeny tree was constructed with RAxML 7.7.2 (Stamatakis, 2006) using the standard algorithm, the PROTGAMMAWAG model of sequence evolution, 1000 bootstrap replicates, and random seed number (12345).

Statistics of JGI genome assemblies (i.e. N50, number of genes and scaffolds, genome size) were obtained from JGI MycoCosm (<https://genome.jgi.doe.gov/programs/fungi/index.jsf>). Genome completeness with single-copy orthologues was calculated using BUSCO v.3.0.2 with default parameters (Sima *et al.*, 2015). The coverage of TEs in genomes was calculated and visualized using our set of custom R scripts TINGO (Morin *et al.*, 2019). Secreted proteins were predicted as described previously by Pellegrin *et al.* (2015). The count and ratio of total (occurrences in genomes) and secreted CAZymes, lipases, proteases, and SSPs (<300 amino acids) and the predicted plant cell-wall-degrading enzymes (PCWDEs) and microbe cell-wall-degrading enzymes (MCWDEs) were calculated, visualized, and compared (note that some enzymes may belong to both PCWDE and MCWDE categories). Global trends of ecological groups were evaluated using nonmetric multidimensional scaling with the count of total and secreted CAZymes using metaMDS in R package VEGAN (Oksanen *et al.*, 2016). The output files generated were combined and visualized with the set of custom R scripts PRINGO (Miyachi *et al.*, 2020a) incorporating R packages GGPlot2, GGTREE and EGG (Wickham, 2009; Auguie, 2017; Yu *et al.*, 2017).

Gene family evolution

ORTHOFinder v.2.4.0 (Emms & Kelly, 2019) and CAFE v.4.2 (Han *et al.*, 2013) were used for the computational analysis of gene family evolution. The proteomes from the 13 fungi (Table S1), plus *Cenococcum geophilum* and *Oidiodendron maius* as external group, were aligned to each other with BLASTP, and the orthogroups were inferred by the MCL algorithm (Enright *et al.*, 2002) within the ORTHOFINDER pipeline.

The previously built phylogenetic tree was rooted and made ultrametric with R package APE (Paradis & Schliep, 2019). We estimated the species divergence times using the mcmctree method implemented in PAML v.4.8. (Yang, 2007). The independent-rates clock model, a WAG substitution model and approximate likelihood calculation were used. The substitution rates of each gene were estimated by CODEML under a global clock model. We set the time unit to 100 Myr and applied uniform priors on two calibration points with lower and upper hard bounds: the Tuberales divergence time estimation between 134 and 179 Ma (Bonito *et al.*, 2013) and the Pezizomycetes–Leotiomyceta divergence time estimation between 400 and 487 Ma (Prieto & Wedin, 2013). The orthogroups, together with the dated ultrametric tree, were used by CAFE to estimate an error model accounting for genome assembly errors (caferor command) and the birth–death parameter λ . Such a model and λ value were finally used to infer expansions/contractions for the protein families analysed.

Pfam domain enrichment/depletion analysis

The top 100 Pfam domains (<https://pfam.xfam.org/>, v.32.0) identified in the gene repertoires of the 13 fungal species were selected. The occurrence frequency values were transformed into *z*-scores, which are a measure of relative enrichment and depletion. The hierarchical clustering was done with a Euclidian distance metric and average linkage clustering method. Data were visualized and clustered using MULTIEXPERIMENT VIEWER (<http://mev.tm4.org>). The Mann–Whitney–Wilcoxon test was performed to compare each Pfam family between the four desert truffles (*T. claveryi*, *T. boudieri*, *T. nivea*, and *Kalahariturber pfeilii*) and the rest of Pezizomycetes.

MAT and sexual-related genes identification

MAT 1-1-1 genes were identified by the JGI annotation pipeline in the genomes of *T. nivea* and *K. pfeilii*. To search for all putative *MAT* genes in *T. claveryi*, *T. boudieri*, *T. nivea* and *K. pfeilii* genomes, a local BLASTP was performed on the JGI website using the *MAT 1-1-1* and *MAT 1-2-1* genes of filamentous ascomycetes as queries and the default parameters. The genomic regions surrounding the *MAT* locus in these species were then scanned to search for conserved genes. Full coding sequences of *MAT 1-1-1* and the HMG domain of *MAT 1-2-1* genes were aligned using MAFFT 7.221 (Katoh & Standley, 2013), and maximum-likelihood analysis was done in IQ-TREE (Nguyen *et al.*, 2014) using an ultrafast bootstrap with 1000 replicates, random seed number (567943) and the best-fit model (VT + I), chosen according to Bayesian information criterion, determined with MODELFINDER (Kalyaanamoorthy *et al.*, 2017). Phylogeny trees were visualized using MEGA X (Kumar *et al.*, 2018). Genes related to pheromone biosynthesis and signalling were also searched in *T. claveryi* and *T. nivea* by BLASTP using orthologous genes already identified in other Pezizomycetes as queries (Murat *et al.*, 2018).

Biological material and experimental conditions for transcriptome analysis

Vegetative mycelium of strain T7 from *T. claveryi* was grown in Erlenmeyer flasks containing MMNo medium (Arenas *et al.*, 2018). The cultures were shaken at 75 rpm and maintained in the dark at 23°C for 3 months, then mycelium was collected, snap frozen in liquid N₂, and stored at –80°C until further analysis.

For host plant production, *H. almeriense* seeds were collected in Zarzadilla de Totana (Murcia, Spain) and germinated as described in Morte *et al.* (2008). Owing to the erratic growth and difficulties for *in vivo* mycorrhiza formation by *T. claveryi* mycelium (Arenas *et al.*, 2018), spore inoculation is the standard and preferred method for mycorrhizal plant production (Morte *et al.*, 2008, 2012, 2017). After 2 months of growth, 36 plants were transferred to 300 ml individual pots, where they were inoculated with approximately 10⁵ *T. claveryi* spore solution obtained from mature ascocarps from the same location stated earlier. Twelve plants were left uninoculated to act as nonmycorrhizal control plants (NMPs). Three months after spore inoculation, the mycorrhizal status of plants was assessed by using a stereomicroscope (Gutiérrez *et al.*, 2003), and 12 of the inoculated plants were subjected to drought stress. Soil moisture was maintained over 75% for the well-watered mycorrhizal plants (WWMPs) and NMPs, whereas it was lowered to 40% and maintained until shoot water potential was < –2 MPa in drought-stressed mycorrhizal plants (DSMPs). Shoot water potential Ψ was measured in the following way: 5 cm long plant apices were covered in dark during 1 h, cut, and immediately placed in a pressure chamber (Soil Moisture Equipment Corp., Santa Barbara, CA, USA) according to Scholander *et al.* (1965). Once DSMPs reached Ψ < –2 MPa, plant biomass was calculated and Chl content was estimated (Morte & Andriano, 2014). Secondary and tertiary roots containing apical tips from six plants per treatment were rinsed to remove soil and harvested. They were immediately stained with trypan blue and observed under an Olympus BH2 microscope (Tokyo, Japan), to calculate mycorrhizal colonization, mycorrhizal intensity, and intracellular colonization from six replicates per treatment. One hundred secondary and tertiary 2–5 mm root sections per sample were observed under the microscope and were classified as mycorrhizal or nonmycorrhizal depending on the presence/absence of *T. claveryi* mycorrhizal structures. The results are given as percentage of the total sections observed. To calculate the mycorrhiza formation intensity, each *T. claveryi* mycorrhizal structure observed was given a score of 1 (< 10% of mycorrhizal structures observed in one segment), 2 (< 50%), 3 (\geq 50%), or 4 (> 90%), according to the amount of mycorrhiza observed in a single section, similar to Derkowska *et al.* (2008). Then, the mycorrhiza formation intensity was calculated as the weighted average divided by 4. Each *T. claveryi* mycorrhizal colonization was also classified as intercellular or intracellular based on the presence of a well-differentiated intercellular Hartig net or intracellular hyphae detected inside root cells. To support these observations, another portion of roots was observed under a

stereomicroscope and some mycorrhizal root morphotypes were selected and processed to make semi-thin sections of 0.5 μ m, using a Leica UC6 ultramicrotome (Vienna, Austria), then stained with toluidine blue and observed under an Olympus BH2 microscope. Secondary and tertiary roots, from three replicates per treatment, were immediately snap frozen in liquid N₂ and stored at –80°C for further RNA extraction.

Transcriptome analysis of *Terfezia claveryi*

Total RNA was extracted from three biological replicates of *T. claveryi* free-living mycelium (FLM) and from mycorrhizal roots of WWMPs, DSMPs, and NMPs with the same methodology as reported earlier herein. RNA sequencing (RNA-seq) was performed using 2 × 75 bp Illumina HiSeq2500 sequencing after messenger RNA (mRNA) library construction by Lifesequencing SL (Valencia, Spain). Raw reads were analysed using FASTQC for quality (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>), paired, trimmed, and mapped against *T. claveryi* reference transcripts (<https://mycocosm.jgi.doe.gov/Tercla1/Tercla1.home.html>) using CLC Genomics WORKBENCH 11 (Qiagen, Courtaboeuf, France). For mapping, the minimum length fraction was 0.9, the minimum similarity fraction 0.9, and the maximum number of hits for a read was set to 10. Raw read counts were normalized (with the function *counts* in DESEQ2), and differential transcription levels of the contigs (Bonferroni adjusted $P < 0.05$) were calculated with DESEQ2 (Love *et al.*, 2014). The RNA-seq data were consistent according to the quality assessment (Figs S1, S2), permitting our transcriptomic analyses. Condition-specific differentially expressed genes ($\log_2(\text{fold-change}) > 2$ or < -2 , false discovery rate FDR < 0.05) were identified using a custom R script, SHIN+GO (Miyachi *et al.*, 2016, 2017, 2018, 2020b). Symbiosis-induced and repressed genes were queried against the 13 genomes analysed using BLASTP, and they were classified according to eukaryotic orthologous groups and secretome categories (i.e. SSPs and secreted CAZymes) as described in Pellegrin *et al.* (2015).

De novo assembly of the plant transcriptome

A *de novo* transcriptome assembly for *H. almeriense* was performed with sequencing reads from NMP, WWMP, and DSMP samples using MEGAHIT v.1.1.3 (Li *et al.*, 2015). Transcripts from each condition (NMP, WWMP, and DSMP) were mapped onto the assembled contigs with BOWTIE2 v.2.3.0 (Langmead & Salzberg, 2012), and reads mapped onto the contigs were counted with SAMTOOLS v.1.7 (Li *et al.*, 2009). Only the contigs with > 200 bases were selected for further steps, and then they were annotated using BLAST on the nonredundant National Center for Biotechnology Information (NCBI) sequence database (March 2018 version) and JGI MycoCosm annotation data (July 2018 version) using DIAMOND v.0.9.19 (Buchfink *et al.*, 2014) with the parameters *more-sensitive max-target-seqs 1 max-hsps 1 evalue 0.00001*. Taxonomic annotations for each contig were selected based on the higher bit scores for the best hits obtained from the NCBI or JGI databases.

The annotated contigs were further separated into three categories (plants, fungi, metazoa) according to their annotations, and unknown contigs were removed. Contigs showing sequence coverage >80% with a known sequence on NCBI or JGI databases and with average reads >5 combined from all conditions were selected for further analyses. A summary of the assembly and the contig selection process is shown in Table S2. Raw read counts were normalized (with the function *counts* in DESEQ2) and differential transcription levels of the contigs (Bonferroni adjusted $P < 0.05$) were calculated with DESEQ2 (Love *et al.*, 2014). Condition-specific differentially transcribed genes ($\log_2(\text{fold-change}) > 2$ or < -2 , $\text{FDR} < 0.05$) were identified with the SHIN + GO pipeline (Miyachi *et al.*, 2016, 2017, 2018, 2020b).

Results

Desert truffles' genome features

As expected, the phylogenomic analyses placed both *T. claveryi* and *T. nivea* in the Pezizaceae family, within the Pezizales order, and close to the other sequenced desert truffles; that is, *T. boudieri* and *K. pfeilii* (Fig. S3). BUSCO score (*T. claveryi*, 83.1%; *T. nivea*, 86.5%) and RNA-seq read mapping (*c.* 96%, see Info page on JGI Genome portals) indicated that the assembly and annotation are of good quality. Genome size for *T. claveryi* (74.5 Mb) and *T. nivea* (63.1 Mb) is in the same range as other ectomycorrhizal Pezizaceae genomes analysed so far (70.0 ± 8.0 Mb), but significantly smaller than the genomes of ectomycorrhizal Tuberaceae (124.3 ± 19.8 Mb) (Table S3). *Terfezia claveryi* and *T. nivea* genome assemblies contain 12 765 and 10 591 genes, respectively, and the average gene content from Pezizaceae was not significantly different from Tuberaceae (Table S3). *Terfezia claveryi* and *T. nivea* assemblies contain 48.3% and 38.3% of TE sequences, respectively (Fig. S3), and this is significantly lower in Pezizaceae than in Tuberaceae (Table S3). The total and secreted CAZyme profiles of the four desert truffles are very similar, but they are different from Tuberaceae and non-mycorrhizal Pezizomycetes (Fig. S4). The number of total and secreted CAZymes is similar in the Pezizaceae desert truffles and the Tuberaceae truffles, but each family showed substantial differences in their CAZyme family distribution, such as fewer MCWDEs and a significantly lower number of genes targeting pectin and chitin in desert truffles (Fig. 1; Table S3).

Desert truffle gene families

A total of 143 orthologous gene family expansions and 1474 contractions were inferred for the common ancestor of desert truffles. The number of orthologous gene families gained in desert truffle species relative to their most recent common ancestor ranges from 300 to 900 families. On the other hand, *K. pfeilii* presents a strikingly low number of orthologous gene family losses (nine) compared with the other desert truffles (Fig. 2a). Most of the expanded genes from desert truffles encode proteins with no known functions or functions related to protein–protein

interactions, signalling networks, stress, and/or cell death (Tables S4, S5).

In the hierarchical clustering of the abundance of the different Pfam protein domains annotated in the genomes of desert truffles and other Pezizomycetes, the desert truffles clustered together (Fig. S5). Only two gene families have an enriched number of genes, whereas 18 families have a lower gene content in desert truffles. Among the latter, it is worth noting the significant reduction in the gene copy number of several genes related to transport, like Major Facilitator Superfamily (MFS; 53.75 vs 77.56 on average, desert truffle vs nondesert truffles Pezizomycetes, respectively), ABC transporters (11.25 vs 14.88 on average, desert truffle vs nondesert truffles Pezizomycetes, respectively), or sugar transporters (20.50 vs 30.67 on average, desert truffle vs nondesert truffles Pezizomycetes, respectively) (Fig. 2b).

Genes involved in sexual reproduction in desert truffles

Putative *MAT* genes were identified in *T. claveryi*, *T. nivea*, *T. boudieri* and *K. pfeilii*. The *MAT 1-1-1* gene, coding for a transcription factor with an α -box domain, was located on scaffold 39 in *T. claveryi* assembly, scaffold 83 in *T. nivea*, and scaffold 10 in *K. pfeilii*, whereas the opposite mating type gene, *MAT 1-2-1*, was found on scaffold 37 in the *T. boudieri* assembly. Only in the *K. pfeilii* genome were both *MAT* genes found to also be present, on scaffold 14, the *MAT 1-2-1* gene. This gene showed the typical HMG domain, maintained the intron position characteristic of *MAT 1-2-1* genes (Fig. S6), and clustered together with Pezizaceae *MAT 1-2-1* genes in a phylogenetic analysis (Fig. S7). The genomic regions flanking the *MAT* genes of all desert truffles showed synteny with other ascomycetes, highlighting a putative DNA lyase (*APN*) next to all desert truffle *MAT* loci except for *K. pfeilii* *MAT 1-2* locus (Fig. 3, S8). Most of the key genes related to pheromone processing, perception, and signalling were also identified in *T. claveryi* and *T. nivea* as per BLASTP search (Table S6). Most of the genes related to the pheromone pathway were not differentially expressed between *T. claveryi* FLM and mycorrhizas, regardless of the fact that these structures were collected from WWMPs or DSMPs.

Symbiosis-induced genes in *Terfezia claveryi*

When comparing the transcriptomic profiles of the mycorrhizal roots and FLM, 751 genes were found to be upregulated (5.89% of the total predicted genes, $\log_2(\text{fold-change}) > 2$, $\text{FDR} < 0.05$; Table S7), whereas 303 were found to be downregulated (2.37% of the total predicted genes, $\log_2(\text{fold-change}) < -2$, $\text{FDR} < 0.05$; Table S8). Among the highly upregulated genes, there were several genes that may be of importance for the establishment and development of functional mycorrhiza, such as SSPs, several transcription factors, N metabolism enzymes (e.g. nitrate assimilation cluster, cyanase), transporters (aquaporin, amino acid, or sugar transporters, among others), PCWDEs (expansins, cellulose-acting copper-dependent lytic polysaccharide monoxygenases (AA9), and carbohydrate esterase (CE8)),

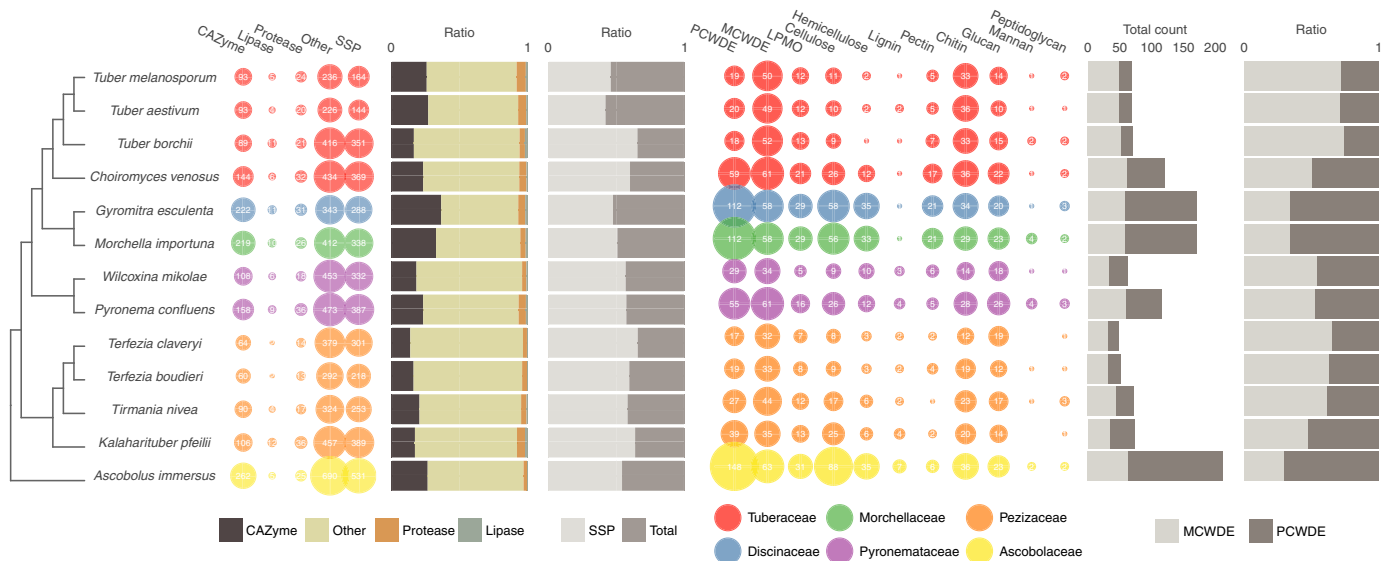


Fig. 1 Secretomic profiles of 13 genomes. Left bubble plot shows the number of secreted genes for carbohydrate-active enzymes (CAZymes), lipases, proteases, and others (i.e. all secreted proteins not in these first three groups). SSP group refers to the number of small secreted proteins (SSPs; < 300 amino acids). The size of bubbles corresponds to the number of genes. Colours in the plots represent different taxonomic families. Left bar plots represent the ratio of CAZymes, lipases, and proteases to all secreted proteins (left) and the ratio of SSPs among the entire secretome (right). Right bubble plot shows the number of plant cell-wall-degrading enzymes (PCWDEs) and microbial cell-wall-degrading enzymes (MCWDEs; fungal cell walls and bacterial membranes), including lytic polysaccharide monoxygenase (LPMO), substrate-specific enzymes for cellulose, hemicellulose, lignin, and pectin (plant cell walls); chitin, glucan, mannan (fungal cell walls); peptidoglycan (bacterial membranes). Right bar plots show the total count of genes including PCWDEs and MCWDEs (left) and the ratio of PCWDEs to MCWDEs (right).

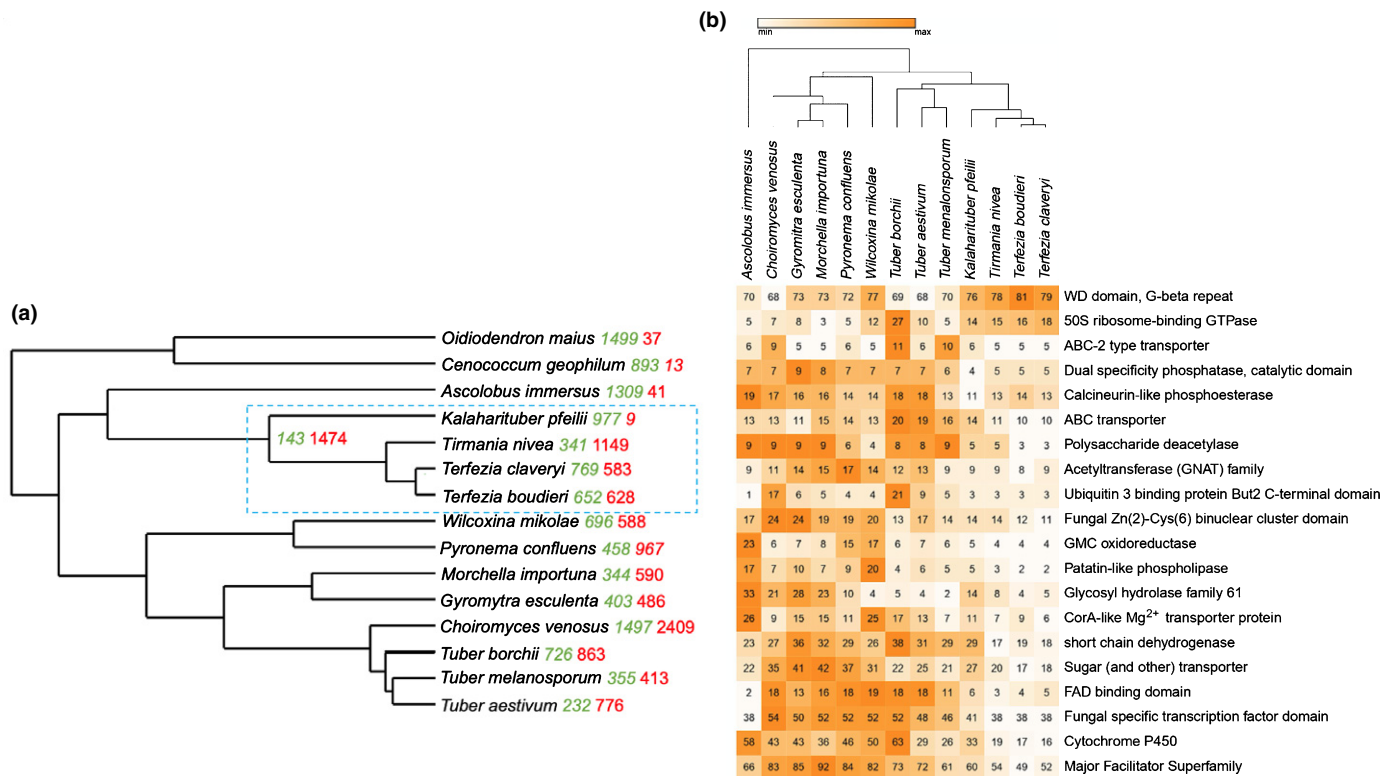


Fig. 2 Gene family distribution in desert truffles. (a) Gene expansion and contraction along a time-calibrated phylogeny tree of 13 Pezizomycetes. Green and red numbers represent significantly ($P < 0.01$) expanded or contracted, respectively, gene family counts. (b) The heatmap depicts a double hierarchical clustering of the significantly enriched or depleted (Mann–Whitney–Wilcoxon test, $P < 0.05$) Pfam families in the desert truffles Pezizaceae compared with the other nine Pezizomycetes analysed. Numbers represent gene counts of each family. The top most frequent 100 domains analysed can be consulted in the Supporting Information Fig. S5.

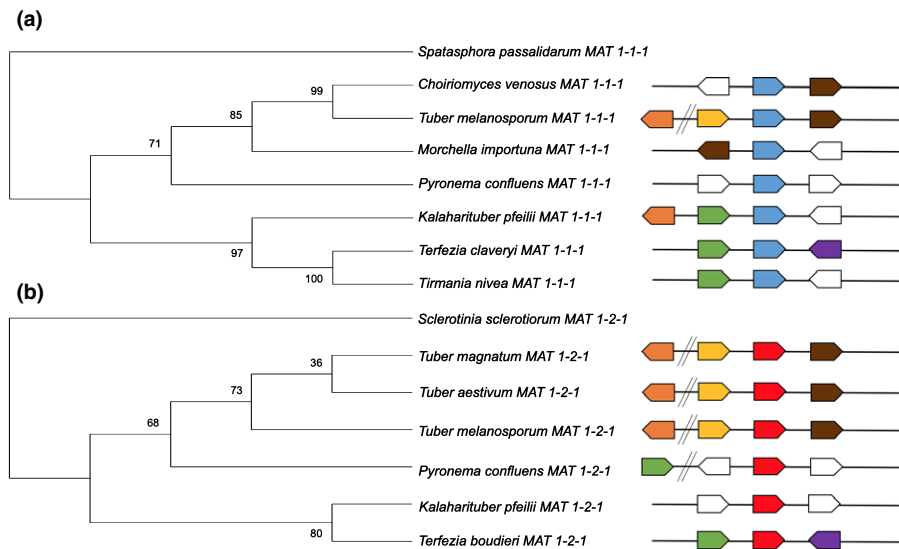


Fig. 3 Phylogenetic tree of MAT sequences from desert truffles and schematic organization of the mating-type loci. Phylogeny tree was constructed by maximum-likelihood method. Coloured arrows represent conserved genes among species. Blue, MAT 1-1-1; red, MAT 1-2-1; green, DNA lyase (APN); purple, Sam decarboxylase; orange, Cyclooxygenase 13 (COX13); yellow, mitochondrial ribosomal protein of the small subunit (RSM22); brown, homologue to GSTUMT2000009628001 from *Tuber melanosporum*; white, nonconserved coding regions. Full coding sequences from (a) MAT 1-1-1 and HMG domain from (b) MAT 1-2-1 proteins were aligned and phylogenetically analysed as explained in the Materials and Methods section.

tryptophan metabolism, and auxin-related proteins (L-amino acid oxygenases, amidases, auxin efflux carrier; Table S7).

In *T. claveryi*, 33% of the symbiosis upregulated genes are conserved among the 13 fungi analysed (clusters VI and VII) (Fig. 4). In these clusters, CAZymes (e.g. AA9 and CE8), genes related to metabolism (e.g. oxidoreductases), cellular processes, and signalling (e.g. peptidases and heat shock proteins) are enriched in comparison with the whole gene repertoire (Fisher's exact test; Table S9). On the other hand, 20% of the upregulated genes are species specific and have no sequence similarity (cluster V). Clusters I to IV (47% of the upregulated genes) contain genes with sequence similarity with genes belonging to all or some of the desert truffles. SSPs are enriched in the whole set of upregulated genes and particularly in clusters I and II compared with the whole gene repertoire (Fisher's exact test; Table S9). Twelve out of the 15 SSPs shared by desert truffles (clusters I and II) also share a substantial level of genome synteny (Fig. S10).

Fungal colonization and gene regulation under drought conditions

No significant differences were found between DSMPs and WWMPs in plant biomass or Chl content (Table S10). *Terfezia claveryi* colonization in DSMPs was greater and more intense (Table 1) than in WWMPs. A greater number of intracellular colonizations (up to 41.9%) was observed in DSMPs vs WWMPs (2.7%) (Fig. 5; Table 1). This morphological observation was corroborated by the higher number of *T. claveryi* RNA reads in DSMPs vs WWMPs *de novo* metatranscriptomes (see next subsection).

A total of 288 and 329 fungal genes (2.26% and 2.58%, respectively, of the entire gene set) were up or downregulated in DSMPs vs WWMPs, respectively ($\log_2(\text{fold-change}) > 2$ or < -2 ,

respectively, $\text{FDR} < 0.05$; Tables S11, S12). Up to 26 fungal genes with a putative function in stress responses (catalase, thioredoxin, cupredoxins, superoxide dismutases, several heat shock proteins) were upregulated.

Under drought conditions, a significant downregulation of secreted CAZymes was found (Fisher's exact test, $P = 0.0007167$ for PCWDEs and $P = 0.004683$ for MCWDEs) compared with the whole gene repertoire. Seven PCWDEs targeting cellulose (AA9, GH131 and GH5_5), hemicellulose (GH43), and pectin (CE8, GH28 and PL4_1) and seven MCWDEs targeting chitin (AA11, CBM18, GH128, GH18 and GH132) and glucans (GH16 and GH17) were downregulated during drought stress conditions, whereas only a GH76 targeting mannan and a GH35 targeting hemicellulose were upregulated (Fig. 6).

Plant co-assembly and gene regulation in mycorrhizal symbiosis

For the plant metatranscriptome analysis, 135 414 transcript contigs were assembled, and we selected 27 346 contigs based on stringent criteria for further transcriptomic analysis (see Materials and Methods section; Table S2). Within each treatment, $> 80\%$ of the reads were either from *T. claveryi* or the host plant with varied proportion of *T. claveryi* (Fig. S11): DSMPs was the treatment with more *T. claveryi* reads (21% vs 12% in WWMPs and 0.4% in NMPs; Table S13). The 0.4% of reads belonging to *T. claveryi* from NMPs can be attributed to slight nursery and/or laboratory contaminations, since no mycorrhizal structures were detected when observed under the optical microscope (Table 1). On average, 72% of the total contigs were from plant origin, as per BLAST results.

From the 19 696 plant contigs identified in *de novo* assembly, 163 were upregulated and 72 were downregulated in

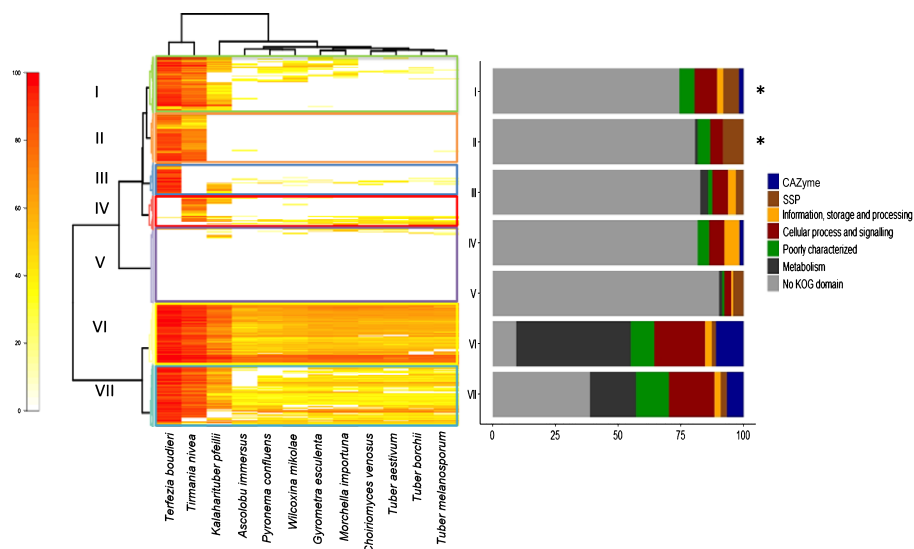


Fig. 4 Sequence conservation and functional analysis of symbiosis-induced transcripts. The heatmap depicts a double hierarchical clustering of the symbiosis-upregulated *Terfezia claveryi* genes (rows, $\log_2(\text{fold-change}) > 2$, FDR-corrected < 0.05 ; Supporting Information Table S7). Symbiosis-induced genes were queried against the genomes of the 13 genomes using BLASTP (see Fig. S9 for the heatmap of symbiosis-downregulated genes). Homologues are coloured from yellow to red depending on the percentage of similarity. Clusters of genes with coordinated expression are numbered and highlighted with different colours. Right of the heatmap, the percentages of putative functional categories are given for each cluster as bar plots. Clusters significantly enriched in small secreted proteins (SSPs) are marked with an asterisk (Fisher's exact test $P < 0.05$). CAZyme, carbohydrate-active enzymes; KOG, eukaryotic orthologous groups.

Table 1 Mycorrhizal colonization of *Terfezia claveryi* \times *Helianthemum almeriense* plants under different treatments.

	Mycorrhizal colonization (%)	Mycorrhizal intensity (out of 4)	Intracellular colonization (%)
Well-watered mycorrhizal plant	21.34 \pm 4.83 a	1.52 \pm 0.19 a	2.73 \pm 1.88 a
Drought-stressed mycorrhizal plant	45.37 \pm 6.62 b	2.20 \pm 0.24 b	41.91 \pm 9.98 b
Nonmycorrhizal plant	nd	nd	nd

Mean \pm SE is presented for each variable. Data were submitted to Mann–Whitney–Wilcoxon test, and different letters represent significant differences ($P < 0.05$). nd, not detected.

WWMPs vs NMPs ($\log_2(\text{fold-change}) > 2$ and < -2 , respectively; Tables S14, S15), whereas 255 plant genes were upregulated and 611 downregulated ($\log_2(\text{fold-change}) > 2$ and < -2 , respectively; Tables S16, S17) in DSMPs vs WWMPs. Among the regulated genes during symbiosis in well-watered conditions, there were several ABC and MFS transporters, pathogen defence-related proteins (thaumatin, peroxidases), and oxidoreductases (e.g. multicopper oxidases, cytochrome P450). When comparing DSMPs with WWMPs, we found a majority of abiotic and biotic stress-response genes (e.g. heat shock proteins, spermidine synthases, thaumatin), transporters (e.g. sugar and amino acid transporters, MFS), carbohydrate-metabolism-related genes (mainly glycoside transferases and glycoside hydrolases), and several GH3 auxin-responsive genes were regulated.

In mycorrhizal plants vs NMPs, regardless of the host plant growing conditions, 20 and 30 genes were up and downregulated, respectively. They represent a core set of symbiosis-regulated *H. almeriense* genes (Fig. 7). Among the upregulated genes, those coding for proteins involved in carbohydrate and lipid metabolism and in transport were the most represented (30% of the total), whereas two genes coding for thaumatin-like pathogen-related proteins showed a further increase under stress. Conversely, among the set of downregulated genes, the majority were of unknown function, two coded for glycoside hydrolase, and a few showed a further decrease in DSMPs vs WWMPs. Among this subset of genes were one gene each for alternative oxidase (AOX) and pectinesterase and two for Pirins and protein kinases.

Discussion

Specific features of the desert truffle genomes

Although desert truffles share many of their genomic features with other Pezizomycetes species, they also exhibit several species-specific characteristics, as shown by our genomic analyses, coupled to the morpho-anatomical and transcriptomic analyses of *T. claveryi* \times *H. almeriense* mycorrhizas. A large genome size and a high number of orphan SSPs, together with a reduced set of PCWDEs, are among the genomic traits shared by *T. claveryi* and *T. nivea* with other mycorrhizal fungi (Kohler *et al.*, 2015; Miyauchi *et al.*, 2020a). Conversely, the expansion of large sets of gene families with unknown Pfam domains are desert-truffle-specific traits, which may be linked to the specific processes that distinguish these fungi from other Pezizomycetes (e.g. different

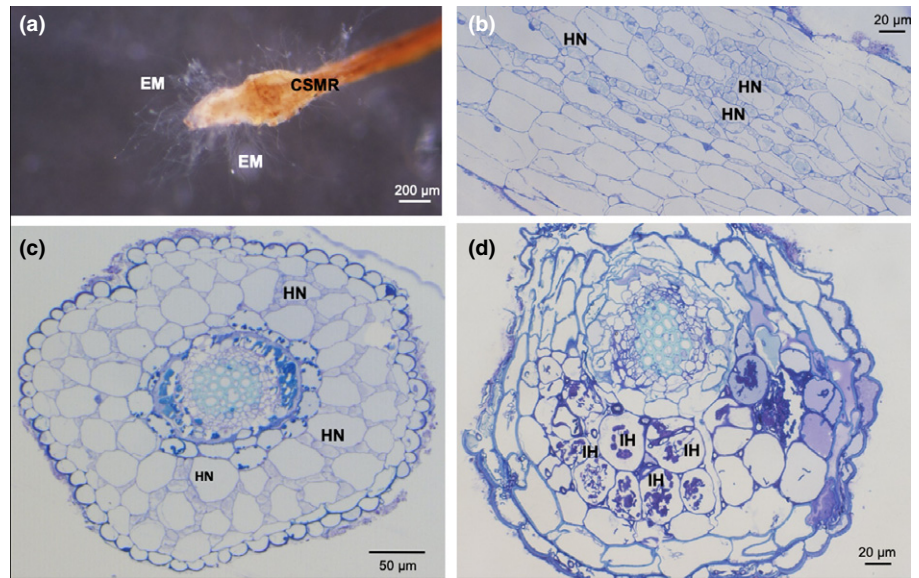


Fig. 5 *Terfezia claveryi* × *Helianthemum almeriense* mycorrhiza. (a) Club-shaped mycorrhizal root (CSMR), one of the four morphotypes described by Gutiérrez *et al.* (2003), with extraradical mycelium (EM). (b) Longitudinal section of a well-watered mycorrhiza (WWMP) with several hyphas forming a Hartig net (HN). (c) Cross-section of a WWMP with several hyphas forming an HN. (d) Cross-section of a drought-stressed mycorrhizal plant with majority of intracellular hyphas (IH) forming clumps.

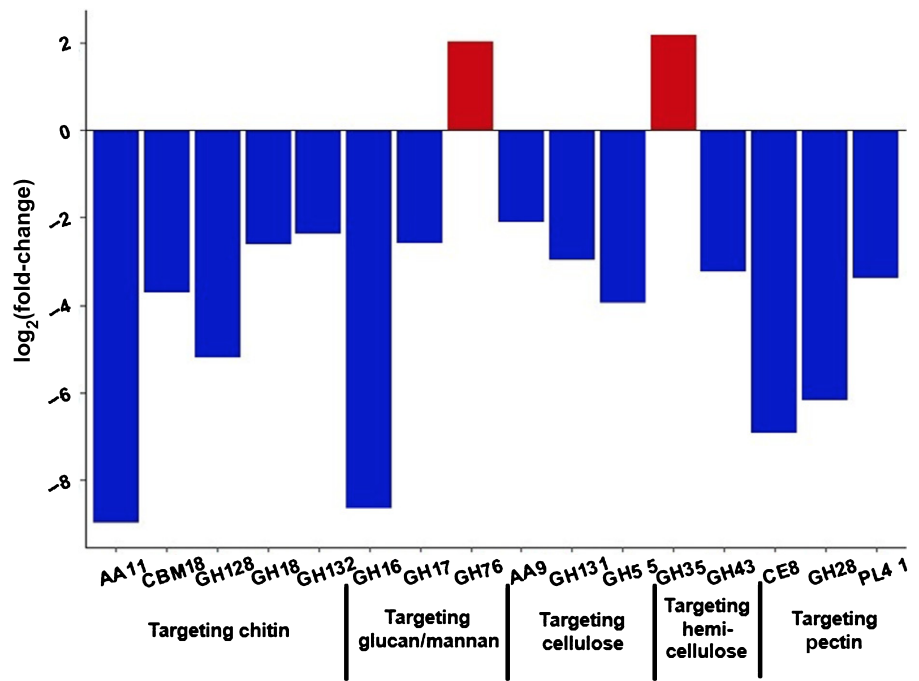


Fig. 6 Regulation of *Terfezia claveryi* plant cell-wall-degrading enzymes and microbial cell-wall-degrading enzymes under drought stress in mycorrhizal symbiosis. Red, upregulation; blue, downregulation.

host specificity, formation of EEMs, adaptation to arid climates). The significant reduction of the number of genes related to metabolite transport, such as MFS, ABC, or sugar transporters, is also a pattern that typifies both *Terfezia* spp. and *T. nivea* with respect to other Pezizomycetes. A similar trend was observed in the mutualistic endophyte *Piriformospora indica*, a well-adapted species to desert environments known to confer several benefits to its hosts under these conditions (Zuccaro *et al.*, 2011). These desert-truffle-specific signatures could be related to unknown mechanisms aimed to optimize water and nutrient resources, ultimately leading to adaptation to arid conditions.

The reproductive modes in desert truffles

Genome sequencing is the only way to characterize the reproductive mode(s) of desert truffles, owing to our inability to master mating mechanisms under laboratory conditions. The occurrence of a single *MAT* gene (either *MAT1-1-1* or *MAT1-2-1*) in the genome of the sequenced strains of *T. claveryi*, *T. nivea* and *T. boudieri* is a signature that these fungi are heterothallic, whereas the presence of both *MAT* genes in *K. pfeilii* genome suggests that this species, differently from all the other symbiotic Pezizomycetes sequenced thus far (Murat *et al.*, 2018), might

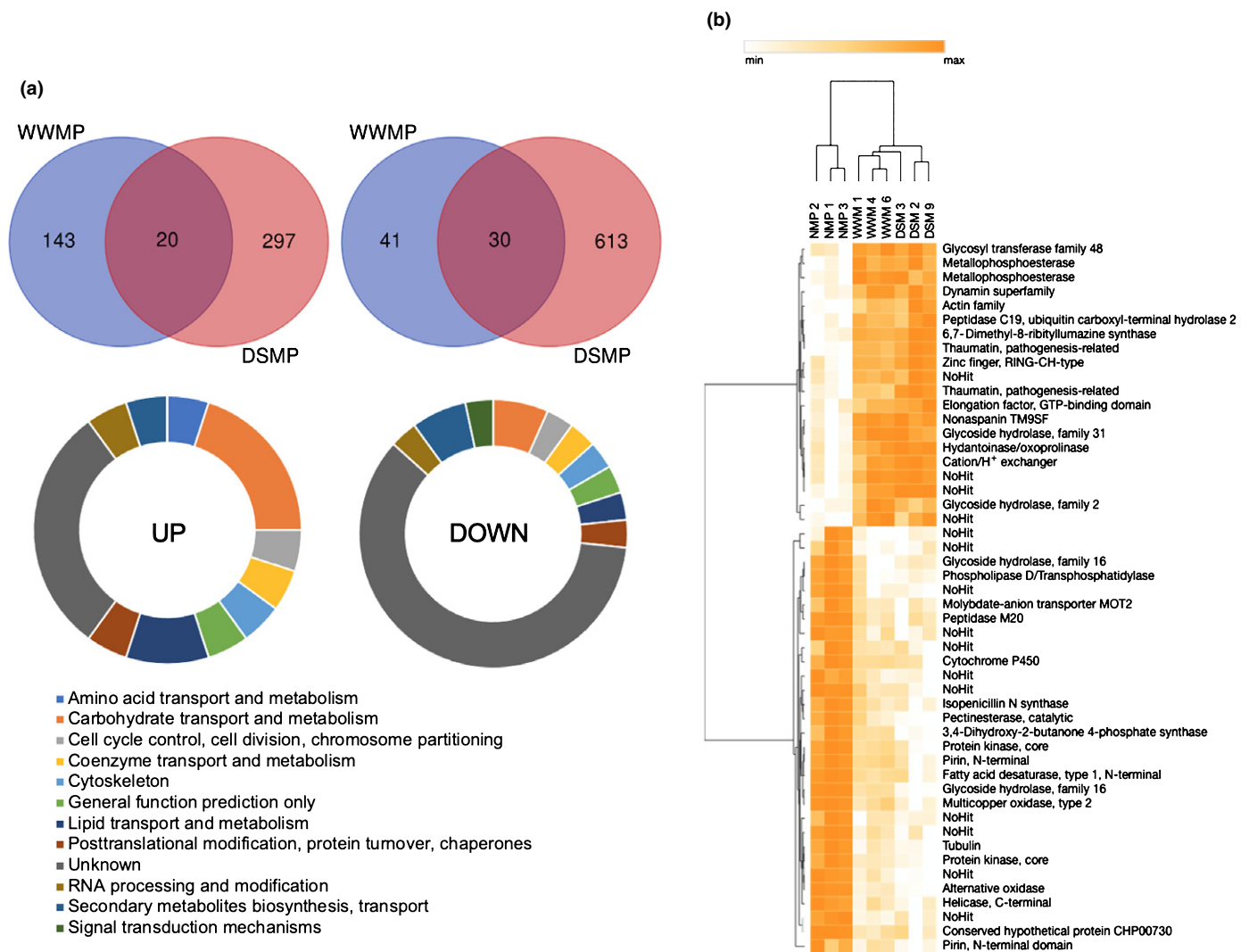


Fig. 7 Core set of plant genes regulated in symbiosis. (a) Venn diagram representing the core upregulated (left) and downregulated (right) genes ($\log_2(\text{fold-change}) > 2$ or < -2 , respectively) and their eukaryotic orthologous groups classification. (b) The heatmap shows a double hierarchical clustering with the number of normalized transcripts by DESeq2 of core genes in three different conditions: nonmycorrhizal plant (NMP 1, 2 and 3), well-watered mycorrhiza (WWM 1, 4 and 6), and drought-stressed mycorrhiza (DSM 2, 3 and 9) replicates.

self-fertilize. Future studies will be needed to verify whether both *K. pfeilii* *MAT* genes are functional and the conditions that promote their expression. The characterization of the structure and organization of the *MAT* locus and its surrounding genomic regions in the desert truffles provides crucial information to optimize the cultivation strategies for these fungi, model the genomic rearrangements within and around this locus, and trace the evolution of the breeding systems within Pezizomycetes.

Genus and species-specific small secreted proteins

Mutual recognition of symbionts and development of mycorrhizal structures rely on a finely tuned exchange of signalling molecules between the partners, which are seen in the symbiosis of plants with Rhizobiaceae, arbuscular mycorrhiza, and ectomycorrhiza (Perret *et al.*, 2000; Bonfante & Genre, 2015; Martin, 2016). Recent studies showed that symbiosis-induced effector

proteins, such as *Laccaria bicolor* MiSSP7 and MiSSP7.6, interact with transcriptional regulators and dampen the local defence responses of *Populus* (Plett *et al.*, 2011, 2014; Kang *et al.*, 2020). Among the symbiosis upregulated genes of *T. claveryi*, we identified several mycorrhiza-induced SSPs (MiSSPs) showing sequence conservation and synteny within the desert truffles (*T. claveryi*, *T. boudieri*, *T. nivea* and *K. pfeilii*). Along with conserved MiSSPs, there are some proteins shared only between *T. claveryi* and its most closely related species (*T. boudieri*, *T. nivea*), and six MiSSPs are unique to *T. claveryi*. The three species *T. claveryi*, *T. boudieri* and *T. nivea* are known to form mycorrhizal symbiosis with the same host species (Moreno *et al.*, 2014). The presence of a common set of MiSSPs suggests the existence of conserved mechanism(s) among the desert truffles to interact with their hosts. At the same time, specific *T. claveryi* MiSSPs induced upon the symbiosis with *H. almeriense* suggest the possible existence of species-specific mechanisms. Further

functional analyses would be required to determine the involvement of these effectors in EEM formation and their specificity among different desert truffles and/or towards different host species.

Helianthemum almeriense transcriptomic features related to symbiosis

As the complete genome of *H. almeriense* is lacking, its transcriptome was assembled *de novo* using our metatranscriptome data set. This represents the first transcriptomics studies in Cistaceae, a family with a prominent role in the configuration of Mediterranean xeric landscapes (Ellul *et al.*, 2002). We annotated 19 696 transcript contigs, but a core set of only 50 genes is strongly regulated in mycorrhizal conditions, regardless of whether or not plants were under water stress. Upregulated genes related to carbohydrate and lipid metabolism may reflect the metabolic changes produced by the symbiotic state and the exchange of nutrients between the plant and the fungus (Wang *et al.*, 2017). The upregulation of thaumatin-like genes, which are known to play a role in pathogen defence (Liu *et al.*, 2010), could be either a direct defence response to limit *T. claveryi* colonization or reflect an enhanced state of defence against pathogens provoked by *T. claveryi* colonization, as it happens in arbuscular mycorrhizal plants (Pozo *et al.*, 2009). Among the downregulated core genes, it is worth noting the *alternative oxidase (AOX)* gene. The AOX pathway is an alternative to the cytochrome oxidase (COX) pathway for electron transport in the respiration process, and it is uncoupled from ATP production. AOX function is particularly important during stress when it is involved in the optimization of respiratory metabolism, which facilitates the continuous turnover of the tricarboxylic acid and acts to prevent the excess generation of reactive oxygen species (ROS) (Li *et al.*, 2013; Vanlerberghe, 2013). The decrease of *AOX* mRNA levels in *T. claveryi* mycorrhizal roots concurs with previous studies that show suppression on AOX activity induced by arbuscular mycorrhizal colonization, a condition in which the COX pathway seems to be the predominant one (Liu *et al.*, 2015; Del-Saz *et al.*, 2018) (see Notes S1 for additional discussion on this issue).

Intracellular colonization under dry conditions

Terfezia claveryi colonizes *H. almeriense* by differentiating symbiotic structures, which differ strikingly from ectomycorrhizal structures. The extent of root colonization, but also the ability to form either inter or intracellular structures, is driven by environmental conditions. As previously observed (Navarro-Ródenas *et al.*, 2013; Marqués-Gálvez *et al.*, 2019), drought stress enhances the mycorrhizal colonization of *H. almeriense* roots by *T. claveryi*. This morphological observation is supported by the higher number of RNA reads corresponding to *T. claveryi* found in dry conditions compared with well-watered conditions. Intriguingly, we observed a shift in the mycorrhizal structures formed under drought conditions. Approximately 40% of the observed mycorrhizal roots entailed intracellular hyphal

structures, whereas *T. claveryi* does not generally penetrate into host cells under well-watered conditions. Pectin-targeting enzymes are thought to play a key role in the formation of the intercellular Hartig net (Veneault-Fourrey *et al.*, 2014). We observed that fungal genes coding for pectin targeting enzymes (CE8, GH28 and PL4_1) were downregulated under the drought condition, suggesting that the activity of these enzymes might be dispensable when the intracellular colonization is favoured. Whereas in ericoid mycorrhiza the intracellular penetration is mainly attributed to PCWDEs because of their increased expression in mycorrhizal symbiosis (Martino *et al.*, 2018), the generalized downregulation of PCWDEs shown in this work suggests that this may not be the case for EEM desert truffles.

The intracellular colonization of desert truffles under drought stress could depend more on the plant side than on the fungal symbiotic partner, as it has been suggested in arbuscular mycorrhizal associations (Tisserant *et al.*, 2013). Our transcriptomic data from *H. almeriense* show some CAZymes upregulated under drought stress. However, for most of them, the lack of full sequences makes it difficult to predict the signal peptides to infer their specific role in plant cell-wall remodelling. The role of plant hormones should also be taken into account. Auxin is already known to determine inter or intracellular colonization in *T. boudieri* × *C. incanus* *in vitro* (Zaretsky *et al.*, 2006a); and in the present study, certain auxin-related genes from both fungal and plant origins were upregulated under dry conditions. This hormone could act in enhancing lateral root development during stress (Korver *et al.*, 2018), thus increasing the surface available for fungal colonization and/or in loosening and remodelling plant cell walls and, possibly, facilitating the accommodation of the intracellular hyphae under drought stress, due to the promotion of 'acid-growth' (Majda & Robert, 2018).

Since intracellular fungal colonization appears to be driven by water or nutritional stresses, it is difficult to infer whether differentially expressed genes under drought stress are master regulators of the switch in mycorrhizal types or are simply part of the mycorrhiza response to drought stress. Moreover, the lack of drought-stressed nonmycorrhizal control plants in this experiment also leaves open the possibility that differentially expressed genes under drought might be a constitutive plant response to drought. For instance, the massive upregulation of fungal stress-response genes, in particular of several ROS scavengers, may have a dual role. It could help in ROS alleviation and, hence, contribute to mycorrhiza adaptation to dry conditions. At the same time, it could serve as a mechanism to increase fungal colonization by decreasing the oxidative burst that is normally produced by plants to limit fungal expansion in the roots (Salzer *et al.*, 1999; Baptista *et al.*, 2007; Marqués-Gálvez *et al.*, 2019). Future research aimed at decoupling the ecto/endo switch from environmental stresses is required to shed more light on the specific mechanisms regulating the host cell intracellular colonization by ectendomycorrhizal fungi. The functional analyses of plant and fungal genes highlighted in this work will also help to improve our understanding of the interaction between desert truffles and their host plants. Coupled to the knowledge of the mating systems of desert truffles gained through the present study, these

future analyses will be of pivotal importance to promote efficient truffle farming in desiccated areas.

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Author contributions

AM and FMM conceived, managed, and coordinated the project. AM, ANR, FMM and A Kohler planned and managed the research. IVG, KWB and A Kuo supervised the sequencing and annotation of the genomes. JEMG, SM, FP, FMM, AM, EM, LA, ANR, FA and A Kohler carried out the experimental set-up, bioinformatic analysis, and analysis and interpretation of the data. JEMG, SM and FP drafted and wrote the manuscript. MPG, ANR, FMM, A Kohler and AM contributed to sections of the manuscript. All the authors read and improved the manuscript.

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Data availability

Full genomes, predicted genes, and transcripts of *T. claveryi* T7 and *T. nivea* G3 can be found at the following JGI portals: <https://genome.jgi.doe.gov/Tercla1/Tercla1.home.html> and <https://genome.jgi.doe.gov/Tirniv1/Tirniv1.home.html>, respectively. RNA-seq data from *T. claveryi* × *H. almeriense* mycorrhiza and drought stress assay are available at Gene Expression Omnibus (GEO) with the accession nos. GSE154490 and GSE155042.

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Supporting Information

Additional Supporting Information may be found online in the Supporting Information section at the end of the article.

Fig. S1 *Terfezia claveryi* distribution and density of normalized log₂ transformed read counts of 9247 genes from nine replicates, three per treatment.

Fig. S2 Correlation of *Terfezia claveryi* transcriptomes among nine replicates, three per treatment.

Fig. S3 Phylogenomic tree with general features and statistics.

Fig. S4 CAZyme-based clusters of 13 fungi.

Fig. S5 Presence and abundance of different Pfam domain-containing proteins in 13 Pezizomycetes.

Fig. S6 Alignment of the MAT 1-2-1 proteins of *K. pfeilii* with other Pezizomycotina species.

Fig. S7 Phylogenetic tree of putative HMG domain containing proteins from *Kalaharituber pfeilii* and *Tuber melanosporum*, together with known MAT 1-2-1 proteins from *Tuberaceae* and *Pezizaceae*.

Fig. S8 Schematic representation of the second MAT locus identified only in *Kalaharituber pfeilii* and the genome synteny with other desert truffles in the regions framing the MAT locus.

Fig. S9 Sequence conservation and functional analysis of symbiosis-downregulated transcripts.

Fig. S10 Genome synteny of SSPs shared by desert truffles.

Fig. S11 Kingdom contig distribution from reads of the meta-transcriptome of *T. claveryi* × *H. almeriense* roots.

Methods S1 *Terfezia claveryi* and *Tirmania nivea* genome sequencing.

Notes S1 Additional discussion on *Helianthemum almeriense* alternative oxidase (AOX) genes downregulation in mycorrhizal plants under drought stress.

Table S1 Genomic features of *Terfezia claveryi* and *Tirmania nivea* together with the rest of the fungi used for genome comparison.

Table S2 Summary of the number of contigs during the selection process for de novo plant, fungi and metazoan assembly.

Table S3 Differences in genomic and secretomic features between *Tuberaceae* and *Pezizaceae* mycorrhizal members used in this study.

Table S4 Most significantly expanded orthogroup gene families according to CAFE analysis for *Terfezia claveryi* (Tercla, green), *Terfezia boudieri* (Terbo, red), *Tirmania nivea* (Tirniv, blue) and *Kalaharituber pfeilii* (Kalpfe, yellow).

Table S5 Most significantly contracted orthogroup gene families according to CAFE analysis for *Terfezia claveryi* (Tercla, green), *Terfezia boudieri* (Terbo, red), *Tirmania nivea* (Tirniv, blue) and *Kalaharituber pfeilii* (Kalpfe, yellow).

Table S6 Genes involved in pheromone biosynthesis and signalling in desert truffles, and its expression in *T. claveryi*.

Table S7 Upregulated *T. claveryi* genes (log₂ fold-change > –2) in mycorrhizal roots vs FLM.

Table S8 Downregulated *T. claveryi* genes (log₂ fold-change > –2) in mycorrhizal roots vs FLM.

Table S9 Fisher's exact test summary results for *Terfezia claveryi* mycorrhiza differentially expressed genes clusters.

Table S10 Plant growth under different water conditions.

Table S11 Upregulated (\log_2 fold-change > 2) *T. claveryi* genes comparing mycorrhizal roots drought stressed with control well-watered mycorrhizal roots.

Table S12 Downregulated (\log_2 fold-change < -2) *T. claveryi* genes comparing mycorrhizal roots drought stressed with control well-watered mycorrhizal.

Table S13 Fungal reads present in plant de novo co-assembly.

Table S14 Upregulated (\log_2 fold-change > 2) *H. almeriense* genes comparing mycorrhizal roots with non mycorrhizal roots.

Table S15 Downregulated (\log_2 fold-change < -2) *H. almeriense* genes comparing mycorrhizal roots with non mycorrhizal roots.

Table S16 Upregulated (\log_2 fold-change > 2) *H. almeriense* genes comparing mycorrhizal roots drought stressed with control well-watered mycorrhizal.

Table S17 Downregulated (\log_2 fold-change < -2) *H. almeriense* genes comparing mycorrhizal roots drought stressed with control well-watered mycorrhizal roots.

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