De novo gene birth, horizontal gene transfer and gene duplication as sources of new gene families associated with the origin of symbiosis in Amanita

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Abstract

By introducing novel capacities and functions, new genes and gene families may play a crucial role in ecological transitions. Mechanisms generating new gene families include de novo gene birth, horizontal gene transfer and neofunctionalization following a duplication event. The ectomycorrhizal (ECM) symbiosis is a ubiquitous mutualism and the association has evolved repeatedly and independently many times among the fungi, but the evolutionary dynamics enabling its emergence remain elusive. We developed a
phylogenetic workflow to first understand if gene families unique to ECM *Amanita* fungi and absent from closely related asymbiotic species are functionally relevant to the symbiosis, and then to systematically infer their origins. We identified 109 gene families unique to ECM *Amanita* species. Genes belonging to unique gene families are under strong purifying selection and are upregulated during symbiosis, compared to genes of conserved or orphan gene families. The origins of seven of the unique gene families are strongly supported as either *de novo* gene birth (two gene families), horizontal gene transfer (four), and gene duplication (one). An additional 34 families appear new because of their selective retention within symbiotic species. Among the 109 unique gene families, the most upregulated gene in symbiotic cultures encodes an ACC deaminase, an enzyme capable of downregulating the synthesis of the plant hormone ethylene, a common negative regulator of plant-microbial mutualisms.

**Keywords**

convergent evolution, gene family evolution, mycorrhizal fungi, phylogenomics, mutualism

**Significance statement**

Mutualisms between fungi and plants appear complex but have evolved repeatedly and independently many times. This convergent evolution is typically explained by gene
loss from fungi, but at the origin of a symbiosis new genes also appear in fungi: where do these new genes come from and what do they do? By systematically querying the origin of genes unique to symbiotic *Amanita* fungi and not found in close relatives we discover *de novo* gene birth, horizontal gene transfer, and gene duplication as the sources of a set of highly selected new genes upregulated during symbiosis.

**Introduction**

Evolutionary novelties are novel properties or features of organisms facilitating adaptation (Mayr 1963; Pigliucci 2008). The concept of an evolutionary novelty can connect dramatic changes in morphologies or phenotypes with ecological transitions in niche. New gene families, without apparent homologies in ancestors, may be considered as genetic evolutionary novelties because they are heritable features potentially shaping adaptations and niche transitions (Villanueva-Cañas *et al.* 2017). New gene families are thought to have three principal sources (Long *et al.* 2013; Andersson *et al.* 2015): *de novo* gene birth, horizontal gene transfer (HGT) and gene duplication (Figure 1). However, genes as evolutionary novelties remain understudied and the functions of many young gene families are unknown.

*De novo* gene birth involves the formation of protein-coding exons from ancestral non-coding loci. Newly evolved exons are typically shorter and bear weaker signatures of purifying selection compared to existing genes (Carvunis *et al.* 2012; Ruiz-Orera *et al.* 2015).
Often, *de novo* genes are identified by the absence of homologous genes in protein databases; hypothetical *de novo* genes are confirmed by aligning their DNA sequences against putatively homologous, non-coding sequences found in closely related species (Cai *et al.* 2008; Knowles & McLysaght 2009). A robust example of a *de novo* gene birth is the *BSC4* gene of *Saccharomyces cerevisiae*. Gene expression data suggest *BSC4* is not a pseudogene, and the sequence homologies between *BSC4* and syntenic but non-coding regions in closely related species confirm the *de novo* nature of this gene (Cai *et al.* 2008).

As mechanisms mediating the emergence of evolutionary novelties, HGT and gene duplication are better understood compared to *de novo* gene birth. HGT genes are often identified when the topologies between a species phylogeny and the phylogeny of a putative HGT gene family are inconsistent (Keeling & Palmer 2008; Husnik & McCutcheon 2018). Because HGT typically involves the movement of genes into distantly related lineages, HGT genes may have distinct properties compared to surrounding genes and preserve a degree of the donor genome's properties (Keeling & Palmer 2008). HGT is well documented in bacteria, while one of the most famous examples of HGT among eukaryotes is the HGT from a fungus to aphids (tribe Macrosiphini) (Moran & Jarvik 2010). The event enabled aphids to synthesize carotenoids (Moran & Jarvik 2010).
Gene duplication introduces paralogs of redundant sequence into a genome. Because they are copies, paralogs can escape the functional constraints of the original gene and undergo positive selection for new functions (Ohno 1970; Zhang 2003). A neofunctionalized gene copy will be more diverged from the originating gene compared to a copy which retains the same function (Assis & Bachtrog 2013). The relative timing of duplication events can be inferred by reconciling the gene tree with the species tree (Bansal et al. 2012; Jacox et al. 2016). A clear example of gene duplication and neofunctionalization involves the duplications of olfactory receptor genes in insects (Saad et al. 2018). Duplications created redundant paralogs and the paralogs evolved the ability to bind new ligands (Saad et al. 2018).

While de novo gene birth, HGT and gene duplication can each give rise to new gene families, their relative influence on genomes remains enigmatic. Nonetheless, these mechanisms have clearly shaped niche transitions. For example, the plant pathogen Pyrenophora tritici-repentis acquired its ToxA virulence gene through HGT from Stagonospora nodorum, enabling P. tritici-repentis to emerge as a devastating pathogen of wheat (Friesen et al. 2006). The transition of tetrapods from water to land was mediated by the duplication and neofunctionalization of HOX genes, resulting in the evolution of limbs from fins (Soshnikova et al. 2013). The evidence for de novo gene birth as a driver of niche transition is indirect. One potential example involves cnidarians (e.g. Hydra, jellyfish, coral, etc.); several genes involved in unique predatory behaviors
are cnidarian-specific, suggesting their origin is \textit{de novo} (Milde \textit{et al.} 2009). While associations between new genes and niche transitions have been explored in multiple systems, few have taken a whole genome approach. Genomics may enable the discovery of all genes, including previously unknown genes, associated with a transition event.

Ectomycorrhizal (ECM) symbioses stand among the most robust examples of niche transitions in nature, having evolved independently multiple times across the fungal kingdom (Matheny \textit{et al.} 2006; Tedersoo \textit{et al.} 2010; Bittleston \textit{et al.} 2016). ECM symbioses are mutualistic associations between fungi and plants and enable the exchange of nutrients and photosynthetically derived carbon. The associations can be identified by a morphological feature termed the Hartig net, which appears as hyphal growth between plant cortical cells (Smith & Read 2008). Research on ECM niche transitions focuses on gene loss, and gene loss appears to characterize diverse origins of the symbiosis (Kohler \textit{et al.} 2015; Peter \textit{et al.} 2016; Hess \textit{et al.} 2018; Murat \textit{et al.} 2018). Although the dynamic of gene loss may explain the repeated emergence of ECM symbiosis across distinct lineages, it does not resolve the mechanisms underpinning the evolution of the association (e.g. how ECM fungi suppress or endure plant immune responses). Gene gain is more rarely the focus of ongoing work, but gene gain may enable the formation of symbiotic structures and exchange of resources. For example, small secreted proteins (SSPs) appear to play a crucial role in fungal-plant communication and SSPs have a larger repertoire in at least some ECM species.
compared to asymbiotic species (Plett, Daguerre et al. 2014; Kohler et al. 2015). Other studies have identified additional gene gains associated with the transition from a saprotrophic to ECM niche, for example expansions in cytochrome P450 and berberine bridge enzyme gene families (Hess et al. 2018). Although multiple lines of evidence suggest a role for new genes in transitions to the ECM niche, the origins of these genes remain unknown.

The fungal genus *Amanita* is an emerging evolutionary model and ideal system to test for connections between evolutionary novelty and gene gain. A single, well-resolved niche transition marks the origin of ECM *Amanita* from asymbiotic ancestral lineages (Wolfe et al. 2012; Hess & Pringle 2014; Hess et al. 2018). While genomic restructuring within ECM *Amanita* does involve the loss of plant cell wall degrading enzymes (Wolfe et al. 2012; Hess et al. 2018), the presence of gene families found only in ECM *Amanita* suggests they may also play a role in mediating the niche transition (Hess & Pringle 2014; Hess et al. 2018). By taking a closer look at these novel gene families, we aim to decipher the genetic underpinnings of the ECM symbiosis. We hypothesize novel genes enabled new functions within the emerged ECM lineage and seek to understand their sources.

Our aims are to (1) explore whether gene families unique to ECM *Amanita* function in the symbiosis and (2) identify the putative origins of these gene families. We developed a phylogenetic workflow to investigate the properties and origins of unique
gene families, defined as genes only found in and shared by species of ECM Amanita. Analyses of transcriptomes and tests for selection support the hypothesis that unique gene families shaped the formation of the mutualism. Our workflow suggests all three gene acquisition processes were at play during the niche transition in Amanita, but HGT gave rise to the majority of new genes that retain enough signal for us to infer their origins.

Materials and Methods

Genome sequencing and annotation

The genomes of five Amanita species and one Volvariella species, including three ECM fungi (A. muscaria var. guessowii, A. brunnescens and A. polypiramis) and three asymbiotic fungi (A. inopinata, A. thiersii and V. volvacea), were used to identify gene families unique to symbiotic species. Genome sequencing and annotation is fully described in Hess & Pringle (2014) and Hess et al. (2018). Data of four of the genomes are available through NCBI's GenBank (Acc. JNHV02000000, JNHW02000000, JNHY02000000, JNHZ02000000) and all data and developed bioinformatic pipelines are available at https://doi.org/10.5061/dryad.g63c748.

ECM-specific orthologous gene family calling
We first identified homologous gene families among each of the six genomes using FastOrtho implementing MCL ver. 11.294 (van Dongen 2000) and BLASTp ver. 2.7.1 (Altschul et al. 1990) with default parameters. To investigate if the inflation value parameter affected results, we ran FastOrtho five additional times with different inflation values ranging from 1.2 to 6. We defined gene families unique to ECM Amanita as families for which homologs are present in all three ECM Amanita but not in any of the three asymbiotic fungi. A similar approach was used in Hess et al. (2018), but the resulting estimates are different from ours because different parameters were used.

**Identifying selection pressures on gene families unique to ECM Amanita**

We next sought to understand which gene families possess signals of purifying selection (dN/dS < 1). The putative protein sequences of all genes from each gene family were first aligned with MAFFT ver. 7.149b (Katoh et al. 2002) and then trimmed with trimAl ver. 1.4.rev15 (Capella-Gutiérrez et al. 2009) using default parameters. A phylogeny for each gene family was built with RAxML ver. 7.2.8 (Stamatakis 2006), using the trimmed protein alignment, and applying gamma rate heterogeneity and the best substitution model (either JTT, LG or WAG) as determined by AICc values calculated by ProtTest ver. 3.4 (Darriba et al. 2011). The DNA sequences of coding regions (CDS) in each gene family were also aligned based on a codon substitution matrix, using PRANK ver. 140603 (Loytynoja & Goldman 2005). Protein phylogenies and CDS alignments were
used to test the alternative hypothesis of dN/dS bias away from neutral selection using the codeml program implemented in PAML ver. 4.8 (model = 0, CodonFreq = 3, fix_kappa = 0, fix_omega = 0 vs 1) (Yang 2007).

The differential expression of gene families unique to ECM Amanita

To test if one of the ECM Amanita, A. muscaria, preferentially expresses genes unique to ECM species during symbiosis, we compared expression patterns of genes in each of three categories: (1) genes conserved across the six species (n=5264), (2) ECM-Amanita-unique genes (n=272) and (3) orphan genes (n=4989) (genes only found in A. muscaria var. guessowii). To assess if the proportion of genes upregulated in symbiosis (=the number of genes upregulated in symbiosis/total number of genes) is higher for ECM-Amanita-unique genes compared to conserved or orphan genes, we first identified all genes upregulated in symbiotic cultures. We retrieved the expression read count table generated from both symbiotic root tips and axenic cultures of A. muscaria var. muscaria from the JGI genome portal (project ID: 1025043) (Kohler et al. 2015). Because the expression data (used for training JGI’s genome annotation and generating expression table) are taken from A. muscaria var. muscaria but the genome assembly was generated from A. muscaria var. guessowii, we mapped our genome annotation (trained with transcriptome from A. muscaria var. guessowii) to the expression data by finding the best hit of each gene in our genome annotation to the
gene sequences used in the expression data with BLASTp (E-value = E-3). A Wald test was performed to screen for differentially expressed genes using the R package DESeq2 (Love et al. 2014). Upregulated genes were defined using FDR adjusted P-values < 0.01 and an expression level fold-change > 2, 4 or 8. The proportion of upregulated genes was compared across the three categories of gene families using Fisher’s exact test and an FDR correction for the P-values. We also performed the same analyses to assess if the genes belonging to any of the three categories mentioned above are upregulated in axenic culture.

**Identifying origins of gene families unique to ECM Amanita**

**Overview**

Before describing our workflow in greater detail in the sections below, we outline our basic approach: In each family unique to ECM *Amanita*, we selected the longest gene from the *A. muscaria var. guessowii* genome as a query to find homologous genes in an in-house, curated proteome database designed to represent the diversity of the three domains of life (see below and Supplementary file 1, 2) (Staehlin et al. 2016), using BLAST. We considered genes with no hits as candidates for de novo gene birth. Next, to identify HGT events from the remaining genes, we compared gene trees to the species tree to look for incompatibilities. If a gene was found in a monophyletic clade without *Amanita* species, the gene was considered as a candidate for HGT. After excluding
candidates for *de novo* gene birth and HGT, we identified potentially duplicated genes by looking for orthologs in asymbiotic species and ECM-paralogs in ECM *Amanita* (we define ECM-paralogs as the paralogs derived from duplications coinciding with the niche transition). If both orthologs and ECM-paralogs were found, the gene was considered as a candidate for duplication. Finally, we attributed genes whose homologs are only absent from asymbiotic *Amanita* and *Volvariella* to the phenomenon of selective retention (multiple independent deletion events in the asymbiotic lineages but not in ECM *Amanita*).

**De novo gene birth detection**

To identify putative homologs of the gene families unique to ECM *Amanita*, we curated an in-house genome database with 354 fungal (Supplementary file 1; last accessed: May 2015), 1153 prokaryotic (Staehlin *et al.* 2016), and 88 plant genomes (Supplementary file 2; last accessed: October 2017). Then, we compared the protein sequence representing the longest *A. muscaria* var. *guessowii* gene of each ECM unique gene family to the database using uBLAST implemented in uSearch ver. 8.0.1517 (Edgar 2010). To maximize the probability of finding potentially homologous sequences, we screened using a conservative E-value of E-3 and minimum identity of 0.25. If matches did not return sequences of all three ECM *Amanita* for a given gene probe, results were considered inconclusive and these genes were discarded from the analyses entirely.
When results consisted of only the three ECM species, and no other hits, gene families remained in consideration as possible *de novo* gene families. The probe sequences were aligned to the NCBI GenBank ([https://www.ncbi.nlm.nih.gov/genbank/](https://www.ncbi.nlm.nih.gov/genbank/)) and UniProt ([http://www.uniprot.org/](http://www.uniprot.org/)) databases using BLASTp with the same E-value cutoff of E-3 and an additional filter for low complexity regions to check for homologs in either database. Sequences without matches were considered as candidates for *de novo* gene birth.

Next, to explore whether identified gene families are potentially derived from non-coding sequence, we identified the syntenic block of each target gene by matching five upstream and five downstream genes across the three ECM genomes. The same upstream and downstream genes were identified in the three asymbiotic genomes to locate syntenic blocks and putatively homologous, non-coding sequences. The putatively homologous sequences were aligned to candidate *de novo* genes to explore synteny with MAFFT. We further used HISAT2 Galaxy Version 2.1.0 (Kim *et al*. 2015) with default parameters to map the Illumina transcriptomic raw reads sequenced from mycelia of asymbiotic species cultured in litter (*Amanita*) or potato dextrose broth (*V. volvacea*) (Acc. SRR089758, SRR619832, SRR7694628) (Bao *et al*. 2013; Hess *et al*. 2018) onto the three asymbiotic reference genomes to understand if the homologous regions of *de novo* genes in asymbiotic lineages are expressed.
**Horizontal gene transfer detection**

After excluding gene families categorized as stemming from *de novo* gene birth as well as gene families for which no strong conclusion could be made, we sought to identify gene families derived from HGT events using the putative homologs in our curated genome database. We first reconstructed a crude phylogeny for each gene to identify potential HGT events. We took uBLAST results and used OrthoMCL ver. 1.4 (Li *et al.* 2003) with an inflation value of 1.5 to cluster the uBLAST results returned for each gene and identify the genes in the same cluster with any target gene, eliminating highly diverged sequences. Refined uBLAST results were aligned with MAFFT and trimmed with trimAl using default parameters. (TrimAl failed to trim Family 12764, resulting an empty alignment, and so this family was not considered further). Preliminary phylogenies of the aligned and trimmed protein sequences were then constructed using FastTree ver. 2.1.7 (Price *et al.* 2010) and compared to the fungal taxonomy (Spatafora *et al.* 2017): genes of putative HGT families should nest within clades unrelated to *Amanita* (e.g. within the Ascomycota).

We next generated more accurate gene phylogenies for downstream analyses only with those gene families tentatively identified as resulting from HGT. Phylogenies were generated using a subset of sequences: for each putative HGT event, we identified branches with bootstrap support of > 90% housing between 100 to 350 sequences and including at least one gene from each of the three ECM *Amanita* species. These datasets
were aligned and trimmed again and used to generate new trees using RAxML (Stamatakis 2006) with best evolutionary models identified by ProtTest based on AICc values.

To rigorously reject the null hypothesis of vertical inheritance of genes, we compared our unconstrained trees with vertical-inheritance-constrained trees by using AU tests to identify the best phylogenetic model for each putative HGT family (Shimodaira 2002). We manually constructed the constraint trees by enforcing the null hypotheses of vertical inheritance (either by enforcing monophyletic Agaricales including *Amanita*, or monophyletic putative donor group) in Mesquite (Maddison & Maddison 2017) (Supplementary file 3). Each constraint tree was used to reconstruct a RAxML phylogeny. To test if the unconstrained phylogenies (suggesting HGT) were strongly favored over constrained phylogenies (indicative of divergence in accordance with speciation), we used the per-site likelihood values of both the original unconstrained and constrained phylogenies to perform AU tests implemented using CONSEL ver. 1.2 (Shimodaira & Hasegawa 2001).

Finally, we explored the gene structure (the intron sites) of genes associated with putative HGT events. We hypothesize the structures of putative HGT genes will be more similar to the genes from putative donors than to genes from Agaricales. To compare putative HGT genes with homologous genes from putative donors and homologous genes from Agaricales, we used GenePainter (Hammesfahr *et al.* 2013) to visualize
YAML-formatted gene structures generated with Webscipio (Hatje et al. 2011) and protein alignments from MUSCLE ver. 3.8.31 (Edgar 2004).

**Gene duplication**

We next tested whether remaining gene families (gene families not resulting from either *de novo* gene birth or HGT) might result from duplication and the subsequent rapid evolution of paralogs. We took a two-step approach: first, we looked for orthologs of putative rapidly evolving genes in asymbiotic species; next, we identified ECM-paralogs (paralogs derived from gene duplications coinciding with the niche transition). The closest asymbiotic ortholog to the new gene family served as an outgroup, allowing us to distinguish new gene families from ECM-paralogs among ECM species.

To identify orthologs in asymbiotic species, we generated a more robust phylogeny using RAxML with the uBLAST results generated for the HGT analysis. If the MCL-reduced dataset housed more than 800 sequences, the dataset was further trimmed using the FastTree trimming method described above; if the MCL-reduced dataset contained 30 to 800 sequences, the MCL-reduced dataset was used; if the MCL-reduced dataset consisted of fewer than 30 sequences, the entire dataset was used to reconstruct the phylogeny. To generate phylogenies we used MAFFT, trimAl, ProtTest and RAxML as above. Next, to identify potential orthologs in asymbiotic species, we first
rooted each phylogeny using its midpoint and then split each phylogeny (of each gene family) into single gene trees with TreeKO algorithm implemented in etetoolkit (Huerta-Cepas et al. 2010; Marcet-Houben & Gabaldón 2011). The TreeKO algorithm splits the phylogenies into multiple single gene trees by trimming off branches that represent duplication events until every species only has a single gene in any single gene tree, and therefore genes in each single gene tree can be treated as orthologs (Marcet-Houben & Gabaldón 2011). We analyzed single gene trees to test if each tree houses (1) only sequences from ECM species or (2) sequences from both ECM and asymbiotic species. Scenario (1) would suggest no orthologous genes can be found in asymbiotic species so selective retention in ECM species (multiple deletions in asymbiotic species) is the more parsimonious explanation of why these families being identified as families unique to ECM Amanita. Scenario (2) suggests the presence of orthologs in asymbiotic species.

Finally, to detect clear signals of gene duplications originating with the niche transition, we sought to identify ECM-paralogs associated with the transition to ECM niche. We first identified all nodes between (1) the most recent common ancestor (MRCA) of a given family (determined by FastOrtho), and (2) the MRCA of this family and the most phylogenetically proximate asymbiotic ortholog(s). We then identified ECM Amanita homologs diverging from the abovementioned intermediary nodes, and labeled these homologs as ECM-paralogs. If ECM-paralogs are present and no homolog
from other species clusters with ECM-paralogs, we consider the origin of the family to be gene duplication and look for a bootstrap value \( \geq 80 \) supporting the ECM-paralogs.

To test if the duplicated genes experienced novel selection pressure, we trimmed off the tips that are not from *Amanita* and *Volvariella* in the phylogeny of each gene family and labeled the branch of duplication as “foreground.” The phylogeny and their CDS alignment from PRANK was then analyzed by aBSREL (Smith et al. 2015), implemented in HYPHY ver. 2.5.1 (Kosakovsky Pond et al. 2005), to test if the duplication branch (foreground branch) has a proportion of codons that has experienced positive selection (\( dN/dS > 1 \)).

**Selective retention in asymbiotic Amanita and Volvariella**

Gene families not fitting into any criterion described above were considered as the results of selective retention (multiple deletions in asymbiotic species). To be stringently considered as a gene family that has undergone selective retention, uBLAST results of genes within a given family must not contain any of the three asymbiotic species.

**Results**

**Number and properties of unique gene families**

The number of gene families found only in the three ECM *Amanita* genomes ranged from 89 to 120. Changing the parameter settings for FastOrtho greatly impacted
the number of gene families identified as unique. The total number of gene clusters increased from 8,694 to 11,436 as the inflation value increased (Supplementary file 4). Because we have no prior knowledge of gene function we decided to use the default inflation value of 1.5, which balances sensitivity and selectivity and fits enzyme family nomenclature according to which reaction an enzyme catalyzes (EC annotation) (Li et al. 2003).

Using this inflation value we identified a total of 9,429 gene families and identified 109 gene families as unique to the three ECM *Amanita* genomes (Supplementary file 5). Among the 109 gene families, 107 are undergoing significant purifying selection ($d_N/d_S < 1$; LRT P-value $< 0.05$). Of the gene families experiencing purifying selection, values of $d_N/d_S$ range from 0.00256 to 0.8504 (Figure 2). The $d_N/d_S$ ratio provides evidence that genes from these gene families encode proteins and are not annotation artifacts. Two gene families had $d_N/d_S$ ratios close to one suggesting either the genes do not code for functional proteins or the genes are under neutral selection (in other words, natural selection does not influence the evolutionary trajectory of these genes).

When using a fold-change cutoff of four, a significantly higher proportion of genes in the 109 unique gene families were upregulated in ECM root tips, compared to orphan genes found only in *A. muscaria* and genes conserved across all six species. In addition, a significantly higher proportion of orphan genes were upregulated in ECM root tips, compared to genes in conserved gene families (Figure 3). However, the difference
between unique and orphan genes was not significant using a fold-change cutoff of two or eight (Figure 3A and C). The difference between unique and conserved genes was significant regardless of the fold-change cutoff. In axenic cultures, unique and orphan were upregulated compared to conserved families when any fold-change cutoff was applied (Supplementary file 6).

**De novo gene birth**

Based on uBLAST searches of our curated genome database, six families were identified as candidates for *de novo* gene birth. However, additional BLAST searches in the GenBank and UniProt databases detected putative homologs for four of these families, leaving only two gene families as candidates for *de novo* gene birth (families 1476 and 3446).

For each of the two gene families, each ECM *Amanita* species has only a single gene copy. The two putative *de novo* gene families have dN/dS ratios significantly lower than one (family 1476: 0.174 and family 3446: 0.198; codeml LRT P-value < 10\(^{-20}\)), suggesting the genes are experiencing strong natural selection. We returned to the transcriptomic data to probe expression patterns of the genes from these two gene families. The gene representing gene family 1476 is expressed constitutively in both symbiotic and axenic cultures (14 to 26 RPKM in each treatment; RPKM = reads per kilobase of exon model per million mapped reads). Transcripts of the gene representing the gene family 3446 are
detected but are not present at levels greater than 1 RPKM in any treatment. While the evidence suggests these are real genes, there is no evidence for the upregulation of either of the two gene families in ECM root tips.

Neither of the two genes has a known function. The lengths of the proteins are 178 (family 1476) and 297 (family 3446) amino acids. Genes from the two gene families have a GC content of 49.9% and 52.0%, respectively, and these GC contents more closely resemble the CDS of conserved genes (49.4%) compared to intergenic regions (46.0%) although each is presumably derived from an intergenic region (Supplementary file 7). In addition, each gene possesses at least one intron (gene 1476 from *A. muscaria* var. *guessowii* has two introns, but genes in the same family from *A. brunnescens* and *A. polypyramis* have only one intron). Introns are less commonly reported in genes derived from *de novo* gene birth, but there may be a bias because most research focuses on recently birthed genes. Our data suggest the evolutionary history of ECM *Amanita* is long enough that these two gene families acquired introns.

Gene family 1476 is located within a non-syntenic region. We hypothesize that this gene family is located in a relatively variable region. In contrast, genes from family 3446 are located within a conserved region across the three species (Figure 4). We attempted to find the homologous non-coding sequence of these genes by aligning homologous regions from ECM and asymbiotic species. However, the pairwise identities (the proportion of aligned nucleotides) of these multiple sequence alignments were low and
ranged from 42.0 to 53.3%. When we searched for evidence of expression of the homologous sequences in asymbiotic species, we found a few raw reads of this region in the transcriptome of all three asymbiotic species: *A. inopinata* (1 read), *A. thiersii* (3 reads) and *V. volvacea* (9 reads) (Supplementary file 8). The low abundance of transcripts from this region leaves open the question of whether this homologous, presumably non-coding region is actually transcribed by the fungus in nature. While no gene is present in these regions in *A. inopinata* and *A. thiersii* according to their annotations, three genes are annotated in the homologous region from *V. volvacea*, and one gene is responsible for its transcriptomic reads. However, these three genes show no homology with gene family 3446.

**Horizontal gene transfer**

Comparing the species tree and crude gene phylogenies generated by FastTree, six gene families emerged as candidates for HGT. After building the RAxML gene phylogenies, one gene family was no longer placed with the putative donor lineage and therefore we did not consider it further. Of the remaining five families, AU tests failed to reject vertical inheritance as a possibility for one of the gene families (7854). In families 11987 and 12806, AU tests rejected the two hypotheses which would suggest vertical inheritance: (1) all genes from Agaricales (including *Amanita*) forming a monophyletic group (P-values < 0.05) and (2) the genes from the putative donor forming a
monophyletic group (P-values < 0.01); these tests strongly suggest this family derived from HGT. In two other families, only one hypothesis suggesting vertical inheritance was rejected. In family 10418, only the first hypothesis was rejected (P-value = 0.001), whereas in family 2813, the second hypothesis was rejected (P-value = 0.01). However we were unable to test for the first hypothesis in family 2813 (no other Agaricales homologs were found) (Figure 5; Supplementary file 9). In summary, we consider each of these four families (2813, 10418, 11987 and 12806) to be the result of HGT. Gene structures provide additional evidence for the HGT of gene family 12806. In this family the ECM *Amanita* genes share more intron sites with homologs from putative donors than with homologs from other Agaricales species (Figure 6).

ECM *Amanita* genes in three of the four HGT families form monophyletic groups in the donor lineage. We hypothesize the genes in family 2813 do not form a monophyletic group because an insufficient phylogenetic signal leads to poorly resolved branches: bootstrap values supporting polyphyly range from 3 to 36. Each of the four families clusters inside Eurotiomycetes, or inside Ascomycota but with Eurotiomycetes as the sister group to the HGT genes. However, there is no evidence that the HGT genes are linked in either the donor lineage or *Amanita* genomes so there is no support for a single transfer of the four genes (e.g. as a gene cluster), which would be more parsimonious.
GO terms were assigned to three HGT families, including GO:0008660 (1-aminocyclopropane-1-carboxylate (ACC) deaminase activity), GO:0009310 (amine catabolic process), GO:0030170 (pyridoxal phosphate binding), GO:0003993 (acid monophosphatase activity), GO:0046872 (metal ion binding), GO:0016787 (hydrolase activity), GO:0016788 (esterase activity) (Supplementary file 5). All HGT gene families have a dN/dS < 1 (0.066 to 0.178) and are expressed. Only the gene from family 10418 (an ACC deaminase) is upregulated in symbiotic cultures, and it is 52-fold overexpressed.

**Gene duplications**

Orthologs of 16 remaining gene families were found in asymbiotic species, and clear evidence of ECM-paralogs is found for four of these 16 families. However, common ancestry with ECM-paralogs is supported by a bootstrap value ≥ 80 for only one family, family 1119 (Figure 7). But only two species (*A. brunnescens* and *A. polypyrampus*) possess these ECM-paralogs, in *A. muscaria* var. guessowii there are no family 1119 ECM-paralogs. The closest asymbiotic ortholog of this family is clustered with some but not all of the ECM-paralogs by FastOrtho, which suggests a divergence after duplication. We hypothesized that this family experienced positive selection shortly after gene duplication. However, we failed to reject the null hypothesis of an absence of positive selection (ABSREL LRT P-value = 0.063). In addition, these genes are not located in

http://mc.manuscriptcentral.com/gbe
syntenic regions and so we are unable to test if family 1119 is the more recent ECM-paralog.

This family is transcribed and annotated as a membrane protein (GO:0016020) (Supplementary file 5). In the transcribed product, no signal peptide was predicted by SignalP ver. 5.0 (Almagro Armenteros et al. 2019), 10 transmembrane domains were found by TMHMM ver. 2.0 (Krogh et al. 2001), and the product was predicted to be located on Golgi apparatus membranes by DeepLoc ver. 1 (Almagro Armenteros et al. 2017). HMMER 3.3 (Potter et al. 2018) predicted the family belongs to the DUF6 (domain of unknown function 6 or EamA) family. The limited evidence of positive selection and the ability to differentiate the functions of the newly duplicated genes and their paralogs prevents any inference of neofunctionalization following duplication for this family.

Selective retention

Thirty-four gene families unique to ECM Amanita appear to be the result of selective retention (equivalent to multiple deletions in asymbiotic lineages). Based on uBLAST and an E-value < E-3, these gene families lack homologs in the three asymbiotic species. False positive signals of selective retention may result from either our choice of E-value cutoff, HGT from unconsidered lineages, or false negatives in annotation. However, our estimate remains conservative because some selectively retained gene
families may house paralogs in asymbiotic species that diverged before the origin of the ECM symbiosis, and these families are not considered here.

Discussion

**The origins of gene families found only in symbiotic Amanita**

New gene families shape genome evolution and can drive adaptation to novel niches (Friesen et al. 2006; Milde et al. 2009). Among ECM Amanita, the genes of unique gene families (gene families found in the three ECM species but not found in closely related asymbiotic fungi) are upregulated in ECM root tips compared to the genes of conserved or orphan gene families, suggesting the new gene families acquired during the niche transition function during symbiosis. Most of the genes of unique gene families have also undergone strong natural selection.

We discovered evidence for the precise origins of 41 families. Thirty-four families are inferred as unique because of selective retention, and the other seven appear as truly new gene families, derived from either *de novo* gene birth (two), HGT (four) or divergence after gene duplication (one). The low number of *de novo* gene families is not surprising. The high turnover rate of *de novo* genes in any genome results in a low preservation of *de novo* families (Palmieri et al. 2014). Although the mechanism(s) driving HGT in fungi remain elusive, accumulating evidence suggests HGT is a key to evolutionary innovation in the fungal kingdom (Soanes & Richards 2014). Finding four
gene families derived from HGT suggests HGT was also critical to the changes in *Amanita* ecology, but this discovery might also reflect the better preservation of the phylogenetic signal of HGT over time, compared to signals from *de novo* gene birth or gene duplication. Unexpectedly, using our workflow we only identified one new gene family derived from gene duplication, contrary to expectations that gene duplication is a common source of new genes in fungi and other eukaryotes (Ohno 1970; Zhang 2003; Wisecaver *et al.* 2014). The discrepancy may result from the use of a Markov clustering algorithm (MCL) to identify new families, as the algorithm can consolidate paralogs into a single gene family if they have not undergone rapid divergence (Li *et al.* 2003), but the discrepancy may also result from a lack of phylogenetic support for the monophyly of paralogs. Changing the parameters used with MCL may influence how clusters are identified (Li *et al.* 2003). For example, while we identified 109 families as unique to ECM genomes Hess *et al.* (2018) identified 171 unique gene families. We used no match cutoff or identity cutoff and used an inflation value of 1.5, while Hess *et al.* (2018) used a match cutoff of 60%, an identity cutoff of 30%, and an inflation value of 3. Parameter choice involves a balance: choosing more stringent clustering results in greater numbers of clusters (Hess *et al.* 2018) and may lead to the identification of greater numbers of duplicated gene families. However, whether these gene families are functionally diverged enough to be judged as novel is an open question, and choosing higher cluster tightness can break orthologs into different families (Li *et al.* 2003). In summary,
the number of gene origins inferred reflects the number of gene family birth events, rates of gene turnover, the decay of phylogenetic signal and the choice of clustering algorithm parameters.

Research on the origins of new genes has focused on orphan genes found in single species and not gene families found across closely related species. Most of these orphan genes are more recent than the niche transition in *Amanita* (the origin of symbiosis among *Amanita* fungi dates to 80 million years ago (Varga et al. 2019)). By focusing on younger genes, these studies take advantage of opportunities to trace homologies and syntenies to elucidate the molecular mechanisms of the emergence of new genes (Donoghue et al. 2011; Arendsee et al. 2019). Moreover, HGT and duplication events are generally left out of their foci.

Other studies on gene family origins have used two major approaches to identify different evolutionary events. The first approach estimates the gene turnover rate with the gene counts of a family in different species, typically using gain-and-death (GD) or birth-death-and-innovation (BDI) models (Librado et al. 2012). The second approach uses a species tree/gene tree reconciliation method to identify duplication, gene loss and HGT (the DTL scenario) (Bansal et al. 2012; Jacox et al. 2016); other algorithms also incorporate incomplete lineage sorting (DTLI) (Stolzer et al. 2012). Because innovation events in the BDI model account for both *de novo* gene birth and HGT (Karev et al. 2002), this first approach does not provide information on the different mechanisms mediating
the origins of new gene families. On the other hand, the second approach does not account for \textit{de novo} gene birth (Tofigh \textit{et al.} 2011). In addition, methods using the DTL scenario require a well-curated species tree for not only the species of interest but also the putative donors for HGT (Bansal \textit{et al.} 2012; Jacox \textit{et al.} 2016) and these are not always available. To account for \textit{de novo} gene birth, HGT and gene duplication simultaneously, and to avoid a reliance on a well-curated species tree, our workflow includes (1) gene clustering to identify new gene families, (2) the search for homologs in a curated genome database, (3) phylogenetic analyses designed for different origin hypotheses, followed by (4) integration of additional support from analyses of e.g. gene structure, composition, and synteny (Gluck-Thaler & Slot 2015).

\textbf{De novo gene families have lost some properties of \textit{de novo} genes}

Interest in \textit{de novo} gene birth is growing. However, misidentification of \textit{de novo} genes can result from either the rapid evolution of an extant gene or a poorly resolved comparison between a putative \textit{de novo} gene and an incomplete genome database (Moyers & Zhang 2015). Casola (2018) suggests investigating four gene features to avoid misidentification and enable recognition of real \textit{de novo} gene families: the absence of homologs in other taxa, a lack of conserved domains, conserved synteny, and substitutions enabling genes encoding proteins (e.g. generating a start codon). We successfully identified the first three features for at least one family (3446). We were not
able to detect the last feature (substitutions enabling genes encoding proteins) because of the dissimilarity between the putative \textit{de novo} genes and their homologous non-coding sequences. Because \textit{de novo} gene families can evolve from sequences encoding long non-coding RNAs (Schlötterer 2015), we searched for transcripts of non-coding sequences orthologous to putative \textit{de novo} gene families in asymbiotic species. However, the low number of raw reads we discovered prevents firm conclusions as to whether or not these regions are transcribed by the fungi in nature.

The genes of \textit{de novo} gene families are reported to possess several distinctive characteristics, including short gene lengths (around 300 to 400 bp) and few to no introns (Wu & Knudson 2018). Sequences may be GC poor or have GC content similar to conserved genes, depending on the species (Palmieri \textit{et al.} 2014; Wu & Knudson 2018). However, the two \textit{de novo} gene families we identified have a structure and composition consistent with the conserved gene families in \textit{Amanita} genomes. Combined with the absence of similarities between \textit{de novo} gene families and their homologous non-coding sequences, these observations lead us to conclude that these gene families have had sufficient time to become ameliorated and now resemble other coding genes (Marri & Golding 2008).

\textit{De novo} gene families of animals and plants have low gene expression levels and higher tissue-specific expression compared to older genes (Schlötterer 2015), especially genes expressed in animal testes (Begun \textit{et al.} 2007). One of the two \textit{de novo} families we
identified has a higher expression rate in axenic mycelium compared to ECM root tips whereas the other is not differentially expressed so these genes are unusual compared to the genes of the other 109 gene families. We hypothesize that these gene families are not directly involved in the plant-fungal interaction, but with other processes shared among ECM *Amanita* species.

**Genes horizontally transferred to ECM *Amanitas* are from Ascomycota, and we hypothesize ACC deaminase (family 10418) was directly involved in the transition to symbiosis in *Amanita***

As with identification of *de novo* gene birth, strategies to identify HGT using only similarity as a criterion are also problematic (Guindon & Perrière 2001). Keeling and Palmer (2008) have suggested gene-species phylogenetic incongruence as the gold standard for HGT detection. Based on that gold standard, we identified four unique gene families originating from HGT. In two families, we successfully rejected the null hypotheses that (1) genes from Agaricales and *Amanita* are monophyletic and (2) the genes from donor group itself are monophyletic, but we failed to reject one of the null hypotheses for the other two families. The failure to reject all null hypotheses may be caused by accumulated substitutions in transferred genes, or too few taxa in our curated genome database. In addition, one gene family showed a high similarity of exon/intron structure to its putative donor, providing further evidence for HGT. These four gene
families are not the first record of HGT to ECM *Amanita*. We previously reported that genes of carbohydrate esterases family 1 (CE1s) were transferred to ECM *Amanita* from bacteria (Chaib De Mares et al. 2015), but these genes are not found in *A. polypyriramis* and thus do not fit our current definition of gene families unique to ECM *Amanita*.

Interestingly, all four HGT families are inferred to have been transferred from Ascomycota and specifically from Eurotiomycetes. Multiple HGTs to a single lineage from the same donor have been reported before and are usually enabled by their transfer as a gene cluster or syntenic block (Slot 2017), but "highways" of HGT between lineages have also been inferred (Qiu et al. 2016). In the case of HGT to ECM *Amanita*, no evidence of physical linkage in either *Amanita* or the putative donors was found. The lack of synteny suggests either these genes were transferred independently or the genes migrated into different genomic locations after HGT.

HGT genes may facilitate adaptation to new niches among the fungi (Soanes & Richards 2014). Of the four gene families transferred to ECM *Amanita*, we consider the family encoding ACC deaminase as the best candidate for driving niche transition because of its expression patterns and putative function. The gene from this family is upregulated in ECM root tips more than 52-fold, which is the highest fold difference among all genes from the 109 gene families. ACC deaminase can remove the amine group from an ACC and produce α-ketobutyrate (Honma & Smmomura 1978). ACC is the immediate precursor of ethylene, and ACC deaminase can therefore inhibit the
ethylene signaling pathway, a negative regulatory pathway of ECM symbioses (Plett, Khachane et al. 2014). In fact, the ACC deaminase knockouts of bacteria involved in a similar mutualistic system, nitrogen fixing rhizobia, are less capable of nodulation compared to the wild types (Nascimento et al. 2016). In addition, in arbuscular mycorrhizal fungi, the SP7 gene also inhibits the ethylene signaling pathway (Kloppholz et al. 2011). We hypothesize the ACC deaminases of ECM Amanita reduce the concentration of ACC and ethylene in ECM roots during symbiosis, and the reduction of ethylene in ECM root tips enables the lateral branching of ECM roots and formation of the Hartig net. Moreover, our finding may provide new evidence of a molecular convergence among mycorrhizal and root-nodulating associations. Lastly, we note ACC deaminase genes are also found in other ECM Amanita genomes not included in our analyses, for example A. bisporigera, A. phalloides and A. jacksonii etc. (van der Nest et al. 2014; Pulman et al. 2016).

**No significant evidence for positive selection on newly duplicated genes**

Many established algorithms are available to detect gene duplication. However, because we are specifically interested in potential duplication events coinciding with niche transition, we chose to detect the orthologs and ECM-paralogs of each of our identified unique gene families. Using this strategy, the only family duplicated during niche transition has a closest ortholog in A. thiersii and ECM-paralogs in A. brunnescens...
and *A. polypyramis*. Multiple evolutionary scenarios can explain the phylogenetic topology (e.g. duplication before the MRCA of *A. inopinata* and ECM *Amanita*, followed by deletion of both homologs in *A. inopinata*), but a single deletion in *A. inopinata* and duplication along the branch leading to niche transition is the most parsimonious explanation. We are also able to detect homologs in *A. inopinata* and *V. volvacea*, but these homologs are from diverged phylogenetic clusters, suggesting the family possesses a dynamic evolutionary background.

Gene duplication provides functional redundancy and paralogs often experience novel selection pressure, undergoing neofunctionalization (Saad *et al.* 2018). An extreme case suggests asymmetric evolutionary rates between two paralogs (or ohnologs to be precise) in yeast (Byrne & Wolfe 2007). However, in the new gene family derived from duplication in ECM *Amanita*, we failed to detect signals of positive selection along the branch leading to the new gene family cluster. We hypothesize synonymous substitutions have reached saturation at selected sites and the substitutions have removed the trace of positive selection (Gharib & Robinson-Rechavi 2013). However, it is also possible the family emerged as a result of non-selective events.

The gene family stemming from a recent duplication encodes two DUF6 domains in the form of a 5 + 5 transmembrane protein (each “5” is one DUF6 domain). Many proteins with this configuration are transporters, e.g., O-acetylserine/cysteine export proteins and nucleotide sugar transporters (Västermark *et al.* 2011), but at least one
gene, *PecM*, is involved in the degradation of pectin and cellulose (Rouanet & Nasser 2001). Although this gene family was identified as a new family, conserved in ECM *Amanita*, there is no evidence for its differential expression. We hypothesize that if this family is not solely the result of stochastic evolution, this gene family could be involved in transmembrane transport, substrate degradation by absorptive hyphae or controlled post-transcriptionally.

**Conclusion**

We developed a workflow for identifying the origins of new gene families unique to symbiotic fungi and not found in closely related free-living fungi. Among the 109 new gene families present in ECM *Amanita*, two, four and one families appear derived from *de novo* gene birth, HGT and gene duplication, respectively, but 34 families only appear new due to selective retention in symbiotic species. The genes of gene families unique to ECM *Amanita* are upregulated during symbiosis and are likely functionally relevant to the symbiosis. The horizontally transferred gene encoding ACC deaminase is potentially crucial to the mutualistic relationship, possibly regulating the immune response in plant symbionts. Our findings suggest a new possibility for ECM evolution: the transition to ECM niche in fungi is not only driven by gene loss but by coincident new gene acquisition as well.
Data availability: Scripts and additional supplementary data are deposited in Dryad under accession doi:10.5061/dryad.s4mw6m94f.

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References

Almagro Armenteros JJ et al. 2019. SignalP 5.0 improves signal peptide predictions using deep neural networks. Nat. Biotechnol. 37:420–423. doi: 10.1038/s41587-019-0036-z.

Almagro Armenteros JJ, Sønderby CK, Sønderby SK, Nielsen H, Winther O. 2017. DeepLoc: prediction of protein subcellular localization using deep learning. Bioinformatics. 33:3387–3395. doi: 10.1093/bioinformatics/btx431.

Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search tool. J. Mol. Biol. 215:403–410. doi: 10.1016/S0022-2836(05)80360-2.
Andersson DI, Jerlström-Hultqvist J, Nasväll J. 2015. Evolution of new functions de novo and from preexisting genes. Cold Spring Harb. Perspect. Biol. 7:a017996. doi: 10.1101/cshperspect.a017996.

Arendsee Z et al. 2019. Fagin: synteny-based phylostratigraphy and finer classification of young genes. BMC Bioinformatics. 20:440. doi: 10.1186/s12859-019-3023-y.

Assis R, Bachtrog D. 2013. Neofunctionalization of young duplicate genes in Drosophila. Proc. Natl. Acad. Sci. 110:17409–17414. doi: 10.1073/pnas.1313759110.

Bansal MS, Alm EJ, Kellis M. 2012. Efficient algorithms for the reconciliation problem with gene duplication, horizontal transfer and loss. Bioinformatics. 28:i283–i291. doi: 10.1093/bioinformatics/bts225.

Bao D et al. 2013. Sequencing and comparative analysis of the straw mushroom (Volvariella volvacea) genome. PLoS One. 8: e58294. doi: 10.1371/journal.pone.0058294.

Begun DJ, Lindfors HA, Kern AD, Jones CD. 2007. Evidence for de novo evolution of testis-expressed genes in the Drosophila yakuba/Drosophila erecta clade. Genetics. 176:1131–1137. doi: 10.1534/genetics.106.069245.

Bittleston LS, Pierce NE, Ellison AM, Pringle A. 2016. Convergence in multispecies interactions. Trends Ecol. Evol. 31:269–280. doi: 10.1016/j.tree.2016.01.006.

Byrne KP, Wolfe KH. 2007. Consistent patterns of rate asymmetry and gene loss indicate widespread neofunctionalization of yeast genes after whole-genome duplication. Genetics. 175:1341–1350. doi: 10.1534/genetics.106.066951.
Cai J, Zhao R, Jiang H, Wang W. 2008. De novo origination of a new protein-coding gene in *Saccharomyces cerevisiae*. Genetics. 179:487–496. doi: 10.1534/genetics.107.084491.

Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. 2009. TrimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. Bioinformatics. 25:1972–1973. doi: 10.1093/bioinformatics/btp348.

Carvunis AR *et al.* 2012. Proto-genes and de novo gene birth. Nature. 487:370–374. doi: 10.1038/nature11184.

Casola C. 2018. From de novo to “de nono”: the majority of novel protein-coding genes identified with phylostratigraphy are old genes or recent duplicates. Genome Biol. Evol. 10:2906–2918. doi: 10.1093/gbe/evy231.

Chaib de Mares M *et al.* 2015. Horizontal transfer of carbohydrate metabolism genes into ectomycorrhizal *Amanita*. New Phytol. 205:1552–1564. doi: 10.1111/nph.13140.

Darriba D, Taboada GL, Doallo R, Posada D. 2011. ProtTest 3: fast selection of best-fit models of protein evolution. Bioinformatics. 27:1164–1165.

van Dongen S. 2000. Graph clustering by flow simulation. doi: 10.1016/j.cosrev.2007.05.001.

Donoghue MT, Keshavaiah C, Swamidatta SH, Spillane C. 2011. Evolutionary origins of Brassicaceae specific genes in *Arabidopsis thaliana*. BMC Evol. Biol. 11:47. doi: 10.1186/1471-2148-11-47.
Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and high
throughput. Nucleic Acids Res. 32:1792–1797. doi: 10.1093/nar/gkh340.

Edgar RC. 2010. Search and clustering orders of magnitude faster than BLAST.
Bioinformatics. 26:2460–2461. doi: 10.1093/bioinformatics/btq461.

Friesen TL et al. 2006. Emergence of a new disease as a result of interspecific virulence
gene transfer. Nat. Genet. 38:953–956. doi: 10.1038/ng1839.

Gharib WH, Robinson-Rechavi M. 2013. The branch-site test of positive selection is
surprisingly robust but lacks power under synonymous substitution saturation and
variation in GC. Mol. Biol. Evol. 30:1675–1686. doi: 10.1093/molbev/mst062.

Gluck-Thaler E, Slot JC. 2015. Dimensions of horizontal gene transfer in eukaryotic
microbial pathogens. PLoS Pathog. 11:e1005156. doi: 10.1371/journal.ppat.1005156.

Guindon S, Perrière G. 2001. Intragenomic base content variation is a potential source of
biases when searching for horizontally transferred genes. Mol. Biol. Evol. 18:1838–1840.
doi: 10.1093/oxfordjournals.molbev.a003972.

Hammesfahr B, Oдрonitz F, Mühlhausen S, Waack S, Kollmar M. 2013. GenePainter: a fast
tool for aligning gene structures of eukaryotic protein families, visualizing the
alignments and mapping gene structures onto protein structures. BMC Bioinformatics.
14:77. doi: 10.1186/1471-2105-14-77.
Hatje K et al. 2011. Cross-species protein sequence and gene structure prediction with fine-tuned WebScipio 2.0 and Scipio. BMC Res. Notes. 4:265. doi: 10.1186/1756-0500-4-265.

Hess J et al. 2018. Rapid divergence of genome architectures following the origin of an ectomycorrhizal symbiosis in the genus *Amanita*. Mol. Biol. Evol. 35:2786–2804. doi: 10.1093/molbev/msy179.

Hess J, Pringle A. 2014. The natural histories of species and their genomes: asymbiotic and ectomycorrhizal *Amanita* fungi. Adv. Bot. Res. 70:235–257. doi: 10.1016/B978-0-12-397940-7.00008-2.

Honma M, Smmomura T. 1978. Metabolism of 1-aminocyclopropane-1-carboxylic acid. Agric. Biol. Chem. 42:1825–1831. doi: 10.1080/00021369.1978.10863261.

Huerta-Cepas J, Dopazo J, Gabaldón T. 2010. ETE: a python environment for tree exploration. BMC Bioinformatics. 11:24. doi: 10.1186/1471-2105-11-24.

Husnik F, McCutcheon JP. 2018. Functional horizontal gene transfer from bacteria to eukaryotes. Nat. Rev. Microbiol. 16:67–79. doi: 10.1038/nrmicro.2017.137.

Jacox E, Chauve C, Szöllosi GJ, Ponty Y, Scornavacca C. 2016. EcceTERA: comprehensive gene tree-species tree reconciliation using parsimony. Bioinformatics. 32:2056–2058. doi: 10.1093/bioinformatics/btw105.
Karev GP, Wolf YI, Rzhetsky AY, Berezovskaya FS, Koonin E V. 2002. Birth and death of protein domains: a simple model of evolution explains power law behavior. BMC Evol. Biol. 2:18. doi: 10.1186/1471-2148-2-18.

Katoh K, Misawa K, Kuma K, Miyata T. 2002. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Res. 30:3059–3066. doi: 10.1093/nar/gkf436.

Keeling PJ, Palmer JD. 2008. Horizontal gene transfer in eukaryotic evolution. Nat. Rev. Genet. 9:605–618. doi: 10.1038/nrg2386.

Kim D, Langmead B, Salzberg SL. 2015. HISAT: a fast spliced aligner with low memory requirements. Nat. Methods. 12:357–360. doi: 10.1038/nmeth.3317.

Kloppholz S, Kuhn H, Requena N. 2011. A secreted fungal effector of Glomus intraradices promotes symbiotic biotrophy. Curr. Biol. 21:1204–1209. doi: 10.1016/j.cub.2011.06.044.

Knowles DG, McLysaght A. 2009. Recent de novo origin of human protein-coding genes. Genome Res. 19:1752–1759. doi: 10.1101/gr.095026.109.

Kohler A et al. 2015. Convergent losses of decay mechanisms and rapid turnover of symbiosis genes in mycorrhizal mutualists. Nat. Genet. 47:410–415. doi: 10.1038/ng.3223.

Kosakovsky Pond SL, Frost SDW, Muse SV. 2005. HyPhy: hypothesis testing using phylogenies. Bioinformatics. 5:676–679. doi: 10.1093/bioinformatics/bti079.
Krogh A, Larsson B, Von Heijne G, Sonnhammer ELL. 2001. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. J. Mol. Biol. 305:567–580. doi: 10.1006/jmbi.2000.4315.

Li L, Stoeckert CJ, Roos DS. 2003. OrthoMCL: identification of ortholog groups for eukaryotic genomes. Genome Res. 13:2178–2189. doi: 10.1101/gr.1224503.

Librado P, Vieira FG, Rozas J. 2012. BadiRate: estimating family turnover rates by likelihood-based methods. Bioinformatics. 28:279–281. doi:
10.1093/bioinformatics/btr623.

Long M, VanKuren NW, Chen S, Vibranovski MD. 2013. New gene evolution: little did we know. Annu. Rev. Genet. 47:307–333. doi: 10.1146/annurev-genet-111212-133301.

Love MI, Huber W, Anders S. 2014. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biol. 15:550. doi: 10.1186/s13059-014-0550-8.

Loytynoja A, Goldman N. 2005. From the cover: an algorithm for progressive multiple alignment of sequences with insertions. Proc. Natl. Acad. Sci. 102:10557–10562. doi: 10.1073/pnas.0409137102.

Maddison WP, Maddison DR. 2017. Mesquite: a modular system for evolutionary analysis. Version 1.0. doi: http://mesquiteproject.org.

Marcet-Houben M, Gabaldón T. 2011. TreeKO: a duplication-aware algorithm for the comparison of phylogenetic trees. Nucleic Acids Res. 39:e66. doi: 10.1093/nar/gkr087.
Marri PR, Golding GB. 2008. Gene amelioration demonstrated: the journey of nascent genes in bacteria. Genome. 51:164–168. doi: 10.1139/G07-105.

Matheny PB et al. 2006. Major clades of Agaricales: a multilocus phylogenetic overview. Mycologia. 98:982–995. doi: 10.3852/mycologia.98.6.982.

Mayr E. 1963. Animal species and evolution. Cambridge (MA): Harvard University Press. doi: 10.2307/1440881.

Milde S et al. 2009. Characterization of taxonomically restricted genes in a phylum-restricted cell type. Genome Biol. 10:R8. doi: 10.1186/gb-2009-10-1-r8.

Moran NA, Jarvik T. 2010. Lateral transfer of genes from fungi underlies carotenoid production in aphids. Science. 328:624–627. doi: 10.1126/science.1187113.

Moyers BA, Zhang J. 2015. Phylostratigraphic bias creates spurious patterns of genome evolution. Mol. Biol. Evol. 32:258–267. doi: 10.1093/molbev/msu286.

Murat C et al. 2018. Pezizomycetes genomes reveal the molecular basis of ectomycorrhizal truffle lifestyle. Nat. Ecol. Evol. 2:1956–1965. doi: 10.1038/s41559-018-0710-4.

Nascimento FX, Brígido C, Glick BR, Rossi MJ. 2016. The role of rhizobial ACC deaminase in the nodulation process of leguminous plants. Int. J. Agron. 2016. doi: 10.1155/2016/1369472.

van der Nest MA et al. 2014. Draft genomes of Amanita jacksonii, Ceratocystis albifundus, Fusarium circinatum, Huntiella omanensis, Leptographium procerum, Rutstroemia.
sydowiana, and Sclerotinia echinophila. IMA Fungus 5:472-486. doi: 10.5598/imafungus.2014.05.02.11.

Ohno S. 1970. Evolution by gene duplication. New York: Springer-Verlag. doi: 10.1007/978-3-642-86659-3.

Palmieri N, Kosiol C, Schlötterer C. 2014. The life cycle of Drosophila orphan genes. Elife. 3:e01311. doi: 10.7554/eLife.01311.001.

Peter M et al. 2016. Ectomycorrhizal ecology is imprinted in the genome of the dominant symbiotic fungus Cenococcum geophilum. Nat. Commun. 7:12662. doi: 10.1038/ncomms12662.

Pigliucci M. 2008. What, if anything, is an evolutionary novelty? Philos. Sci. 75:887–898. doi: 10.1086/594532.

Plett JM, DAGUERRE Y, et al. 2014. Effector MiSSP7 of the mutualistic fungus Laccaria bicolor stabilizes the Populus JAZ6 protein and represses jasmonic acid (JA) responsive genes. Proc. Natl. Acad. Sci. 111:8299–8304. doi: 10.1073/pnas.1322671111.

Plett JM, Khachane A, et al. 2014. Ethylene and jasmonic acid act as negative modulators during mutualistic symbiosis between Laccaria bicolor and Populus roots. New Phytol. 202:270–286. doi: 10.1111/nph.12655.

Potter SC et al. 2018. HMMER web server: 2018 update. Nucleic Acids Res. 46:W200–W204. doi: 10.1093/nar/gky448.
Pulman JA, Childs KL, Sgambelluri RM, Walton JD. 2016 Expansion and diversification of the MSDIN family of cyclic peptide genes in the poisonous agarics Amanita phalloides and A. bisporigera. BMC Genomics 17:1038. doi: 10.1186/s12864-016-3378-7

Price MN, Dehal PS, Arkin AP. 2010. FastTree 2 - approximately maximum-likelihood trees for large alignments. PLoS One. 5:e9490. doi: 10.1371/journal.pone.0009490.

Qiu H, Cai G, Luo J, Bhattacharya D, Zhang N. 2016. Extensive horizontal gene transfers between plant pathogenic fungi. BMC Biol. 14:41. doi: 10.1186/s12915-016-0264-3.

Rouanet C, Nasser W. 2001. The PecM protein of the phytopathogenic bacterium Erwinia chrysanthemi, membrane topology and possible involvement in the efflux of the blue pigment indigoidine. J. Mol. Microbiol. Biotechnol. 3:309–318.

Ruiz-Orera J, Verdaguer-Grau P, Villanueva-Cañas JL, Messeguer X, Albà MM. 2018. Translation of neutrally evolving peptides provides a basis for de novo gene evolution. Nat. Ecol. Evol. 2:890–896. doi: 10.1038/s41559-018-0506-6.

Saad R, Cohanim AB, Kosloff M, Privman E. 2018. Neofunctionalization in ligand binding sites of ant olfactory receptors. Genome Biol. Evol. 10:2490–2500. doi: 10.1093/gbe/evy131.

Schlötterer C. 2015. Genes from scratch - the evolutionary fate of de novo genes. Trends Genet. 31:215–219. doi: 10.1016/j.tig.2015.02.007.

Shimodaira H. 2002. An approximately unbiased test of phylogenetic tree selection. Syst. Biol. 51:492–508. doi: 10.1080/10635150290069913.
Shimodaira H, Hasegawa M. 2001. CONSEL: for assessing the confidence of phylogenetic tree selection. Bioinformatics. 17:1246–1247. doi: 10.1093/bioinformatics/17.12.1246.

Slot JC. 2017. Fungal gene cluster diversity and evolution. Adv. Genet. 100:141–178. doi: 10.1016/bs.adgen.2017.09.005.

Smith MD et al. 2015. Less is more: an adaptive branch-site random effects model for efficient detection of episodic diversifying selection. Mol. Biol. Evol. 32:1342–1353. doi: 10.1093/molbev/msv022.

Smith S, Read D. 2008. *Mycorrhizal symbiosis*. 3rd ed. Cambridge (MA): Academic Press. doi: 10.1016/B978-0-12-370526-6.X5001-6.

Soanes D, Richards TA. 2014. Horizontal gene transfer in eukaryotic plant pathogens. Annu. Rev. Phytopathol. 52:583–614. doi: 10.1146/annurev-phyto-102313-050127.

Soshnikova N, Dewaele R, Janvier P, Krumlauf R, Duboule D. 2013. Duplications of Hox gene clusters and the emergence of vertebrates. Dev. Biol. 378:194–199. doi: 10.1016/j.ydbio.2013.03.004.

Spatafora JW et al. 2017. The fungal tree of life: from molecular systematics to genome-scale phylogenies. Microbiol. Spectr. 5:5. doi: 10.1128/microbiolspec.FUNK-0053-2016.

Staehlin BM, Gibbons JG, Rokas A, O’Halloran TV, Slot JC. 2016. Evolution of a heavy metal homeostasis/resistance island reflects increasing copper stress in enterobacteria. Genome Biol. Evol. 8:811–826. doi: 10.1093/gbe/evw031.
Stamatakis A. 2006. RAxML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. Bioinformatics. 22:2688–2690. doi: 10.1093/bioinformatics/btl446.

Stolzer M et al. 2012. Inferring duplications, losses, transfers and incomplete lineage sorting with nonbinary species trees. Bioinformatics. 28:i409–i415. doi: 10.1093/bioinformatics/bts386.

Tedersoo L, May TW, Smith ME. 2010. Ectomycorrhizal lifestyle in fungi: global diversity, distribution, and evolution of phylogenetic lineages. Mycorrhiza. 20:217–263. doi: 10.1007/s00572-009-0274-x.

Tofigh A, Hallett M, Lagergren J. 2011. Simultaneous identification of duplications and lateral gene transfers. IEEE/ACM Trans. Comput. Biol. Bioinforma. 8:517–535. doi: 10.1109/TCBB.2010.14.

Vakirlis N et al. 2018. A molecular portrait of de novo genes in yeasts. Mol. Biol. Evol. 35:631–645. doi: 10.1093/molbev/msx315.

Varga T et al. 2019. Megaphylogeny resolves global patterns of mushroom evolution. Nat. Ecol. Evol. 3:668–678. doi: 10.1038/s41559-019-0834-1.

Västermark Å, Almén MS, Simmen MW, Fredriksson R, Schiöth HB. 2011. Functional specialization in nucleotide sugar transporters occurred through differentiation of the gene cluster EamA (DUF6) before the radiation of Viridiplantae. BMC Evol. Biol. 11:123. doi: 10.1186/1471-2148-11-123.
Villanueva-Cañas JL et al. 2017. New genes and functional innovation in mammals. Genome Biol. Evol. 9:1886–1900. doi: 10.1093/gbe/evx136.

Wisecaver JH, Slot JC, Rokas A. 2014. The evolution of fungal metabolic pathways. PLoS Genet. 10:e1004816. doi: 10.1371/journal.pgen.1004816.

Wolfe BE, Tulloss RE, Pringle A. 2012. The irreversible loss of a decomposition pathway marks the single origin of an ectomycorrhizal symbiosis. PLoS One. 7:e39597. doi: 10.1371/journal.pone.0039597.

Wu B, Knudson A. 2018. Tracing the de novo origin of protein-coding genes in yeast. MBio. 9:4. doi: 10.1128/mBio.01024-18.

Yang Z. 2007. PAML 4: phylogenetic analysis by maximum likelihood. Mol. Biol. Evol. 24:1586–1591. doi: 10.1093/molbev/msm088.

Zhang J. 2003. Evolution by gene duplication: an update. Trends Ecol. Evol. 18:292–298. doi: 10.1016/S0169-5347(03)00033-8.
Figures

Fig. 1. Three mechanisms of gene family emergence. A: *de novo* gene birth; B: HGT; C: gene duplication. Blue: species tree; Red: new gene families (arrow indicates origin); Black: homologous genes.
Fig. 2. Frequency distribution of the dN/dS ratios of the 109 gene families unique to ECM *Amanita*. Most gene families (98.2%) have experienced purifying selection with dN/dS ratios significantly lower than one (P-value < 0.05).
Fig. 3. The ratio of the number of genes upregulated in ECM symbiosis to the total number of genes for conserved (n=5264), unique (n=272) or orphan families (n=4989). Different fold cutoffs (two, four and eight) are used to define upregulation in subplots A, B and C. Fisher exact test P-value $< 10^{-4}$ for all cutoffs. Letters mark significant (adjusted P-value $< 0.05$).
Fig. 4. Syntenic structure between scaffold 17 of *A. muscaria var. guessowii* (ECM) and the scaffold 77 of *A. thiersii* (asymbiotic) highlighting the location of one gene (AmumRNA.3446.1; purple) putatively identified as deriving from *de novo* gene birth.

Light blue: scaffolds; Dark blue: exons; Black: introns; Brown: homologous genes (determined by BLASTx); Grey: non-alignable regions. Units: kbp.
Fig. 5. Protein trees of the four putative HGT gene families. A: family 2813 (with unknown function); B: family 10418 (ACC deaminase); C: family 11987 (metal binding acid phosphatase); D: family 12806 (with unknown function).
Fig. 6. Gene structures of the HGT family 12806 (with unknown function). The intron structures of ECM *Amanita* genes deriving from HGT are more similar to their putative donors’ homologs than to homologs belonging to other Agaricales members.
Fig. 7. Phylogeny of gene duplication in family 1119 (DUF6). Branches with bootstrap values $\geq 80$ are labeled. Arrow head: duplication generating family 1119.
Conserved Unique Orphan
Ratio of the number of genes upregulated in ECM symbiosis to the total number of genes

A. fold-cutoff: 2
B. fold-cutoff: 4
C. fold-cutoff: 8

Gene origin
Conserved Unique Orphan

a b c a b a b
