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Comparative genomics of pyrophilous fungi reveals a link between fire events and developmental genes

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Summary

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Forest fires generate a large amount of carbon that remains resident on the site as dead and partially 'pyrolysed' (i.e. burnt) material that has long residency times and constitutes a significant pool in fire-prone ecosystems. In addition, fire-induced hydrophobic soil layers, caused by condensation of pyrolysed waxes and lipids, increase post-fire erosion and can lead to long-term productivity losses. A small set of pyrophilous fungi dominate post-fire soils and are likely to be involved with the degradation of all these compounds, yet almost nothing is currently known about what these fungi do or the metabolic processes they employ. In this study, we sequenced and analysed genomes from fungi isolated after Rim fire near Yosemite National Park in 2013 and showed the enrichment/expansion of CAZymes and families known to be involved in fruiting body initiation when compared to other basidiomycete fungi. We found gene families potentially involved in the degradation of the hydrophobic layer and pyrolysed organic matter, such as hydrophobic surface binding proteins, laccases (AA1_1), xylanases (GH10, GH11), fatty acid desaturases and tannases. Thus, pyrophilous fungi are important actors to restate the soil's functional capabilities.

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Introduction

The frequency of massive, high-severity wildfires is increasing in the western United States and regions around the world due to fuel accumulations from long-term fire suppression strategies and from the lengthened fire season due to global warming (Westerling *et al.*, 2006). These fires have direct, adverse effects on soil carbon (C) stocks through combustion, but they have indirect, positive effects on soil carbon stocks through the production of pyrolysed organic matter (PyOM) (González-Pérez *et al.*, 2004). PyOM is chemically heterogeneous material; its composition and structure are determined by both the initial organic matter and the temperature at which it is produced (Whitman *et al.*, 2013; Gul *et al.*, 2015).

Another soil-chemical result of high intensity fires is the production of hydrophobic soils. These soils are thought to result from the deposition of partially pyrolysed waxes and lipids that are volitalized at high temperatures and condense at slightly lower temperatures in the upper layers of soil (Atanassova *et al.*, 2014). The production of hydrophobic soils correlates with higher risk of erosion and can therefore reduce site productivity for decades (Certini, 2005; Mainwaring *et al.*, 2013). Hydrophobic layers can persist for weeks, months or even years after a fire.

It seems likely that soil microbes have some role in the degradation of these unique post-fire soil carbon compounds, although this possibility has not yet been explored. While microbes are known to be able to mineralize a small fraction of PyOM, much of the PyOM is highly aromatic and relatively slow to decompose, and thus results in long-term C storage (Debano, 2000; González-Pérez et al., 2004). Nevertheless, the initial post-fire microbial community may determine the rates at which decomposition of PyOM occurs either by their direct actions or by their capture of more labile carbon compounds that could otherwise prime the decomposition of more recalcitrant forms of PyOM. Microbial decomposition of hydrophobic layers is another possibility that has not yet been explored, but it would be surprising if these energy-rich, concentrated layers were not targeted by microbes.

Pyrophilous (fire-loving) fungi are a well-known, ecological guild of fungi that are restricted to post-fire environments and fruit abundantly in months immediately

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following fire (Petersen 1970, Hughes *et al.*, 2020). Because of their predicable occurrence after fire, they almost certainly need to be able to interact with post-fire soil chemistry. Here we report on the genomic content of four such pyrophilous that are members of the Agaricomycetes. All four fungi are saprobic (i.e. they live on dead material), yet little is known about what such fungi feed on in these post-fire environments. All of these fungi were found and isolated into culture after the 2013 Rim fire near Yosemite Park in California, United States, and they, or closely related species, have been reported from other fires around the world (Bruns *et al.*, 2020).

Although many pyrophilous fungi are members of the Ascomycota, we have focused initially on those that are mushroom-forming fungi (Basidiomycota, Agaricomycetes) because these classes of fungi are important wood-degraders involved in global carbon cycling. In the last years, the Agaricomycetes have been of great interest in comparative genomic studies regarding organic matter degradation and as complex multicellular organisms that produce fruiting bodies relevant in agriculture and medicine. The genome sequencing of these fungi showed important components of lignocellulose decomposition and a repertoire of developmental genes involved in fruiting body development in mushroom-forming fungi (Ohm et al., 2010; Sakamoto et al., 2011; Sipos et al., 2017; Almási et al., 2019; Krizsán et al., 2019).

The genomes of Lyophyllum atratum, Coprinellus angulatus, Pholiota molesta and Crassisporium funariophilum were sequenced and reported here for the first time, and we link the environmental changes caused by fires events to fruiting body development and degradation of PyOM through a comparative genomics approach. Understanding the role of pyrophilous fungi in post-fire soils is essential, as they are likely to affect the fate of carbon storage in these environments, and may affect the productivity and recovery of burnt soil.

Results

Genome features and phylogeny of pyrophilous genomes

Genomes of pyrophilous species were sequenced using long-read Pacific Biosciences (PacBio) technology, assembled into 41–701 scaffolds with length ranging from 37–93 Mbp and annotated with predicted 13,637–25,937 gene models. BUSCO (Seppey *et al.*, 2019) analysis showed 93.9–98.7% completeness using the Agaricales database (Table 1).

During the assembly and annotation process, *L. atratum*, *C. angulatus* and *P. molesta*, secondary scaffolds were detected, suggesting that those genomes are functionally diploid (number of bases in secondary scaffolds >20%).

This is expected in DNA isolated from dikaryon fungi and sequenced using PacBio technology (Kües, 2000; Rhoads and Au, 2015). For all analyses of this work, we moved the secondary scaffolds and their respective annotation into a separate set and used the primary sets for a more accurate comparison. The primary and secondary tracks are available at MycoCosm (https://mycocosm.jgi.doe.gov) (Grigoriev et al., 2014).

We reconstructed the species phylogeny from 1134 single-copy orthologs (1 604 504 amino acid characters) using 29 Agaricomycetes taxa: 25 Agaricales, three Polyporales, and one Boletales (Fig. 1A). With high bootstrap support (> 90%), the tree shows a topology expected of an Agaricomycetes phylogenetic tree (Almási et al., 2019) and a broad distribution of pyrophilous species across the tree, with C. funariophilum placed outside, but near the family Strophariaceae. Pholiota molesta and C. funariophilum show similar genomic metrics when compared with other related white-rot fungi. However, C. angulatus, on the Psathyrellaceae clade, shows a slightly larger genome size and a higher number of gene models when compared with the other 29 fungi (Fig. 1B). Finally, L. atratum, the only species from the Lyophyllaceae family, follows a similar pattern of other white-rot fungi (Fig. 1B).

As seen in Fig. 1C, the core gene set has a low variation (mean ± standard deviation = 4392 ± 369 genes) among the 29 genomes in this study. On the other hand, the number of unique genes varies greatly, ranging from 410 genes (O. olearius) to 7294 (C. micaceus) and correlates with the proteome size ($R^2 = 0.78$). Regarding these unique genes (unassigned to clusters), pyrophilous genomes show numbers (3393 ± 2052) on par with other Agaricales genomes (3490 ± 1753). Enrichment analysis of PFAM domains on pyrophilous unique genes (P < 0.05, Fisher's exact test, Benjamini Hochberg adjusted P values, abbreviated as FET) showed eight domains enriched, seven of them in C. angulatus (Supplemental file 1) which have the largest genome in our dataset. Among these enriched PFAM domains, there are two cytosol aminopeptidases (PF00883 and PF02789), two ribonucleotide reductase (PF02867 and PF00317), two mannosyltransferases (PF16192, PF02366), a MIR domain (PF02815) and a carboxyl transferase (PF02626). Interestingly, 23 domains are more frequent (frequency > 50%, Supplemental file 1) in the unique dataset than in the rest of the genome with a high diversity of functions reflecting the metabolic repertoire present in pyrophilous fungi. To infer functions enriched in pyrophilous fungi, we analysed all PFAM domain counts across 29 fungal Agaricomycetes genomes, which revealed 48 significantly overrepresented domains and 21 underrepresented (FET, $P \le 0.05$) (Supplemental file 2).

Comparing the functional annotation normalized by proteome size (Fig. 1D), there is a trend of basal nodes of the tree showing higher counts, except for the small

Table 1. Summary statistics of pyrophilous genomes.

Species	Assembly size (Mbp)	No. of contigs	Coverage	N50	L50 (Mbp)	# genes	BUSCO (%)
Lyophyllum atratum CBS 144462	64.90	171	30.2X	15	1.21	15 046	97.7
Coprinellus angulatus CBS 144469	93.57	517	63.55X	56	0.48	25 937	96.6
Pholiota molesta CBS 144467	66.04	701	65.79X	65	0.25	17 825	93.9
Crassisporium funariophilum CBS 144457	37.12	41	120.54X	7	2.16	13 637	98.7

BUSCO represents the percentage of complete BUSCO models using the Agaricales dataset.

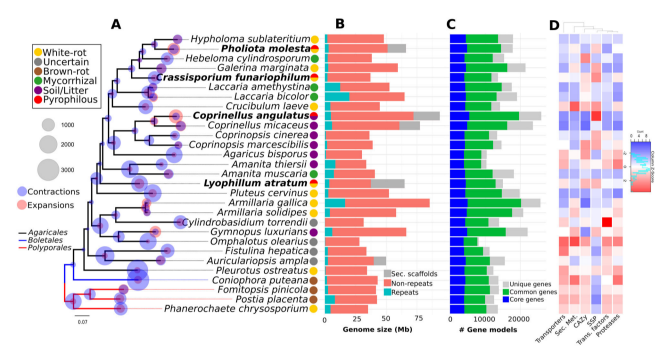


Fig. 1. Genome features of 29 Agaricomycetes genomes.

- A. Maximum likelihood tree showing the phylogenetic relationship based on 1134 single-copy orthologs of 29 Agaricomycetes fungi and their respective nutritional mode. Pyrophilous species are represented in bold. All support values are >90%. Circles in the nodes represent all familywide gene expansions (red) and contractions (blue) using CAFE.
- B. Genome size in Mbp. showing the distribution of repeats and non-repeats content.
- C. Gene model counts of each genome (only primary alleles are shown) divided into core genes (present in all genomes), common genes (present in two or more genomes) and unique genes (exclusively found on that genome).
- D. Column z-score heatmap of functional annotations normalized by total proteome size.

secreted proteins (with signal peptide and < 300 amino acids, abbreviated as SSP), which follows the opposite trend. Among different broad classes of annotation used in this study, the pyrophilous genomes do not show any clear distinctions. The only feature shared by them is the high number of SSP, but this expansion is not exclusive to pyrophilous fungi. SSPs are associated with several ecophysiological features of mushroom-forming fungi linked to developmental stages (Almási et al., 2019; Krizsán et al., 2019). We found two SSP domains - Hydrophobin (PF01185) and Hydrophobic surface binding protein (PF12296) – being enriched (FET, $P \le 0.05$) and having a strong negative correlation with proteome size in pyrophilous $(R^2 = -0.971 \text{ and } R^2 = -0.861 \text{ respectively})$ when compared with all genomes ($R^2 = 0.354$ and $R^2 = 0.147$ respectively) (Supplemental files 2 and 3). Other domains, like cutinase (PF01083) and alpha-L-arabinofuranosidase (PF6964), follow a similar pattern (Supplemental file 2).

Few CAZymes are correlated with pyrophilous lifestyle

To access the enrichment of CAZymes between different trophic modes, we performed a FET with FDR correction $(P \le 0.05)$ followed by contrast analysis (Felsenstein, 1985) and phylo-PCA (Revell, 2009) (Fig. 2A-C respectively). In pyrophilous genomes, six CAZy families are enriched: endo- α -1,4-polygalactosaminidase (GH114), gluco/chitooligosaccharide oxidase (AA7 dist), acetyl xylan

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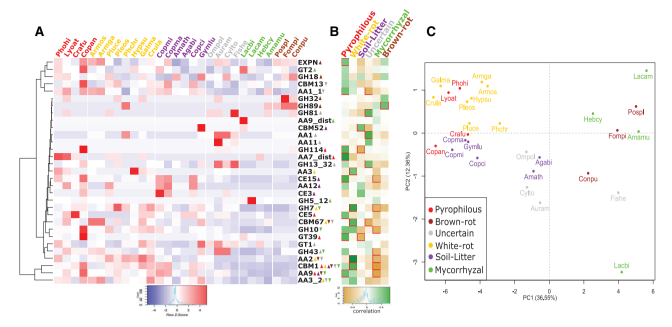


Fig. 2. CAZy families enriched on pyrophilous genomes (FDR < 0.05).

A. Heatmap (blue-red) of CAZy counts (z-score normalization). The arrows on each CAZy row represents whether that family is overrepresented (up-arrow) or underrepresented (down-arrow).

B. The heatmap (brown-green) shows the correlation of each cluster with trophic mode using the program contrast from phylip package (Felsenstein, 1985). Red outlines represent correlation values $\geq |0.4|$.

C. Phylogenetic principal component analysis of CAZy families enriched at least one nutritional mode.

esterase/cutinase (CE5), α-mannosyltransferase (GT39), lytic polysaccharide monooxygenases (LPMO/AA9) and cellulose-binding module (CMB1). There is a clear imbalance of gene counts for most families, except for AA9 and CBM1, which are enriched in all pyrophilous genomes. The families GH114 and GT39 are enriched mainly in C. angulatus, family AA7 dist in P. molesta and L. atratum, and CE5, mainly in C. funariophilum and L. atratum. In addition, we used the contrast program (Felsenstein, 1985) to correlate gene counts with nutritional mode (Fig. 2B). We highlighted families with correlation value > |0.4| using this threshold. We found eight families enriched in pyrophilous genomes: all six described earlier, distantly related to plant expansins (EXPN), laccase (AA1_1), and 4-O-methylglucuronoyl methylesterase (CE15). To visualize the enriched CAZy families in different trophic modes, we applied phylogenetically informed principal component analysis (phylo-PCA). Since pyrophilous fungi have other trophic modes (Fig. 1A), they clustered together with similar white-rot and soil/litter decomposers, and no clear separation was found (Fig. 2C).

Enriched clusters involved in the degradation of the hydrophobic zone and fruiting body development

Since the fungi isolated from post-fire soil are polyphyletic and show different trophic modes, the prediction of gene families related to this environment is not straightforward. In order to access orthologous clusters (or gene families) for these 29 fungi proteomes, the program Orthofinder v2.3.0 with default parameters (Emms and Kelly, 2015) was used to generate the count table used as input for Wilcoxon rank-sum test and contrast analysis for pyrophilous and non-pyrophilous fungi (Fig. 3). Comparing the distribution of protein count within each cluster of pyrophilous and other nutritional modes, 94 clusters were found statistically significant (rank-sums *P*-value ≤0.05) and contrast correlation ≥0.4 (Fig. 2A and B respectively). Among the clusters enriched on pyrophilous genomes, 70 showed at least one PFAM domain or presence of functional annotation (e.g. signal peptide), and 14 different clusters of SSP were enriched.

These families show a high diversity of functions and some are associated with fungal cell remodelling like α -mannosyltransferase (GT39), endo-1,3- β -glucosidase (GH17), chitinase (GH18) and exo- β -1,3-glucanase (GH55); transcription factors and transcriptional regulators: TFIIF, fungal Zn(2)-Cys(6), homeodomain, protein kinases, F-box proteins; and 14 SSP clusters. Comparing these results with developmentally regulated genes in other Agaricales genomes (Muraguchi *et al.*, 2015; Almási *et al.*, 2019; Krizsán *et al.*, 2019), we found that 67% of the enriched clusters are involved in at least one developmental stage (black circles in Fig. 3B and D). To confirm the specificity of these clusters to the pyrophilous lifestyle, we used phylo-PCA, showing a clear separation of pyrophilous genomes on the first two principal components (Fig. 3C).

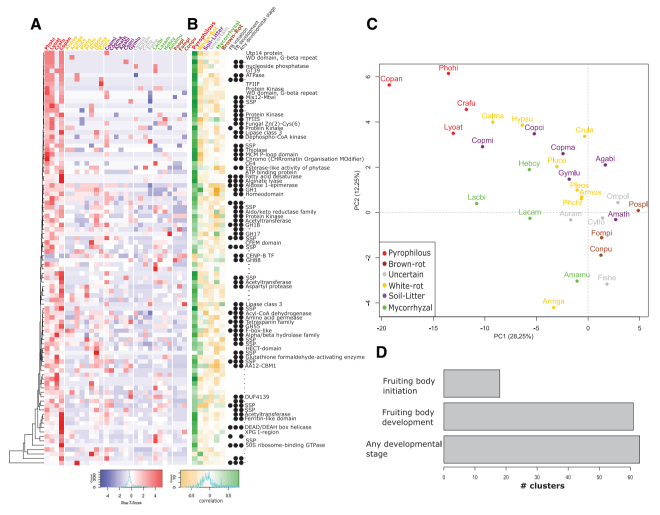


Fig. 3. Orthologous clusters enriched (P-value of Wilcoxon test $P \le 0.05$) on pyrophilous fungi.

- A. The heatmap (blue-red) shows z-score of gene copy number on each cluster detected using Orthofinder counts.
- B. The heatmap (yellow-green) shows the correlation of each cluster with trophic mode using the program contrast from phylip package (ref). Genes involved in developmental stages are shown as a black circle.
- C. Phylogenetic PCA of the same clusters showing a separation of pyrophilous species on both principal components.
- D. The number of clusters enriched and involved in developmental stages by the condition.

To perform an in-depth analysis of these developmentally regulated genes, we used proteinortho v6 tool (Lechner et al., 2011) using -synteny option to get as close as one-to-one gene orthologs with all four pyrophilous genomes and three Agaricales genomes with transcriptome data available (Krizsán et al., 2019) (Supplemental file 4). We decided to use Agaricales genomes since all pyrophilous fungi analysed here belong to this order. As seen in Fig. 4, about half of the gene space found differentially expressed in Coprinosis cinerea, Armillaria ostovae and Schizophyllum commune on developmental stages are shared with at least one pyrophilous genome. Among those genes, we can see that most of them are shared with all four pyrophilous genomes (UpSet diagram on Fig. 4A and B), and on average, 17.7% and 19.8% are expanded in pyrophilous genomes on the conditions fruiting body initiation and fruiting body development (Krizsán *et al.*, 2019) respectively.

The nodes of the phylogenetic tree on Fig. 1A show the number of gene families expanded and contracted (family-wide P < 0.01) using CAFE program (De Bie et al., 2006). Among the pyrophilous fungi, only *C. angulatus* and *P. molesta* showed a higher proportion of expansions/contractions (3.62 and 1.77 respectively). Looking at the rapidly evolving families (Viterbi P < 0.01) (Supplemental file 5), all pyrophilous genomes show a proportion of expansions/contractions >1, totalling 408 expanded and 100 contracted families. In addition, those CAFE expanded families are not only related to fruiting body development but potentially involved with the degradation of the hydrophobic layer – condensed waxes and lipids formed after the fire (Keiluweit et al., 2010; Bruns et al., 2020) – and

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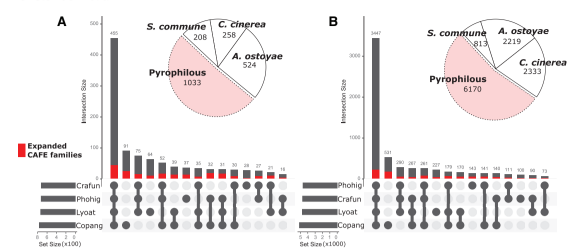


Fig. 4. Clusters involved in different developmental stages (Krizsán *et al.*, 2019) in Agaricales genomes and shared with at least one pyrophilous genome (pink on the pie chart). Each upset plot shows the distribution of respective pyrophilous slice. CAFE expanded clusters are shown in red for each intersection.

- A. Clusters that showed over four-fold upregulation at fruiting body initiation.
- B. Clusters that showed over four-fold expression dynamics (up- or downregulation) across the range of fruiting body stages.

pyrolysed biomass that is enriched in aromatics. Genes such as a laccase (AA1_1), xylanases (GH10, GH11), a fatty acid desaturase and a tannase and detoxification: DNA repair protein XPG_1, hsp20, p450, among others (Supplemental file 5) are examples.

Discussion

In this study, for the first time we sequenced, annotated and analysed four genomes from basidiomycetes/ Agaricales fungi isolated from post-fire soil and compared with 25 other fungi with different known nutritional modes in basidiomycetes (Riley et al., 2014; Almási et al., 2019). For this study, we considered pyrophilous as a nutritional mode. However, they show a similar arsenal of enzymes to degrade cellulose when compared with white-rot and soil/litter decomposer fungi (supplemental file 6). The purpose of this separation was to highlight that they are restricted to fruiting on post-fire soil and to explore whether the altered soil chemistry of post-fire soils correlates with changes in gene content. Their position in the phylogenetic tree follows what we expected based on the taxonomy and pyrophilous fungi are broadly distributed across different families rather than being members of a single clade; however, it is essential to note that while other ascomycetes fungi are often found in post-fire soil (El-Abyad and Webster, 1968; Bruns et al., 2020) (e.g., Pyronema spp.), we restricted the analysis to basidiomycetes.

We pinpointed 94 orthologous gene families enriched in pyrophilous genomes, 48 overrepresented PFAM domains, and another 508 rapidly evolving families in at least one pyrophilous fungal species. Among these gene

families and domains, we see several cell-wall modifying enzymes, various SSP, kinases, and transcription factors (TFs). Agaricomycetes-specific phylostrata of developmentally regulated genes are particularly enriched in F-box genes, TFs, and kinases, which indicates an increased rate of origin for these genes in mushroomforming fungi (Krizsán et al., 2019). We can see this pattern even more pronounced in pyrophilous fungi when compared with other Agaricomycetes (Fig. 3, supplemental file 2). This suggests that developmental processes in pyrophilous fungi can be even more complex than is expected from a regular mushroom-forming fungus.

Hydrophobins are a particularly important class of SSP, playing a role in development and morphogenesis in the majority of the filamentous fungi, influencing spore properties and fitness-related traits (Bayry *et al.*, 2012; Cai *et al.*, 2020). Interestingly, there is a slight enrichment of this domain on pyrophilous genomes (PF01185, P = 0.045 – Supplemental file 2), and a negative correlation of hydrophobin domain counts compared to total proteome size (pyrophilous $R^2 = -0.94$; all genomes $R^2 = 0.35$), suggesting the retention of these domains during its evolutionary history (Supplemental file 3). A similar pattern was found for hydrophobic surface binding protein (PF12296), cutinase (PF01083) and alpha-L-arabinofuranosidase (PF06964). The latter two are CAZymes that might be involved in the enzymatic arsenal to degrade PyOM.

Carbohydrate-active enzyme is an essential class of proteins in fungi since it is involved in the nutrient acquisition through degradation of biomass and cell wall transformation, among other specific functions (Lombard *et al.*, 2014). Our analysis results reveal that CAZy families involved with the degradation of plant biomass (a mix

of with PvOM in the context of this study), and cell wall remodelling were enriched in pyrophilous genomes. Some examples are CAZvmes active in the fungal cell wall glucan and chitins, like GH17, GH18, GH55, GH71 and GH88, all developmentally regulated (Krizsán et al., 2019). The degradation of PvOM and the hydrophobic layer are crucial to recovering the soil properties. Enzymes belonging to auxiliary activities (AA) such as laccases (AA1), peroxidases (AA2), and lytic polysaccharide monooxygenases (AA9) are candidates to perform this degradation. Laccases and peroxidases are well known to be active in phenolic compounds and highly recalcitrant environmental pollutants (Novotný et al., 2004: Viswanath et al., 2014). And finally, enzymes active on cellulose/ hemicellulose such as GH1, CBM1-containing protein, GH114 and GH10/GH11 were also enriched/expanded on pyrophilous fungi suggesting a role of these families processing the pyrolysed biomass. Interestingly, classic cellulases families like GH5, GH6, GH7, GH12 and GH45 were not enriched on pyrophilous fungi, which might be related to the low accessibility to cellulose due to the carbonization of plant biomass on post-fire environments.

Soil is an excellent insulator, and as a result, there is a steep drop in temperature with depth during a fire (Massman, 2012; Bruns et al., 2020). Peak soil temperature is predictable and drops as the log of soil depth. That means that within the span of only 15 cm, temperatures can change from over 500 °C at the surface to pre-fire ambient temperatures below. Pyrophilous fungi are likely to survive as spores or sclerotia at soil depths with intermediate temperatures that kill most of their competitors and allow them to rapidly colonize an open, post-fire niche (Bruns et al., 2020). All four of our sampled pyrophilous fungi were common fruiters after the 2013 Rim fire in California, and they are common throughout the Northern Hemisphere in prescribed fires and wildfire events (Bruns et al., 2020). Interestingly, a significant number of gene families involved in fruiting body development were found enriched on these genomes (Figs. 3 and 4).

Environmental conditions play a crucial role in the decision of whether a fruiting body will be formed. The optimal environmental for fruiting body development is often induced after drastically altering the environmental circumstances, and there is no universal set of conditions that leads to fructification in all fungi (Kües and Liu, 2000). Some environmental changes known to trigger this process are drop in temperature, nutrient depletion, shifts in light condition, pH, humidity, salinity and CO2 concentration. It seems counterintuitive, but fire events alter all these environmental parameters, and they can match the required triggers for fruiting body initiation, though not necessarily in the short term. Drop in temperature and CO2 concentration are good examples: not surprisingly, fire causes a rise in temperature and CO₂

levels, which cools down after it extinguishes. Therefore, these two artificial changes can work as a trigger; for instance. A. bisporus requires at least a drop of 5°C to start mushroom formation (Kües and Liu, 2000).

Another component is the formation of black carbon, commonly known as charcoal (González-Pérez et al., 2004). which can influence mushroom formation as well. Experiments with charcoal suggest that it induces fruiting body development by eliminating inhibitory compounds present in the substrate (De Groot et al., 1998). It is known that pH influences the fruiting body initiation, requiring neutral to slightly acidic pH (6.5-7) to trigger this process (Kües and Liu. 2000). The ashes formed burning increases soil pH (Certini, 2005), and considering that Yosemite Park is acidic with pH ranging from 4.5-6.5 (USDA, 2007), this rise in pH could be another factor inducing mushroom formation. The onset of fruiting body development correlates with the nutritional exhaustion of the growth substrates (Kües and Liu, 2000). The C/N ratios of soil after burning are usually lower than in the original soils, a phenomenon frequently cited in several types of post-fire soils (Almendros and Leal, 1990; González-Pérez et al., 2004), and nutrient depletion is an essential trigger for fruiting body initiation (Sakamoto, 2018; Almási et al., 2019).

It is important to note that pyrophilous fungi fruits are found only in burned habitats and are abundant in the first weeks, months or years after fire (Bruns et al., 2020, Hughes et al., 2020, Petersen, 1970), not necessarily right after the event. At first glance, the causal relationship between fire events and fruiting body development seems circumstantial. However, in this study, we sequenced and analysed genomes from fungi isolated after Rim fire at Yosemite Park and showed the enrichment/expansion of families known to be involved in fruiting body initiation when compared to other basidiomycete fungi. We found gene families potentially involved in the degradation of the hydrophobic layer. This is important because the hydrophobic layer can persist for several years and dramatically increases post-fire soil erosion, resulting in losses in productivity and long-term C storage (Mainwaring et al., 2013). Thus, pyrophilous fungi may be important actors to restate the soil's functional capabilities.

Methods

Isolation of fungi on post-fire soil and nucleotide extraction

All four fungi were isolated from tissue explants from inner pileus tissue of young mushrooms. Mushroom caps were ripped open to expose inner, uncontaminated tissue, and small pieces of these tissues were cut out with sterile scalpels, dipped into 30% H₂O₂ for 10 to 15 s and rinsed in sterile water before being pushed into 2% malt

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extract agar (MA) containing 50 mg/l Streptomycin sulphate and 50 mg/l chloramphenicol. After mycelium grew out into the agar, explants were transferred to 2% MA plates lacking antibiotics and maintained on this media.

All samples were grown in Petri dishes on cellophane over cornmeal yeast malt agar media at 25° C for 3–14 days, depending on the growth rate, and were allowed to experience light and dark cycles. The samples were ground by mortar and pestle in liquid nitrogen and frozen at -80° C.

RNA extraction was performed using the 'Fleming method' (Sessitsch et al., 2002; Fleming et al., 1998) modified by using as a lysis buffer a solution of 50% dH₂O and 50% Buffer RLT from the Qiagen RNeasy Mini Kit (Qiagen, Germantown, MD, USA) and precipitated by lithium chloride. DNA was extracted using the Qiagen Monarch Genomic DNA Purification Kit (Qiagen, Germantown, MD, USA). DNA and RNA samples were quantified by Qubit fluorospectrometer (Thermo Fisher Scientific, Waltham, MA, USA), and the quality of samples was assessed by 260:230 and 260:280 ratios using Nanodrop spectrophotometers (Thermo Fisher Scientific, Waltham, MA, USA). DNA and RNA were visualized via gel electrophoresis to confirm integrity. DNA fragments were size-selected using AmPure XP bead cleanup kit (Beckman Coulter, Indianapolis, IN, USA) or Blue Pippin (Sage Science, Beverly, MA, USA).

Genome sequencing and assembly

The genomes in this study (Lyophyllum atratum CBS 144462, Coprinellus angulatus CBS 144469, Pholiota molesta CBS 144467 and Crassipodium funariophilum CBS 144457) were sequenced using PacBio AMPure Bead Size Selection. Unamplified libraries were generated using the PacBio standard template preparation protocol for creating >10 kb libraries. Five microgram of gDNA (10 μg for Blue Pippin protocol) was used to generate each library, and the DNA was sheared using Covaris g-TubesTM to generate sheared fragments of >10 kb in length. The sheared DNA fragments were then prepared using PacBio SMRTbell template preparation kit, where the fragments were treated with DNA damage repair, had their ends repaired so that they were blunt-ended, and 5' phosphorylated. PacBio hairpin adapters were then ligated to the fragments to create the SMRTbell template for sequencing. The SMRTbell templates were then purified using exonuclease treatments and size-selected using AMPure PB beads. For Blue Pippin protocol, the SMRTbell templates were size-selected using the Sage Science BluePippin instrument with a 4 kb or 7 kb lower cutoff depending on DNA quality. In both protocols, PacBio Sequencing primer was then annealed to the SMRTbell template library, and sequencing polymerase

was bound to them using Sequel Binding kit 2.0. The prepared SMRTbell template libraries were then sequenced on a Pacific Biosystem's Sequel sequencer using v3 sequencing primer, 1 M v2 SMRT cells and v2.0 (2.1 for Blue Pippin protocol) sequencing chemistry with 6- and 10-h sequencing movie run times.

The genomes were assembled with Falcon version 1.8.8 (Chin et al., 2016) to generate an initial assembly. Lyophyllum atratum CBS 144462 passed through an extra step to remove contaminant data using kmer matching with bbtools bbduk.sh (k = 25 mm = f mkf = 0.05) (https://sourceforge.net/projects/bbtools/). The mitochondria were assembled separately from the Falcon pre-assembled reads (preads) using an in-house tool (assemblemito.sh), used to filter the preads and polished with Arrow version SMRTLink v5.0.1.9578 (https://github.com/PacificBiosciences/Genomic Consensus). A secondary Falcon assembly was generated using the mitochondria-filtered preads, improved with finisherSC v. 2.1 (Lam et al., 2015), and polished with Arrow version SMRTLink v5.0.1.9578.

Transcriptome sequencing and assembly

For transcriptomes, stranded cDNA libraries were generated using the Illumina Truseq Stranded RNA LT kit. mRNA was purified from 1 µg of total RNA using magnetic beads containing poly-T oligos. mRNA was fragmented and reversed transcribed using random hexamers and SSII (Invitrogen) followed by secondstrand synthesis. The fragmented cDNA was treated with end-pair, A-tailing, adapter ligation and eight cycles of PCR. The prepared library was quantified using KAPA Biosystem's next-generation sequencing library qPCR kit and run on a Roche LightCycler 480 real-time PCR instrument. The quantified library was then multiplexed with other libraries, and the pool of libraries was then prepared for sequencing on the Illumina HiSeq sequencing platform utilizing a TruSeq paired-end cluster kit v4, and Illumina's cBot instrument to generate a clustered flow cell for sequencing. Sequencing of the flow cell was performed on the Illumina HiSeg 2500 sequencer using HiSeq TruSeq SBS sequencing kits, v4, following a 2 x 150-bp indexed run recipe. For Lyophyllum atratum CBS 144462, the pool of libraries was prepared for sequencing on the Illumina NovaSeq 6000 sequencing platform using NovaSeg XP v1 reagent kits, S4 flow cell, following a 2 x 150-bp indexed run recipe.

Raw fastq file reads were filtered and trimmed using the JGI QC pipeline. Using BBDuk (https://sourceforge.net/projects/bbmap/), raw reads were evaluated for artefact sequence by kmer matching (kmer = 25), allowing one mismatch and the detected artefact was trimmed from the 3' end of the reads. RNA spike-in reads, PhiX reads and reads containing any Ns were removed.

Quality trimming was performed using the Phred trimming method set at Q6. Finally, following trimming, reads under the length threshold were removed (minimum length 25 bases or one-third of the original read length whichever is longer). Filtered fastg files were used as input for de novo assembly of RNA contigs. Reads were assembled into consensus sequences using Trinity (v2.3.2) (Grabherr et al., 2011). Trinity partitions the sequence data into many individuals de Bruijn graphs, each representing the transcriptional complexity at a given gene or locus and then processes each graph independently to extract full-length splicing isoforms and to tease apart transcripts derived from paralogous genes. Trinity was run with the --normalize reads (In-silico normalization routine) and --jaccard_clip (Minimizing fusion transcripts derived from gene dense genomes) options.

Genome annotation

All genomes were annotated using the JGI Annotation Pipeline (Grigoriev et al., 2014), which combines several gene predictions and annotation methods with transcriptomics data and integrates the annotated genomes into MycoCosm (https://mycocosm.jqi.doe.gov), a webbased fungal resource for comparative analysis (Grigoriev et al.. 2014). Completeness of genome annotation was accessed using BUSCO v4.0.6 (Seppey et al., 2019) using the agaricales odb10 database.

Phylogenetic analysis

Single-copy orthologs were identified in MCL clusters of the 29 Agaricomycetes, including brown rotters, ectomycorrhizal, saprotrophs/litter decomposers/organic matter degraders. white rotters and fungi of uncertain ecology. Protein sequences in the clusters were aligned using MAFFT (Katoh et al., 2019). Ambiguously aligned regions were removed using the -automated1 tag of Trimal (Capella-Gutiérrez et al., 2009). Maximum-likelihood (ML) inference was performed in igtree v1.6.9, using 5000 ultrafast bootstrap and partition model. We considered each gene as a partition (ModelFinder to select the best model) and used the partition-resampling strategy with -sampling option (Nguyen et al., 2015).

Identification of orthologous clusters

Orthogroups were identified using OrthoFinder v2.3.7 (Emms and Kelly, 2015). Two analyses were performed, one to delimit orthogroups across the 29 Agaricomycetes species and the second using proteinortho v6.0.10 (Lechner et al., 2011) with -synteny option to find a near one-to-one orthologous shared by pyrophilous genomes and other Agaricales with transcriptome during fruiting body development: A. ostoave. C. cinerea, and S. commune (Krizsán et al., 2019).

Functional annotation and principal component analysis

Functional annotation was done using INTERPROSCAN v5.35-74.0. CAZymes were annotated using the CAZy annotation pipeline (Lombard et al., 2014). Gene count heatmaps were created using the 'heatmap.2' function of gplots-R using z-score normalization. Hierarchical clustering with Euclidean distance and averaged-linkage clustering was carried out on copy-number using the 'hclust' function in R. SSPs were defined as proteins with <300 amino acids, having a signal peptide, an extracellular localization and no transmembrane domain using TMHMM 2.0 (Krogh et al., 2001). Proteins <300 amino acids long were subjected to signal peptide prediction through signalP5.0 (Almagro Armenteros et al., 2019) with the 'eukaryotic' option.

Phylogenetic PCA was performed on CAZyme copy numbers using the phyl.pca (Revell, 2009) function from PHYTOOLS. A copy number matrix normalized by proteome size (EMI 15273 Supplemental file 1) and the ML species tree were used as input. Independent contrasts were calculated under the Brownian motion model and the parameter mode = 'cov'.

Enrichment and CAFE analysis

The enrichment of PFAM domains was tested using FET and corrected for multiple testing by the Benjamini-Hochberg method in R. To identify orthologous clusters signal in pyrophilous fungi; we used Wilcoxon-test method in R. For both cases, $P \le 0.05$ was considered significant. To access gene family expansions and contraction analysis, we used CAFE v4.2.1 (De Bie et al., 2006), a tool that uses a stochastic birth and death process to model the evolution of gene family sizes over a phylogeny. We used the auto-estimation of λ values based on our dataset and a Viterbi $P \le 0.01$ to be considered a rapidly evolving family (Supplemental file 5).

Species abbreviations

Hypholoma sublateritium FD-334 SS-4 - Hypsu; Pholiota molesta CBS 144467 - Phohi; Hebeloma cylindrosporum h7 - Hebcy; Galerina marginata CBS 339.88 - Galma; Crassisporium funariophilum CBS 144457 - Crafu; Laccaria amethystina LaAM-08-1 -Lacam; Laccaria bicolor - Lacbi; Crucibulum laeve CBS 166.37 - Crula: Coprinellus angulatus CBS 144469 -Copan; Coprinellus micaceus FP101781 - Copmi; Coprinopsis cinerea - Copci; Coprinopsis marcescibilis CBS121175 - Copma; Agaricus bisporus var. bisporus

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H97 – Agabi; Amanita thiersii Skay4041 – Amath; Amanita muscaria Koide BX008 – Amamu; Lyophyllum atratum CBS 144462 – Lyoat; Pluteus cervinus NL-1719 – Pluce; Armillaria gallica 21–2 – Armga; Armillaria solidipes 28–4 – Armos; Cylindrobasidium torrendii FP15055 – Cylto; Gymnopus luxurians FD-317 M1 – Gymlu; Omphalotus olearius – Ompol; Fistulina hepatica ATCC 64428 – Fishe; Auriculariopsis ampla NL-1724 – Auram; Pleurotus ostreatus PC15 – Pleos; Coniophora puteana RWD-64-598 SS2 – Conpu; Fomitopsis pinicola FP-58527 SS1 – Fompi; Postia placenta MAD-698-R-SB12 – Pospl; Phanerochaete chrysosporium RP-78 – Phchr.

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Data availability statement

Genome assemblies and annotations are available at MycoCosm (https://mycocosm.jgi.doe.gov) (Grigoriev et al., 2014) and have been deposited at DDBJ/ENA/GenBank under the following accessions *L. atratum* CBS 144462 (JACGCN000000000), *C. angulatus* CBS 144469 (JACGCI000000000), *C. funariophilum* CBS 144457 (JACGCL000000000) and *P. molesta* CBS 144467 (JACGCK000000000).

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

Supplemental file 1. Enrichment analysis of unique genes on pyrophilous genomes.

Supplemental file 2. Enrichment analysis of PFAM counts from 29 Agaricomycetes genomes and rank-sums analysis.

Supplemental file 3. Hydrophobin correlation with proteome size and gene tree.

Supplemental file 4. Developmentally regulated genes on pyrophilous fungi.

Supplemental file 5. MCL clusters and rapidly evolving families detected using CAFE software.

Supplemental file 6. Distribution of CAZy families and substrate classification.