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Genome and secretome of *Chondrostereum purpureum* correspond to saprotrophic and phytopathogenic life styles

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Abstract

The basidiomycete Chondrostereum purpureum (Silverleaf fungus) is a saprotroph and plant pathogen commercially used for combatting forest "weed" trees in vegetation management. However, little is known about its lignocellulose-degrading capabilities and the enzymatic machinery that is responsible for the degradative potential, and it is not yet clear to which group of wood-rot fungi it actually belongs. Here, we sequenced and analyzed the draft genome of C. purpureum (41.2 Mbp) and performed a quantitative proteomic approach during growth in submerged and solid-state cultures based on soybean meal suspension or containing beech wood supplemented with phenol-rich olive mill residues, respectively. The fungus harbors characteristic lignocellulolytic hydrolases (GH6 and GH7) and oxidoreductases (e.g. laccase, heme peroxidases). High abundance of some of these genes (e.g. 45 laccases, nine GH7) can be explained by gene expansion, e.g. identified for the laccase orthogroup ORTHOMCL11 that exhibits a total of 18 lineage-specific duplications. Other expanded genes families encode for proteins more related to a pathogenic lifestyle (e.g. protease and cytochrome P450s). The fungus responds to the presence of complex growth substrates (lignocellulose, phenolic residues) by the secretion of most of these lignocellulolytic and lignin-modifying enzymes (e.g. alcohol and aryl alcohol oxidases, laccases, GH6, GH7). Based on the genetic and enzymatic constitution, we consider the 'marasmioid' fungus C. purpureum as a 'phytopathogenic' white-rot fungus (WRF) that possesses a complex extracellular enzyme machinery to accomplish efficient lignocellulose degradation during both saprotrophic and phytopathogenic life phases.



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Introduction

Wood-degrading organisms play an important role in carbon and nitrogen cycling. The most efficient ones are filamentous fungi, which can ecologically be categorized into brown-rot, white-rot and soft-rot fungi, colonizing compact wood (logs, branches, stumps) and degrading all polymeric cell wall components [1, 2]. Ascomycetous soft-rot fungi (SRF) erode the secondary wall or form discrete cavities within the cell wall where they break down mainly cellulose and hemicelluloses but have little or no effects on the wood lignin and the middle lamellae [3, 4]. Basidiomycetous brown-rot fungi (BRF) degrade primarily the polysaccharide components of wood and leave a partially modified lignin framework behind, whereas basidiomycetous white-rot fungi (WRF) decompose efficiently all cell wall components. The rate and extent of lignin, cellulose, and hemicellulose removal varies among white-rot species [5, 6]. They are considered to decompose the three wood components via the synergistic action of extracellular hydrolases and oxidoreductases classified within the CAZy database [7]. The class of Agaricomycetes includes numerous well-known WRF (wood and litter decomposers) and BRF, e.g. within the orders Polyporales (Trametes versicolor, Phlebia radiata, Bjerkandera adusta, Irpex lacteus), Gloeophyllales (Gloeophyllum trabeum) and Agaricales (Agaricus bisporus, Stropharia *coronilla*, *Agrocybe praecox*) [7].

Despite the crustous morphology of its fruiting body (basidiocarp), Chondrostereum purpureum (Pers. ex Fr.) Pouzar, belongs to the order Agaricales. It is a wood-decomposing fungus commonly found on broad-leaved trees in temperate and boreal vegetation zones. The fungus is saprotrophic mainly during the initial decomposition phase affecting felled trees or stumps. However, it also occurs as a secondary parasite causing the silverleaf disease in orchard trees [8, 9]. Spores can penetrate dead wood of living trees or on wounded wood where they grow and pair with other spores forming heterokaryotic mycelia that grows deeper into the wood [10]. When the hyphae of *C. purpureum* spread within a stump, tree vessels are occluded [11]. Induced dehydration combined with fungal toxins (sterpurenes, sesquiterpene metabolites) strengthens the adverse effects of the fungus in preventing the resprouting of stumps [11, 12]. The fungus consumes carbohydrates and during that process, it also decomposes lignin with the help of a large set of extracellular enzymes [1, 13-15]. When the decay process has penetrated deep enough into the wood, the ability of a stump to produce new sprouts diminishes and the stump dies. Later, C. purpureum is quickly replaced by other fungi [16]. Due to these abilities, C. purpureum has been used as a biocontrol agent against sprouting and root suckering of tree species such as red alder (Alnus rubra), black cherry (Prunus serotina), white birch (Betula papyrifera) and aspen (Populus spp.) [9, 17, 18].

Fruiting bodies of *C. purpureum* have a characteristic laminar-like shape and therefore, it was for a long time taxonomically assigned within the Polyporales (Meruliaceae; [18]). Matheny et al. [19] analyzed a larger phylogeny of agaric species, which resulted in the affiliation of *C. purpureum* within the Agaricales order, more precisely in the family of Cyphellaceae within the 'marasmioid clade' (comprising seven fungal families). However, other recent reports have claimed no strict consensus about the phylogenetic position of *C. purpureum*, being still included either within Polyporales or Agaricales [9, 20].

Beside the discrepancy in the phylogenetic affiliation of *C. purpureum*, there is only little known concerning its lignocellulolytic enzyme machinery, by which the fungus accomplishes its saprotrophic life style with that strong tendency to pathogenicity. In this context, it is yet not clear to which type of wood-rot fungi *C. purpureum* actually belongs. Next generation sequencing has enormously accelerated studies in fungal genomics and evolution, for example, the genomes of over 50 basidiomycetous fungi have been made available in 2014, including those of numerous Polyporales species [21]. Riley and coworkers [21] were able to separate

white-rot (and white-rot like fungi) *vs.* brown-rot species based on the content of lignocellulose decomposing genes. Kohler et al. [22] found dramatic lignocellulose-related gene losses in brown-rot and ectomycorrhizal species compared to white-rot species, and Hess et al. [23] observed a similar tendency within the genus *Amanita*. Floudas et al. [24], and recently Nagy and coworkers [25], used comparative genomics to trace back the origins of lignocellulose decay capabilities. Sipos et al. [26] used a similar approach to elucidate the 'signature' of the pathogenicity in the genus *Armillaria* and found a large repertoire of plant cell wall degrading enzymes and pathogenicity factors, which are seemingly involved in the severe tree pathogenicity of several *Armillaria* species.

Several recent studies have evaluated the gene inventory of saprotrophic fungi often in combination with transcriptomic and secretomic analyses. Thus, the secretomes of ligno-cellulose-degrading fungi were studied when growing on different complex media often based on lignocellulosic materials [27]. Among them were basidiomycetous fungi causing typical white or brown-rot in deadwood (e.g. *Phanerochaete chrysosporium*, [28–30]; *Pleurotus ostrea-tus*, [27] or *Serpula lacrymans*, [2]; *Postia placenta*, [31]), or so far unclassified/unspecific types of wood-rot (e.g. *Schizophyllum commune*, [24]) as well as phytopathogenic wood-decay fungi (i.e. facultative parasites such as *Heterobasidion irregulare*, [32] or *Armillaria mellea*, [33]).

From the biotechnological perspective, wood-degrading fungi and their enzymes are promising tools for the bioconversion of natural lignocellulose-containing polymers into renewable resources and feedstocks, e.g. chemicals and biofuels [34]. In that context, the fermentation of unused lignocellulosic by-products or their extractives, given in large quantities by various industrial processes manufacturing agricultural or forestal products, offers a promising approach to convert cheap, underutilized materials (e.g. rape straw, grain or olive mill residues) into useful final goods (e.g. base chemicals, fibers or fertilizers). An important preparing step of such biotechnological fermentation processes is the biological pretreatment of lignocellulosic materials, which reduces substances inhibiting fermentation (i.e. toxic, persistent and protecting plant ingredients like lignin, suberin, tannins and phenolics) via their enzymatic degradation and transformation [34–36]. For example, dry olive mill residue (DOR), a byproduct of a two-phase extraction process during olive oil manufacturing, is rich in organic matter and nutritionally relevant substances, which makes it attractive for an agronomic use. However, DOR contains significant amounts of phytotoxic ingredients. For that reason, fungal pretreatments of DOR to get rid of the toxic compounds have been intensively studied during the last years [37]. It was demonstrated that fungus-treated DOR enhances the growth of tomato plants and hence is applicable as a valuable organic fertilizer [38-40].

From the eco-physiological point of view, the addition of agricultural by-products to fungal cultures may reflect growth conditions in a complex environment close to nature and could therefore stimulate the secretion of enzymes required for degradation, 'digestion' and detoxification of lignocelluloses [38, 41, 42]. On the other hand, to our best knowledge, not much is known about the effects of natural phenolics, tannins and humic substances in leaf-litter, organic soil fractions, compost or agricultural wastes (e.g. DOR) on the protein expression profiles of fungi [43, 44].

Against this ecological and biotechnological background, it has been our intention–besides the general analysis of the *C. purpureum* genome–to analyze the secretomes of this fungus when growing on natural substrates (beech and birch wood), on the agricultural by-product DOR and on nutrient-rich soy medium. This approach may both help to deepen our understanding of the physiology and ecology of *C. purpureum* and to develop new fungus-based technologies.

Materials and methods

Dry olive mill residue (DOR)

DOR was obtained from the Sierra Sur olive oil company in Granada, Spain (2011–2012 harvest). It was sieved, autoclaved in three cycles and stored at 4°C before use. The aqueous extract of DOR (abbreviated as ADOR) was produced by a 1:2 (w:v) DOR:water extraction process lasting 8 h under orbital shaking (170 rpm) and subsequent centrifugation and filtration through glass fiber filters (Whatman no. 5) [41].

Fungal cultivation

The *C. purpureum* strain used was obtained from the German Collection of Microorganisms and Cell Cultures (DSMZ, Braunschweig, Germany) where it is deposited under DSM 4894. Pre-cultures were incubated at 24°C on 2% malt extract agar (MEA) over two weeks to obtain fresh inoculum. Solid-state fermentations (SFFs) with the fungus were performed in 250-mL Erlenmeyer flasks. Each flask contained 4 g of beech wood (BW) and 14 mL of distilled water. After autoclaving BW two times for 20 min at 121°C, the sterile wood was inoculated with 9 mL of a homogenized suspension from four fully overgrown agar plates in 80 mL sterile tap water. After the fungal mycelium had grown for one week at 25°C, half of the flasks were mixed with DOR (50% w:w) for preparing the DOR supplemented beech wood cultures (BWD). Sampling occurred weekly over an incubation time of seven weeks. To obtain the extracellular enzymes as well as the complete secreted protein profile, the harvested cultures were extracted with distilled water (1:5 w:v) by shaking on a rotatory shaker at 150 rpm for two hours. Extracts for label-free proteome analyses were prepared after seven weeks of cultivation by aqueous extraction, centrifugation and concentration *via* lyophilization.

To identify differences in the enzyme secretion pattern in dependence of the composition of the liquid media, submerged fermentation (SF) with *C. purpureum* was performed in 500 mL round-bottomed flasks using either 200 mL of complex soybean meal suspension (SM) or 200 mL of synthetic KIRK-medium (KM); the latter was prepared as described by [1]. SM was prepared with distilled water in a 3% ratio (w:v). To both culture media, ADOR (5% (v:v); ASKM & ASSM) as well as birch wood (1:2 (w:v); BSKM & BSSM) were supplemented after four days of fungal growth, respectively (Table 1).

The liquid cultures were incubated at 25°C and 100 rpm on a rotatory shaker. Every second or third day, samples were taken (1.5 mL) from the culture liquids until the end of the

purpureum.				
Medium	Composition			
Solid state fermentation (SSF)				
BW	beech wood (1:3 with distilled water, w:v),			
BWD	beech wood (1:3 with distilled water, w:v), plus DOR (50%, w:w)			
Submerged fermentation (SF)				
КМ	KIRK-medium			
SM	Soybean meal medium (3% suspension in distilled water, w:v),			
ASKM	KM, plus ADOR (5%, v:v)			
ASSM	SM, plus ADOR (5%, v:v)			
BSKM	KM, plus birch wood (1:2, w:v)			
BSSM	SM, plus birch wood (1:2, w:v)			

Table 1. Composition of solid and liquid media used for the analysis of the proteomes of Chondrostereum
purpureum.

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experiment on day 16 and used for enzymatic measurements. After the end of the experiment, culture liquids were concentrated by ultrafiltration, lyophilized and used for label-free proteome analyses. All treatments were carried out in triplicate.

Enzymatic activity measurements

Manganese peroxidase (MnP) activity was determined as described by [40] by following the formation of Mn³⁺-malonate complexes ($\varepsilon_{270} = 11.95 \text{ mM}^{-1} \text{ cm}^{-1}$) spectrophotometrically in the presence of MnCl₂ (0.5 mM) and H₂O₂ (0.1 mM). To distinguish between activities of laccase and manganese-independent peroxidase (MiP, including lignin, versatile and generic peroxidase) a sequential assay was performed that based on the oxidation of ABTS (0.3 mM; $\varepsilon_{420} = 36 \text{ cm}^{-1} \text{ mM}^{-1}$) with and without H₂O₂ addition (0.1 mM); peroxidase activity was corrected by the calculated Lac activity [45–47]. Unspecific peroxygenase activity (UPO) was measured as reported by Ullrich et al. [48] using veratryl alcohol at 310 nm ($\varepsilon_{310} = 9.3 \text{ mM}^{-1} \text{ cm}^{-1}$) and pH 7.0 in the presence of 1 mM H₂O₂. The mean of triplicate measurements was calculated and expressed in international units (U). An international unit is defined as the amount of enzyme that forms 1 µmol of product or converts 1 µmol of substrate per minute under assay conditions. Enzymatic activities detected in SSF cultures were expressed in U g⁻¹ and those determined in SF cultures as U L⁻¹.

DNA isolation and genome sequencing

Genomic DNA was purified from a dikaryotic strain of C. purpureum. High quality RNA-free DNA was obtained using DNeasy Plant Maxi Kit (QIAGEN). The obtained gDNA (a total of 1 μg) was fragmented with Ion Shear Plus Reagent to obtain a 200-basepair-read library. The reaction was performed at 37°C during 4 min. Fragmented gDNA was purified with the Agencourt AMPure XP Kit and the fragment size was checked with the Agilent 2100 Bioanalyzer. Adapters were ligated and blunt-end was nick-repaired with the Ion Plus Fragment Library Kit, then the ligated DNA was again purified. Subsequently, fragments of 250 bp were sizeselected on an E-Gel SizeSelect agarose gel. Fragment size was again checked using the Agilent 2100 Bioanalyzer and a PCR amplification was not required. Library was diluted to a final concentration of 26 pM with the aid of Bioanalyzer to calculate the dilution factor. Templatepositive ISPs containing clonally amplified DNA fragments were obtained using the Ion One-Touch 200 Template Kit v2 according to the manufacturer protocol. The quality of the unenriched template-positive ISPs was assessed using a Qubit 2.0 fluorometer and the Quality Control of the Ion Sphere kit. The percentage of Templated ISPs was 16%. ISP enrichment was performed with the aid of Ion OneTouch ES. The enriched template-positive ISPs were sequenced using an Ion Torrent Personal Genome Machine (PGM) (Life Technologies; Grand Island, NY, USA) according to the manufacturer's protocols provided for a 318v2 chip.

Assembly and genome annotation

All reads obtained from polyclonal and low quality ISPs were excluded. The assembly was performed according to the procedure described by Kellner et al. [49]. Only raw-reads with a read length between 120–250 bp were considered. They were assembled using MIRA 4 [50] with an accurate sensibility and a minimum of 50 reads per contig. The obtained contigs were reassembled with the Geneious R8 *de novo* assembler to filter for duplicate contigs. To assess the completeness of the assembled genome, we used CEGMA v2.5 [51] and the quality of the assembly was calculated using QUAST [52]. To verify the assembly quality and to calculate the coverage (empirical: number of reads * read length / assembly size), a remapping approach was performed. All reads were mapped against the assembly using the Geneious assembler (sensitivity: low/fastest) to analyze coverage and their uniformity.

Ab initio gene prediction was performed with Augustus [53], using Laccaria bicolor as a reference organism. No transcript variants were selected. The contigs with higher coverage after remapping were chosen for searching the rRNA genes' cistron. The functional annotation of the protein-coding genes after prediction with AUGUSTUS was carried out by a bulk blastp search against a non-redundant database (nr) obtained from GenBank. The output file (xml) with the BLAST results was imported into the Blast2GO platform, which was used for creating a C. purpureum gene database as well as for mapping and annotating of the BLAST results and displaying gene ontology terms (GO), which were merged with InterPro motifs after an Inter-ProScan. The latter step helped in functional annotation of the genome and finding of relevant proteins in the total set of 13,739 predicted gene models. Furthermore, it provided the annotated protein database for subsequent proteomic analyses. The sequences of interest, such as unspecific peroxygenase (UPO), dye-decolorizing peroxidase (DyP), laccase (Lac) and class II peroxidase (POD), were also identified and annotated manually by BLAST searches against the created proteine database and the genome contigs using gene models as references. A comprehensive analysis of CAZy genes was performed by custom BLAST searches and by using the dbCAN webserver (settings: HMMER search, E-Value <1e-4, coverage >0.3) [54]. The raw data and genome assembly is accessible at National Center for Biotechnology Information under LBLO00000000 (BioProject PRJNA281625).

Phylogenetic analysis

To confirm broad phylogenetic placement of *C. purpureum* using genome-scale data, 35 published Agaricomycete genomes [21, 26, 44, 55–60] as well as the newly sequenced *C. purpureum* were used to infer a species tree. To this end, predicted proteomes of all species were clustered into gene families using the FastOrtho (http://enews.patricbrc.org/fastortho/) implementation of the OrthoMCL software [61] with default search and clustering parameters. This resulted in a dataset of 43,749 clusters of which 1,368 were single copy and present in at least 30 species, and consequently, selected for phylogenetic analysis. Amino acid (aa) sequences constituting each cluster were aligned using Canopy (https://github.com/chunxiangli/Canopy) with the PRANK aligner [62] and three iterations of alignment and guide tree building. The resulting alignments were trimmed to remove poorly aligning regions using the software TrimAl [63] with the '-automated1' algorithm. ParGenes [64] was used to infer the best evolutionary model and a gene tree for each of the 1,368 clusters using the following parameters to RAxML-NG: '-s 10 -p 10 -b 100' [65]. Genes with strong phylogenetic signal were identified based on average bootstrap support (BS) of >50% and a minimum alignment length of at least 150 aa.

This resulting high quality set of genes was then used to infer a species tree using two complementary approaches: i) a partitioned Maximum Likelihood approach, encompassing a total of 84,248 sites and ii) ASTRAL-III, a super tree approach based on the multispecies coalescence [66]. For i) PartitionFinder2 [67] was used to identify the best-fit partitioning scheme using linked branch lengths, AICc-based model selection, the rclusterf search algorithm and the following models of protein evolution: LG, LG+G, LG+I+G, WAG, WAG+G, WAG+I+G, JTT, JTT+G, JTT+I+G, LG4X, LG4M+G. RAxML-NG [65] was run using the partition model, encompassing 112 partitions, the Majority Rule Extended automatic stopping criterion for bootstrap trees and scaled branch lengths. To infer the coalescent species tree (ii), gene trees of the high quality gene set were modified to remove nodes with less than 70% BS and ASTRA-L-III was run using default parameters.

Evolutionary analyses of gene content

Genome-wide duplication and loss rates across the 36 studied genomes were inferred using CAFÉ v4.1 [68]. To this end, an ultrametric tree was estimated using R8s v1.81 [69], with the penalized likelihood algorithm and root age arbitrarily set to 350 MYA. As input tree, we used the topology of the ASTRAL tree but with branch lengths estimated as substitution per site using RAxML-NG with the partitioned model (see above). OrthoMCL clusters were filtered to remove clusters with more than 100 gene copies in any of the species, as well as families represented in less than five species. This resulted in a final dataset of 12,032 clusters with wide taxonomic distribution. CAFÉ was run estimating a single gain (λ) and loss (μ) parameter across the tree. Gene families with significantly slower or faster rates of evolution were identified using branch-specific P-values reported by CAFÉ with a threshold of 0.01.

For target gene families of interest (GH6, GH7 and Lac), we also implemented a full phylogenetic approach for inferring gene duplications and losses. Gene trees for each target cluster were aligned as above. TreeFix v.1.1.10 [70] was run for species-tree aware error correction of each gene tree using the following options: nquickiter = 100 and niter = 1000, and the best-fit model determined by ParGenes [64], if available and WAG+G otherwise. Corrected gene trees were then reconciled with the species tree using NOTUNG 2.9 [71] in the 'phylogenomics' mode.

Furthermore, a principal component analysis was performed for the number of core lignocellulolytic enzyme encoding genes (according to Riley et al. [21] using PAST 3.22 [72]).

Proteome analysis of the C. purpureum strain

Lyophilized secreted protein fractions of fungal culture liquid (3 mg) were resuspended in 4 mL SDS buffer (1.25% SDS, 0.1 M TRIS, 0.3% DTT), a spatula of glass beads were added and the solution was incubated for one hour at room temperature under shaking. Afterwards, the FASTPrep (5.5 m/s, 45 sec, 3 cycles) protocol was applied followed by further steps including three cycles of freezing and thawing (freezing in liquid nitrogen, thawing in 60°C water bath), the addition of 0.6 mL of 10% (w/v) SDS solution and two cycles of ultrasonic treatment using an ultrasonic disintegrator (ultrasonic processor UP50H equipped with ultrasonic probe MS7, Hilscher Inc., Germany; 4°C, 2 min/80% amplitude/80% power, break of 2.5 min between cycles). Phenol extraction was applied to the sample supernatant by adding 4 mL phenol solution (10 g mL⁻¹ in ddH₂O) and incubating under shaking at 500 rpm at room temperature for one hour. The mixture was centrifuged at 4°C and 12,000 \times g for 10 min to achieve phase separation (Sorvall RC 6 plus, Thermo Fisher Scientific, Waltham, MA, USA). The lower phase was mixed with a fivefold volume of ice cold 100 mM ammonium acetate in methanol. Precipitation was performed overnight at -20°C followed by centrifugation at 12,000 \times g. The supernatant was removed and the pellet was air-dried. The resulting protein pellets were separated on SDS-PAGE with a 15°µL sample buffer. Each sample lane was then cut into one gel band and prepared for proteolytic cleavage. Protein lysate was reduced (2.5 mM DTT for one hour at 60°C) and alkylated (10 mM iodacetamide for 30 min at 37°C). Proteolysis was performed overnight using trypsin (Promega, Madison, WI, USA) with an enzyme/substrate ratio of 1:25 at 37°C. Peptide lysates were extracted from the gel and desalted using SOLAµ (Thermo Scientific) [73].

The peptide lysates were separated on a UHPLC system (Ultimate 3000, Dionex/Thermo Fisher Scientific, Idstein, Germany). Peptides were first trapped for 3 min on a C18-reverse phase trapping column (Acclaim PepMap 100, 75 μ m × 2 cm, particle size 3 μ M, nanoViper, Thermo Fisher Scientific), followed by separation on a C18-reversed phase analytical column (Acclaim PepMap 100, 75 μ m × 25 cm, particle size 3 μ M, nanoViper, Thermo Fisher

Scientific) using a two-step gradient (90 min from 4% to 30% B, then 30 min from 30% to 55% B; A: 0.1% formic acid in MS-grade water; B: 80% acetonitrile, 0.1% formic acid in MS-grade water) with a solvent flow-rate of 300 nL min⁻¹ and a column temperature of 35° C.

Mass spectrometry was performed on a Q Exactive HF mass spectrometer (Thermo Fisher Scientific, Waltham, MA, USA) with a TriVersa NanoMate (Advion, Ltd., Harlow, UK) source in LC chip coupling mode with the following settings: MS resolution 120,000, MS automatic gain control (AGC) target 3,000,000 ions, maximum injection time for MS 80 ms, intensity threshold for MS/MS of 17,000 ions, dynamic exclusion 30 sec, TopN = 20, isolation window 1.6 *m/z*, MS/MS resolution 15,000, MS/MS AGC target 50,000 ions, maximum injection time for MS/MS 120 ms.

Proteome Discoverer (v1.4.1.14, Thermo Scientific) was used for protein identification and the acquired MS/MS spectra were searched with the Sequest HT algorithm against the protein-coding database of *C. purpureum* (containing 13,739 protein-coding gene entries). Enzyme specificity was selected to trypsin with up to two missed cleavages allowed using 10 ppm pep-tide ion tolerance and 0.05 Da MS/MS tolerances. Oxidation (methionine) and carbamylation (lysine and arginine) were selected as variable modifications and carbamidomethylation (cysteine) as a static modification. Only peptides with a false discovery rate (FDR) <1% calculated by Percolator and peptide rank = 1 were considered as identified. Relative protein abundances were calculated based on the normalized spectral abundance factor (NSAF) [74].

Statistical analysis

Secretome samples were triplicated for each treatment. Hotelling's T2 was performed to find statistically significant differences between treatments and principal component analysis (PCA) using non-linear iterative partial least squares algorithms (NIPALS) was used to determine the main trend in the data set and to compare the samples replicates and treatments. The software used for statistical analyses were SPSS Statistics 21, Unscrambler X10.2 and R [75].

Results

Assembly and quality assessment of *Chondrostereum purpureum* draft genome

The genome of *C. purpureum* was assembled from 3,971,460 quality filtered reads obtained from the Ion Torrent PGM System. The average fragment size was 168 bp and a final 41.2 Mbp-sized draft genome organized in 3,435 contigs was obtained after the assemblies using MIRA and Geneious R8 (Table 2).

A remapping approach resulted in a uniform coverage (S1 Fig), and the average coverage was 15.2x. From the subset of 248 ultra-conserved CEGMA CEGs considered, we found 229 (92.3%) complete and 237 (95.6%) partial proteins sequences. In sum, a number of 13,739 protein-coding gene models was predicted. For 99% of the sequences, a BLAST hit was obtained and 30%, i.e. 4,016 sequences of the gene models contained a GO term associated. Considering the 'biological process domain', 480 of the predicted models with a GO were related to the metabolism of *C. purpureum* (S2 Fig). An assignment of gene sequences to enzymes and their classification according to EC nomenclature is given in S3 Fig.

Classification of the C. purpureum genome and proteome

The *C. purpureum* genome comprised an extensive repertoire of CAZymes characteristic for wood-rot fungi, including hydrolytic enzymes attacking cellulose and hemicelluloses (e.g. almost 50% GHs, CEs & CBMs; Fig 1) as well as multiple oxidative 'auxiliary' activities acting

207,970
390
3,435
23,869
655
98
13,739
15,468
1,522
47.5

Table 2. Statistical assembly of the C. purpureum genome.

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on lignin or lignin-derived aromatics (e.g. class II heme peroxidase, Lac, DyP and UPO; <u>S4</u> Fig).

Many of these proteins were secreted by the fungus during its growth both in synthetic and complex media (1,151 and 981 in solid-state and liquid media, respectively), especially key enzymes involved in lignocellulose decomposition (Figs 2 & 3). Overall, CAZy enzymes involved in lignocellulose degradation represented between 46 and 53% as well as 35 and 46% NSAF in the *C. purpureum* secretome on solid (BW and BWD, respectively) and liquid cultures (KM- and SM-based, respectively), respectively (S5 & S6 Figs). During liquid cultivation, distinct secretion patterns were observed (S7 Fig). Altogether 67 GHs, seven CEs, eight PLs, 28 AAs and ten CBMs showed significant differences between the different cultivations.

Glycoside Hydrolases. The analysis of the putative *C. purpureum* enzymes classified in the genome using dbCAN indicated that almost half of them were glycoside hydrolases (GHs, 49.5%) (Fig 1). The most abundant GHs were found in the functionally diverse families GH16 and GH5 with 43 and 25 sequences, respectively. In addition, a large set of genes from other families encoding cellulolytic (e.g. GH6, GH7, GH12), hemicellulolytic (e.g. GH10, GH11, GH30) and pectinolytic (e.g. GH43, GH28, GH53) proteins are present in the C. purpureum genome and most of them were expressed by the fungus during the different liquid and solid cultivations (GH6, 7, 10, 12, 28 & 43; S7 Fig). The GH class was with 34% and 32% NSAF and with a number of 112 and 124 different genes the most abundant group in the secretome obtained from the BW and BWD cultures of the fungus. Among them, GH6 and GH7 gene products (e.g. g8516 & g6636, respectively) were strongly expressed (~0.9 & 0.5%, respectively) in the BW cultures compared to the other lignocellulolytic enzymes. In the liquid cultures, a high percentage of GHs (~44% NSAF of the overall set of secreted proteins) was determined in SM and BSSM compared to the other liquid media (23–29% NSAF). High relative abundances were specifically found for the GH6 class (e.g. g13545), with 12%, 8% and 14% NSAF in SM, ASSM and BSSM, respectively. Together with the gene product g8516 (<3.9% NSAF), both were the most abundant proteins secreted by C. purpureum in all soybean-containing media (S8 Fig). The expressed CBM1 modules found in the fungal secretome pertain to a β -glucanase (GH131; g3194) and an exoglucanase (GH7; g9459); the latter occurred with a relatively high abundance mainly in BW (0.6% NSAF) (Fig 2).

Other CAZymes. CBMs are necessary for the functioning of most GHs; they were found to be present in the *C. purpureum* genome in form of 14 different families (e.g. CBM1, 13, 18, 67) and could be identified in the secretome especially in BW (~0.6% NSAF for *g9459*) and soybean-based media (0.1–0.3% NSAF for *g1461*).



Fig 1. C. purpureum CAZy genes classification (except auxiliary activities of oxidoreductases from AA1 to AA9).

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Carbohydrate esterases (CE) accounted for approx. 22% of the predicted CAZy gene models with hydrolytic activities. Among them were putative pectinolytic enzymes like CE4, 8, 9 and 15, while the CE10 family (containing carboxyl and aryl esterases) was the most abundant one with 57 predicted genes. Some of the CE families were expressed at higher levels (e.g. CE4, 8, 9; <u>S7 Fig</u>) when the fungus was growing in liquid media containing either soybean, wood or ADOR components. An acetylxylan esterase belonging to the CE1 family associated with a CBM1 module (*g1461*) was expressed by the fungus in soybean and wood cultures (Fig 3).

The addition of phenolic DOR to the solid-state cultures seemingly stimulated the secretion of polysaccharide lyases (PL). The relative amount of PL was found to be three-fold higher (9.6–2.9% NSAF) in BWD than in BW (S5 Fig). Also non-CAZy hydrolases such as peptidases (25.4–11.9% S5 Fig) showed higher relative protein amounts after DOR addition. Interestingly,

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	Gene		Putative	Me	dia
	2010		function	BW BWD	
1		g2048	UPO	0.000	0.122
		g5789	DyP	0.083	0.041
		g6352	AA3_1 (CDH)	0.850	0.947
		g12139		0.521	0.456
		g10622	AA3_3	0.042	0.037
		g5697	(AlcOx.)	0.242	0.061
		g1416		0.049	0.030
		g11873		0.033	0.052
		g10495	AA1_1	0.004	0.018
C		g6136	(Lac)	0.000	0.011
-ignin		g1744		0.023	0.100
60		g9579	AA5	0.007	0.098
		g9433		0.032	0.075
		g9062	(CRO)	0.037	0.109
		g5969		0.000	0.033
		g2075	(GlyOx.)	0.000	0.017
		g12545	AA3_2	0.083	0.172
		g8332		0.098	0. 375
		g8331	(AAO)	0.034	0.037
		g7252	(GLO)	0.032	0.050
		g11636		0.023	0.011
		g9185	AA2	0.058	0.074
		g8056		0.070	0.008
		g9543		0.014	0.005
		g8686		0.011	0.011
		g6045		0.000	0.018
		g4746		0.067	0.106
Se		g4608	AA9	0.000	0.030
0		g3824	(LPMO)	0.085	0.160
lu		g3183		0.167	0. 357
e B		g2570		0.051	0.133
		g2420		0.072	0.255
ne		g2388		0.106	0.105
Crystalline cellulo		g1818		0.163	0 .327
sta		g7415	GH7	0.207	0.4 99
<u> </u>		g6636	GH	0.451	0. 363
J .		g8516	СНЕ	0.923	0 .279
		g4675	GH6	0.040	0.074
		g9459*		0.636	0.125
		g3194 **	СВМ	0.027	0.009
		g1461#		0.076	0.140

Lignin

Fig 2. Distribution of the lignocellulose-degrading enzymes secreted by *C. purpureum* (after seven weeks of cultivation). Relative abundance (% NSAF) of the proteins found in solid state cultures containing beech-wood (BW) or beech-wood and DOR (BWD) is given (Table 1). CBM1 of *GH7, **GH131 (β -glucanase) and [#]acetylxylan esterase.

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the secretion level of the peptidase enzyme subclass (EC 3.4.x) was also remarkably increased (up to 15% NSAF) when the fungus grew in ASSM compared to SM without ADOR supplementation (8% NSAF) (S6 Fig).

Oxidoreductases. Altogether 153 oxidoreductases with relevance for lignocellulose decomposition were found in the *C. purpureum* genome and 81 of them were expressed during the different cultivations (Table 3).

Among them are typical enzymes involved in lignocellulose breakdown caused by woodrot fungi. Besides AA1 (multicopper oxidases, e.g. Lac) and AA3 (GMC, e.g. cellobiose dehydrogenase or alcohol oxidase), a high number of AA genes were found to encode LPMO

		Gene	Putative			Med	ia		
			function	км	ASKM	BSKM	SM	ASSM	BSSM
	[g6938		0.037	0.000	0.000	0.000	0.000	0.000
		g5554	UDO	0.042	0.072	0.000	0.048	0.124	0.034
		g4785	UPO	0.000	0.073	0.000	0.000	0.022	0.000
		g2048		0.000	0.117	0.000	0.000	0.000	0.000
		g6591	DyP	0.000	0.000	0.000	0.000	0.000	0.077
		g6352	AA3_1 (CDH)	0.216	0.086	0.107	0.182	0.130	0.199
		g13239		0.000	0.000	0.000	0.026	0.000	0.000
		g12972		0.019	0.042	0.014	0.015	0.015	0.000
		g12139		0.190	0.125	0.264	0.290	0.250	0.157
		g10622	AA3_3	0.144	0.431	0.411	0.167	0.223	0.172
		g9260	(AlcOx.)	0.000	0.118	0.000	0.026	0.000	0.013
		g3473		0.000	0.014	0.000	0.009	0.005	0.010
		g2782		0.000	0.021	0.000	0.000	0.000	0.000
		g1416		0.216	0.332	0.207	0.076	0.180	0.039
		g11873		0.088	0.096	0.074	0.027	0.037	0.020
		g11793		0.000	0.015	0.021	0.000	0.000	0.000
		g10758		0.000	0.000	0.012	0.022	0.000	0.000
		g10495		0.018	0.099	0.044	0.007	0.000	0.000
		g10095		0.000	0.071	0.023	0.026	0.031	0.000
		g9564		0.000	0.224	0.000	0.000	0.000	0.000
.⊆		g8701		0.000	0.008	0.000	0.000	0.016	0.000
L		g8577		0.038	0.022	0.042	0.000	0.000	0.000
Ë.		g7276		0.000	0.119	0.137	0.022	0.000	0.040
_		g7032		0.000	0.037	0.000	0.000	0.000	0.000
		g6961	AA1_1	0.000	0.000	0.000	0.000	0.020	0.000
		g6664	(Lac)	0.249	0.447	0.071	0.031	0.010	0.000
		g6292	(Lac)	0.042	0.107	0.000	0.000	0.000	0.000
		g6136		0.000	0.000	0.027	0.000	0.000	0.000
		g6000		0.000	0.156	0.143	0.011	0.030	0.066
		g5841		0.000	0.025	0.049	0.000	0.000	0.000
		g5639		0.013	0.384	0.239	0.058	0.079	0.061
		g5532		0.031	0.026	0.014	0.000	0.016	0.000
		g5369		0.000	0.038	0.040	0.000	0.013	0.000
		g3879		0.000	0.029	0.020	0.000	0.009	0.000
		g2393		0.078	0.148	0.135	0.033	0.016	0.018
		g1744		0.133	0.391	0.197	0.042	0.071	0.048
		g1645		0.290	0.284	0.266	0.120	0.058	0.065
		g9433		0.024	0.085	0.151	0.027	0.049	0.015
		g9062	AA5	0.000	0.021	0.006	0.012	0.012	0.008
		g8520	(CRO)	0.000	0.014	0.000	0.000	0.000	0.000
		g304		0.029	0.013	0.000	0.009	0.028	0.010
		g8503		0.643	0.115	0.068	0.292	0.751	0.621
		g5474	(GlyOx.)	0.173	0.023	0.036	0.113	0.231	0.127
	L	g2075		0.106	0.021	0.030	0.070	0.110	0.072

		Gene	Putative			Med	ia		
	,		function	KM	ASKM	BSKM	SM	ASSM	BSSM
		g12545		0.127	0.383	0.523	0.160	0.166	0.133
	1	g8332	AA3_2	0.000	0.345	0.376	0.170	0.151	0.101
	1	g8331	(AAO)	0.000	0.213	0.372	0.096	0.072	0.039
Ē		g1551		0.000	0.021	0.019	0.000	0.000	0.000
		g7252	(GLO)	0.128	0.158	0.097	0.139	0.145	0.113
_		g5232		0.000	0.016	0.008	0.000	0.000	0.000
		g5379	AA2 (GP)	0.000	0.051	0.000	0.000	0.000	0.000
	ļ	g5041	(MnP)	0.000	0.012	0.000	0.015	0.018	0.016
		g13314		0.000	0.192	0.021	0.026	0.074	0.085
		g9543		0.000	0.000	0.000	0.013	0.016	0.020
	1	g8686		0.000	0.000	0.000	0.010	0.016	0.012
		g7832		0.000	0.016	0.000	0.000	0.025	0.000
		g4746		0.000	0.000	0.000	0.035	_	
		g4575		0.000	0.000	0.000	0.018		0.027
	1	g3824	AA9	0.000	0.000	0.000	0.091		
	1	g3200	(LPMO)	0.000	0.031	0.000	0.033		0.000
		g3183	(1.1.1.0)	0.000		0.000	0.186		0.142
b		g2570		0.000	0.000	0.000	0.031	0.012	0.023
SC		g2420		0.000	0.029	0.000	0.025	0.045	0.018
Ĕ		g2388		0.000	0.000	0.000	0.046		0.049
=		g2289		0.000	0.000	0.000	0.000	0.026	0.011
S		g1818		0.000	0.109	0.000	0.022	0.083	0.017
e		g195		0.000	0.000	0.000	0.000	0.008	0.015
Crystalline cellulose		g7992		0.000		0.021	0.000	0.007	0.000
a	1	g7415		0.067			0.275	0.307	0.219
'st	1	g6636		0.062	0.060	0.097	0.174	0.150	0.190
2		g6241	GH7	0.104	0.024	0.126	0.000		0.000
0		g5468		0.015	0.019	0.000	0.000	0.000	0.000
		g3092		0.387	0.305	0.461	0.295	0.182	0.299
		g2102		0.000	0.000	0.000	0.031	0.000	0.013
		g1181		0.000	0.000	0.013	0.016	0.022	0.000
		g13545		0.280	0.609	0.200	0.121	0.829	0.136
		g8516		0.353	0.342	0.280	0.368	0.196	0.394
		g8515	GH6	0.072	0.028		0.044	0.043	0.086
		g4675		0.250	-	0.000	0.283	0.211	0.233
		g4112		0.098		0.166	0.258	0.352	0.197
		g4111		0.054	0.000	0.108	0.132	0.207	0.070
		g9459*	CBM	0.022	0.015	0.043	0.069	0.032	0.054
	L	g1461 #		0.039	0.052	0.036	0.310	0.140	0.227

Fig 3. Distribution of the lignocellulose-degrading enzymes secreted by *C. purpureum* (after 16 days of cultivation). Relative abundance (% NSAF, >0.01) of the proteins found in liquid cultures of synthetic KM, ASKM, BSKM and complex SM, ASSM and BSSM (<u>Table 1</u>). To obtain an appropriate resolution, high NSAF values (>1%) were divided by the factor 10 (blue) or 100 (red). CBM1 of *GH7 and *acetylxylan esterase.

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(AA9) and more than half of them were expressed with NSAF >0.01% (17 out of 31 LPMO genes). During growth on wood (BW and BWD), LPMO made up the largest number of proteins among the lignocellulose-modifying enzymes. Some of them had a higher relative abundance in media supplemented with phenol-rich olive mill residues (e.g. in BWD with 0.36, 0.26, 0.33% NSAF for *g3183*, *g2420*, *g1818*, respectively and in ASKM with 0.19, 0.05, 0.11% NSAF for *g13314*, *g3183*, *g1818*, respectively), or in soybean-based liquid media (Figs 2 & 3). Whereas none of them was present in synthetic KM without any additive, 15 genes encoding for LPMO were expressed in soybean-based medium (e.g. 0.097–0.186% NSAF for *g3183*) and seven of these LPMOs showed significant differences during cultivation (S7 Fig).

Four class II peroxidases were found in the genome of *C. purpureum* (S9 Fig). Among them two MnPs (g2700 & g5041), the key enzyme type of incipient lignin degradation by white-rot fungi, were identified according to the conserved manganese-binding aa residues (D34, D39 & D179; S10 Fig). The presence of MnP was confirmed by respective activity measurements in both solid and liquid media (<10 to 30 U L⁻¹; S9 and S10 Figs) and by detection of an expressed short MnP protein (g5041; Fig 3) mainly in soybean-based medium but also in ASKM. Due to the absence of typical manganese-binding aa residues and LiP-characteristic tryptophan (at position W171 in *P. chrysosporium*, [76]), the other two sequences (g5379, g10149) of class II peroxidases seemingly represent generic peroxidases (GPs) that typically oxidize phenolics. One of the GPs was found to be expressed exclusively in ASKM (S10 Fig).

In addition to class II peroxidases, sequences encoding for seven DyPs and eight HTPs/ UPOs were predicted in the genome. Due to the presence of two specific amino acid motifs, PCP and EHD [77], six complete UPO sequences could be assigned to the 'short' UPOs. The other two sequences were incomplete but phylogenetic analysis related them also to the clade

Proteins	Total	Expressed ^a
UPO ^b	8	5
DyP ^b	7	2
AA1	47	27
Lac	45	25
FeOx Fet3	1	1
Fungal pigment oxidase (MCO)	1	1
AA2 class II peroxidase (POD)	4	2
MnP	2	1
GP	2	1
AA3 GMC	36	16
AA4 VAO	3	0
AA5 GLX and CRO	12	9
AA7 GOO	3	3
AA8 iron reductase domain	2	0
AA9 LPMO	31	17

Table 3. Oxidoreductases found in the C. purpureum genome and expressed in fungal cultures.

Abbreviations: unspecific peroxygenase (UPO), dye-decolorizing peroxidase (DyP), CAZy classified auxiliary activities (AA): AA2 class II peroxidases (manganese peroxidase, MnP and generic peroxidase, GP) as well as AA1 (laccase, Lac), ferroxidase (FeOx) and fungal pigment multicopper oxidase (MCO), AA3 (glucose-methanol-choline oxidoreductase, GMC), AA4 (vanillyl alcohol oxidase, VAO), AA5 (glyoxal oxidase, GLX) and cooper radical oxidase (CRO), AA7 (glucooligosaccharide oxidase, GOO), AA8 and AA9 (lytic polysaccharide monooxygenase, LPMO) enzymes.

^aNumber of expressed genes in the secretomes obtained from different culture media.^bNon CAZy protein families.

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of 'short' UPOs. The fungus secreted one DyP (g5789) in BW and BWD, whereas one 'short' UPO (g2048) was exclusively found in BWD, the presence of which was confirmed by a corresponding enzymatic activity 0.5 U g⁻¹ after six weeks of cultivation (Figs 2 & S11). In SF, another DyP gene (g6591) and four further 'short' UPO genes (g4556, g4785, g5554, g6938) were expressed by the fungus. UPO activities were also detectable in all liquid cultures with values up to ~10 U L⁻¹ (S12 Fig). For the sake of completeness, three genes of intracellular class I peroxidases–two ascorbic acid peroxidases (APXs) and one cyctochrome c peroxidase (CcP)–were found in the *C. purpureum* genome as well (S9 Fig). The expression of CcP (g2498) was confirmed for BW cultures on the protein level (NSAF <0.01%).

A remarkable finding has been the high number of Lac genes (45, including 38 full-length sequences) predicted in the genome of *C. purpureum*. Phylogenetic analysis of the whole MCO family (including Lac, ferroxidase and pigment oxidase) revealed 40 full-length sequences and 14 partial sequences, which frequently were split on the end of contigs, i.e. showing only the N- or C terminus in an alignment (S13 Fig). Considering the 40 full-length genes, 38 can be defined as Lac *sensu stricto*. Highest identity between two full-length Lacs was 97% (S14 Fig) and more than half of all Lacs were expressed (in total 25, S13 Fig) with NSAF >0.01% in liquid and solid state cultures (23 and four proteins, respectively; Figs 2 and 3). Moreover, moderate Lac activities were detected during cultivations (up to ~10 U L⁻¹; S11 and S12 Figs). The secretion of some of the Lacs was obviously stimulated in the presence of phenol-rich olive mill residues (e.g. *g1744* in SSF and *g9564*, *g6664*, *g5639*, *g1744* in KM with DOR and birch wood supplemented ASKM and BSKM, respectively). One MCO sequence turned out to belong to a Fet3 ferroxidase cluster (FeOX; *g718*) and another one to a fungal pigment MCO cluster (*g10148*; S13 Fig).

Further CAZy AA representatives (AA3-9) predicted in the genome rank among enzymes supporting peroxide-dependent biocatalysts acting on lignin and aromatics (i.e. oxidases that supply class II POD, UPO and DyP) or interacting with carbohydrates (GMC, GLX, LPMO); some of them occur with high relative abundance (4 to 36 genes; Table 3 & S4 Fig). The number of predicted proteins producing hydrogen peroxide (AA3 & AA5) was 48. Among this enzyme group, the only available CDH gene (AA3_1; g6352), encoding an enzyme involved in Fenton-based chemistry and LPMO activation, was expressed by C. purpureum in all culture media tested (Figs 2 and 3). Interestingly, the respective CDH gene product was in the solidstate cultures one of the most abundant proteins (NSAF ~0.9%) among the lignocellulose-relevant enzymes (Fig 2). From the AA3_2 group (GMCs), 24 genes encoding for glucose and aryl alcohol oxidases were predicted and the fungus expressed up to six of them in all media tested (Figs 2 and 3). In KM, the addition of DOR or birch wood (ASKM and BSKM) led to a higher relative abundance of some of the AAOs (e.g. g12545, g8332, g8331), which were also present in the soybean medium with and without additives. Furthermore, ten sequences of alcohol oxidases (AA3 3) and one pyranose oxidase gene (AA3 4) were identified in the genome (S4 Fig). Several genes of the former groups (e.g. g12139) were expressed both in SSF and SF. There is indication that the AA3_3 gene g12139 could be constitutive, since it showed also high relative abundance in KM, in which eligible inducing compounds (e.g. aromatics from wood or DOR) were not present (Fig 3). The number of genes encoding for copper radical oxidases (CRO and GLX, AA5) was twelve, and some of them were expressed in SSF and SF with NSAF >0.01% (e.g. *g9579*, *g9433*, *g9062*, *g2075*). Eventually, four genes encoding for benzoquinone reductase (AA6) were found in the genome indicating that the enzymatic basis of Fenton-based attack on lignocellulose is realized in C. purpureum.

Taxonomy. A phylogenetic analysis that comprised of 609 single copy genes from 36 basidiomycete genomes placed *C. purpureum* in a clade close to *Moniliophthora perniciosa* (Marasmiaceae) and four *Armillaria* species (Physalacriaceae), which all represent distinct





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pathogens within the Agaricales (Fig 4). Complementary approaches using either ML reconstruction or a summary method based on the multispecies coalescent resulted in nearly identical, well resolved trees (Fig 4 and S15 Fig), with the exception of the branching order of *C. purpureum* and *M. perniciosa*. Currently only few genomes of the so called 'marasmioid clade' [19] are known and larger numbers of taxa in this clade will be required for accurate resolution of the relationship between *C. purpureum* and *M. perniciosa*. In analyses with a more dense taxon sampling but less phylogenetically relevant genes (two to six genes), a possible phylogenetic position to the Cyphellaceae was shown [19, 78]; the genes found in the assembled genome were identical to the previously analyzed *C. purpureum* genes in the study of Matheny and coworkers.

Gene content evolution. Inference of gene duplication (λ) and loss (μ) rates yielded global estimates of 9.27e-4 and 2.59e-3 events per unit branch length, respectively. A total of 25 orthogroups showed significant deviation from these genomic background rates on the *C. purpureum* branch (P <0.01), and in any case, this entailed expansions rather than contractions (Table 4). Approximately two thirds of significantly expanded families were attributable to transposable elements based on annotation with PFAM domains or BLAST searches. The most pronounced expansion among structural genes was ORTHOMCL11, encoding a family of Lac, in the case of which we inferred a total of 18 lineage-specific duplications using the model-based approach. Significantly expanded families also included a putative non-ribosomal peptide synthase, two families of proteases and a family of cytochrome P450s.

Since both Lac and GH7 cellobiohydrolase were found in unusually high numbers in the *C. purpureum* genome, we also included a focused phylogenetic analysis of orthogroups encoding genes in these families (Fig 4). Among annotated Lac, 39 were found spread across five orthogroups, ORTHOMCL11 being the largest (Fig 4 and S1 Fig). Thirteen Lacs did not cluster into orthogroups, of which ten constituted partial gene models, suggesting that FastOrtho clustering mitigates the problem of fragmentation for downstream evolutionary analyses. All

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OrthoGroup	PFAM domains	Putative Function
ORTHOMCL2869[+4]	Plavaka (PF18759)	Likely transposable element
ORTHOMCL7313[+2]	None (RT based on BLAST)	Likely transposable element
ORTHOMCL383[+20]	None (Gag-Pol based on BLAST)	Likely transposable element
ORTHOMCL194[+21]	KDZ (PF18758)	Likely transposable element
ORTHOMCL305[+16]	CxC2 (PF18803)	Likely transposable element
ORTHOMCL7306[+15]	Transposase_21 (PF02992)	Likely transposable element
ORTHOMCL7397[+2]	None	Hypothetical protein
ORTHOMCL477[+16]	RVT_1 (PF00078)	Likely transposable element
ORTHOMCL480[+2]	KDZ (PF18758)	Likely transposable element
ORTHOMCL7175[+2]	None (FAR1 based on BLAST)	Likely transposable element
ORTHOMCL3789[+21]	None	-
ORTHOMCL88[+3]	RVT_1 (PF00078)	Likely transposable element
ORTHOMCL7296[+2]	None	-
ORTHOMCL404[+5]	PIF1 (PF05970)	Helicase—possibly TE
ORTHOMCL601[+2]	DUF4470 (PF14737)	-
ORTHOMCL330[+7]	Retrotrans gag (PF03732)	Likely transposable element
ORTHOMCL75[+4]	AMP-binding (PF00501)	Non-ribosomal peptide synthetase (NRPS)
ORTHOMCL812[+3]	None	-
ORTHOMCL526[+5]	None (RT from BLAST)	Likely transposable element
ORTHOMCL727[+3]	RVT_2 (PF07727)	Likely transposable element
ORTHOMCL80[+6]	Helitron-like N (PF14214)	Likely transposable element
ORTHOMCL86[+3]	Peptidase_M36 (PF02128)	Fungalysin metalloprotease
ORTHOMCL192[+5]	Peptidase_S8 (PF00082)	Subtilisin like protease
ORTHOMCL11[+18]	Cu-oxidase (PF00394)	Laccase
ORTHOMCL43[+4]	P450 (PF00067)	Cytochrome P450

Table 4. Significantly expanded orthogroups in C. purpureum.

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annotated GH7s were found in ORTHOMCL56 (Fig 4). Phylogenetic analyses confirmed model-based inference, showing an exceptionally large lineage-specific expansion of Lac in *C. purpureum* mainly driven by ORTHOMCL11. Similarly, five of the eight GH7 copies likely arose on the *C. purpureum* branch (based on six duplications and one loss; Fig 4). Patterns for GH6 were complex, since members of this family were spread among two different orthogroups, with two out of five copies constituting singletons (S16 Fig).

To classify the position of *C. purpureum* within genome-sequenced wood decomposing fungi, a principal component analysis was performed using the core lignocellulolytic genes (Fig 5). The axis 1 explained 44.3% of the variation and axis 2 explained 32%. Brown-rot fungi formed a distinct group, whereas white-rot and the so called 'white-rot-like' species (*Botryobasidium botryosum, Jaapia argillacea* and *S. commune*; all containing no class II peroxidase genes) formed a far more variable group (Fig 5). The lignocellulolytic gene content placed *C. purpureum* close to *Stereum hirsutum*, *M. perniciosa* and three *Armillaria* species. Conspicuous differences in the gene repertoire among the ecological classifications 'white-rot' (without the pathogenic species), 'white-rot-like', 'pathogenic white-rot' (*Armillaria* species, *M. perniciosa* and *C. purpureum*) and 'brown-rot' were found (S4 Fig). For example, class II peroxidases (AA2) occurred in average with seven genes in pathogenic WRF, twelve in the remaining WRF, and none in WRF-like and BRF. Class AA3_2 occurred on average with 52 genes in pathogenic WRF, 18 in WRF-like and 13 in BRF. Class AA1_1, Lac, occurred on average with 27 genes in pathogenic WRF, nine in the remaining WRF, one in WRF-like and four in BRF (S4 Fig).



Component 1

Fig 5. Principal component analysis of main CAZy lignocellulolytic gene content in the analysed genomes (published data from JGI & NCBI). A variance-covariance matrix of the values in (S4 Fig) were used. Agabi, *Agaricus bisporus*; Agrae, *Agrocybe aegerita* [60]; Armce, *Armillaria cepistipes*; Armga, *Armillaria gallica*; Armme, *Armillaria mellea*; Armso, *Armillaria solidipes*, Armos, *Armillaria ostoyae*; Aursu, *Auricularia subglabra*; Bjead, Bjerkandera adusta; Botbo, Botryobasidium botryosum; Cersu, Ceriporiopsis subvermispora; Chopu, Chondrostereum purpureum; Conpu, Coniophora puteana; Copci, Coprinopsis cinereus; Dacsp, Dacryopinax sp.; Dicsq, Dichomitus squalens; Fomme, Fomitiporia mediterranea; Fompi, Fomitopsis pinicola; Galma, Galerina marginata; Glotr, Gloeophyllum trabeum; Hetan, Heterobasidion annosum; Jaaar, Jaapia argillacea; Mycch, Mycena chlorophos [79]; Monpe, Moniliophthora perniciosa; Phaca, Phanerochaete carnosa; Phchr, Phanerochaete chrysosporium; Phlbr, Phlebia brevispora; Pleos, Pleurotus ostreatus; Pospl, Postia placenta; Punst, Punctularia strigosozonata; Pycci, Pycnoporus cinnabarinus; Schco, Schizophyllum commune; Serla, Serpula lacrymans; Stehi, Stereum hirsutum; Trave, Trametes versicolor; Volvo, Volvariella volvacea and Wolco, Wolfiporia cocos.

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Discussion

Phylogenetic position of C. purpureum

C. purpureum has been placed in the Cyphellaceae within the so called 'marasmioid clade' by Matheny and coworkers [19], and the current databases (MycoBank, Index Fungorum) follow this taxonomic affiliation. A follow-up study using a six gene approach could not further resolve the positioning of *C. purpureum* [78]. In our approach using more than 600 single copy genes, it has not been expected to reach further resolution, since only a 'handful' of genomes is currently being available from the 'marasmioid clade'. Nevertheless, the identified genes of *C. purpureum* perfectly match with those of both studies of Matheny et al. [19, 78], and hence support its current phylogenetic position. Thus, the *C. purpureum* genome presented here will contribute to future studies dealing with the phylogeny of the marasmioid clade by using comparative genomics.

Repertoire of genes and secreted proteins related to lignocellulose degradation in comparison to other wood-rot fungi

In addition to the *C. purpureum* genome, we provide insights into one of the few existing secretomes of fungi that can switch between phytopathogenic and saprotrophic life-styles. To obtain a broad proteomic data set, the fungus was cultured in liquid synthetic and plant-based

complex media as well as in solid media containing wood and/or olive-wastes to follow enzyme secretion under different degradation and detoxification strategies.

The size of the *C. purpureum* genome (46.5 Mbp) is within the typical range of other fungi's genomes, especially basidiomycetes (40–50 Mbp; [80]). On the other hand, the fungus secreted a high number of proteins (981 to 1,151 in dependence of the culture medium) in comparison to other basidiomycetes [27]. For example, following numbers of secreted proteins were reported for some other basidiomycetes: 243 for *Ceriporiopsis subvermispora* [81], 293 for *A. mellea* [33], 356 and 413 for *P. chrysosporium* and *P. placenta*, respectively [82] as well as 508 for *P. ostreatus* [27] when growing on lignocellulose-containing or synthetic media. An equally large number of secreted proteins (with almost 800) was reported for the secretome of *P. chrysosporium* when growing on the wood of three different poplar genotypes [83]. However, it should be taken into consideration that steady improvements in mass spectrometry may have influenced these data sets [84].

The fungus *C. purpureum* possesses in the genome and secretes diverse hydrolases and oxidoreductases that are involved in the degradation and modification of lignocelluloses and are typical for saprotrophic fungi. *C. purpureum* pertains to the ecological group of 'pathogenic white-rot' fungi like *Armillaria* spp. or *M. perniciosa* as indicated by a PCA considering core lignocellulolytic genes. *Sensu lato, C. purpureum* belongs to the WRF ecotype, incorporating key enzymes of lignin attack like manganese peroxidase.

Glycoside hydrolases. Although cellobiohydrolases of the families GH6 and GH7 are known to correlate with white-rot, the number of six genes belonging to family GH6 in the *C. purpureum* genome is one of the highest reported so far, considering all available data on WRF and BRF (<5 genes). The number of GH7 genes was nine and thus ranges in the middle of reported numbers for other WRF (e.g. one and 16 genes for *Heterobasidion annosum* and *P. ostreatus*, respectively). In this context, the fungus houses the highest number of GH7s in its clade and six lineage-specific duplications in this family (Fig 4). This is contrary to BRF, in which genes encoding these enzymes are rarely present and only few of them have one or two respective enzymes (GH6 or GH7) available. The presence of cellulases (families GH6 and GH7) in the secretome of *C. purpureum* is also a characteristic that it shares with other WRF (e.g. *P. chrysosporium, C. subvermispora*; [81, 83]), for which these enzymes were shown to be major proteins secreted in wood-containing media. These enzymes act on 'bulky' polysaccharide backbones and are seemingly up-regulated in *C. purpureum* when it grows in soybean-based medium for more than two weeks.

It has been accepted that seven CAZy families preferably target complex hemicelluloses (e.g. GH10, GH11, GH30), and eleven families the even more heterogeneous pectins (e.g. GH43, GH28) [21, 85]. Most of these enzymes were found to be present in the *C. purpureum* genome except families GH11 and GH26. Highly abundant GHs found in the *C. purpureum* secretomes of liquid cultures containing wood or ADOR belong to the families GH3 (*g3528*), GH5 (*g6251, g7580*) and GH10 (*g13143*). These proteins were also reported to be over-produced in other WRF in the presence of lignocellulosic substrates [27]. GH5 represents a large protein family that contains a range of enzymes acting on β -linked oligo- and polysaccharides [64], and 17 out of 25 genes present in the genome were expressed by *C. purpureum*, predominantly in soybean-based medium.

Pectinolytic hydrolases expressed by *C. purpureum* in SF (e.g. GH28; CE4, 8, 9 and 15) play probably a role during the invasive step of fungal plant pathogens as proposed for typical representatives such as *Botrytis cinerea* and *Fusarium oxysporum* [14, 21, 86]. The GH28 family comprises endo- and exo-[rhamno]galacturonases, essential for pectinolysis by both parasitic and saprotrophic fungi [87]. Genes encoding such enzymes that disintegrate the middle lamel-lae of plant cell walls, are highly abundant in facultative parasites (i.e. WRF that are both

necrotrophic and biotrophic such as the tree pathogen *A. mellea* that contains 17 respective genes in its genome) and are just moderately represented in exclusively saprotrophic fungi (e.g. dead-wood dwellers such as *C. subvermispora* with six and *P. chrysosporium* with four respective genes; [33, 87]).

The expressed CE proteins of *C. purpureum*-found in the secretome of *A. mellea* as wellare also involved in the degradation of pectins, e.g. by members of the CE8 family, which cleave methyl esters. CE4 is a chitin deacetylase and acts on the acetyl group of *N*-acetyl glucosamine [33]. It may cooperate with fungal chitinase, e.g. GH18 of *C. purpureum*, to deacetylate chitin to chitosan oligomers that do not longer elicit a plant defence response, and thus enabling the fungus to invade living plants/trees without any appreciable resistance [88]. The high abundance of pectinolytic enzymes and the presence of chitin-degrading enzymes in the genome and secretome of *C. purpureum* implies that they are part of its pathogenic system causing the silver-leaf disease ([89, 90], originally described to be caused by 'Stereum' purpureum [91]).

Oxidoreductases. The number of LPMO genes (31) is in the upper range of that of other WRF like B. adusta (Meruliaceae, 28 genes; [44]) or P. ostreatus (Pleurotaceae, 29 genes; [21]) and leaf-litter decomposers such as Coprinopsis cinerea (Psathyrellaceae, 34 genes; [56]). Even in 'atypical' wood-rot fungi, high numbers of LPMO genes can be found (e.g. 32 genes in B. botryosum), while in BRF, the number of LPMO genes generally goes below ten [21]. LPMOs are thought to be involved in the oxidative cleavage of recalcitrant wood polysaccharides, particularly in that of crystalline cellulose [92] but may have also completely different roles, for example, during chitin development in ancient arthropods [93]. The expression levels of LPMOs reported for wood-degrading fungi differ considerably. Thus, during growth on woody substrates, almost half of the existing 15 genes were expressed in *P. chrysosporium* [94]; Phlebia gigantea expressed six out of 15 genes [95], C. subvermispora five out of nine genes [81] and P. radiata seven out of twelve [96]. On the other hand, only low or even no expression of LPMOs was observed for Wolfiporia coccos [97] and P. ostreatus [27] under comparable conditions. LPMOs are capable of enhancing oxidative attack on crystalline cellulose by cooperating with CDH [98]. The high abundance of LPMO in SF of C. purpureum points to a certain LPMO-stimulating effect by plant secondary ingredients and hemicellulose derivatives [39, 40, 99]. Furthermore, it could be an indication for their joint action with CDH that functions as an activating electron donor for LPMO and thereby improving the incipient degradation of crystalline cellulose [98]. Moreover, CDH has been proposed to play a role in the generation of hydroxyl radicals (HO•) via Fenton chemistry and thereby may also contribute to the degradation of cellulose as well as the modification of lignin [100]. CDH is uniformly present as a single gene copy in all WRF but absent in the majority of BRF [21, 22, 100]. High expression of CDH in relation to lignocellulose degradation (e.g. of aspen wood or wheat straw) was reported for P. chrysosporium [94], I. lacteus [101], P. gigantea [95], C. subvermispora [81] and P. radiata [96]. In contrast to these findings, CDH was not detectable in secretomes of P. ostreatus grown on poplar wood or wheat straw.

The presence of high-redox potential class II PODs (i.e. MnPs) in the genome of *C. purpureum* fulfills a well-known characteristic of all WRF and is a prerequisite for the efficient degradation of lignin. BRF, on the other hand, are completely lacking these biocatalysts similar as some 'uncategorized' (atypical or 'white-rot-like') wood-rot fungi [21]. Ligninolytic PODs can occur in various combinations of MnP, VP and LiP (and there are several subtypes of MnPs with somewhat different substrate spectra; [58, 102]. The number of MnP genes in white-rot fungi ranges from two in *A. bisporus* to 13 and 16 in *C. subvermispora* and *Fomitoporia mediteranea*, respectively [21, 81]. *C. purpureum* with its two MnP encoding genes rather groups in the WRF clade with few MnP genes in this classification system. As an agaric WRF, *C*. *purpureum* is lacking LiP genes; the occurrence of LiP is seemingly restricted to the evolutionary old order of Polyporales, for example, to species like *T. versicolor*, *P. chrysosporium*, *P. radiata* or *B. adusta* [58].

Among the two 'short-type' MnPs of *C. purpureum*, only one was expressed and only in liquid culture. Similarly, the necrotrophic WRF *H. irregulare* possesses six short MnPs out of seven class II PODs but none of them showed strong expression when the fungus grew on spruce heartwood [32]. For *P. gigantea*, a similar observation was made; from seven genes encoding MnPs, the expression level in the corresponding secretome was rather low during growth on acetone-extracted and non-extracted pine wood [95]. A possible explanation for the low expression level of MnP genes in *C. purpureum* could be related to the fungus' affiliation to the group of 'agaric' WRF, which seemingly possess a simpler ligninolytic enzyme system compared to 'polyporous' WRF. *Volvariella volvacea* (Rice straw mushroom) with its two MnPs can serve as a typical example of such fungi [57]. Possibly, lignin degradation by fungi with small numbers of class II peroxidases may be supported by other oxidative enzymes and/ or mechanisms. Furthermore, MnPs are known to be secreted during early stages of fungal lignocellulose degradation (e.g. second to fourth week [103]) so that they may have already disappeared at the time point when the *C. purpureum* secretome was analyzed (after six weeks).

Interestingly also HTPs, with UPOs as their most prominent representatives, were found to be present in the *C. purpureum* genome. In general, these enzymes are widespread among the whole fungal kingdom including all phyla of true fungi (Eumycota) and a few 'pseudofungal' stramenopiles ('Oomycota') [77]. However, it is not much known on their physiological functions and on the natural substances that regulate their expression [104]. *C. purpureum* expressed five out of six UPOs, which all belonged to the phylogenetically older 'short' type genes. Because of their catalytic versatility, it is conceivable that UPOs may be involved in extracellular detoxification reactions, e.g. of low molecular weight compounds typically found in living plants, wood, and soil (secondary metabolites, tannins, phytoalexins, microbial toxins, xenobiotics, etc.). In addition to triggering UPO production by soybeans and their ingredients in different fungi (e.g. by soybean meal, [48]; soybean peptone, [105]; glycinin or conglycinin, [106]), recently published data indicate that DOR and its aqueous extract (ADOR) have stimulating effects on UPO secretion too [38, 107].

Two of six genes encoding for DyPs were expressed by *C. purpureum*, but both only in beech-wood supplemented soybean medium and in solid-state cultures; in the latter case, the corresponding NSAF (%) was higher in beech-wood (BW) than in respective solid cultures supplemented with DOR (BWD). It cannot be ruled out that both types of peroxide-using enzymes, UPOs and DyPs, are partially overlapping with class II PODs in their ability to oxidize lignin structures, and thus, they should be included in the pool of lignin-modifying enzymes [46–48, 108, 109].

Peroxide generating enzymes (e.g. AA3 and AA5) constitute another important component of the ligninolytic system of wood-rot fungi, since H_2O_2 is required as the oxidant (electron acceptor) in peroxidative (and peroxygenating) reactions. The high abundance of respective genes/proteins in *C. purpureum*'s genome/secretome reflects this fact and indirectly points to the importance of peroxidase-based processes during wood decay [110, 111].

There are other oxidoreductases that are thought to be involved in lignin modification. The most prominent ones are Lacs being abundant in most WRF (though they are lacking in the model organisms *P. chrysosporium* and *B. adusta*; [5, 112]) and they also occur in some BRF [21]. Taking into account all information from fungal genomes, *C. purpureum* has, with 45 sequences in total, the largest number of Lac genes so far reported for a fungus. It exceeds the 23 to 25 Lac genes of the phylogenetically related, phytopathogenic *Armillaria* species, which were hitherto the 'record holders' in terms of the number of Lac genes [113, 114]. Birth-death

model based analyses of gene content evolution pinpoint Lacs as some of the most rapidly evolving gene families in *C. purpureum*, showing significant lineage-specific expansion in this species (Table 4). This was confirmed by sequence-based analyses, in the course of which we inferred 37 lineage-specific duplications in five families (Fig 4). High numbers of Lac genes were also found in the genomes of the biotrophic (parasitic) fungi *H. annosum* (14 genes, [21]), *H. irregulare* (18 genes, [115]) and *M. perniciosa* (24 genes, [55]). Remarkably, the high abundance of Lac genes was adequately reflected by the expression patterns of *C. purpureum* cultures comprising 25 Lac proteins in total. This has led to the assumption that Lacs are not only involved in fruiting body development [116] and mycelial growth [117] but may be also active in both parasitic and saprotrophic life styles of the fungus (e.g. during plant tissue invasion or cell wall modification; [55]). Such role of Lacs in virulence (e.g. by detoxifying phenolic compounds during host-defense) has already been postulated for *A. mellea* and *H. annosum* [32, 33, 115], and also for phytopathogenic Ascomycota like *Collectorichum* spp. and *Sclerotinia* spp. [118, 119] [94, 95] and even for human pathogens like *Talaromyces marneffei* [120].

A further genomic feature that may be associated with *C. purpureum*'s phytopathogenic life style is the significant expansion of non-ribosomal peptide synthetases (NRPSs), proteases and cytochrome P450 (Table 4), all of which were also found to be expanded in pathogenic *Armillaria* spp. highlighting the dual ecology of this genus [26]. The whole repertoire of genes–with expansions, for example, in Lacs, GH7 and proteases combined with relative high abundances of pathogenicity and lignocellulose-related proteins (e.g. Lacs, GH7, AA3_3) in different culture media–is, from our point of view, indicative for the switching lifestyle of *C. purpureum* and suggests its adaptability to changing environmental conditions. It confirms the affiliation of *C. purpureum* to the group of facultative parasitic agarics among the WRF (along with *Armillaria* spp. and *M. perniciosa*), which are notorious for their changing lifestyles.

Conclusion

C. purpureum is a member of the order Agaricales, family Cyphellaceae within the current taxonomic classification. According to its genome, the fungus has a diverse genetic repertoire of heme peroxidases (including MnPs, HTPs/UPOs and DyPs), LPMOs, Lacs as well as of H₂O₂generating oxidases. The presence of a single CDH, CBMs as well as of diverse cellulolytic and hemicellulolytic hydrolases perfectly matches the eco-physiological classification of C. purpureum as a WRF. Secretomic analyses revealed that the fungus realizes production and secretion of all these enzymes degrading and modifying lignocelluloses during growth in different complex solid and liquid media. Some of these media facilitated higher protein abundances in the presence of special medium components such as soybean meal (mainly with respect to GH6, seven LPMOs and one UPO, g5554), milled wood or phenol-rich agro-wastes (mainly for CBMs, PLs and LPMOs). These findings, along with the close relation of C. purpureum to other wood-rot agarics with both phytopathogenic and saprotrophic life styles, implies a particular role of these enzymes in transformation and detoxification of plant secondary metabolites, in a way of self-protection and manipulation of the environment in order to overcome plant resistance mechanisms. At the same time, they are responsible for the disintegration of biopolymers in the fungus' micro-environment and thus contribute to its acquisition of nutrients. Not least, in that context, C. purpureum seems to be a suitable candidate for further biotechnological studies regarding the disintegration and modification of recalcitrant materials.

Supporting information

S1 Fig. Remapping of raw reads to the assembly. Examples are given for few full length laccase, GH6 and GH7 genes. Coverage is given on the top in blue, tracks indicate intro-exon and CDS structure and the red arrow indicates the track with the non-synonymous SNP positions. (TIF)

S2 Fig. Gene sequences of *C. purpureum* classified according Gene Ontology domains: (left) biological process, (middle) molecular function and (right) cellular component. (TIF)

S3 Fig. Assignment of gene sequences of *C. purpureum* to main enzyme classes according to the EC nomenclature.

(TIFF)

S4 Fig. Lignocellulose-degrading enzymes in different basidiomycetous wood-decay fungi (modified according to Riley et al., 2014; published data from JGI & NCBI), yellow-WRF, grey-BRF, blue-unresolved wood-rot fungi. Agabi, Agaricus bisporus; Agrae, Agrocybe aegerita (Gupta et al. 2018); Armce, Armillaria cepistipes; Armga, Armillaria gallica; Armme, Armillaria mellea; Armso, Armillaria solidipes, Armos, Armillaria ostoyae; Aursu, Auricularia subglabra; Bjead, Bjerkandera adusta; Botbo, Botryobasidium botryosum; Cersu, Ceriporiopsis subvermispora; Chopu, Chondrostereum purpureum; Conpu, Coniophora puteana; Copci, Coprinopsis cinereus; Dacsp, Dacryopinax sp.; Dicsq, Dichomitus squalens; Fomme, Fomitiporia mediterranea; Fompi, Fomitopsis pinicola; Galma, Galerina marginata; Glotr, Gloeophyllum trabeum; Hetan, Heterobasidion annosum; Jaaar, Jaapia argillacea; Mycch, Mycena chlorophos (Tanaka et al. 2014); Monpe, Moniliophthora perniciosa; Phaca, Phanerochaete carnosa; Phchr, Phanerochaete chrysosporium; Phlbr, Phlebia brevispora; Pleos, Pleurotus ostreatus; Pospl, Postia placenta; Punst, Punctularia strigosozonata; Pycci, Pycnoporus cinnabarinus; Schco, Schizophyllum commune; Serla, Serpula lacrymans; Stehi, Stereum hirsutum; Trave, Trametes versicolor; Volvo, Volvariella volvacea and Wolco, Wolfiporia cocos; *generic peroxidase, ! probably a generic peroxidase. (PDF)

S5 Fig. Relative protein abundance (% NSAF) of enzymes produced by *C. purpureum* **during solid-state fermentation (SSF).** Beech-wood (BW, inner ring) and beech-wood plus DOR (BWD, outer ring). CAZy proteins are highlighted in bold letters. Organelle proteins include ribosomal, peroxisomal and vacuolar proteins without defined catalytic properties. Values are the mean of three replicates. (TIF)

S6 Fig. Relative protein abundance (% NSAF) of enzymes produced by *C. purpureum* **during (submerged) liquid fermentation (SF).** (A) KM (inner ring) ASKM, (middle ring) and BSKM (outer ring) and (B) SM (inner ring), ASSM (middle ring) and BSSM (outer ring). CAZy proteins are highlighted in bold letters. Organelle proteins include ribosomal, peroxisomal and vacuolar proteins without defined catalytic properties. Values are the mean of three replicates. (TIF)

S7 Fig. Heat map of the secreted CAZymes of *C. purpureum* **that show significant differences during SF in KM, SM, ASKM, ASSM, BSKM and BSSM.** Differences between treatments were corroborated with Hotelling's T2 test. Abundance is demonstrated by the normalized spectral abundance factor (% NSAF). GHs are shown in the upper side while the rest of the CAZymes (including AAs) are given in the lower side. (PDF)

S8 Fig. Principal component analysis (PCA) bi-plot of the *C. purpureum* secretome from SSF cultures. (left; BW and BWD loadings are highlighted in red) and SF cultures (right; SM, ASSM,

BSSM, KM, ASKM and BSKM loadings are highlighted in red) using NIPALS algorithms. (PDF)

S9 Fig. Neighbor-Joining phylogenetic tree of 54 class I and II peroxidase protein sequences (manganese, lignin, versatile and generic peroxidases (MnP, LiP, VP, GP), ascorbate peroxidases (APX) and cytochrome c peroxidases (CcP)). The sequences of *C. purpureum* are marked by a "g" and in bold. Numbers with asterisks indicate proteins detected in the secretomes; only complete sequences were considered. Sequences were aligned with Clustal W and Jukes-Cantor distance models were used. (TIF)

S10 Fig. Alignment of a versatile peroxidase of *Pleurotus eryngii* (2BOQ_A) with the class II peroxidases of *C. purpureum*. The three acidic amino acid residues typical for the manganese-binding sites of MnPs are indicated. (TIF)

S11 Fig. Time course of extracellular oxidoreductase production by *C. purpureum* during solid-state fermentation (SSF) of cultures containing beech wood (BW, left) and beech wood supplemented with olive-mill residues 'DOR' (BWD, right); manganese-dependent peroxidase activities (MnP, squares), unspecific peroxygenase (UPO, circles) and laccase activities (Lac, triangles) and pH (dashed line). (PDF)

S12 Fig. Time course of extracellular oxidoreductase production by *C. purpureum* during SF in cultures containing (a) KIRK medium (KM) and (b) soybean meal suspension (SM), (c) KM-ADOR (ASKM), (d) SM-ADOR (ASSM), (e) KM-birch wood (BSKM) and (f) SM-birch wood (BSSM); manganese-dependent peroxidase activities (MnP, squares), unspecific peroxygenase (UPO, circles) and laccase activities (Lac, triangles). (PDF)

S13 Fig. Neighbor-Joining phylogenetic tree of 67 multicopper oxidases (MCO) protein sequences (laccase (Lac), ferroxidase (FeOX), ascorbate oxidase (ASC). The sequences of *C. purpureum* (38 full length and 3 partial sequences (from C-terminal)) are marked by an "g" and in bold. Numbers with asterisks indicate that the proteins were found in the secretome. Sequences were aligned by Clustal W and Jukes-Cantor distance model were used. (TIF)

S14 Fig. Alignment of all full-length and partial MCO genes. The alignment was performed using ClustalW with parameters: BLOSUM cost matrix, gap opening cost = 10, gap extension cost = 0.1.

(TIF)

S15 Fig. Maximum likelihood phylogeny. Maximum Likelihood phylogeny based on 609 genes across 36 genomes. Numbers indicate bootstrap support at each node. (PDF)

S16 Fig. Numbers of inferred duplications across all orthogroups encoding GH6 cellobiohydrolases. The middle section indicates the number of orthogroups housing genes in each of the respective classes and their copy number in each species. Agabi, *Agaricus bisporus*; Agrae, *Agrocybe aegerita*; Armce, *Armillaria cepistipes*; Armga, *Armillaria gallica*; Armme, *Armillaria mellea*; Armso, *Armillaria solidipes*, Armos, *Armillaria ostoyae*; Aursu, *Auricularia subglabra*; Bjead, *Bjerkandera adusta*; Botbo, *Botryobasidium botryosum*; Cersu, *Ceriporiopsis subvermispora*; Chopu, *Chondrostereum purpureum*; Conpu, *Coniophora puteana*; Copci, *Coprinopsis* cinereus; Dacsp, Dacryopinax sp.; Dicsq, Dichomitus squalens; Fomme, Fomitiporia mediterranea; Fompi, Fomitopsis pinicola; Galma, Galerina marginata; Glotr, Gloeophyllum trabeum; Hetan, Heterobasidion annosum; Jaaar, Jaapia argillacea; Mycch, Mycena chlorophos; Monpe, Moniliophthora perniciosa; Phaca, Phanerochaete carnosa; Phchr, Phanerochaete chrysosporium; Phlbr, Phlebia brevispora; Pleos, Pleurotus ostreatus; Pospl, Postia placenta; Punst, Punctularia strigosozonata; Pycci, Pycnoporus cinnabarinus; Schco, Schizophyllum commune; Serla, Serpula lacrymans; Stehi, Stereum hirsutum; Trave, Trametes versicolor; Volvo, Volvariella volvacea and Wolco, Wolfiporia cocos. (PDF)

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References

- 1. Kirk TK, Farell RL. Enzymatic "combustion": the microbial degradation of lignin. Annu Rev Microbiol. 1987; 41:465–505. https://doi.org/10.1146/annurev.mi.41.100187.002341 PMID: 3318677
- Eastwood DC, Floudas D, Binder M, Majcherczyk A, Schneider P, Aerts A, et al. The plant cell walldecomposing machinery underlies the functional diversity of forest fungi. Science. 2011; 333:762–5. https://doi.org/10.1126/science.1205411 PMID: 21764756
- Blanchette RA, Nilsson T, Daniel G, Abad A. Biological degradation of wood. In: Rowell RM, Barbour RJ, editors. Archaeological wood: properties, chemistry, and preservation; advances in chemistry series. American Chemical Society; 1990. p. 141–74.
- 4. Eriksson KE. Biotechnology in the pulp and paper industry. Wood Sci Technol. 1990; 24:79–101.
- 5. Hatakka A. Lignin-modifying enzymes from selected white-rot fungi: production and role from in lignin degradation. FEMS Microbiol Rev. 1994; 13:125–35.
- Kersten P, Cullen D. Extracellular oxidative systems of the lignin-degrading Basidiomycete Phanerochaete chrysosporium. Fungal Genet Biol. 2007; 44:77–87. https://doi.org/10.1016/j.fgb.2006.07.007 PMID: 16971147

- Lundell TK, Makela MR, Hilden K. Lignin-modifying enzymes in filamentous basidiomycetes: ecological, functional and phylogenetic review. J Basic Microbiol. 2010; 50:5–20. https://doi.org/10.1002/ jobm.200900338 PMID: 20175122
- Wall RE. The fungus Chondrostereum purpureum as a silvicide to control stump sprouting in hardwoods. North J Appl For 1990; 7:17–9.
- Hamberg L, Vartiamäki H, Hantula J. Breeding increases the efficacy of *Chondrostereum purpureum* in the sprout control of birch. PLoS ONE. 2015; 10:e0117381. https://doi.org/10.1371/journal.pone. 0117381 PMID: 25674794
- Hamberg L, Hantula J. The efficacy of six elite isolates of the fungus Chondrostereum purpureum against the sprouting of European aspen. J Environ Manage. 2016; 171:217–24. <u>https://doi.org/10.1016/j.jenvman.2016.02.016</u> PMID: 26899306
- 11. Butler EJ, Jones SG (1949) Silver leaf disease of plum, *Stereum purpureum* (Fr.) Fr. Plant Pathology. MacMillian & Co., London, pp. 763–767.
- Strunz GM, Bethell R, Dumas MT, Boyonoski N. On a new synthesis of sterpurene and the bioactivity of some related *Chondrostereum purpureum* sesquiterpene metabolites. Can J Chem. 1997; 75:742– 53.
- 13. Leatham GF. The ligninolytic activities of *Lentinus edodes* and *Phanerochaete chrysosporium*. Appl Microbiol Biotechnol. 1986; 24:51–8.
- ten Have R, Teunissen PJM. Oxidative mechanisms involved in lignin degradation by white-rot fungi. Chem Rev. 2001; 101:3397–413. PMID: <u>11749405</u>
- Shamoun SF, Wall RE. Characterization of Canadian isolates of *Chondrostereum purpureum* by protein content, API ZYM and isozyme analyses. Eur J For Path. 1996; 26:333–42.
- 16. Rayner ADM. Fungal colonization of hardwood stumps from natural sources. II. Basidiomycetes. Trans Brit Mycol Soc. 1977; 69:303–12.
- Bellgard SE, Johnson VW, Than DJ, Anand N, Winks CJ, Ezeta G, et al. Use of the silverleaf fungus *Chondrostereum purpureum* for biological control of stump-sprouting, riparian weedy tree species in New Zealand. Australas Plant Pathol. 2014; 43:321–6.
- Hamberg L, Malmivaara-Lämsä M, Löfström I, Hantula J. Effects of a biocontrol agent Chondrostereum purpureum on sprouting of Sorbus aucuparia and Populus tremula after four growing seasons. BioControl. 2014; 59:125–37.
- Matheny PB, Curtis JM, Hofstetter V, Aime MC, Moncalvo J-M, Z-W. G, et al. Major clades of Agaricales: a multilocus phylogenetic overview. Mycologia. 2006; 98:982–95. PMID: 17486974
- Djeddour D, Shaw R. Classical biocontrol of weeds in Europe–are we pushing against an open door. In: Impson FAC, Kleinjan CA, Hoffmann JH, editors. Proceedings of the XIV International Symposium on Biological Control of Weeds. 2014; p. 129–34.
- Riley R, Salamov AA, Brown DW, Nagy LG, Floudas D, Held BW, et al. Extensive sampling of basidiomycete genomes demonstrates inadequacy of the white-rot/brown-rot paradigm for wood decay fungi. Proc Natl Acad Sci USA. 2014; 111:9923–8. <u>https://doi.org/10.1073/pnas.1400592111</u> PMID: 24958869
- Kohler A, Kuo A, Nagy LG, Morin E, Barry KW, Buscot F, et al. Convergent losses of decay mechanisms and rapid turnover of symbiosis genes in mycorrhizal mutualists. Nat Genet. 2015; 47:410–5. https://doi.org/10.1038/ng.3223 PMID: 25706625
- Hess J, Skrede I, Chaib De Mares M, Hainaut M, Henrissat B, Pringle A. Rapid divergence of genome architectures following the origin of an ectomycorrhizal symbiosis in the genus Amanita. Mol Biol Evol. 2018; 35:2786–804. https://doi.org/10.1093/molbev/msy179 PMID: 30239843
- Floudas D, Binder M, Riley R, Barry K, Blanchette RA, Henrissat B, et al. The Paleozoic origin of enzymatic lignin decomposition reconstructed from 31 fungal genomes. Science. 2012; 336:1715–9. https://doi.org/10.1126/science.1221748 PMID: 22745431
- Nagy LG, Riley R, Tritt A, Adam C, Daum C, Floudas D, et al. Comparative genomics of early-diverging mushroom-forming fungi provides insights into the origins of lignocellulose decay capabilities. Mol Biol Evol. 2016; 33:959–70. https://doi.org/10.1093/molbev/msv337 PMID: 26659563
- Sipos G, Prasanna AN, Walter MC, O'Connor E, Bálint B, Krizsán K, et al. Genome expansion and lineage-specific genetic innovations in the forest pathogenic fungi *Armillaria*. Nat Ecol Evol. 2017; 1:1931–41. https://doi.org/10.1038/s41559-017-0347-8 PMID: 29085064
- 27. Fernández-Fueyo E, Ruiz-Dueñas FJ, López-Lucendo MF, Pérez-Boada M, Rencoret J, Gutiérrez A, et al. A secretomic view of woody and nonwoody lignocellulose degradation by *Pleurotus ostreatus*. Biotechnol Biofuels. 2016; 9:49. https://doi.org/10.1186/s13068-016-0462-9 PMID: 26933449
- 28. Vanden Wymelenberg A, Gaskell J, Mozuch M, Kersten P, Sabat G, Martinez D, et al. Transcriptome and secretome analyses of *Phanerochaete chrysosporium* reveal complex patterns of gene

expression. Appl Environ Microbiol. 2009; 75:4058–68. https://doi.org/10.1128/AEM.00314-09 PMID: 19376920

- Vanden Wymelenberg A, Sabat G, Martinez D, Rajangam AS, Teeri TT, Gaskell J, et al. The *Phanero-chaete chrysosporium* secretome: database predictions and initial mass spectrometry peptide identifications in cellulose-grown medium. J Biotechnol. 2005; 118:17–34. <u>https://doi.org/10.1016/j.jbiotec.</u> 2005.03.010 PMID: 15888348
- 30. Vanden Wymelenberg A, Minges P, Sabat G, Martinez D, Aerts A, Salamov A, et al. Computational analysis of the *Phanerochaete chrysosporium* v2.0 genome database and mass spectrometry identification of peptides in ligninolytic cultures reveal complex mixtures of secreted proteins. Fungal Genet Biol. 2006; 43:343–56. https://doi.org/10.1016/j.fgb.2006.01.003 PMID: 16524749
- Martinez D, Challacombe J, Morgenstern I, Hibbett D, Schmoll M, Kubicek CP, et al. Genome, transcriptome, and secretome analysis of wood decay fungus *Postia placenta* supports unique mechanisms of lignocellulose conversion. Proc Natl Acad Sci USA. 2009; 106:1954–9. https://doi.org/10.1073/pnas.0809575106 PMID: 19193860
- Yakovlev IA, Hietala AM, Courty P-E, Lundell T, Solheim H, Fossdal CG. Genes associated with lignin degradation in the polyphagous white-rot pathogen *Heterobasidion irregulare* show substrate-specific regulation. Fungal Genet Biol. 2013; 56:17–24. <u>https://doi.org/10.1016/j.fgb.2013.04.011</u> PMID: 23665189
- Collins C, Keane TM, Turner DJ, O'Keeffe G, Fitzpatrick DA, Doyle S. Genomic and proteomic dissection of the ubiquitous plant pathogen, *Armillaria mellea*: toward a new infection model system. J Proteome Res. 2013; 12:2552–70. https://doi.org/10.1021/pr301131t PMID: 23656496
- Kuhar S, Nair LM, Kuhad RC. Pretreatment of lignocellulosic material with fungi capable of higher lignin degradation and lower carbohydrate degradation improves substrate acid hydrolysis and the eventual conversion to ethanol. Can J Microbiol. 2008; 54:305–13. https://doi.org/10.1139/w08-003 PMID: 18389003
- Hofrichter M, Fritsche W. Depolymerization of low-rank coal by extracellular fungal enzyme systems. III. *In vitro* depolymerization of coal humic acids by a crude preparation of manganese peroxidase of the white-rot fungus *Nematoloma frowardii* b19. Appl Microbiol Biotechnol. 1997; 47:566–71.
- **36.** Hatakka A, Hammel KE. Fungal biodegradation of lignocelluloses. Mycota, Industrial applications. Springer Berlin Heidelberg; 2010. p. 319–40.
- 37. Reina R, Garcia Sánchez M, Liers C, García-Romera I, Aranda E. An overview of fungal applications in the valorization of lignocellulosic agricultural by-products: The case of two-phase olive mill wastes. In: Prasad R, editor. Mycoremediation and Environmental Sustainability. Fungal Biology. Springer; 2018. p. 213–38.
- Reina R, Liers C, Ocampo JA, Garcia-Romera I, Aranda E. Solid state fermentation of olive mill residues by wood- and dung-dwelling Agaricomycetes: effects on peroxidase production, biomass development and phenol phytotoxicity. Chemosphere. 2013; 93:1406–12. <u>https://doi.org/10.1016/j.chemosphere.2013.07.006 PMID</u>: 23920362
- Aranda E, Sampedro I, A. Ocampo J, García-Romera I. Phenolic removal of olive-mill dry residues by laccase activity of white-rot fungi and its impact on tomato plant growth. Int Biodeterior Biodegradation. 2006; 58:176–9.
- Sampedro I, Romero C, Ocampo JA, Brenes M, García I. Removal of monomeric phenols in dry mill olive residue by saprobic fungi. J Agric Food Chem. 2004; 52:4487–92. <u>https://doi.org/10.1021/</u> if0400563 PMID: 15237956
- Aranda E, Sampedro I, Ocampo JA, García-Romera I. Contribution of hydrolytic enzymes produced by saprophytic fungi to the decrease in plant toxicity caused by water-soluble substances in olive mill dry residue. Appl Microbiol Biotechnol. 2004; 64:132–5. <u>https://doi.org/10.1007/s00253-003-1368-6</u> PMID: 12811426
- Girard V, Dieryckx C, Job C, Job D. Secretomes: the fungal strike force. Proteomics. 2013; 13:597– 608. https://doi.org/10.1002/pmic.201200282 PMID: 23349114
- **43.** Carabajal M, Kellner H, Levin L, Jehmlich N, Hofrichter M, Ullrich R. The secretome of *Trametes versi-color* grown on tomato juice medium and purification of the secreted oxidoreductases including a versatile peroxidase. J Biotech. 2013; 168:15–23.
- Morin E, Kohler A, Baker AR, Foulongne-Oriol M, Lombard V, Nagy LG, et al. Genome sequence of the button mushroom *Agaricus bisporus* reveals mechanisms governing adaptation to a humic-rich ecological niche. Proc Natl Acad Sci USA. 2012; 109:17501–6. <u>https://doi.org/10.1073/pnas.</u> 1206847109 PMID: 23045686
- 45. Wolfenden BS, Willson RL. Radical-cations as reference chromogens in kinetic studies of ono-electron transfer reactions: pulse radiolysis studies of 2,2-azinobis-(3-ethylbenzthiazoline-6-sulphonate). J Chem Soc, Perkin Transactions 2. 1982:805–12.

- Liers C, Bobeth C, Pecyna MJ, Ullrich R, Hofrichter M. DyP-like peroxidases of the jelly fungus Auricularia auricula-judae oxidize nonphenolic lignin model compounds and high-redox potential dyes. Appl Microbiol Biotechnol. 2010; 85:1869–79. https://doi.org/10.1007/s00253-009-2173-7 PMID: 19756587
- Liers C, Pecyna MJ, Kellner H, Worrich A, Zorn H, Steffen KT, et al. Substrate oxidation by dye-decolorizing peroxidases (DyPs) from wood- and litter-degrading agaricomycetes compared to other fungal and plant heme-peroxidases. Appl Microbiol Biotechnol. 2013; 97:5839–49. <u>https://doi.org/10.1007/</u> s00253-012-4521-2 PMID: 23111597
- Ullrich R, Nüske J, Scheibner K, Spantzel J, Hofrichter M. Novel haloperoxidase from the agaric basidiomycete Agrocybe aegerita oxidizes aryl alcohols and aldehydes. Appl Environ Microbiol. 2004; 70:4575–81. https://doi.org/10.1128/AEM.70.8.4575-4581.2004 PMID: 15294788
- Kellner H, Pecyna MJ, Buchhaupt M, Ullrich R, Hofrichter M. Draft genome sequence of the chloroperoxidase-producing fungus *Caldariomyces fumago* Woronichin DSM1256. Genome Announce. 2016; 4:e00774–16.
- Chevreux B (2005) MIRA: An automated genome and EST assembler. PhD thesis. Ruprecht-Karls University, Heidelberg, Germany.
- Parra G, Bradnam K, Korf I. CEGMA: a pipeline to accurately annotate core genes in eukaryotic genomes. Bioinformatics. 2007; 23:1061–7. <u>https://doi.org/10.1093/bioinformatics/btm071</u> PMID: 17332020
- 52. Gurevich A, Saveliev V, Vyahhi N, Tesler G. QUAST: quality assessment tool for genome assemblies. Bioinformatics. 2013; 29:1072–5. https://doi.org/10.1093/bioinformatics/btt086 PMID: 23422339
- Stanke M, Morgenstern B. AUGUSTUS: a web server for gene prediction in eukaryotes that allows user-defined constraints. Nucleic Acids Res. 2005; 33:W465–7. <u>https://doi.org/10.1093/nar/gki458</u> PMID: 15980513
- Yin Y, Mao X, Yang J, Chen X, Mao F, Xu Y. dbCAN: a web resource for automated carbohydrateactive enzyme annotation. Nucleic Acids Res. 2012; 40:W445–W51. <u>https://doi.org/10.1093/nar/ gks479 PMID: 22645317</u>
- 55. Mondego JMC, Carazzolle MF, Costa GGL, Formighieri EF, Parizzi LP, Rincones J, et al. A genome survey of *Moniliophthora perniciosa* gives new insights into Witches' Broom Disease of cacao. BMC Genomics. 2008; 9:548. https://doi.org/10.1186/1471-2164-9-548 PMID: 19019209
- 56. Stajich JE, Wilke SK, Ahren D, Au CH, Birren BW, Borodovsky M, et al. Insights into evolution of multicellular fungi from the assembled chromosomes of the mushroom *Coprinopsis cinerea* (*Coprinus cinereus*). Proc Natl Acad Sci USA. 2010; 107:11889–94. <u>https://doi.org/10.1073/pnas.1003391107</u> PMID: 20547848
- Bao D, Gong M, Zheng H, Chen M, Zhang L, Wang H, et al. Sequencing and comparative analysis of the straw mushroom (*Volvariella volvacea*) genome. PLoS ONE. 2013; 8:e58294. <u>https://doi.org/10. 1371/journal.pone.0058294 PMID: 23526973</u>
- Ruiz-Duenas FJ, Lundell T, Floudas D, Nagy LG, Barrasa JM, Hibbett DS, et al. Lignin-degrading peroxidases in Polyporales: an evolutionary survey based on 10 sequenced genomes. Mycologia. 2013; 105:1428–44. https://doi.org/10.3852/13-059 PMID: 23921235
- Levasseur A, Lomascolo A, Chabrol O, Ruiz-Dueñas FJ, Boukhris-Uzan E, Piumi F, et al. The genome of the white-rot fungus *Pycnoporus cinnabarinus*: a basidiomycete model with a versatile arsenal for lignocellulosic biomass breakdown. BMC Genomics. 2014; 15:486. <u>https://doi.org/10.1186/1471-2164-15-486 PMID: 24942338</u>
- 60. Gupta DK, Rühl M, Mishra B, Kleofas V, Hofrichter M, Herzog R, et al. The genome sequence of the commercially cultivated mushroom Agrocybe aegerita reveals a conserved repertoire of fruiting-related genes and a versatile suite of biopolymer-degrading enzymes. BMC Genomics. 2018; 19:48. https://doi.org/10.1186/s12864-017-4430-y PMID: 29334897
- Li L. OrthoMCL: Identification of ortholog groups for eukaryotic genomes. Genome Res. 2003; 13:2178–89. https://doi.org/10.1101/gr.1224503 PMID: 12952885
- Löytynoja A, Goldman N. Phylogeny-aware gap placement prevents errors in sequence alignment and evolutionary analysis. Science. 2008; 320:1632–5. <u>https://doi.org/10.1126/science.1158395</u> PMID: 18566285
- Capella-Gutiérrez S, Silla-Martínez JM, Gabaldón T. trimAl: a tool for automated alignment trimming in large-scale phylogenetic analyses. Bioinformatics. 2009; 25:1972–3. https://doi.org/10.1093/ bioinformatics/btp348 PMID: 19505945
- 64. Morel B, Kozlov AM, Stamatakis A. ParGenes: a tool for massively parallel model selection and phylogenetic tree inference on thousands of genes. Bioinformatics. 2018.
- Kozlov A, Darriba D, Flouri T, Morel B, Stamatakis A. RAxML-NG: A fast, scalable, and user-friendly tool for maximum likelihood phylogenetic inference. bioRxiv. 2018:447110.

- 66. Zhang C, Rabiee M, Sayyari E, Mirarab S. ASTRAL-III: polynomial time species tree reconstruction from partially resolved gene trees. BMC Bioinformatics. 2018; 19:153. https://doi.org/10.1186/s12859-018-2129-y PMID: 29745866
- Lanfear R, Senfeld T, Frandsen PB, Wright AM, Calcott B. PartitionFinder 2: New methods for selecting partitioned models of evolution for molecular and morphological phylogenetic analyses. Mol Biol Evol. 2016; 34:772–3.
- Han MV, Thomas GWC, Lugo-Martinez J, Hahn MW. Estimating gene gain and loss rates in the presence of error in genome assembly and annotation using CAFE 3. Mol Biol Evol. 2013; 30:1987–97. https://doi.org/10.1093/molbev/mst100 PMID: 23709260
- **69.** Sanderson MJ. r8s: inferring absolute rates of molecular evolution and divergence times in the absence of a molecular clock. Bioinformatics. 2003; 19:301–2. PMID: <u>12538260</u>
- Wu Y-C, Rasmussen MD, Bansal MS, Kellis M. TreeFix: statistically informed gene tree error correction using species trees. Syst Biol. 2012; 62:110–20. <u>https://doi.org/10.1093/sysbio/sys076</u> PMID: 22949484
- Stolzer M, Lai H, Xu M, Sathaye D, Vernot B, Durand D. Inferring duplications, losses, transfers and incomplete lineage sorting with nonbinary species trees. Bioinformatics. 2012; 28:i409–i15. https://doi. org/10.1093/bioinformatics/bts386 PMID: 22962460
- 72. Hammer Ø, Harper DAT, Ryan PD. PAST: paleontological statistics software package for education and data analysis. Palaeontol Electron. 2001; 4.
- 73. Reina R, Kellner H, Jehmlich N, Ullrich R, García-Romera I, Aranda E, et al. Differences in the secretion pattern of oxidoreductases from *Bjerkandera adusta* induced by a phenolic olive mill extract. Fungal Genet Biol. 2014;72.
- Zybailov B, Mosley AL, Sardiu ME, Coleman MK, Florens L, Washburn MP. Statistical analysis of membrane proteome expression changes in *Saccharomyces cerevisiae*. J Proteome Res. 2006; 5:2339–47. https://doi.org/10.1021/pr060161n PMID: 16944946
- 75. Team RC. R: A language and environment for statistical computing. R foundation for statistical computing: Vienna, Austria. ISBN 3-900051-07-0.
- 76. Choinowski T, Blodig W, Winterhalter KH, Piontek K. The crystal structure of lignin peroxidase at 1.70 Å resolution reveals a hydroxy group on the Cβ of tryptophan 171: A novel radical site formed during the redox cycle. J Mol Biol. 1999; 286:809–27. https://doi.org/10.1006/jmbi.1998.2507 PMID: 10024453
- Hofrichter M, Kellner H, Pecyna MJ, Ullrich R. Fungal unspecific peroxygenases: heme-thiolate proteins that combine peroxidase and cytochrome P450 properties. Adv Exp Med Biol. 2015; 851:341– 68. https://doi.org/10.1007/978-3-319-16009-2_13 PMID: 26002742
- Matheny PB, Wang Z, Binder M, Curtis JM, Lim YW, Nilsson RH, et al. Contributions of rpb2 and tef1 to the phylogeny of mushrooms and allies (Basidiomycota, Fungi). Mol Phylogenet Evol. 2007; 43:430–51. https://doi.org/10.1016/j.ympev.2006.08.024 PMID: 17081773
- Tanaka Y, Kasuga D, Oba Y, Hase S, Sato K. Genome sequence of the luminous mushroom Mycena chlorophos for searching fungal bioluminescence genes. Luminescence. 2014; 29:47–8.
- Mohanta TK, Bae H. The diversity of fungal genome. Biol Proced Online. 2015; 17:8. https://doi.org/10.1186/s12575-015-0020-z PMID: 25866485
- Hori C, Gaskell J, Igarashi K, Kersten P, Mozuch M, Samejima M, et al. Temporal alterations in the secretome of the selective ligninolytic fungus *Ceriporiopsis subvermispora* during growth on aspen wood reveal this organism's strategy for degrading lignocellulose. Appl Environ Microbiol. 2014; 80:2062–70. https://doi.org/10.1128/AEM.03652-13 PMID: 24441164
- 82. Vanden Wymelenberg A, Gaskell J, Mozuch M, BonDurant SS, Sabat G, Ralph J, et al. Significant alteration of gene expression in wood decay fungi *Postia placenta* and *Phanerochaete chrysosporium* by plant species. Appl Environ Microbiol. 2011; 77:4499–507. <u>https://doi.org/10.1128/AEM.00508-11</u> PMID: 21551287
- Gaskell J, Marty A, Mozuch M, Kersten PJ, Splinter BonDurant S, Sabat G, et al. Influence of *Populus* genotype on gene expression by the wood decay fungus *Phanerochaete chrysosporium*. Appl Environ Microbiol. 2014; 80:5828–35. https://doi.org/10.1128/AEM.01604-14 PMID: 25015893
- Shishkova E, Hebert AS, Coon JJ. Now, more than ever, proteomics needs better chromatography. Cell Syst. 2016; 3:321–4. https://doi.org/10.1016/j.cels.2016.10.007 PMID: 27788355
- van den Brink J, de Vries RP. Fungal enzyme sets for plant polysaccharide degradation. Appl Microbiol Biotechnol. 2011; 91:1477–92. https://doi.org/10.1007/s00253-011-3473-2 PMID: 21785931
- Di Pietro A, Roncero MI. Cloning, expression, and role in pathogenicity of pg1 encoding the major extracellular endopolygalacturonase of the vascular wilt pathogen *Fusarium oxysporum*. Mol Plant Microbe Interact. 1998; 11:91–8. https://doi.org/10.1094/MPMI.1998.11.2.91 PMID: 9450333

- Sprockett DD, Piontkivska H, Blackwood CB. Evolutionary analysis of glycosyl hydrolase family 28 (GH28) suggests lineage-specific expansions in necrotrophic fungal pathogens. Gene. 2011; 479:29– 36. https://doi.org/10.1016/j.gene.2011.02.009 PMID: 21354463
- Cord-Landwehr S, Melcher RL, Kolkenbrock S, Moerschbacher BM. A chitin deacetylase from the endophytic fungus *Pestalotiopsis* sp. efficiently inactivates the elicitor activity of chitin oligomers in rice cells. Sci Rep. 2016; 6:38018. https://doi.org/10.1038/srep38018 PMID: 27901067
- Simpson RM, Hekezen RV, Lune FV, Brewster D, Christeller JT, Spiers AG. Extracellular enzymes of Chondrostereum purpureum, causal fungus of silverleaf disease. N Z Plant Prot. 2001; 54:202–8.
- 90. Ogawa S, Shimizu T, Ohki H, Araya T, Okuno T, Miyairi K. Expression, purification, and analyses of glycosylation and disulfide bonds of *Stereum purpureum* endopolygalacturonase I in *Pichia pastoris*. Protein Expr Purif. 2009; 65:15–22. https://doi.org/10.1016/j.pep.2008.12.014 PMID: 19174189
- Hamada S, Toda K, Ogawa S, Kubota K, Miyairi K. Characterization of the effects of C-terminal prosequence on self-inactivation of *Stereum purpureum* endopolygalacturonase I. FEMS Microbiol Lett. 2015; 362:fnv134. https://doi.org/10.1093/femsle/fnv134 PMID: 26293910
- Busk PK, Lange L. Classification of fungal and bacterial lytic polysaccharide monooxygenases. BMC Genomics. 2015; 16:368. https://doi.org/10.1186/s12864-015-1601-6 PMID: 25956378
- Sabbadin F, Hemsworth GR, Ciano L, Henrissat B, Dupree P, Tryfona T, et al. An ancient family of lytic polysaccharide monooxygenases with roles in arthropod development and biomass digestion. Nat Commun. 2018; 9:756. https://doi.org/10.1038/s41467-018-03142-x PMID: 29472725
- Adav SS, Chao LT, Sze SK. Quantitative secretomic analysis of *Trichoderma reesei* strains reveals enzymatic composition for lignocellulosic biomass degradation. Mol Cell Proteomics. 2012; 11: M111.012419.
- Hori C, Ishida T, Igarashi K, Samejima M, Suzuki H, Master E, et al. Analysis of the *Phlebiopsis gigan*tea genome, transcriptome and secretome provides insight into its pioneer colonization strategies of wood. PLoS GENET. 2014; 10:e1004759. <u>https://doi.org/10.1371/journal.pgen.1004759</u> PMID: 25474575
- Kuuskeri J, Häkkinen M, Laine P, Smolander OP, Tamene F, Miettinen S, et al. Time-scale dynamics of proteome and transcriptome of the white-rot fungus *Phlebia radiata*: growth on spruce wood and decay effect on lignocellulose. Biotechnol Biofuels. 2016; 9:192. <u>https://doi.org/10.1186/s13068-016-0608-9 PMID: 27602055</u>
- 97. Gaskell J, Blanchette RA, Stewart PE, BonDurant SS, Adams M, Sabat G, et al. Transcriptome and secretome analyses of the wood decay fungus *Wolfiporia cocos* support alternative mechanisms of lignocellulose conversion. Appl Environ Microbiol. 2016; 82:3979–87. https://doi.org/10.1128/AEM. 00639-16 PMID: 27107121
- Kracher D, Scheiblbrandner S, Felice AK, Breslmayr E, Preims M, Ludwicka K, et al. Extracellular electron transfer systems fuel cellulose oxidative degradation. Science. 2016; 352:1098–101. <u>https:// doi.org/10.1126/science.aaf3165</u> PMID: 27127235
- Reina R, Ullrich R, García-Romera I, Liers C, Aranda E. Integrated biovalorization of wine and olive mill by-products to produce enzymes of industrial interest and soil amendments. Span J Agric Res. 2016; 14.
- Langston JA, Shaghasi T, Abbate E, Xu F, Vlasenko E, Sweeney MD. Oxidoreductive cellulose depolymerization by the enzymes cellobiose dehydrogenase and glycoside hydrolase 61. Appl Environ Microbiol. 2011; 77:7007–15. https://doi.org/10.1128/AEM.05815-11 PMID: 21821740
- 101. Salvachua D, Martinez AT, Tien M, Lopez-Lucendo MF, Garcia F, de Los Rios V, et al. Differential proteomic analysis of the secretome of *Irpex lacteus* and other white-rot fungi during wheat straw pre-treatment. Biotechnol Biofuels. 2013; 6:115. https://doi.org/10.1186/1754-6834-6-115 PMID: 23937687
- 102. Hilden K, Martinez AT, Hatakka A, Lundell T. The two manganese peroxidases Pr-MnP2 and Pr-MnP3 of *Phlebia radiata*, a lignin-degrading basidiomycete, are phylogenetically and structurally divergent. Fungal Genet Biol. 2005; 42:403–19. https://doi.org/10.1016/j.fgb.2005.01.008 PMID: 15809005
- 103. Hofrichter M, Vares T, Kalsi M, Galkin S, Scheibner K, Fritsche W, et al. Production of manganese peroxidase and organic acids and mineralization of ¹⁴C-labelled lignin (¹⁴C-DHP) during solid-state fermentation of wheat straw with the white rot fungus *Nematoloma frowardii*. Appl Environ Microbiol. 1999; 65:1864–70. PMID: 10223971
- 104. Hofrichter M, Ullrich R. Oxidations catalyzed by fungal peroxygenases. Curr Opin Chem Biol. 2014; 19:116–25. https://doi.org/10.1016/j.cbpa.2014.01.015 PMID: 24607599
- 105. Gröbe G, Ullrich R, Pecyna MJ, Kapturska D, Friedrich S, Hofrichter M, et al. High-yield production of aromatic peroxygenase by the agaric fungus *Marasmius rotula*. AMB Express. 2011; 1:31. <u>https://doi.org/10.1186/2191-0855-1-31</u> PMID: 21988939

- 106. Pecyna MJ (2016) Molekularbiologische Charakterisierung von Häm-Thiolat- und DyP-type Peroxidasen ausgewählter Basidiomyceten. PhD thesis. Dresden University of Technology, International Institut Zittau; 25.10.2016. 241 p.
- 107. Reina R, Liers C, García-Romera I, Aranda E. Enzymatic mechanisms and detoxification of dry olivemill residue by *Cyclocybe aegerita, Mycetinis alliaceus* and *Chondrostereum purpureum*. Int Biodeterior Biodegradation. 2017; 117:89–96.
- 108. Kinne M, Poraj-Kobielska M, Ralph SA, Ullrich R, Hofrichter M, Hammel KE. Oxidative cleavage of diverse ethers by an extracellular fungal peroxygenase. J Biol Chem. 2009; 284:29343–9. <u>https://doi.org/10.1074/jbc.M109.040857</u> PMID: 19713216
- 109. Karich A, Ullrich R, Scheibner K, Hofrichter M. Fungal unspecific peroxygenases oxidize the majority of organic EPA priority pollutants. Front Microbiol. 2017; 8:1463. https://doi.org/10.3389/fmicb.2017. 01463 PMID: 28848501
- Hernandez-Ortega A, Ferreira P, Martinez AT. Fungal aryl-alcohol oxidase: a peroxide-producing flavoenzyme involved in lignin degradation. Appl Microbiol Biotechnol. 2012; 93:1395–410. https://doi. org/10.1007/s00253-011-3836-8 PMID: 22249717
- Kersten P, Cullen D. Copper radical oxidases and related extracellular oxidoreductases of wooddecay Agaricomycetes. Fungal Genet Biol. 2014; 72:124–30. https://doi.org/10.1016/j.fgb.2014.05. 011 PMID: 24915038
- 112. Liers C, Arnstadt T, Ullrich R, Hofrichter M. Patterns of lignin degradation and oxidative enzyme secretion by different wood- and litter-colonizing basidiomycetes and ascomycetes grown on beech-wood. FEMS Microbiol Ecol. 2011; 78:91–102. https://doi.org/10.1111/j.1574-6941.2011.01144.x PMID: 21631549
- 113. Kilaru S, Hoegger PJ, Kues U. The laccase multi-gene family in *Coprinopsis cinerea* has seventeen different members that divide into two distinct subfamilies. Curr Genet. 2006; 50:45–60. https://doi.org/10.1007/s00294-006-0074-1 PMID: 16775746
- 114. Rühl M, Majcherczyk A, Kües U. Lcc1 and Lcc5 are the main laccases secreted in liquid cultures of Coprinopsis cinerea strains. Antonie Leeuwenhoek. 2013; 103:1029–39. <u>https://doi.org/10.1007/</u> s10482-013-9883-7 PMID: 23340718
- Kuo HC, Detry N, Choi J, Lee YH. Potential roles of laccases on virulence of *Heterobasidion annosum* s.s. Microb Pathog. 2015; 81:16–21. https://doi.org/10.1016/j.micpath.2015.03.004 PMID: 25757691
- Pezzella C, Lettera V, Piscitelli A, Giardina P, Sannia G. Transcriptional analysis of *Pleurotus ostrea*tus laccase genes. Appl Microbiol Biotechnol. 2013; 97:705–17. <u>https://doi.org/10.1007/s00253-012-</u> 3980-9 PMID: 22395908
- 117. Zhang J, Chen H, Chen M, Ren A, Huang J, Wang H, et al. Cloning and functional analysis of a laccase gene during fruiting body formation in *Hypsizygus marmoreus*. Microbiol Res. 2015; 179:54–63. https://doi.org/10.1016/j.micres.2015.06.005 PMID: 26411895
- 118. Lin SY, Okuda S, Ikeda K, Okuno T, Takano Y. LAC2 encoding a secreted laccase is involved in appressorial melanization and conidial pigmentation in *Collectotrichum orbiculare*. Mol Plant Microbe Interact. 2012; 25:1552–61. https://doi.org/10.1094/MPMI-05-12-0131-R PMID: 22934563
- 119. Coman C, Mot AC, Gal E, Parvu M, Silaghi-Dumitrescu R. Laccase is upregulated via stress pathways in the phytopathogenic fungus *Sclerotinia sclerotiorum*. Fungal Biol. 2013; 117:528–39. https://doi.org/ 10.1016/j.funbio.2013.05.005 PMID: 23931118
- 120. Williamson PR. Role of laccase in the virulence of *Talaromyces marneffei*: A common link between AIDS-related fungal pathogens? Virulence. 2016; 7:627–9. https://doi.org/10.1080/21505594.2016. 1198867 PMID: 27282335