



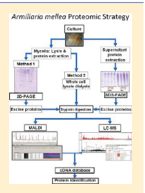
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Genomic and Proteomic Dissection of the Ubiquitous Plant Pathogen, Armillaria mellea: Toward a New Infection Model System

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Supporting Information

ABSTRACT: Armillaria mellea is a major plant pathogen. Yet, no large-scale "-omics" data are available to enable new studies, and limited experimental models are available to investigate basidiomycete pathogenicity. Here we reveal that the A. mellea genome comprises 58.35 Mb, contains 14473 gene models, of average length 1575 bp (4.72 introns/gene). Tandem mass spectrometry identified 921 mycelial (n = 629 unique) and secreted (n = 183 unique) proteins. Almost 100 mycelial proteins were either species-specific or previously unidentified at the protein level. A number of proteins (n = 111) was detected in both mycelia and culture supernatant extracts. Signal sequence occurrence was 4-fold greater for secreted (50.2%) compared to mycelial (12%) proteins. Analyses revealed a rich reservoir of carbohydrate degrading enzymes, laccases, and lignin peroxidases in the A. mellea proteome, reminiscent of both basidiomycete and ascomycete glycodegradative arsenals. We discovered that A. mellea exhibits a specific killing effect against Candida albicans during coculture. Proteomic investigation of this interaction revealed the unique expression of defensive and potentially offensive A. mellea proteins (n = 30). Overall, our data reveal



new insights into the origin of basidiomycete virulence and we present a new model system for further studies aimed at deciphering fungal pathogenic mechanisms.

KEYWORDS: Armillaria mellea, basidiomycete proteomics, basidiomycete genomics, Next Generation Sequencing, fungal proteomics, glycosidases, Candida

INTRODUCTION

Basidiomycete fungi are attracting increased interest because of their plant pathogenic properties, along with an ability to produce enzymes, in particular glycoside hydrolases, which can degrade recalcitrant carbon sources, thereby increasing the range of substrates available for bioethanol production. Although the genomes of a number of basidiomycete species have been sequenced and a number of proteomic studies have been carried out, one genera in particular, Armillaria, has received little attention to date.^{2–4} Yet, Armillaria spp. are among the most serious plant root pathogens and cause major economic losses following infection of tree species of both agronomic importance and those grown for timber production purposes.⁵ The genera consists of at least 40 species, including A. mellea, A. gallica, A. tabescens, and A. borealis, with the former species attracting the most attention.

In particular, A. mellea has been studied because of its virulence, bioluminescent properties, ability to produce unusual natural products and from a strain typing perspective, for definitive identification of the Armillaria spp. responsible for timber crop infection. 6 Recently, a novel infection model system for A. mellea has been established which deployed confocal microscopy and quantitative PCR for the estimation of fungal biomass produced during grape rootstock infection.⁷ This approach has facilitated a more quantitative assessment of fungal infection of plant roots, in addition to distinguishing between susceptible and resistant plant varieties. Armillaria spp. also produce uncommon secondary metabolites, such as sesquiterpene aryl esters, diketopiperzines and sphingolipids. 8,9 Recent work has begun to explore the biomedical application of such metabolites.5

Next generation sequencing has dramatically accelerated research into basidiomycete genomics, for example the genomes of over 50 basidiomycetes are currently available, including the maize and teosinte pathogen Ustilago maydis,2 the ectomycorrhizal fungus, Laccaria bicolor,3 white rots (Phanerochaete chrysosporium on Schizophyllum commune and the multicellular species Coprinopsis cinerea. 12

In initial basidiomycete proteomics studies, 13 2-D PAGE coupled with MALDI-ToF or LC-MS/MS led to identification of 25 extracellular proteins from P. chrysosporium, mainly those involved in carbohydrate polymer degradation, and noted difficulties with peptide identification caused by protein glycosylation. Soon after, these techniques were used to assess the effects of vanillin, an intermediate in lignin degradation on the same fungus and revealed up- and down-regulation of the citric acid and glyoxylate cycles, respectively. 14,15 A proteomic

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approach was also used to investigate the differential secretome composition of P. chrysosporium grown under ligninolytic conditions, where peroxidases were mainly detected. A greater diversity of fungal glycoside hydrolases was identified during culture in biopulped material including cellulases, hemicellulases, putative glucan 1,3- β -glucosidases and α -glycosidases. This work, which was the first differential secretome characterization of P. chrysosporium, highlighted the power of high-throughput proteomic analysis to reveal culture-dependent proteome remodelling due to substrates used for fungal culture. Although no genome sequences are available, 16 proteomic analyses were also carried out on two economically important edible basidiomycetes, Sparassis crispa and Hericium erinaceum, yielding identification of 77 and 121 proteins from both species, respectively, 21 of which were found to be common in both species. L. bicolor is an ectomycorrhizal basidiomycete, which exists in either a saprotrophic phase in soil or an extended mutualistic interaction with tree roots. 17 A recent proteomic study of the L. bicolor secretome in liquid culture identified 224 proteins by a combined electrophoretic fractionation and mass spectrometry-based shotgun approach. 18 This work significantly extended knowledge of the secretome of this saprophytic organism and identified multiple glycoside hydrolases and proteases, hypothesized to be involved in either plant colonisation or basidiomycete defense. Of interest, among proteins of unknown function, an induced small secreted protein, probably a putative effector, has been detected.¹⁸ Functional assignment of unknown proteins produced by basidomycetes is set to become one of the major challenges in fungal biology in the coming years.

However, despite such advances in other basidiomycete species, to date, only 16 distinct *A. mellea* proteins are curated in GenBank including an ATP synthase subunit 6, GAPDH isoform, putative zinc finger/binuclear cluster transcriptional regulator, RNA polymerase (two subunits), a laccase-like multicopper oxidase Lcc2 and a metalloendopeptidase. This is especially problematic since so little is known at the molecular level about this plant pathogen. Here, we reveal the first extensive genome sequence analysis of *A. mellea*, combined with a detailed proteomic characterization. In addition, we propose a novel infection model system that utilizes *Candida albicans* as a target to investigate *A. mellea* virulence.

■ EXPERIMENTAL PROCEDURES

All chemicals were purchased from Sigma-Aldrich Chemical Co. Ltd. (U.K.), unless otherwise stated.

Fungal Culture

A. mellea (Vahl:Fr) Kummer DSM 3731 was purchased from DSMZ (Braunschweig, Germany) and maintained on malt extract agar (50 g/L) (Oxoid CM0059). Liquid cultures (200 mL) were undertaken in 500 mL Erlenmeyer flasks using potato dextrose broth (PDB) (2.4% (w/v)) or minimal media (5 mM ammonium tartarate as nitrogen-source, trace elements according to Schrettl and colleagues and containing 1% (w/v) of one of the following carbon-sources: lignin, xylan, cellulose, rutin, yeast extract). Ethanol (0.15% (v/v)) was added to liquid cultures before inoculation with mycelia and incubation at 25 °C in darkness for 2-6 weeks. Mycelia from liquid cultures were harvested by filtration through miracloth, washed with distilled water, dried thoroughly with absorbent paper, and put immediately into liquid nitrogen and stored at -70 °C. Culture supernatants were stored at -20 °C. Malt extract agar (MEA)

plates were inoculated with A. mellea mycelia and incubated at 25 °C in darkness for 2–6 weeks. Plates were wrapped in parafilm and stored at 4 °C. Coculture experiments involving A. mellea and Candida albicans were carried out on PDB agar. C. albicans (5 μ L; 10⁸ cells/mL) was applied in quadruplicate around central mycelial cultures of A. mellea. After 52 days coculture, C. albicans plugs were collected for subsequent growth (via reculture on PDB agar) and viability assessment (via fluorescein diacetate (FDA)²¹), in the absence of A. mellea. Briefly, FDA (2 μ g/100 μ L) and propidium iodide (PI: 0.6 μ g/30 μ L) were added to 2 \times 10⁸ C. albicans cells for 30 min in the dark. After sequential washing with phosphate buffered saline (PBS) and 20% (w/v) bovine serum albumin (BSA) in PBS, fluorescence was determined using an Olympus Fluoview 1000 confocal microscope. Viability was also assessed by replating serial dilutions of monoculture and coculture-derived C. albicans on MEA. Proteins of A. mellea produced during coculture were extracted by gently grinding mycelia and agar (25 mL) to avoid mycelial lysis with 50 mM KH₂PO₄ pH 7.0. After overnight agitation and filtration, proteins were TCA precipitated, resolubilized in 50 μ L of 10 mM ammonium hydrogen carbonate containing 10% (v/v) acetonitrile and trypsin (65 ng/mL) and digested (37 $^{\circ}\text{C})$ overnight. Resultant peptides were diluted 1/4 in 0.1% (v/v) formic acid (20 μ L) and analyzed directly by LC–MS/MS.

DNA Isolation

A. mellea DNA was extracted using a ZR Fungal/Bacterial DNA Kit (Zymo Research Group) using a modified protocol. Mycelia were ground into a fine powder in liquid N2, and 300 mg were added to tubes containing glass beads (200 mg, 0.4 mm). Lysis solution (750 μ L) was added, specimens were bead-beaten for 2 min at 20 Hz followed by centrifugation at 10000× g for 1 min. Supernatants (400 μ L) were transferred to Zymo-Spin IV Spin Filters and the manufacturer's instructions were then followed. DNA was precipitated by adding 3 M sodium acetate (1/10 vol.)and 3 vol. ice-cold ethanol (100% (v/v)). Following incubation at -20 °C for 3 h, specimens were centrifuged at $13000 \times g$ for 10 min and supernatants removed. Pellets were washed again with ice-cold ethanol (75% (v/v); 180 $\mu L)$ followed by centrifugation at 13000× g for 10 min. After supernatant removal, pellets were air-dried to remove traces of ethanol before resuspending in Milli-Q H_2O (20 μ L). PCR to confirm A. mellea DNA presence was carried out using two different primer pairs; RegF1 and RegR1 primers and AMEL3 and ITS4 primers.^{22,2} products were separated on 1% (w/v) agarose gel.

Genome Sequencing, Assembly and Annotation

A single no-PCR paired-end library was generated from A. mellea genomic DNA according to the method of Kozareva and Turner 24 with an estimated fragment size of 168 \pm 92 bp (inferred by remapping the reads onto the assembly with Maq software²⁵). The library was sequenced on three lanes of an Illumina GAII at 76 bp read length²⁶ producing a total of 6.72 Gb Q20 bases. The reads were assembled with Velvet v0.7.53²⁷ with a kmer of 53 bp. The raw sequence data have been submitted to the European Nucleotide Archive (ENA: http://www.ebi.ac.uk/ ena/) under accession: ERP000894. Gene models were predicted using the ab initio gene finder Augustus.²⁸ Based on a close phylogenetic relationship, L. bicolor was chosen as an appropriate training sequence (Figure 1). A preliminary annotation of predicted gene models was performed using Blast2GO.²⁹ Blast2GO was also used to assign gene ontology (GO)³⁰ terms to all gene models. Signal secretion peptides were predicted using SignalP ver3.0.31 Proteins that may be secreted

Journal of Proteome Research

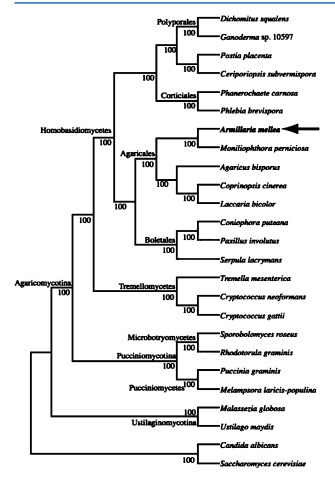


Figure 1. Basidiomycete phylogenetic supertree: Supertree is derived from 5936 individual gene families. Basidiomycotina subphyla, class and orders are shown where applicable. Numbers along branches represent bootstrap support values. Two ascomycete species (*Candida albicans* and *Saccharomyces cerevisiae*) were chosen as outgroups.

via the nonclassical secretion pathway were located using the SecretomeP 2.0 server. *A. mellea* multigene families were located by performing an "all against all" BlastP database search³² and the resulting hits were clustered into families using the mcl algorithm³³ with an inflation value of 2.

The initial genome assembly and predicted protein coding genes are available at http://www.armillariamellea.com.

Phylogenetic Supertree Construction

Our representative genome species tree consisted of 23 Basidiomycete species and 2 Ascomycete outgroups (Supplementary Table 1, Supporting Information). Genome data was obtained from the relevant sequencing centers (Supplementary Table 1, Supporting Information). In total, 8860 homologous gene families (containing 4 or more genes) were identified using a previously described randomized BlastP approach. Gene families were aligned using the multiple sequence alignment software Muscle v3.7 with the default settings. Misaligned or fast evolving regions of alignments were removed with Gblocks. Permutation tail probability (PTP) tests 38,39 were performed on each alignment to ensure that the presence of evolutionary signal was better than random (p < 0.05). We found that 2924 gene families failed the PTP test, these were removed, leaving a data set of 5936 gene families. Optimum models of

protein evolution were selected using Modelgenerator⁴⁰ and these were used to reconstruct maximum likelihood phylogenies in Phyml v3.0.⁴¹ Bootstrap resampling was performed 100 times on each alignment and majority rule consensus (threshold of 50%) trees were reconstructed. Gene families were used to reconstruct supertrees using gene tree parsimony^{42,43} implemented in the software DupTree version 1.48.⁴⁴

Protein Extraction

Mycelial protein extraction was carried out at 4 °C using two different methods. For 2-D PAGE, mycelia were ground in liquid N₂ and 1 g resuspended in 6 mL 10% (w/v) trichloroacetic acid (TCA) (4 °C) and sonicated six times (Bandelin Senoplus HD2200 sonicator, Cycle 6, 10 s, Power 10%). Samples were then centrifuged at 20200× g for 10 min (4 °C), followed by washing in ice cold acetone, twice. Pellets were finally resuspended in IEF buffer (10 mM Tris, 8 M urea, 2 M thiourea, 4% (w/v) CHAPS, 1% (v/v) Triton X-100, 65 mM dithiothreitol (DTT) and 0.8% (w/v) IPG buffer), centrifuged at $14000 \times g$ for 5 min (4 °C) and supernatants used for further analysis. Proteins were quantified using the Bradford assay and 300 μ g were used for IEF using 13 cm strips (pH range 3-10; IPGphorII; GE Healthcare) (Step: 50 V \times 12 h; Step: 250 V \times 0.25 h; Gradient to 5000 V \times 2 h; Step: 5000 V \times 5 h; Gradient to 8000 V \times 2 h; Step: $8000 \text{ V} \times 1 \text{ h}$) followed by SDS-PAGE separation (60 V, 30 min; 120 V, 1.5 h) on PROTEAN Plus Dodeca Cell (Bio-Rad) and Colloidal Coomassie Blue staining. For shotgun analysis, 1 g of mycelial powder was resuspended in 6 mL of 25 mM Tris-HCl, 6 M Guanidine-HCL, 10 mM DTT, pH 8.6 and sonicated five times to generate whole cell lysates. Samples were centrifuged at $20200 \times g$ for 10 min (4 °C). DTT (1 M; 10 μ L/mL lysate) was added to supernatants and incubated at 56 °C for 30 min. Samples were cooled to room temperature and 55 µL of 1 M iodoacetamide per milliliter of lysate was added followed by incubation (in the dark) for 20 min. Samples were then dialyzed twice against cold 100 mM ammonium bicarbonate, (2 × 50 vol.), for at least 3 h with stirring. Precipitates of denatured protein were present after dialysis. One-hundred microliters of denatured protein solution was transferred to an Eppendorf tube and 5 μ L trypsin (Promega V5111) (0.4 μ g/ μ L in 10% (v/v) acetonitrile in 10 mM ammonium bicarbonate) was added. After overnight incubation at 37 °C, the protein precipitates disappeared and protein digests were spin filtered (Agilent Technologies, 0.22 μ m cellulose acetate) prior to LC-MS/MS analysis.

For extracellular protein isolation, a modified version of that described by Fragner et al. 45 was used. Culture supernatants (40 mL) were frozen -20 °C, thawed, centrifuged at $48000 \times g$ for 40min, transferred to clean tubes and frozen in liquid N2. Samples were lyophilized to 4 mL, brought to 10% (w/v) TCA and centrifuged 10000× g for 5 min before washing with ice cold acetone (3 times) with centrifugation steps of 10000× g for 5 min. Protein pellets were solubilized with 100 μ L of 10% (w/v) sodium dodecyl sulfate (SDS), 0.1 M DTT and 5 μ L of SDS-PAGE solubilization buffer and boiled for 10 min prior to SDS-PAGE. For protein extraction of fungal cultures grown on solid phase, MEA plates (25 mL) were briefly ground with a pestle, transferred to containers and brought to 50 mL with 50 mM KH₂PO₄ pH 7.0. 46 Samples were agitated overnight at 4 °C, after which liquid was removed by gravity filtration through Miracloth. Polysaccharides were removed by centrifugation at 48000× g for 30 min, and supernatants were precipitated with 10% (w/v) TCA under agitation overnight at 4 °C. Samples were centrifuged at

 $1700\times g$ for 40 min, pellets were washed (3 times) with 20% (v/v) 50 mM Tris-HCl in ice cold acetone followed by washing in pure ice-cold acetone before air drying and storage -20 °C. Proteins were digested overnight at 37 °C with 50 μ L of trypsin (65 ng/ μ L in 10% (v/v) acetonitrile in 10 mM ammonium bicarbonate). All samples were spin filtered prior to LC–MS/MS analysis.

In-gel digestions (SDS-PAGE bands and 2-D PAGE spots) were carried out according Shevchenko and colleagues with little modification. ⁴⁷ Spots were excised and destained with 10% (v/v) acetonitrile in 100 mM ammonium bicarbonate. Gel pieces were shrunk with 500 μ L of acetonitrile and digested with 50 μ L of trypsin (13 ng/ μ L) at 37 °C overnight before filtration and LC–MS/MS analysis.

LC-MS/MS, Spectral Analysis and Database Searching

An Agilent 6340 ion trap mass spectrometer running ChemStation 0.01.03-SR2 (204) and a ProtiID-Chip150 C₁₈ 150 mm (G4240-62006) chip was used for peptide separation from in-gel protein digests, while a 160 nL 150 mm C18 chip (G4240-62010) was used for tryptic peptides prepared by total protein digestion. Solvent A: 0.1% (v/v) formic acid in water and Solvent B: 0.1% (v/v) formic acid in 90% (v/v) acetonitrile / 10% (v/v) water. For protein identification following SDS- or 2-D PAGE, a flow rate of 0.6 μ L/min with a linear gradient of 5 to 70% Solvent B over 7 min followed by 1 min increase to 100 % Solvent B for 1 min and decrease to 5% Solvent B for 1 min was deployed. For shotgun proteomic studies, a flow rate of 0.4 μ L/ min with non-linear acetonitrile/water + 0.1% (v/v) formic acid gradients, optimised on a sample-specific basis from initial chromatographic analyses, were used. For shotgun proteomic studies, a flow rate of 0.4 μ L/min with a nonlinear acetonitrile/ water +0.1% (v/v) formic acid gradient was optimized on a sample-specific basis from initial chromatographic analyses. Gradients ranging from 20 to 180 min were optimized for identification of the maximum number of proteins per sample: peptide identification from protein spots or bands required 20 min gradients, while shotgun (complex mixtures) gradients were generally up to 180 min duration.

All LC–MS spectra were analyzed using Spectrum Mill software (Rev A.03.03.084 SR4; Agilent Technologies) to identify proteins against the translated A. mellea cDNA database. MS/MS searches were carried out with the following settings: enzymatic trypsin digestion, carbamidomethylation as fixed modification and two variable modifications were set: deamination and oxidation of methionine. The minimum scored peak intensity was set to 70, with precursor mass tolerance ± 2.5 Da and product mass tolerance ± 0.7 . The maximum ambiguous charge was set to 3 with a sequence tag length >3 and minimum detected peaks set to 4. Reversed database scores were calculated.

■ RESULTS AND DISCUSSION

Genome Sequencing and Analysis

PCR confirmed the presence of *A. mellea* DNA (data not shown). The genome of *A. mellea* (Vahl:Fr) Kummer DSM 3731 was sequenced using an Illumina Genome Analyzer II. A total of 6.72 Gb of sequence data was obtained from 3 single read lanes. On the basis of nuclear DNA measurements by RFLP analysis,⁴⁷ this represents 57.25-fold coverage of the *A. mellea* genome. Assembly of the Illumina reads with the Velvet assembler resulted in 58.35 Mb of sequence data in 4377 scaffolds with an N50 size of 36.7 kb (Table 1). The overall GC content of the genome is 49.1%, while in the coding regions it is 53.2%.

Table 1. Genome Assembly Statistics for A. mellea (Vahl:Fr) Kummer DSM 3731

	scaffolds	contigs
N50	36679 bp	5485 bp
Largest	639 705 bp	154 911 bp
Count	4377	15215
Total Length	58 358 340 bp	71 738 977 bp

A total of 14473 putative A. mellea open reading frames (ORFs) were predicted (Table 2) which was subsequently used

Table 2. Gene Statistics for *A. mellea* (Vahl:Fr) Kummer DSM 3731 in Comparison to Closely-related Basidiomycete Species

	A. mellea	L. bicolor	Coprinopsis cinerea
Genome Assembly (Mb)	58.3	64.9	37.5
Number of protein-coding genes	14473	20614	13544
Coding sequence <300 bp	957	2191	838
Average gene length (bp)	1575	1533	1679
Average coding sequence length (bp)	1227.75	1134	1352
Average exon length	217.52	210.1	251
Average intron length	73.62	92.7	75
Average number of introns per gene	4.72	4.44	4.66

as the reference database for proteomic analyses. The average gene length was found to be 1575 bp, similar to what is observed in close relatives of *A. mellea*, such as *L. bicolor* and *Coprinopsis cinerea* (Figure 1 and Table 2). The average exon/intron lengths were also comparable among these species, as were the average number of introns per gene (4.72, 4.44 and 4.66, respectively). Blast2GO successfully assigned a GO term to 9514 (65.7%) of the predicted genes (data not shown). The SignalP webserver identified 1903 proteins with secretion signal peptides, while Phobius predicted 4035 proteins (27.8%) with membrane regions, 2746 (19%) with transmembrane regions and 246 with transmembrane regions plus signal peptide regions. Protein GRAVY⁴⁸ identified 448 proteins with a GRAVY Score >0.5.

Homology searches followed by Markovian clustering inferred that 9939 (68.7%) of the A. mellea ORFs belonged to multigene families. Using the same approach we calculated that 68.5% of L. bicolor genes belong to multigene families. Approximately 18.8% (2721) of the predicted A. mellea genes have no significant homologues in the NCBI database, a figure consistent with other newly sequenced basidiomycete genomes. We also successfully located 240 tRNA genes which compares favorably to L. bicolor which has 297 tRNA genes.

Protein Identification

Given the paucity of A. mellea proteins curated and present in GenBank (n=16), it was necessary to interrogate the translated version of the A. mellea cDNA database with peptide sequence data for complete and accurate de novo protein identification. As a consequence, a total of 921 proteins were identified in both mycelia and culture supernatants from A. mellea grown under a range of different culture conditions. These identifications were based on the detection of 3641 unique peptides, 2839 of which came from mycelia and 802 from the supernatant, respectively, with 197 common to both (Supplementary Tables 2 and 3, Supporting Information).

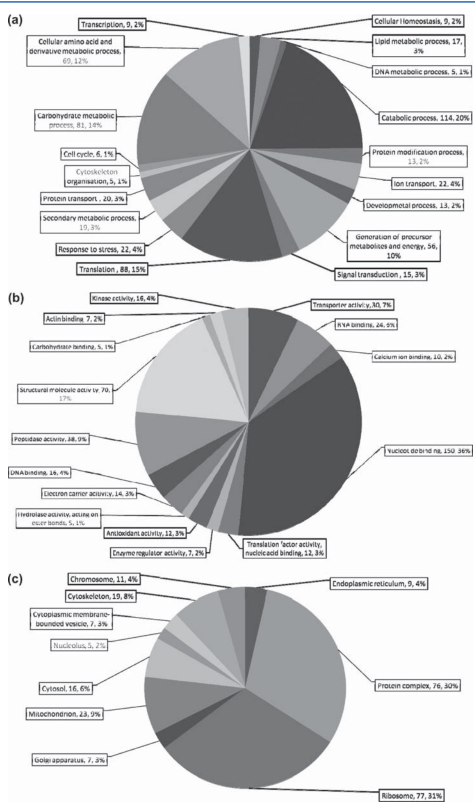


Figure 2. Blast2Go annotation of mycelial protein classification: (a) biological process; (b) molecular function; (c) cellular component. The total number of proteins in each category, as well as the overall %, are given for each subdivision (No., %).

Journal of Proteome Research

Table 3. Unique, Previously Hypothetical and Predicted Proteins from A. mellea Mycelia and Secretome

accession no."	InterPro domains ^b	tM_{r}^{c}	$_{p}I$ dı	coverage (%)	unique peptides	SM score	GRAVY score ^f	TM^g	SigP/ SecP ^h	Method'	Source
Am18468	Civ. bladed betannoneller Talk like	62204.0	4 10	36	01	15 691	-0.272		GiaD	Ē	NS
Am8114	None detected	20938.8	7.77	31	4	77.92	-0.278		SecP	2D	M. SN
Am8753	Macrophage migration inhibitory factor: Tautomerase	20131.3	5.1	15	7	32.87	-0.053		SecP	s	` M
Am14766	Glucose-methanol-choline oxidoreductase, N-terminal; Protein of unknown function DUF427	83438.2	6.5	-	1	16.04	-0.295			1D	SN
Am16713	Monooxygenase, FAD-binding; Aromatic-ring hydroxylase-like; Thioredoxin-like fold	60524.2	6.17	33	14	247	-0.184		SigP	2D	M
Am11784	RNA recognition motif domain; Aldo/keto reductase; Nucleotide-binding, alpha-beta plait; RNA-binding motif	25961.1	6.36	4	6	151.55	-0.458			2D	M
Am14065	Globin, structural domain	24081.8	5.9	17	8	61.66	-0.128		SecP	S	M, SN
Am6826	None detected	6811.8	9.52	24	1	19.01	-0.138		SecP	S	M
Am11329	None detected	38549.9	5.64	23	4	74.13	0.032		SecP	1D	M, SN
Am15909	6-phosphogluconate dehydrogenase, NADP-binding; 6-phosphogluconate dehydrogenase, C-terminal- like; Dehydrogenase, multihelical; Hydroxy monocarboxylic acid anion dehydrogenase, HIBADH- type; NAD(P)-binding domain	39265	6.83	31	^	138.16	-0.089	1	SecP	2D	M
Am19857	PLC-like phosphodiesterase, TIM beta/alpha-barrel domain	33320.7	4.36	4	1	20.33	0.081		SigP	1D	SN
Am11737	None detected	31480.1	9.19	15	4	65.72	-0.316			S	M
Am10761	None detected	104910	5.02	4	2	36.1	-0.605			S	M
Am6759	None detected	9075.6	68.01	19	1	17.65	-0.064		SecP	S	M
Am687	None detected	31486.3	4.74	42	∞	153.64	-0.773		SecP	2D	M
Am12929	Phosphatidylethanolamine-binding protein PEBP	25186.2	6.02	13	7	37.32	0.35		SigP	1D	SN
Am15856	Thioredoxin-like fold	29393.2	5.89	8	1	11.6	-0.132		SigP	S	M
Am10869	None detected	29854.2	4.44	6	-	17.66	0.217	1	SigP	1D	SN
Am7778	Lambda repressor-like, DNA-binding domain	7398.4	6.7	22	1	22.18	-0.027		SecP	S	M
Am13799	HAD-superfamily hydrolase, subfamily IA, variant 3; Phosphoglycolate phosphatase, domain 2; HAD-like domain	36474.5	5.58	^		14.84	-0.364		SecP	s	M
Am18955	Beta-lactamase-like	37813.3	5.59	3	1	12.65	-0.164		SecP	2D	M
Am11484	None detected	13748.5	4.85	13	1	16.43	-0.687		SecP	1D	NS
Am19019	Carboxylesterase, type B; Carboxylesterase type B, conserved site	61132.2	4.65	10	4	63.48	-0.097		SigP	1D	M, SN
Am17323	None detected	6874.9	6.18	19	1	19.82	0.077		SecP	s	M
Am11327	None detected	33136.6	4.5	11	3	51.17	-0.01		SecP	1D	SN
Am18523	Vacuolar protein sorting-associated protein 62	40908	4.45	3	1	17.86	-0.258		SigP	1D	NS
Am16822	None detected	310055	4.05	<u>^</u>	-	13.54	-0.318		SecP	S	M
Am16119	Crotonase, core; Armadillo-like helical	105164	5.71	7	1	21.12	-0.037	7		S	M
Am15204	None detected	20487.3	4.95	13	1	23.43	-0.319		SecP	1D	M, SN
Am9185	None detected	61991	2.67	7	1	17.45	-0.08		SecP	2D	M
Am10375	Six-hairpin glycosidase-like; Protein of unknown function DUF1680	67815.8	5.1	7	1	11.46	-0.156		SecP	1D	NS
Am13156	None detected	72243.6	4.96	25	10	177.07	-0.065		SigP	1D	NS
Am6889	Short-chain dehydrogenase/reductase SDR; Glucose/ribitol dehydrogenase; NAD(P)-binding domain; Short-chain dehydrogenase/reductase, conserved site	26913	6.52	51	6	163.3	0.145		SecP	S	M
Am10376	FAD dependent oxidoreductase; Lanthionine synthetase C-like	158913	5.63	2	2	24.79	-0.143		SecP	1D	NS
Am9103	None detected	9374.6	9.15	43	2	39.82	-0.38		SecP	2D	M
Am6752	NADH:flavin oxidoreductase/NADH oxidase, N-terminal; Aldolase-type TIM barrel	39576	6.03	4	1	22.1	-0.322			1D	SN
Am1857	PLC-like phosphodiesterase, TIM beta/alpha-barrel domain	29991.3	4.21	\$	1	22.58	-0.069	1	SecP	1D	SN
Am16247	NADH:flavin oxidoreductase/NADH oxidase, N-terminal; Aldolase-type TIM barrel	58857	5.18	7	-	17.2	-0.358			1D	SN

Table 3. continued

Table 3. continued

accession no."	${\rm InterPro\ domains}^b$	tM,°	^{p}I dt	coverage (%)	unique peptides	SM score ^e	GRAVY score ^f	$^{2}_{ m CM}$	SigP/ SecP ^h	Method ⁱ	Source
Am12286	Ubiquitin; Ubiquitin conserved site; Ubiquitin supergroup; Ubiquitin subgroup	57329.3	4.22	2	-	13.06	-0.132		SigP	1D	SN
Am12502	None detected	60386.3	6.2	26	10	188.36	-0.174	-		2D	M, SN
Am13522	None detected	12745.6	7.54	10	П	16.59	-0.495			s	M
Am13594	None detected	73800.3	9.33	3	П	20.46	-0.658			S	M
Am14055	Alpha/beta hydrolase fold-1; Peptidase S33 tripeptidyl aminopeptidase-like, C-terminal	55061	4.49	П	П	14.07	-0.051		SecP	1D	SN
Am14569	None detected	30523.7	98.9	8	2	25.97	-0.218			2D	M
Am14848	Aldo/keto reductase; NADP-dependent oxidoreductase domain	49709.4	5.72	s	П	20.05	0.004			1D	M
Am15133	None detected	14436.6	4.91	31	3	52.74	0.168		SigP	1D	M, SN
Am17215	None detected	95309.3	6.93	2	П	16.14	0.153	11		S	M
Am17941	None detected	115634	6.11	-	-	12.04	-0.085	2	SecP	2D	M
Am18255	None detected	90035.9	4.75	2	П	14.5	-0.599			S	M
Am18727	None detected	10573	8.96	18	-	12.61	-0.73		SecP	S	M
Am19502	None detected	18009.9	4.37	34	33	54.75	0.623		SigP	1D	SN
Am19696	Ribonuclease/ribotoxin	15150	4.47	13	-	16.89	-0.209		SecP	1D	SN
Am19883	None detected	54306.3	2.67	s	П	20.27	-0.729			S	M
Am20219	None detected	16248.1	9.58	7	П	14.89	-0.966	П		S	M
Am20301	Histone chaperone domain CHZ	49025.6	5.85	\$	2	22.48	-0.29		SecP	S	M, SN
Am4394	None detected	26796.8	7.7	4	-	18.36	-0.193			S	M, SN
Am5361	None detected	8625.1	8.09	41	2	36.28	-0.36		SecP	2D	M
Am6074	None detected	27554.6	7.72	9	П	17.89	-0.087			S	M
Am9062	None detected	16419.1	6.9	9	-	16.06	-0.066			1D	SN
Am9780	None detected	17457.1	4.13	22	3	50.38	0.52		SigP	1D	SN
Am9972	None detected	73660.6	4.59	7	-	16.89	-0.112		SecP	1D	SN
Am17500	None detected	27349.2	9.13	8	2	26.53	-0.607		SecP	1D	SN
Am8226	High mobility group, HMG-I/HMG-Y; AT hook, DNA-binding motif; AT hook-like	38794	7.62	18	4	62.82	-0.226			S	M
Am1474	None detected	5325.2	6.28	29	1	12.29	0.2		SecP	s	M

Spectrum Mill protein score, ^fGRAVY score, grand average of hydropathy (Negative score indicates hydrophilicity while Positive score indicates hydrophobicity). ⁸TM, number of transmembrane regions. ^hSigp, classical secretion signal peptide; SecP, nonclassical secretion signal. [†]Method: 1D SDS-PAGE prior to LC-MS/MS analysis, 2-D proteins separated in two dimensions prior to LC-MS/MS analysis. [†]Source: M, mycelia; SN, culture supernatant/secretome. ^aAccession number from A. mellea cDNA database. ^bInterPro domains located within protein following Blast2GO analysis. ^ctM₀, theoretical molecular mass. ^atpJ, theoretical isoelectric point. ^eSM score,

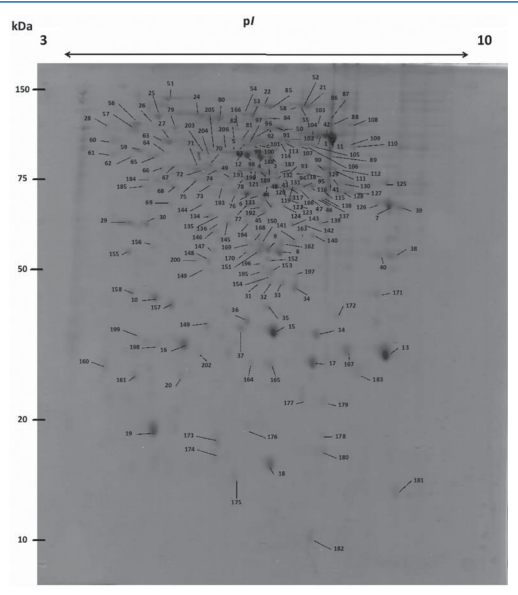


Figure 3. 2-D PAGE reference map of the A. mellea mycelial proteome.

Mycelial Proteins

Mycelial proteins (n = 739) were identified by using the A. mellea database obtained from the translation of the protein-coding genes. Six-hundred twenty-nine proteins were uniquely identified in mycelia (Supplementary Table 2, Supporting Information), while others were also identified in the A. mellea secretome (Supplementary Tables 3 and 4, Supporting Information). Of the mycelia-specific proteins, 509 were uniquely identified by a shotgun approach, with 98 and 9 proteins identified following prior 2-D PAGE and SDS-PAGE fractionation, respectively (Supplementary Figure 1, Supporting Information). Supplementary Table 2, Supporting Information, shows the actual peptide sequence with corresponding Spectrum Mill scores where values >11 were deemed significant according to manufacturer's guidelines. Sequence coverage ranged from 1 to 5%, 1 to 81% and 1 to 72%, respectively, using SDS-PAGE, 2-D PAGE fractionation and shotgun analysis.

Functional analysis undertaken with BLAST2GO and database searches classified proteins according to the biological process (BP) in which they are involved, the molecular function (MF) and the cellular component (CC). GO analysis by BP of mycelial proteins generated 18 categories. The largest categories were catabolic processes (n = 114) followed by translation (n =88), carbohydrate metabolic processes (n = 81), precursor metabolites and energy (n = 56), response to stress (n = 22) and secondary metabolic processes (n = 19) (Figure 2a). MF of mycelial proteins revealed nucleotide binding (n = 150) and structural molecule activity (n = 70) as the largest categories, accounting for 53% of mycelial proteins identified. GO identified 38 proteins in MF as having peptidase activity (Figure 2b). Mycelial proteins CC mainly comprised ribosomal proteins (n =78) and protein complexes (n = 77). There were also 23 proteins classified as mitochondrial (Figure 2c) and 14 proteins that have not been previously annotated and appear to be unique to A. mellea. One-hundred two proteins were detected that were previously reported as hypothetical or predicted (Table 3). As expression of these proteins is now demonstrated, they can be reclassified as Unknown Function Proteins. A search of these

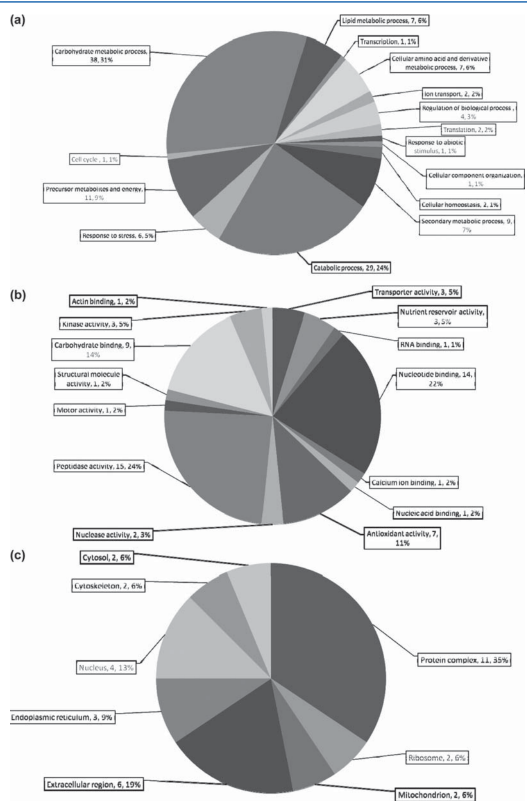


Figure 4. Blast2Go annotation of proteins identified in the secretome. (a) Biological process; (b) molecular function; (c) cellular component. The total number of proteins in each category, as well as the overall %, are given for each subdivision (No., %).

proteins against the InterPro database⁴⁹ suggested putative functions. Overall, 114 known domains were assigned to 55 proteins (Table 3). Eight proteins (Am15909, Am6889,

Am18190, Am14843, Am14848, Am14568, Am13925 and Am17640) were found to contain a $NAD(P)^+$ -binding domain, however no other domain was over-represented in our analysis.

Table 4. Glycoside Hydrolase (GH) Expression Identified from A. mellea Mycelia and Secretome

accession no. ^a	BLAST2GO annotation b	tM _r ^c	$\operatorname{tp} I^d$	coverage (%)	unique peptides	SM score ^e	GRAVY score ^f	TM^g	SigP/ SecP ^h	method ⁱ	source ^j
Am13982	GH	58105.5	5.05	6	3	53.65	-0.151		SigP	1D	SN
Am18619	GH	51878.3	4.63	6	1	17.68	-0.039		SigP	1D	SN
Am19316	GH	49972.4	4.76	2	1	16.09	-0.259		SigP	1D	SN
Am9430	GH Family 1 Protein	16311.1	4.08	6	1	14.64	-0.401		SecP	1D	SN
Am9431	GH Family 1 Protein	26301.4	5.77	7	1	24.73	0.219		SigP	1D, S	M, SN
Am9862	GH Family 1 Protein	65185	5.04	1	2	21.05	-0.156		SecP	1D	SN
Am10289	GH Family 115 Protein	114008	4.55	2	2	40.53	-0.094		SigP	1D	SN
Am16201	GH Family 13 Protein	46456.7	4.55	9	2	38.04	-0.136		SecP	1D	SN
Am17847	GH Family 13 Protein	56284.6	4.94	5	3	58	-0.24		SigP	1D	SN
Am14148	GH Family 15 Protein	87387	4.97	2	1	17.02	0.015		SigP	1D	SN
Am14157	GH Family 15 Protein	61191.1	4.56	11	4	68.88	0.016		SigP	1D	SN
Am13749	GH Family 16 Protein	35677.5	4.32	4	1	19.18	-0.063		SigP	1D	SN
Am17932	GH Family 16 Protein	48410.8	4.69	2	1	17.58	-0.017	1	SigP	2D	M
Am18038	GH Family 16 Protein	39327.9	6.39	17	5	80.3	-0.114		SigP	1D	SN
Am19909	GH Family 16 Protein	147978	9.69	1	2	35.77	-0.157		SecP	S	M
Am11859	GH Family 17 Protein	39604.6	4.44	7	1	22.22	-0.125		SigP	1D	SN
Am18174	GH Family 18 Protein	28184	4.53	15	2	37.1	0.318		SigP	1D	SN
Am18992	GH Family 18 Protein	54832.1	4.96	4	2	28.75	-0.253		SigP	1D	SN
Am6678	GH Family 2 Protein	98191.6	5.67	2	1	16.69	-0.455		0-8-	S	M
Am12096	GH Family 20 Protein	61819.8	4.98	3	1	17.55	-0.067		SigP	1D	SN
Am9564	GH Family 20 Protein	59239.4	4.75	7	3	42.51	-0.083		SigP	1D	SN
Am14135	GH Family 27 Protein	47065	4.7	18	5	91.48	-0.182		0-8-	1D, S	M, SN
Am17468	GH Family 28 Protein	8969.1	9.4	20	1	21.85	0.087		SecP	1D, 5	SN
Am17470	GH Family 28 Protein	44505.2	4.76	7	2	37.48	0.001		SigP	1D	SN
Am12215	GH Family 3 Protein	148863	5.59	2	2	38.48	-0.211		SecP	1D, S	M, SN
Am15538	GH Family 3 Protein	81458.6	4.62	2	1	23.05	-0.021		SigP	1D, S	M, SN
Am4117	GH Family 3 Protein	66127.4	4.54	6	4	59.81	0.014	1	SecP	1D, 5	SN
Am4142	GH Family 3 Protein	110105	4.95	3	3	49.7	-0.151	-	SigP	1D	SN
Am9629	GH Family 3 Protein	94245.5	5.9	2	1	16.83	-0.238		0-8-	S	M
Am19156	GH Family 31 Protein	105123	5.18	1	1	14.1	-0.137	1	SecP	1D	SN
Am5759	GH Family 31 Protein	191108	6.39	2	4	60.82	-0.498	1	5001	1D, S	SN
Am14693	GH Family 37 Protein	71119.8	4.35	6	3	44.78	-0.258	-	SigP	1D, 5	SN
Am14694	GH Family 37 Protein	15385	4.14	7	1	16.41	-0.11		SecP	1D	SN
Am14901	GH Family 38 Protein	116608	6.16	3	1	13.06	-0.311		5001	S	M
Am13752	GH Family 47 Protein	59389.7	4.97	32	15	282.75	-0.154		SigP	1D	SN
Am10113	GH Family 5 Protein	88731.2	4.81	2	1	19.29	-0.446	1	SecP	1D	SN
Am12892	GH Family 5 Protein	64378.6	6.12	12	4	68.82	-0.515	-	5001	1D, 2D, S	M
Am14455	GH Family 5 Protein	77060.8	4.83	1	1	19.42	-0.032	6		1D, 2D, 6	SN
Am15965	GH Family 5 Protein	46113.4	4.8	25	6	108.18	-0.212		SigP	1D	SN
Am17147	GH Family 5 Protein	57264.9	4.97	2	1	13.01	-0.272		SecP	2D	M
Am8313	GH Family 5 Protein	75693.8	6.8	1	1	19.28	0.322	9	SecP	1D	SN
Am19121	GH Family 55 Protein	79789	6.09	2	1	27.82	-0.147		SecP	1D	SN
Am19122	GH Family 55 Protein	62305.6	4.43	2	1	22.56	-0.077		SecP	1D	SN
Am15544	GH Family 61 Protein	22656.6	4.7	9	1	11.98	0.212		SigP	1D	SN
Am15111	GH Family 72 Protein	58531.8	4.34	11	5	84.6	-0.01		SigP	1D, S	M, SN
Am13430	GH Family 74 Protein	85505.4	4.88	2	1	11.49	-0.01		SigP	1D, 3 1D	SN
Am17906	GH Family 74 Protein	71627.9	4.32	1	1	16.07	-0.134		SigP	1D 1D	SN
Am18634	GH Family 79 Protein	38585.2	4.86	19	5	84.99	-0.146 -0.019		SecP	1D 1D	SN
Am18034 Am9625	GH Family 79 Protein GH Family 79 Protein	55806	4.49	5	1	14.53	-0.019 -0.064		SigP	1D 1D	SN
Am17712	GH Family 92 Protein	87085.9	4.49	15	7	112.69	-0.203		SigP	1D 1D	SN
Am9020	GH Family 92 Protein	86824.7	4.98	3	1	15.69	-0.203 -0.425		SigP	1D, 2D	M, SN
Am7125	GH Family 92 Protein GH Family 95 Protein				8		-0.423 -0.138		Sigr	1D, 2D 1D	SN
AIII/123	Gri Failmy 93 Protein	89157.1	4.8	13	٥	125.65	-0.138			110	SIN

[&]quot;Accession number from A. mellea cDNA database. Blast annotation following Blast2GO analysis of proteins identified from cDNA database. thm, theoretical molecular mass. thp, theoretical isoelectric point. SM score, Spectrum Mill protein score. GRAVY score, grand average of hydropathy (Negative score indicates hydrophilicity while Positive score indicates hydrophobicity). TM, number of transmembrane regions. SigP, classical secretion signal peptide; SecP, nonclassical secretion signal. Method: 1D SDS-PAGE prior to LC—MS/MS analysis, 2-D proteins separated in two dimensions prior to LC—MS/MS analysis. Source, M: mycelia; SN: culture supernatant/secretome.

Moreover, no GO function was identifiable for 73 proteins, while 26 showed no BLAST description. Among these are proteins

with conserved domains which indicate a putative process, function or cellular location such as Am12246 containing a

START domain typical of proteins which transport ubiquinone during respiration or Am14390 containing a 60 residue conserved BSD domain of unknown function usually located in BTF2-like transcription factors, Synapse-associated proteins and DOS2-like.⁵⁰

Regarding 2-D PAGE analysis, 206 protein spots were excised, trypsin-digested and subjected to LC-MS/MS. Two hundred twenty five proteins were identified in total. A reference 2-D PAGE map is shown in Figure 3 and information (pI and $M_{\rm r}$) on proteins identified can be found in Supplementary Table 2 (as "2D" under column "Method", Supporting Information). Possible protein isoforms were identified in different spots (e.g., spots 89, 92, 107 (Am10779) and 47, 122, 124, 128, 130 (Am16991). One protein, Am17600, was identified in 17 spots (Nos. 1, 3, 11, 58, 88, 89, 90, 97, 103, 104, 107, 117, 118, 119, 131, 132, 133) probably due to a protease activity or degrading events. BLAST analysis classified this protein as a peroxisomal catalase.

The A. mellea Secretome

A total of 293 proteins were identified in culture supernatants derived from A. mellea, 183 of which were uniquely detected in the secretome, while 111 were also identified in mycelia (Supplementary Table 3, Supporting Information). To overcome the considerable difficulty of precipitating low abundance proteins from liquid culture, A. mellea was grown on agar, which was subsequently gently macerated and the liquid phase subject to TCA precipitation followed by SDS-PAGE fractionation. A nonredundant protein search of the translated A. mellea gene set, identified 226 proteins. Analysis of LC-MS/MS data identified 157 unique proteins, while 37 were uniquely identified by a shotgun strategy without SDS-PAGE fractionation. Sequence coverage ranged from 1 to 55% and 1 to 46% following SDS-PAGE and the shotgun approach, respectively, in the case of supernatant protein analyses (Supplementary Table 3, Supporting Information). Eight proteins were identified in culture supernatants which were A. mellea-specific and included: Am1874, Am2320, Am5632, Am9253, Am11032, Am11414, Am13545 and Am13611. None returned BLAST hits and Am13545 had no InterProScan (IPS) match. IPS revealed lowcomplexity segments in Am11414, Am2320, Am5632 and Am9253. Analysis revealed two SigP hits (Am11032 and Am13611), three SecP signals (Am11414, Am13545 and Am1874) and a TMHMM search predicted Am11032 to be a hydrophobic membrane protein.

Secretome proteins were classified into 15 distinct biological processes. The proportions of proteins involved in catabolic processes (n = 29) and carbohydrate metabolism (n = 38) was greater (55%) than in mycelia (33%) (Figure 4a). Seven supernatant proteins were GO annotated to be involved in lipid metabolism. This classification was absent for mycelial proteins (Figure 2a). GO analysis related to the molecular function of supernatant proteins showed peptidase activity as the largest functional category (n = 15) followed by nucleotide binding proteins (n = 14). The class including carbohydrate binding proteins represented 14% in the secretome, yet in mycelia it accounted for only 1%. Proteins classified as having antioxidant activity represent 11% of the secretome, while only 3% of mycelial proteins were similarly classified (Figures 2b, 4b). Concerning GO annotation of CC, the largest categories in the secretome comprised protein complexes (n = 11), and extracellular proteins (Figure 4c). Several proteins of intracellular origin were also detected.

SignalP analysis predicted signal peptide presence for 12% (n =90) and 50.2% (n = 147) of mycelial and secretome proteins, respectively. Furthermore SecretomeP indicated that 99 secreted proteins (34%) contained a nonclassical secretion signal meaning that less than 20% (55) of those proteins do not contain a secretion signal. GRAVY and Phobius evaluation of hydrophobicity and transmembrane domain presence indicated that 12.7% (n = 93) of mycelial proteins were hydrophobic in nature, 10.5% (n = 77) contained transmembrane regions, and 1.4% (n = 77) 10) contained both a transmembrane region and signal peptide. Six proteins (Am9118, 13744, 6597, 5593, 17215 and 19909) contained greater than 10 transmembrane regions, while 71 proteins contained between 1 and 8 transmembrane domains. With respect to supernatant proteins, 26.3% (n = 77) were hydrophobic in nature, 29 contained predicted transmembrane domains, and 9 of these were also predicted to possess signal sequences.

Glycoside Hydrolases

Although basidiomycetes have been considered as sources of glycoside hydrolases (GH), with potential applications in recalcitrant carbohydrate digestion, no data whatsoever exists on GH complement in A. mellea. Here, we reveal the presence of 226 GH genes in A. mellea (Supplementary Table 5, Supporting Information), and further demonstrate the expression of 52 of these at the protein level, in culture supernatants, under a range of culture conditions (Table 4). Specifically, GH genes belonging to 43 families are present (Supplementary Table 5, Supporting Information). The number of A. mellea GH genes is greater than that identified in either P. chrysosporium (n = 177) or Ceriporiopsis subvermispora (n = 171), both of which are white rot fungi, 51 a finding that may be related to the pathogenic nature of A. mellea.

The A. mellea genome encodes 16 xyloglucan:xyloglucosyltransferases (GH16 family, Supplementary Table 5, Supporting Information). Xyloglucan, a hemicellulose, is a major component of plant cell walls whereby its cellulose binding capacity provides structural support for the cell wall.⁵² GH 16 family enzymes cleave the 1-4 glycosidic backbone of xyloglucan and transfer the segment either to another xyloglucan or to a xyloglucan oligosaccharide (XGO), thus restructuring and modifying the primary cell wall.⁵³ XGOs have been shown to activate plant cellulases and promote plant growth, but plant XGOs do not activate fungal cellulase. 54 The GH16 family proteins also have amylase and cellulase activity. Two enzymes (Am17743 and Am19909) in this class were also implicated in GTPase activity, which may indicate a signaling role. 55 Other GH families with increased or decreased representation in the A. mellea genome compared to lignolytic and cellulolytic fungi are reported in Table 5. Contrary to expectation that lignolytic fungi would have similar frequencies of particular GH family proteins, A. mellea appears to contain more GH28, GH61, and GH76 family members than the other white or brown rot fungi (Table 5). For example, 17 GH28 genes (endo/exo-(rhamno)galacturonases, essential for pectin degradation) are present in the A. mellea genome (Supplementary Table 5, Supporting Information) in comparison to the 6 identified in *L. bicolor* and the 4 present in *P.* chrysosporium. Yet, A. mellea possesses similar GH family occurrence to ascomycete species. Moreover, GH109, an α -Nacetylgalactosaminidase present in A. mellea and the mycorrhizial L. bicolor (Supplementary Table 5, Supporting Information), has previously been also identified in prokaryotes and its crystal structure defined.56

Table 5. Comparison of *A. mellea* with Lignolytic and Cellulolytic Basidiomycetes and Ascomycetes with Respect to Four Glycoside Hydrolase Families Present in the Respective Genomes

		GH :	family	
fungus	28	35	47	76
Armillaria mellea	20	8	12	8
Ceriporiopsis subvermispora	6	1	4	
Phanerochaete chrysosporium	4	3	6	
Schizophyllum commune H4–8	3	4	6	3
Coprinopsis cinerea Okayama 7 (#130)	3		8	
Postia placenta Mad-698-R	7	1	5	
Podospora anserina S mat+		1	9	9
Aspergillus niger CBS 513.88	21	5	5	11
Aspergillus nidulans FGSC A4	9	4	7	7

Chitinases, GH18 (endo- β -N-acetylglucosaminidases) (n=11), are also encoded within the genome, and the expression of two of these was detectable in A. mellea culture supernatants (Am17468 and Am17470) (Supplementary Table 3, Supporting Information). In addition to glycoside hydrolases, 72 glycosyltransferase and 19 polysaccharide lyases have been identified (Supplementary Table 5, Supporting Information).

Carbohydrate-binding Modules (CBM)

Besides glycoside hydrolases, other CBM enzymes were discovered in the A. mellea genome. CBM proteins exhibit three functions encompassing (i) carbohydrate binding and concentration of enzymes on the substrate, (ii) binding to the polysaccharides that are the substrate targets of the catalytic moiety of the enzyme and (iii) substrate degradation.⁵⁷ Three CBM proteins were identified (Am18408, AM9123 and Am11736) by LC-MS/MS analysis, belonging to families 1, 12 and 13, respectively (Table 4). CBM1 is a small module (40 aa) that binds cellulose. CBM12 previously only described in bacteria, comprises 40-60 residues, binds chitin, which is unusual among CBM proteins which have broad polysaccharide specificity. The C-terminus of the chitin binding domain (ChBD) binds very specifically to insoluble chitin but not to chitin oligosaccharides or glucose.⁵⁷ Interestingly, chitinases have been investigated as biotechnological biocontrol for plant pathogenic fungi. Se CBM13 is a 150 residue module that has been identified in galactose-binding plant lectins.⁵⁷ Predominantly found in bacteria, they have also been identified in eukaryotes, including some fungi; CBM13 modules occur in glycoside hydrolases and glycosyltransferases and also bind xylan. **Carbohydrate Esterases**

Compared to some basidiomycete and ascomycete species, *A. mellea* has increased representation of carbohydrate esterase (CE) families (Supplementary Table 5, Supporting Information), specifically CE4, CE8, CE9 and CE14 comprised 94 genes. CE1 is an acetyl xylan esterase (AXE) which depolymerises xylan microfibrils and solubilizes lignocellulosic material and releases fermentable sugars. ⁵⁹ A CE1 enzyme (Am19807) was identified at the protein level in the growth condition we tested (Supplementary Table 2, Supporting Information). The CE4 family exhibits acetyl xylan, chitin, chitooligosaccharide and peptidoglycan deacetylase activity. Four CE4 enzymes (Am18938, Am3987, Am3988, and Am6064) have been identified in *A. mellea* culture in both mycelia and secretome (Supplementary Tables 2 and 3, Supporting Information). These enzymes act on N-acetylglucosamine (GlcNAc) which is present

in bacterial as well as fungal cell walls; have very broad substrate specificity and catalyze the hydrolysis of acetyl groups from xylan and carbon-nitrogen bonds in linear amides. They act in concert with chitinases to deacetylate both chitin and chitosan. 60 Chitin deacetylases (CDA) are secreted during hyphal penetration of plants by Colletotrichum lindemuthianum. Tsigos and Bouriotis hypothesized that chitin of invading pathogens triggers recognition by plant defenses inducing plant chitinase activity which gives rise to formation of lignin and callus. 61 Deacetylases modify plant chitinases thereby neutralizing plant defenses, and modify chitin to chitosan which plant deacetylases are unable to modify, thus enabling the fungus to invade the plant. Pectin methylesterase (PME) are members of the CE8 family and catalyze the demethyloxidation of pectin, a major plant cell wall polysaccharide. PME disrupts pectin structure and makes it liable to subsequent enzymatic depolymerisation. PMEs are important in vegetative and reproductive plant development, but pathogenic fungi can also degrade pectin in the cell wall during plant invasion. 62 One CE8 (Am12061) was identified in A. mellea culture supernatants. CE9 (N-acetylglucosamine 6-phosphate deacetylase) is another gene family detected in A. mellea which is involved in carbohydrate, specifically N-acetyglucosamine, catabolism. Surprisingly, a GlcNAc-6-phosphate deacetylase (DAC1) has been linked to *C. albicans* virulence. He CE14 deacetylase family has a conserved LmbE domain which catalyzes the N-deacetylation of appropriate substrates.⁶⁵ Tanaka et al. identified a deacetylase from Thermococcus kodakarensis KOD1 which degrades diacetylchitobiose (GlcNAc2) to GlcN-GlcNAc which in a second round of deacetylation is deacetylated to GlcN.66 Another function of these enzymes in eukaryotes is GPI anchor modification.⁶⁷

Laccases

The *A. mellea* genome encodes 6 laccases (Am380, Am12382, Am16732, Am16804, Am17018 and Am19478) (Figure 5), all of which contain the canonical L1-L4 signature sequences, characteristic of this multicopper oxidase (MCO) family of enzymes.⁶⁸ Overall, 15 laccase-like enzymes are encoded within the *A. mellea* genome. Laccase Am380 expression was detected in mycelial and supernatant by LC–MS/MS analysis (Supplementary Tables 2 and 3, Supporting Information), while expression of any other laccase was undetectable under the condition tested. Expression of two laccase-like enzymes (Lcc5: Am2859 and Am4666) was also evident in the *A. mellea* secretome (Supplementary Table 3, Supporting Information). Notably, the genome of the white rot fungus *P. chrysosporium* does not encode any laccase,⁵¹ while those of *C. subvermispora*⁵¹ and *L. bicolor*⁶⁹ encode 7 and 9 laccase gene models, respectively.

Peroxidases

Basidiomycete peroxidases are a class of oxidoreductase involved in lignin biodegradation 70,71 In white rot fungi, ligninolytic haem peroxidases have been shown to cleave the main linkage types in lignin due to their high redox-potential and specialized catalytic mechanisms. To date, three ligninolytic haem peroxidases have been isolated and characterized in white-rot fungal species. These include lignin peroxidase (LiP), characterized by its ability to oxidize high redox-potential compounds such as alcohol and methoxybenzenes and veratryl, manganese peroxidase (MnP) which exhibits an ability to oxidize Mn²+ to Mn³+72 and versatile peroxidase (VP) which can oxidize both LiP and MnP substrates. Our genome analysis and subsequent homology modeling infers that A. mellea contains nine putative ligninolytic peroxidases. In an attempt to elucidate the function

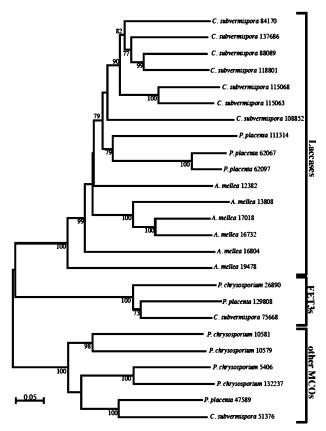


Figure 5. Maximum likelihood phylogenetic tree of laccase protein sequences of *A. mellea*. Sequences were aligned using MUSCLE (v3.6), ⁸⁴ with the default settings. Appropriate protein model of substitution was selected using ModelGenerator. ⁴⁰ One-hundred bootstrap replicates were applied with the appropriate protein model using the software program PHYML (v3.0) ⁸⁵ and summarized using the majority-rule consensus method. Monophyletic laccases, canonical ferroxidases (FET3) and other multicopper oxidases (MCO) are highlighted.

of the A. mellea peroxidases we reconstructed a phylogenetic tree using representative MnP, LiP and VP sequences from C. subvermispora and P. chrysosporium. S1 Our phylogeny infers that the A. mellea peroxidases are the result of a species-specific expansion as they are clustered together in a monophyletic clade (Figure 6). Interestingly the A. mellea-specific clade is clustered beside *C. subvermispora* MnP13, and this gene has previously been described as a short MnP.⁵¹ The length of the *A. mellea* genes is comparable to typical MnPs however (not shown). The phylogeny also infers that the A. mellea genes are more closely related to P. chrysosporium LiP genes than the "typical" C. subvermispora MnP genes (Figure 6). However, close sequence inspection shows that the A. mellea peroxidases do not contain the conserved aromatic substrate oxidation sites required for LiP enzymatic activity.⁷⁵ Furthermore, five of the nine A. mellea peroxidases contain the three acidic residues (two glutamates and an aspartate) required for the Mn²⁺ binding site of MnPs. Based on this observation we hypothesize that at least five of the peroxidases found in A. mellea have MnP activity.

Fungal Coculture

Comparative growth studies of *C. albicans* recovered from either mono- or coculture conditions demonstrates that significant killing (p = 0.0004) of *C. albicans* by *A. mellea* occurs in

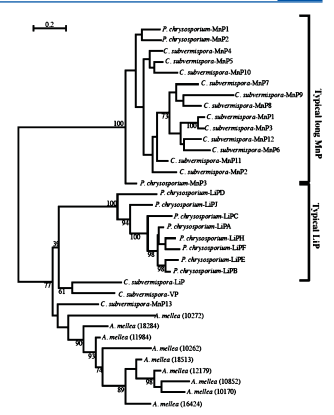


Figure 6. Maximum likelihood phylogenetic tree of peroxidase protein sequences of *A. mellea*. Sequences were aligned using MUSCLE (v3.6), ⁸⁴ with the default settings. Appropriate protein model of substitution was selected using ModelGenerator. ⁴⁰ One hundred bootstrap replicates were used with the appropriate protein model using the software program PHYML (v3.0)⁸⁵ and summarized using the majority-rule consensus method.

cocultures (Figure 7). This observation can be associated with the detection of 30 A. mellea proteins, not identified under any other culture condition (Table 6) including redox-active, degradative and attack-type proteins. Interestingly, Cys-2 peroxiredoxin (Am10593) and thioredoxin (Am7929) were both uniquely detected in fungal cocultures. Cys-2 peroxiredoxin converts peroxides to water, with concomitant formation of intramolecular disulfide linkages. These disulfide linkages are in turn reduced by thioredoxin to regenerate the active form of Cys-2 peroxiredoxin.^{76,77} We speculate that this system may be induced in A. mellea as an antifungal response to C. albicans presence. Alternatively, peroxide dismutation may be required to attenuate products originating from C. albicans cell wall degradation. In this regard, it is notable that a class II chitinase (Am13890) was also uniquely detected in fungal cocultures which may be deployed by A. mellea to degrade Candida cell wall material. Two proteins classified as prohibitins (termed Am14001 and Am20304), which have been proposed to be involved in the negative regulation of cell proliferation, 78,79 have also been identified elsewhere. 80 Intriguingly, Am3423 is classified by PFAM as a GPI-anchored protein. These serinethreonine-rich membrane proteins are anchored by glycosylphosphatidylinositol ligands. Several GPI-anchored proteins have been characterized in fungi as having multiple functions including cell wall organization in A. fumigatus, fruiting body formation in Lentinus edodes, and interaction with MAPK, where

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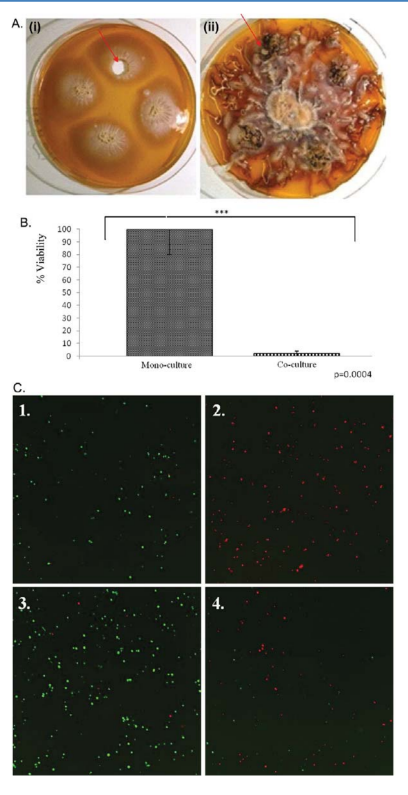


Figure 7. Fungicidal affect of *A. mellea* against *C. albicans* in coculture. (A) Culture plates: (i) *C. albicans* only; (ii) Coculture of *C. albicans* and *A. mellea* for up to 52 days. Reculture of *C. albicans* was carried out from multiple plugs taken from inoculation points (arrows). (B) Comparative viable cell count of recultured *C. albicans* from single and coculture plates shows that coculture with *A. mellea* results in significant (p = 0.0004) killing of *C. albicans*. (C) Fluorescent determination of *C. albicans* viability. Simultaneous FDA (live) and PI (dead) cell staining of *C. albicans*. 1. *C. albicans* live culture (24 h; positive control); 2. *C. albicans* killed by autoclaving (negative control); 3. Monoculture of *C. albicans* and 4. *C. albicans* following coculture with *A. mellea* (A.). Magnification: 20×.

Table 6. Proteins Found Uniquely in Coculture of A. mellea and C. albicans

accession no. ^a	BLAST2GO annotation b	tM _r ^c	$\operatorname{tp} I^d$	coverage (%)	unique peptides	SM score ^e	GRAVY score ^f	TM^g	$\begin{array}{c} \operatorname{SigP/} \\ \operatorname{SecP}^h \end{array}$	method ⁱ	source ^j
Am5344	60s ribosomal protein 110a	12603.7	9.6	10.7	1	11.51	-0.7		SecP	S	SN
Am12506	Aryl-alcohol oxidase	64088.4	4.7	2.7	1	16.33	-0.1		SigP	S	SN
Am17545	Coproporphyrinogen iii oxidase	103512.9	9.5	15.7	4	79.77	-0.1	2.0		S	SN
Am10593	Cys 2 peroxiredoxin	22242.4	5.2	6.5	1	12.87	-0.2		SecP	S	SN
Am9607	Cytochrome c oxidase subunit v	24861.9	9.7	7.5	1	20.74	-0.3	2.0	SecP	S	SN
Am19980	F1F0-ATP syn F	15435.1	10.5	14.5	1	19.06	-0.1	1.0	SecP	S	SN
Am16128	Glycerol-3-phosphate o- acyltransferase	64013.5	9.8	3.3	1	13.13	0.0	3.0		S	SN
Am13890	Glycosyl hydrolase 53 domain- containing protein	21828.7	6.9	14.7	1	18.79	0.0		SigP	S	SN
Am13814	Hypothetical protein	8166.3	5.8	18.7	1	18.59	-0.3			S	SN
Am13829	Hypothetical protein	66976.6	6.0	7.9	2	26.2	-0.3		SecP	S	SN
Am18856	Hypothetical protein	30601.9	9.5	6.1	1	13.69	0.4	4.0	SigP	S	SN
Am12218	Hypothetical protein	56022.9	4.6	4.1	1	14.46	-0.2		SigP	S	SN
Am12353	Iron-sulfur protein subunit	30797.5	8.7	9.7	1	13.5	-0.5		SecP	S	SN
Am19926	Lipoic acid synthase	67338.7	9.6	3.3	1	16.04	-0.2	1.0		S	SN
Am16692	Unknown Function Protein	25812.9	4.6	6.5	1	17.44	-0.4			S	SN
Am13379	Nadh dehydrogenase	61958.6	9.3	3.2	1	15	-0.2	2.0	SecP	S	SN
Am14001	Prohibitin Phb1	26007.0	6.9	7.9	1	20.8	0.1		SigP	S	SN
Am20343	Unknown Function Protein	10943.5	9.1	21.2	1	13.75	-0.3		SecP	S	SN
Am3423	Unknown Function Protein	13693.7	4.6	18.1	1	20.53	0.2		SigP	S	SN
Am6084	Unknown Function Protein	32823.9	5.7	9.8	1	13.81	-0.5		SecP	S	SN
Am20304	Proteolysis and peptidolysis-related protein	22016.4	8.7	6.6	1	15.62	-0.2		SecP	S	SN
Am17796	Secreted protein	46381.6	5.9	12.1	2	36.22	0.0			S	SN
Am18628	Subunit Vib of cytochrome c oxidase	10044.2	5.3	15.1	1	12.99	-0.8		SecP	S	SN
Am15086	Succinate-semialdehyde dehydrogenase	74240.9	8.5	2.0	1	15.07	0.1	4.0	SecP	S	SN
Am16124	Sulfide-quinone oxidoreductase	118017.6	8.7	4.0	2	33.19	-0.4	2.0		S	SN
Am7929	Thioredoxin	13706.0	4.7	15.6	1	15.39	0.1			S	SN
Am14705	Twin-arginine translocation pathway signal	63826.1	5.8	4.9	1	16.77	0.0		SecP	S	SN
Am2793	Ubiquitin domain-containing	37237.5	5.2	6.4	1	16.2	-0.2			S	SN
Am18503	Uracil phosphoribosyltransferase	19530.7	4.9	10.3	1	15.46	0.2		SecP	S	SN
Am14973	Urea hydro-lyase cyanamide hydratase	28060.0	5.7	8.4	1	19.82	-0.2		SecP	S	SN

[&]quot;Accession number from A. mellea cDNA database. "Blast annotation following Blast2GO analysis of proteins identified from cDNA database. "tM_D theoretical molecular mass. "tpI, theoretical isoelectric point. "SM score, Spectrum Mill protein score. "fGRAVY score, grand average of hydropathy. "TM, number of transmembrane regions. "SigP, classical secretion signal peptide; SecP, nonclassical secretion signal. "Method: 1D SDS-PAGE prior to LC-MS/MS analysis, 2-D proteins separated in two dimensions prior to LC-MS/MS analysis. "Source, M: mycelia; SN: culture supernatant/secretome

they may have a signaling role. ⁸¹ Am3423 belongs to the same protein family of DRMIP, Hesp-379, which is a haustorially expressed secreted protein involved in host tissue penetration in *L. edodes*. ⁸² In addition, a number of oxidoreductases, monooxygenases, reductases and dehydrogenases of basidiomycete origin were detected in the fungal cocultures (Table 6). Similarly, metabolite extracts from *A. tabescens* have also been observed to exhibit antifungal activity. ⁸³ Apart from demonstrating the power of proteomic investigations to dissect fungal–fungal interactions, the coculture system we have developed for *A. mellea* and *C. albicans* may represent a model system for the further investigations of *A. mellea* pathogenicity.

CONCLUSIONS

Combined application of Illumina sequencing with LC-MS/MS has, for the first time, revealed new insights into the proteome of the important fungal plant pathogen *Armillaria mellea*, including the identification of multiple glycodegradative enzyme systems which can be useful for recalcitrant carbon compound degradation in biotechnology applications. Moreover, proteo-

mics tools shed light on potent antifungal properties of *A. mellea*, which in part may be mediated by basidiomycete enzymes.

ASSOCIATED CONTENT

Supporting Information

Supplemental tables and figure. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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