

## Original article

# FunSecKB: the Fungal Secretome KnowledgeBase

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The Fungal Secretome KnowledgeBase (FunSecKB) provides a resource of secreted fungal proteins, i.e. secretomes, identified from all available fungal protein data in the NCBI RefSeq database. The secreted proteins were identified using a well evaluated computational protocol which includes SignalP, WolfPsort and Phobius for signal peptide or subcellular location prediction, TMHMM for identifying membrane proteins, and PS-Scan for identifying endoplasmic reticulum (ER) target proteins. The entries were mapped to the UniProt database and any annotations of subcellular locations that were either manually curated or computationally predicted were included in FunSecKB. Using a web-based user interface, the database is searchable, browsable and downloadable by using NCBI's RefSeq accession or gi number, UniProt accession number, keyword or by species. A BLAST utility was integrated to allow users to query the database by sequence similarity. A user submission tool was implemented to support community annotation of subcellular locations of fungal proteins. With the complete fungal data from RefSeq and associated web-based tools, FunSecKB will be a valuable resource for exploring the potential applications of fungal secreted proteins.

**Database URL:** <http://proteomics.yosu.edu/secretomes/fungi.php>

## Introduction

Fungi play an important role in carbon cycling as they use secreted enzymes to break down lignocelluloses and other biopolymers then transporting the resulting products into the cells as their food. The secreted proteins in plant associated fungi play important roles in plant and fungi symbiosis or fungal pathogenicity (1). Fungal secreted proteins also play important roles in the development of fungal diseases in human (2,3). Secreted fungal enzymes have found a wide range of applications in the food, feed, pulp and paper, bioethanol and textile industries (4).

Signal-peptide dependent secreted proteins contain a signal peptide (SP) at the N-terminus that directs the ribosomes to the rough endoplasmic reticulum (ER) for completing polypeptide synthesis (5,6). The signal peptide, typically 15–30 amino acids long and consisting of 15–20 hydrophobic amino acid residues, is cleaved off during

translocation across the membrane. While some proteins without an N-terminal signal peptide can be found in the ER and the Golgi, over 90% of human secreted proteins (7) and ~90% of the *Aspergillus niger* extracellular proteins identified by mass spectrometry contain classical N-terminal signal peptides (8). There are also examples of non-classically secreted proteins in fungi, including the *Saccharomyces cerevisiae* mating pheromone  $\alpha$ -factor (9) and two galectins from *Coprinus cinereus* (10), but it is generally believed that the vast majority of secreted fungal proteins are processed by the classical secretory pathway (8).

The term secretome is often used to refer to the complete set of secreted proteins in an organism (2,11,12). However, the term has also been used to include the set of proteins involved in the secretory pathway (13,14). In the work described here, the secretome only includes the secreted proteins in an organism. Along with an increased

number of species having genomes being completely sequenced, we see an increased number of publications on fungal secretome identification and analysis using both computational and experimental approaches (15). For example, secretomes have been reported in following fungi including *A. niger* (8), *Candida albicans* (16), *Phanerochaete chrysosporium* (17), *Sclerotinia sclerotiorum* (18), *Fusarium graminearum* (19) and *Ustilago maydis* (20). Considering the biological importance of secreted proteins and their potential industrial applications, we developed a knowledgebase of fungal secretomes for identification, annotation and curation of both computationally predicted and experimentally identified fungal secreted proteins. This knowledgebase is designed to serve as a central portal for providing as well as collecting information on fungal secretomes.

## Data collection and database implementation

The fungal protein sequences were retrieved from the NCBI Reference Sequence collection (RefSeq) database (release April, 2010) (<http://www.ncbi.nlm.nih.gov/RefSeq/>). The rationale for choosing the RefSeq protein data set was that RefSeq provides a comprehensive, integrated, non-redundant, well-annotated set of proteins and also the corresponding nucleotide sequences were also linked for these protein sequences in their database (21). The data in the fungal secretome knowledgebase (FunSecKB) were obtained from the following three sources: (i) the features predicted using computational approaches; (ii) subcellular locations annotated in UniProtKB; and (iii) our manual curation with experimental evidence obtained from recent literature.

### Computational methods for prediction of secreted proteins

The fungal protein sequences downloaded from the NCBI RefSeq database were processed using the following programs including SignalP (version 3.0, <http://www.cbs.dtu.dk/services/SignalP/>) (22), Phobius (<http://phobius.binf.ku.dk/>) (23,24), WolfPsort (<http://wolfsort.org/>) (25,26) and TargetP (<http://www.cbs.dtu.dk/services/TargetP/>) (27), for signal peptide and subcellular location prediction. We chose these four predictors because they were previously evaluated favorably and widely used by the fungal secretome research community (8,16,28). TMHMM (<http://www.cbs.dtu.dk/services/TMHMM/>) was used to identify proteins having transmembrane domains (29) and PS-Scan (<http://www.expasy.org/tools/scanprosite/>) was used to scan ER targeting sequence (Prosites: PS00014) (30). With each of the programs, the default parameters for eukaryotes or fungi were used. For SignalP prediction, only entries that were

predicted having a 'mostly likely cleavage site' by SignalP-NN algorithm and a 'signal peptide' by SignalP-HMM algorithm were considered to be true signal peptide 'positives', using the N-terminal 70 amino acids (22). For predicting membrane proteins using TMHMM, the entries having membrane domains not located within the N-terminus (the first 70 amino acids) were treated as real membrane proteins. Protein sequences predicted to have a signal peptide by SignalP were further processed using FragAnchor to identify the glycosylphosphatidyinositol (GPI) anchors (<http://navet.ics.hawaii.edu/~fraganchor/NNHMM/NNHMM.html>) (31). Protein sequences predicted as having a GPI anchor may be attached to the outside of the plasma membrane or may be secreted to be targeted to the cell wall (32).

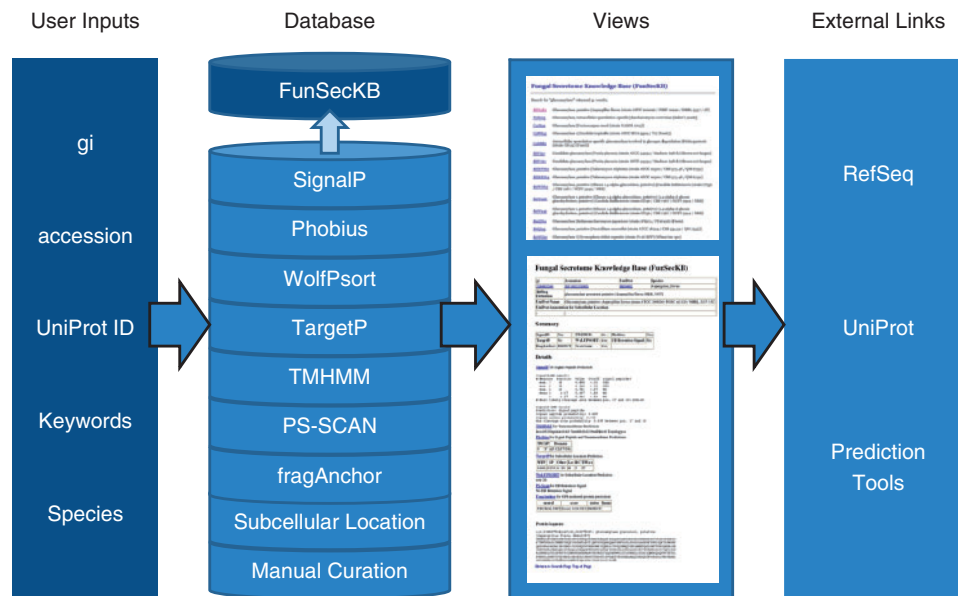
We recently performed the accuracy evaluation of the computational methods, using 241 experimentally identified secreted proteins and 5992 non-secreted proteins in fungi that were retrieved from UniProt/Swiss-Prot data set, and found that the highest prediction accuracy (92.1% in sensitivity and 98.9% in specificity) was achieved by combining SignalP, WolfPsort and Phobius for signal peptide prediction, TMHMM for eliminating membrane proteins, and PS-Scan for removing ER targeting proteins (28). Thus, the secretomes defined in this study include the manually curated secreted proteins along with the proteins predicted as having a signal peptide at their N-terminus by SignalP and Phobius and with a subcellular location predicted as extracellular by WolfPsort, but not having a transmembrane domain or an ER targeting signal. The information provided by TargetP and fragAnchor were also included in the annotation which may be useful for identifying mitochondrial targeted proteins or GPI anchored membrane or cell wall proteins. An overview of the database's features are shown in Figure 1.

### Linking RefSeq proteins to UniProtKB annotation

The fungal protein entries in FunSecKB are linked to the UniProtKB using the mapping information generated in UniProtKB ([ftp://ftp.uniprot.org/pub/databases/uniprot/current\\_release/knowledgebase/idmapping/](ftp://ftp.uniprot.org/pub/databases/uniprot/current_release/knowledgebase/idmapping/)) (33). We also integrated the subcellular location information of fungal proteins annotated in UniProtKB including curated (reviewed, from the UniProtKB/Swiss-Prot data set) and predicted (unreviewed, from the UniProtKB/TrEMBL data set). In addition, we also included manually curated protein entries in UniProtKB/Swiss-Prot data set which could not be mapped to entries in the RefSeq database.

### Manual curation and community annotation

FunSecKB supports community curation of subcellular locations of fungal proteins based on published experimental evidence. A submission form was developed for users to provide subcellular location annotation and the literature



**Figure 1.** Overview of FunSecKB. To search the database users can enter NCBI RefSeq gi or accession number, UniProt accession number, keywords or species. The database consists of information generated using seven prediction tools and subcellular location annotated in UniProtKB and our own manual curation. Users can browse through the results using the web user-interface. Links to external databases and resources are also provided for further exploration. Whole secretome sequences can be downloaded and BLAST utility can be accessed from the database interface.

source to support the annotation. After our curator's validation, these data will be incorporated into the database. Currently we have manually curated more than two hundred secreted proteins from *A. niger* (8). Manual curation is an ongoing process, thus additional secreted proteins will be manually curated and integrated into the database with time.

The information from the above three sources are integrated in the annotation (Figure 1). The annotated entries are linked to the RefSeq database in NCBI and UniProtKB as well as related literature for entries manually curated by our curators or the community. The data will be updated when a new RefSeq data set is released from NCBI (<http://www.ncbi.nlm.nih.gov/RefSeq/>).

## Data access

FunSecKB can be accessed through the database web interface at <http://proteomics.yu.edu/secretomes/fungi.php>. There are three approaches to accessing the data including: (i) search individual proteins using NCBI's RefSeq gi or accession number, UniProt accession number, keyword or by species; (ii) search or download the whole secretome or a subset of manually curated secreted proteins of a species and (iii) search all fungal proteins or fungal secreted proteins using BLAST.

The annotation page contains the summary and the details of subcellular locations predicted by the tools

mentioned above and annotation retrieved from UniProtKB. Each entry is linked to both RefSeq and UniProtKB. The secretome, including predicted and curated secreted proteins from a particular species, can be searched and downloaded by selecting a species from the species list for complete genomes or inputting a species name for others not having a complete genome. The protein sequences of the secretome from a species can be downloaded into a fasta file. Manually curated secreted proteins consist of proteins retrieved from UniProtKB/Swiss-Prot with subcellular locations labeled as 'reviewed' and proteins curated by our curators and the users. The proteins curated by us and by the community are supported by experimental evidence for their subcellular location annotation and the related literature can be found on the same page. The annotation page also contains the primary protein sequence (Figure 1). The database interface provides a link to the BLAST input interface to search through the proteins retrieved from RefSeq: either all fungal proteins or just the fungal secretomes.

## Preliminary data analysis

Currently FunSecKB contains a total of 478 073 fungal protein sequences including 23 878 predicted and/or curated secreted proteins from a total of 118 fungal species. This includes 52 fungal species, with one species having two different varieties, having a complete predicted proteome set.

We performed a preliminary analysis on the 53 complete secretomes of 52 fungal species including 43 Ascomycetes, 7 Basidiomycetes (with *Cryptococcus neoformans* having two varieties) and 2 Microsporidia (Table 1). Overall, fungal species having an expanded genome size encode more proteins in their predicted proteomes ( $r=0.75$ ) (Figure 2a). *Ajellomyces dermatitidis* and *Postia placenta* are two outliers. For the *P. placenta* genome of 69 Mb the RefSeq only has 9083 predicted proteins, however, Martinez *et al.* (2009) reported 17 173 proteins predicted from the *P. placenta* genome (34). Thus the discrepancy may be caused by lagged database update. The reason for the *A. dermatitidis* data is not known.

The proportion of the secretomes in the proteomes in different species varies significantly from <1% in *Encephalitozoon cuniculi* and *Enterocytozoon bieneusi*, two Microsporidia species (unicellular parasites), to >10% in *Magnaporthe grisea*, a rice pathogenic fungus (Table 1). Overall, predicted secretome sizes increase with expanded proteome sizes in fungal species ( $r=0.83$ ) (Figure 2b). We further identified GPI-anchored proteins in the predicted secretome, which represent insoluble portions of secreted proteins that are components of cell walls or attached to the outside of cell membrane. We see that both insoluble and soluble portions are increased with increased proteome size in different fungal species (Figure 2c and 2d).

The functional categorization of predicted secretomes was analyzed using the rpsBLAST tool in the NCBI BLAST package to search the conserved domain database (35). The highly encoded secreted protein families having more than 50 members in the whole database are listed in Table 2. Preliminary functional analysis revealed that the fungal secretomes largely consist of enzymes, particularly hydrolases, which are used to breakdown carbohydrates, lipids, proteins and all other types of organic materials by fungi (Table 2). Furthermore, a total of 10 397 secreted proteins have GO annotations in UniProtKB. Among them, molecular functional classification using GOSlimViewer ([http://agbase.msstate.edu/cgi-bin/tools/goslimviewer\\_select.pl](http://agbase.msstate.edu/cgi-bin/tools/goslimviewer_select.pl)) showed 43% were hydrolases including peptidases (Figure 3) (36). These enzymes have potential applications in biofuel production. The database user interface features an easy to use option to download predicted secretomes from completely sequenced fungal species. This provides a resource for further detailed species specific or interspecies comparative analysis.

## Discussion

While constructing our database, a similar fungal secretome database (FSD, <http://fsd.snu.ac.kr/>) was published by Choi *et al.* (37). However, there are several important differences between the two databases (Table 3). We used RefSeq data while the FSD used only completely sequenced

fungal genome data including some 'work in progress' genomes (37). The prediction methods used for identification of secreted proteins were also different. The FSD used a three-layer hierarchical identification rule based on 9 different programs and considered entries to be secreted proteins as long as any one of the tools predicted it to be secreted, thus the number of secreted proteins were much higher than the number predicted in our database. For example, in *A. niger*, we predicted 832 secreted proteins in the strain CBS 513.88, while Choi *et al.* (37) predicted 1831 secreted proteins in the same strain and 2616 secreted proteins in the ATCC1015 strain in the FSD (37). However, there were only from 691 to 881 proteins which were predicted to be secreted, with 160 of them being confirmed experimentally in the ATCC1015 strain by Tsang *et al.* (8). Thus, we believe the methods used in the FSD significantly over-estimated the number of secreted proteins in fungi. In addition, the search for the FSD is limited to using the sequence locus name and can not be searched with NCBI gi and accession number, UniProt accession number or keywords. There is also not a curation tool available for the community annotation in FSD (37).

In addition to the signal-peptide dependent secreted proteins using the classical ER-Golgi secretory pathway, there are non-classical, signal peptide independent, secretory pathways in all domains of organisms. Mammalian and bacterial leadless secreted proteins have been collected and used to implement the prediction software, SecretomeP, for predicting these proteins (<http://www.cbs.dtu.dk/services/SecretomeP/>) (38,39). The tool has not been trained with fungal-specific data and the accuracy for predicting fungal non-classical secreted protein could not be evaluated, thus we did not include this tool in our data processing. Although the FSD used SecretomeP to predict non-classical secreted proteins, the predicted secreted proteins were not included in the secretome analysis; including them would make the putative secretome >40% of whole proteome (37). Nevertheless, the FunSecKB and the FSD databases could complement each other as different data sources, prediction tools and data access utilities were implemented.

In summary, we constructed FunSecKB to identify, annotate and curate the secreted proteins in fungi. The data can be searched using protein identifiers or keywords, and by species. Most of the secreted proteins are currently predicted by computational tools. However, the community can use the curation module implemented in our site to manually curate subcellular locations of fungal proteins having experimental evidence. The resource described in the work is expected to provide a query and curation system that will help the community to further understand the secretome biology and explore various potential applications of fungal secreted proteins in bio-processing or environmental remediation industries.

Table 1. Summary of genome size, proteome size, secretome size in different fungi

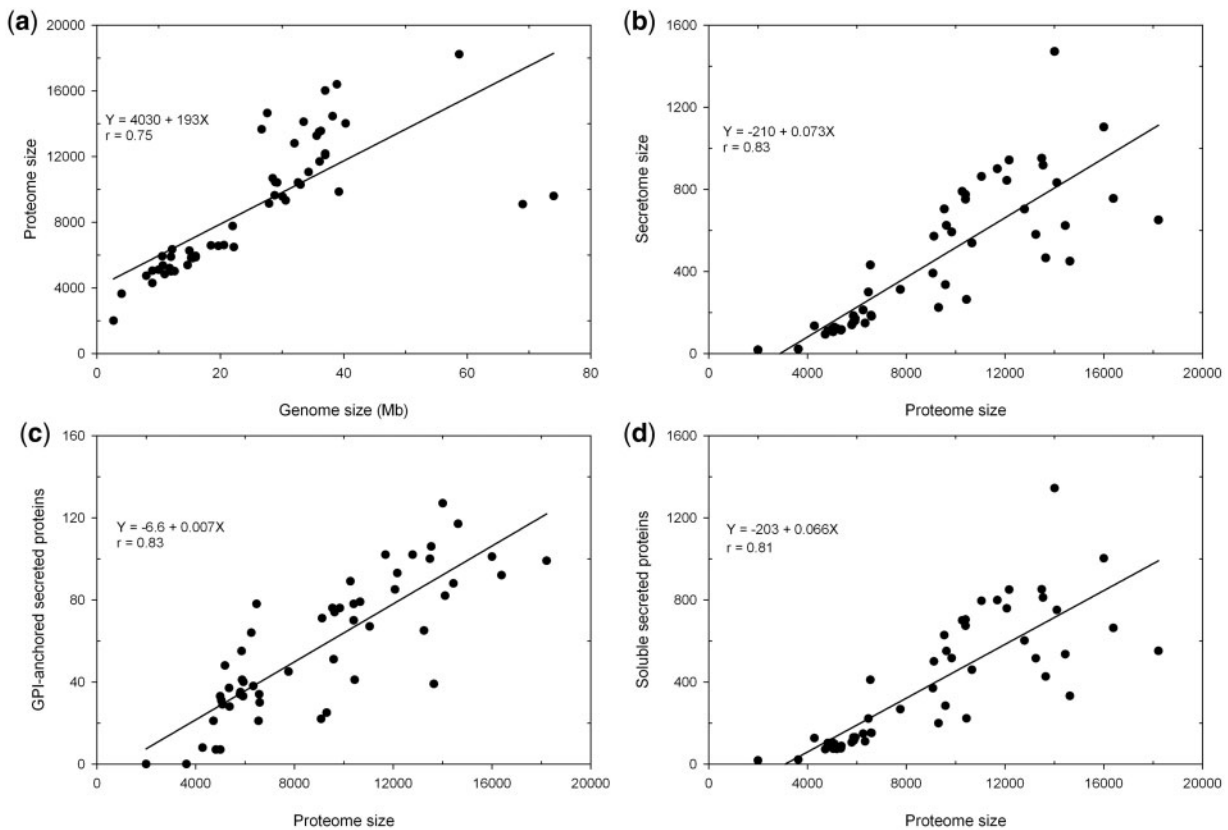
Species	Phylum	Genome (Mb)	Predicted Proteome	Predicted Secretome	Curated Secretome	GPI-anchored Secretome	Soluble Secretome	Secretome (%)	GPI-anchored Portion (%)
<i>Ajellomyces capsulatus</i>	Ascomycota	31	9313	224	0	25	199	2.4	11.2
<i>Ajellomyces dermatitidis</i>	Ascomycota	74	9587	335	0	51	284	3.5	15.2
<i>Ashbya gossypii</i>	Ascomycota	8	4725	93	2	21	72	2.0	22.6
<i>Aspergillus clavatus</i>	Ascomycota	28	9121	571	17	71	500	6.3	12.4
<i>Aspergillus flavus</i>	Ascomycota	36	13 487	951	25	100	851	7.1	10.5
<i>Aspergillus fumigatus</i>	Ascomycota	29	9630	624	58	74	550	6.5	11.9
<i>Aspergillus nidulans</i>	Ascomycota	30	9541	704	29	76	628	7.4	10.8
<i>Aspergillus niger</i>	Ascomycota	34	14 102	832	253	82	750	5.9	9.9
<i>Aspergillus oryzae</i>	Ascomycota	37	12 074	843	28	85	758	7.0	10.1
<i>Aspergillus terreus</i>	Ascomycota	29	10 401	774	23	70	704	7.4	9.0
<i>Botryotinia fuckeliana</i>	Ascomycota	39	16 389	755	4	92	663	4.6	12.2
<i>Candida albicans</i>	Ascomycota	28	14 633	449	41	117	332	3.1	26.1
<i>Candida dubliniensis</i>	Ascomycota	16	5860	184	0	55	129	3.1	29.9
<i>Candida glabrata</i>	Ascomycota	12	5192	121	7	48	73	2.3	39.7
<i>Candida tropicalis</i>	Ascomycota	15	6254	212	1	64	148	3.4	30.2
<i>Chaetomium globosum</i>	Ascomycota	34	11 048	862	1	67	795	7.8	7.8
<i>Clavisporea lusitanae</i>	Ascomycota	16	5936	169	0	40	129	2.8	23.7
<i>Coccidioides immitis</i>	Ascomycota	29	10 440	263	2	41	222	2.5	15.6
<i>Debaryomyces hansenii</i>	Ascomycota	12	6335	148	1	38	110	2.3	25.7
<i>Gibberella zeae</i>	Ascomycota	36	11 690	900	1	102	798	7.7	11.3
<i>Kluyveromyces lactis</i>	Ascomycota	11	5357	113	5	37	76	2.1	32.7
<i>Lachancea thermotolerans</i>	Ascomycota	10	5091	128	0	29	99	2.5	22.7
<i>Lodderomyces elongisporus</i>	Ascomycota	16	5799	139	0	34	105	2.4	24.5
<i>Magnaporthe grisea</i>	Ascomycota	40	14 010	1471	3	127	1344	10.5	8.6
<i>Neosartorya fischeri</i>	Ascomycota	33	10 406	751	21	78	673	7.2	10.4
<i>Neurospora crassa</i>	Ascomycota	39	9844	592	10	76	516	6.0	12.8
<i>Penicillium chrysogenum</i>	Ascomycota	32	12 791	703	5	102	601	5.5	14.5
<i>Penicillium marneffei</i>	Ascomycota	29	10 663	538	0	79	459	5.0	14.7
<i>Phaeosphaeria nodorum</i>	Ascomycota	37	16 002	1103	1	101	1002	6.9	9.2
<i>Pichia guilliermondii</i>	Ascomycota	11	5920	159	0	33	126	2.7	20.8
<i>Pichia pastoris</i>	Ascomycota	9	5040	105	0	31	74	2.1	29.5
<i>Pichia stipitis</i>	Ascomycota	15	5816	144	0	35	109	2.5	24.3

(Continued)

Table 1. Continued.

Species	Phylum	Genome (Mb)	Predicted Proteome	Predicted Secretome	Curated Secretome	GPi-anchored Secretome	Soluble Secretome	Secretome (%)	GPi-anchored Portion (%)
<i>Podospira anserina</i>	Ascomycota	33	10 272	789	1	89	700	7.7	11.3
<i>Pyrenophora tritici-repentis</i>	Ascomycota	37	12 169	942	0	93	849	7.7	9.9
<i>Saccharomyces cerevisiae</i>	Ascomycota	12	5885	156	101	41	115	2.7	26.3
<i>Schizosaccharomyces japonicus</i>	Ascomycota	11	4824	109	0	7	102	2.3	6.4
<i>Schizosaccharomyces pombe</i>	Ascomycota	13	5001	112	43	7	105	2.2	6.3
<i>Sclerotinia sclerotiorum</i>	Ascomycota	38	14 446	623	1	88	535	4.3	14.1
<i>Talaromyces stipitatus</i>	Ascomycota	36	13 252	580	0	65	515	4.4	11.2
<i>Uncinocarpus reesii</i>	Ascomycota	22	7760	312	0	45	267	4.0	14.4
<i>Vanderwaltozyma polyspora</i>	Ascomycota	15	5376	116	0	28	88	2.2	24.1
<i>Yarrowia lipolytica</i>	Ascomycota	22	6472	299	5	78	221	4.6	26.1
<i>Zygosaccharomyces rouxii</i>	Ascomycota	12	4994	120	0	33	87	2.4	27.5
<i>Coprinopsis cinerea</i>	Basidiomycota	36	13 546	917	8	106	811	6.8	11.6
<i>Cryptococcus neoformans (neoformans B-3501A)</i>	Basidiomycota	19	6578	186	0	34	152	2.8	18.3
<i>Cryptococcus neoformans (neoformans JEC21)</i>	Basidiomycota	21	6594	181	0	30	151	2.7	16.6
<i>Laccaria bicolor</i>	Basidiomycota	59	18 215	650	0	99	551	3.6	15.2
<i>Malassezia globosa</i>	Basidiomycota	9	4286	134	0	8	126	3.1	6.0
<i>Moniliophthora perniciosa</i>	Basidiomycota	27	13 649	465	0	39	426	3.4	8.4
<i>Postia placenta</i>	Basidiomycota	69	9083	391	0	22	369	4.3	5.6
<i>Ustilago maydis</i>	Basidiomycota	20	6548	431	2	21	410	6.6	4.9
<i>Encephalitozoon cuniculi</i>	Microsporidia	3	1996	17	2	0	17	0.9	0.0
<i>Enterocytozoon bieneusi</i>	Microsporidia	4	3632	21	0	0	21	0.6	0.0
Other species			998	367	366				
Total			478 073	23 878	1067	3014			





**Figure 2.** Relationship between genome size, proteome size and secretome size in fungi. (a) genome size and proteome size; (b) proteome size and secretome size; (c) proteome size and GPI-anchored secreted proteins and (d) proteome size and soluble secreted proteins.

**Table 2.** Highly encoded secreted protein families in fungi

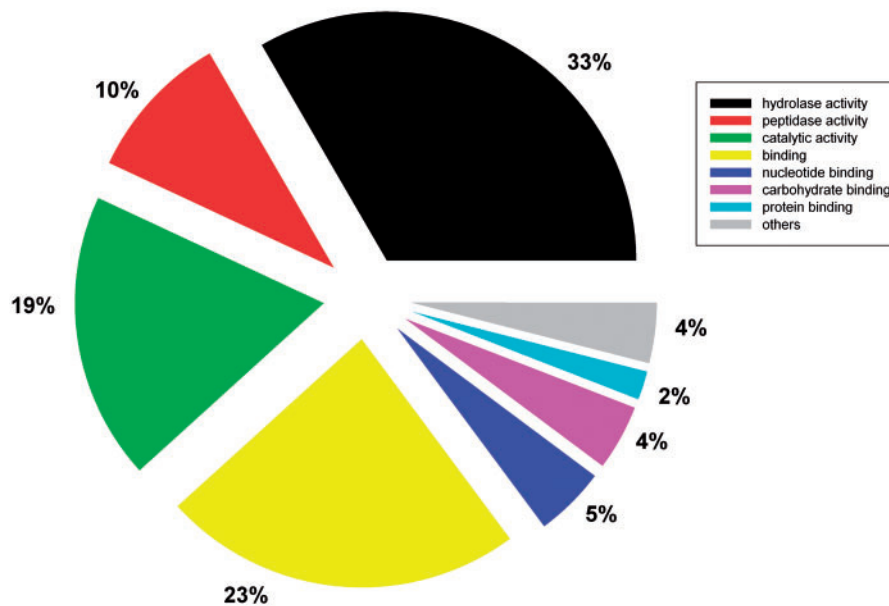
CDD functional domains	Numbers
pfam00135, COesterase, Carboxylesterase	314
pfam03443, Glyco hydro 61, Glycosyl hydrolase family 61	301
COG0277, GlcD, FAD/FMN-containing dehydrogenases	287
cd04077, Peptidases S8 PCSK9 ProteinaseK like: Peptidase S8 family domain in ProteinaseK-like proteins	223
pfam00450, Peptidase S10, Serine carboxypeptidase	215
pfam00295, Glyco hydro 28, Glycosyl hydrolases family 28	207
pfam00067, p450, Cytochrome P450	160
pfam00933, Glyco hydro 3, Glycosyl hydrolase family 3 N terminal domain	156
cd05474, pepsin-like proteinases secreted from pathogens to degrade host proteins	154
COG2303, BetA, Choline dehydrogenase and related flavoproteins	152
pfam01083, Cutinase	139
pfam09362, DUF1996, Domain of unknown function (DUF1996)	136
pfam00264, Tyrosinase, Common central domain of tyrosinase	130
TIGR03388, ascorbase, L-ascorbate oxidase, plant type	128
cd04056, Peptidases S53, Peptidase domain in the S53 family	124
pfam04389, Peptidase M28, Peptidase family M28	122
COG5309, COG5309, Exo-beta-1,3-glucanase	121
pfam04616, Glyco hydro 43, Glycosyl hydrolases family 43	114

(Continued)

Table 2. Continued.

CDD functional domains	Numbers
cd00519, Lipase 3, Lipase (class 3)	106
PRK02106, PRK02106, choline dehydrogenase	100
COG2730, BglC, Endoglucanase	99
pfam00328, Acid phosphat A, Histidine acid phosphatase	98
pfam03856, SUN, Beta-glucosidase (SUN family)	97
pfam07519, Tannase, Tannase and feruloyl esterase	97
smart00656, Amb all, Amb all domain	94
pfam00457, Glyco hydro 11, Glycosyl hydrolases family 11	92
cd06097, Aspergillopepsin like: Aspergillopepsin like, aspartic proteases of fungal origin	91
cd02877, GH18 hevamine Xipl class III	88
pfam00331, Glyco hydro 10, Glycosyl hydrolase family 10	88
pfam01565, FAD binding 4, FAD binding domain	87
pfam03583, LIP, Secretory lipase	87
pfam03659, Glyco hydro 71, Glycosyl hydrolase family 71	87
pfam01185, Hydrophobin, Fungal hydrophobin	85
pfam01532, Glyco hydro 47, Glycosyl hydrolase family 47	79
cd02181, GH16 MLG1 glucanase	78
cd05471, Pepsin-like aspartic proteases, bilobal enzymes that cleave bonds in peptides at acidic pH	77
cd05384, SCP PRY1 like, SCP-like extracellular protein domain, PRY1-like sub-family restricted to fungi	75
cd07203, Fungal Phospholipase B-like; cPLA2 GpIIVA homologs; catalytic domain	71
pfam00840, Glyco hydro 7, Glycosyl hydrolase family 7	71
pfam00150, Cellulase, Cellulase (glycosyl hydrolase family 5)	70
pfam11790, Glyco hydro cc, Glycosyl hydrolase catalytic core	70
pfam01522, Polysacc deac 1, Polysaccharide deacetylase	69
pfam07971, Glyco hydro 92, Glycosyl hydrolase family 92	68
smart00636, Glyco 18, Glycosyl hydrolase family 18	68
cd00842, MPP ASMase, acid sphingomyelinase and related proteins	67
cd03457, intradiol dioxygenase like, Intradiol dioxygenase supgroup	67
pfam03663, Glyco hydro 76, Glycosyl hydrolase family 76	67
pfam05577, Peptidase S28, Serine carboxypeptidase S28	67
pfam12296, HsbA, Hydrophobic surface binding protein A	65
cd02183, GH16 GPI glucanosyltransferase	64
COG0654, 2-polyprenyl-6-methoxyphenol hydroxylase and related FAD-dependent oxidoreductases	63
pfam01055, Glyco hydro 31, Glycosyl hydrolases family 31	62
cd06248, Peptidase M14 Carboxypeptidase A/B-like subfamily	61
pfam02128, Peptidase M36, Fungalysin metallopeptidase (M36)	61
pfam04185, Phosphoesterase, Phosphoesterase family	61
pfam11765, Hyphal reg CWP, Hyphally regulated cell wall protein	60
pfam01328, Peroxidase 2, Peroxidase, family 2	59
pfam01828, Peptidase A4, Peptidase A4 family	58
pfam03198, Glyco hydro 72, Glycolipid anchored surface protein	57
cd01846, Fatty acyltransferase-like subfamily of the SGNH hydrolases, a diverse family of lipases and esterases	56
pfam02102, Peptidase M35, Deuterolysin metalloprotease (M35)	56
pfam00723, Glyco hydro 15, Glycosyl hydrolases family 15	54
pfam00128, Alpha-amylase, Alpha amylase, catalytic domain	53
cd08588, Catalytic domain of Arabidopsis thaliana PI-PLC X domain-containing protein	52
PHA03247, PHA03247, large tegument protein UL36; Provisional	52
pfam01301, Glyco hydro 35, Glycosyl hydrolases family 35	51
pfam11937, DUF3455, Protein of unknown function (DUF3455)	51





**Figure 3.** Molecular functional classification of fungal secreted proteins using GOSlimViewer.

**Table 3.** Comparison of the two independently developed fungal secretome databases

	FSD	FunSecKB
Data source	Fungal genomes	Fungal proteins in RefSeq
Prediction tools	SignalP3.0; SigCleave; SigPred; RPSP; TMHMM2.0c; TargetP1.1b; PsortII; PredictNLS; SecretomeP1.0f	SignalP 3.0; Phobius1.01; WolfPsort0.2; TargetP1.1b, TMHMM2.0c; PS-Scan
Data access	Sequence locus name; BLAST	Keywords, RefSeq gi or accession, UniProt accession; BLAST
Community curation tool	Not available	Available

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