# Evolutionary histories of expanded peptidase families in *Schistosoma mansoni*

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Schistosoma mansoni is one of the three main causative agents of human schistosomiasis, a major health problem with a vast socio-economic impact. Recent advances in the proteomic analysis of schistosomes have revealed that peptidases are the main virulence factors involved in the pathogenesis of this disease. In this context, evolutionary studies can be applied to identify peptidase families that have been expanded in genomes over time in response to different selection pressures. Using a phylogenomic approach, we searched for expanded endopeptidase families in the S. mansoni predicted proteome with the aim of contributing to the knowledge of such enzymes as potential therapeutic targets. We found three endopeptidase families that comprise leishmanolysins (metallopeptidase M8 family), cercarial elastases (serine peptidase S1 family) and cathepsin D proteins (aspartic peptidase A1 family). Our results suggest that the Schistosoma members of these families originated from successive gene duplication events in the parasite lineage after its diversification from other metazoans. Overall, critical residues are conserved among the duplicated genes/proteins. Furthermore, each protein family displays a distinct evolutionary history. Altogether, this work provides an evolutionary view of three S. mansoni peptidase families, which allows for a deeper understanding of the genomic complexity and lineage-specific adaptations potentially related to the parasitic lifestyle.

Key words: phylogenomics - maximum likelihood analysis - homology prediction - functional annotation - proteases - paralogous families - parasite genomics

Schistosomiasis, which is caused by different species from the Schistosoma genus, remains one of the most prevalent tropical neglected diseases, affects 210 million people worldwide, and is responsible for at least 280,000 deaths every year (van der Werf et al. 2003, Steinmann et al. 2006, Han et al. 2009). Schistosoma mansoni is one of the three major species that infect humans and is the causative agent of intestinal and hepatic schistosomiasis mainly in Africa and South America (Han et al. 2009). Measures to control schistosomiasis rely almost entirely on praziquantel<sup>®</sup>, which is the only drug available for mass chemotherapy. Despite the effectiveness of this treatment, re-infection is common and drug-resistant parasites have been found in the laboratory and in the field, which demonstrate the urgent need to develop additional chemotherapeutic agents and effective vaccines (Liang et al. 2003, Pica-Mattoccia & Cioli 2004, Botros & Bennett 2007, Melman et al. 2009).

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sites composed of an aspartic acid and an asparagine, the latter being the P1 residue, the amino acid or molecule, which can be found at a specific location in the cleavage site (Rawlings et al. 2010). In turn, aspartic peptidases have their catalytic centres formed by two aspartate residues that activate a water molecule that mediates the nucleo-

lar analysis of major parasites have identified some key factors involved in parasitic diseases and peptidases as one of the major factors of pathogenicity (McKerrow et al. 2006, Kasný et al. 2009). These enzymes have been implicated in processes that are crucial to the development and survival of helminth parasites, including digestion, invasion from host tissues, activation of inflammation and evasion of the host immune system (McKerrow et al. 2006, Kasný et al. 2009).

Over the past several years, advances in the molecu-

Peptidases (also termed proteases, proteinases or proteolytic enzymes) are hydrolytic enzymes that cleave peptide bonds in proteins. Endopeptidases cleave internal peptide bonds, whereas exopeptidases hydrolyse the amino terminus (aminopeptidases) or carboxy terminus (carboxypeptidases) of different proteins. Enzymatic specificity is determined based on the chemical groups responsible for catalysis in the peptide's active site. Thus, peptidases are classified into one of the following classes: asparagine, aspartic, cysteine, glutamic, metallo, serine, threonine and unknown peptidases (Rawlings & Barrett 1993, Rawlings et al. 2010).

Asparagine peptidases are enzymes that have active

philic attack on the peptide bond (James 2004, Rawlings

et al. 2010). In general, cysteine peptidases have cysteine and histidine residues forming their "catalytic dyad". Meanwhile, other active site residues have been found. Glutamic peptidases have glutamic acid residues as their primary catalytic residues, which are probably the nucleophilic attack mediators involved in the catalysis (Fujinaga et al. 2004, Rawlings et al. 2010). In metallopeptidases, the catalytic mechanism usually involves a single catalytic zinc ion tetrahedrally coordinated by one glutamate and two histidine residues (Rawlings et al. 2010). Serine peptidases have serine residues at their active sites, which together with two other variable amino acids constitute the "catalytic triad" (Hedstrom 2002, Rawlings et al. 2010). Threonine peptidases have threonine residues as their nucleophiles during catalysis. For unknown peptidases, the active site residues have not yet been determined.

Evolutionary analyses have been applied to a broad range of studies, which include the identification of gene/protein families that have expanded in a specific lineage over evolutionary time and possibly indicate the existence of selective pressure (Irving et al. 2003, Sargeant et al. 2006, Nahum & Pereira 2008, Robinson et al. 2008, Wu et al. 2009, Huzurbazar et al. 2010). The availability of faster and more powerful computers combined with the development of automated pipelines has enabled the investigation of such evolutionary processes through the reconstruction of phylogenetic trees for the complete set of proteins encoded in a genome (known as phylome). The results obtained by this analysis provide a broad view of the evolution of an organism's genome and proteome, which allows for a deeper understanding of genomic complexity and lineage-specific adaptations (Huerta-Cepas et al. 2007, 2010b).

In a previous study, we described the reconstruction of the S. mansoni phylome to improve gene/protein functional annotation and provide insights into parasite's biology (phylomedb.org). By applying an automated pipeline, we also identified lineage-specific gene duplications, which may have led to a potential diversification of several protein families that are relevant for host-parasite interactions, such as tetraspanins, fucosyltransferases and sperm-coating protein-like proteins. Here, we explore the S. mansoni phylome data to analyse three endopeptidase families that expanded in this lineage since its diversification from 15 other metazoan species with the aim of contributing to the available knowledge of parasite biology and host-parasite interactions from an evolutionary perspective. The members of these families include leishmanolysins (metallopeptidase M8 family), cercarial elastases (serine peptidase S1 family) and cathepsin D proteins (aspartic peptidase A1 family).

The present paper is centred on two main research questions: (i) Did any peptidase families expand in the *S. mansoni* genome/proteome and if so, which ones? (ii) What are the evolutionary histories of these peptidase families? To address these questions, we used a so-called species-overlap algorithm (Huerta-Cepas et al. 2007) to detect lineage-specific duplications that occurred during the evolution of the parasite's genome. We also integrated information on sequence alignments, phylogenetic trees, protein architecture and the conservation of critical resi-

dues to characterise these proteins. Our results indicate that each peptidase family has a unique evolutionary history within/across the analysed species. Furthermore, our data support the hypothesis that gene duplication events followed by divergence is the main mechanism shaping the evolution of *S. mansoni*-specific paralogous groups.

The analysis of the evolutionary histories of these three *S. mansoni* families is relevant to functional genomics, evolutionary biology, medicine and biotechnology, especially taking into account the importance of *S. mansoni* peptidases in the development of schistosomiasis and that they have been described as promising vaccine and drug targets (McKerrow et al. 2006, Abdulla et al. 2007, Kasný et al. 2009).

### **MATERIALS AND METHODS**

Organisms and sequence data - The dataset of species selected for analysis includes eight invertebrates (Nematostella vectensis, Caenorhabditis elegans, Caenorhabditis briggsae, S. mansoni, Drosophila melanogaster, Anopheles gambiae, Bombyx mori and Strongylocentrotus purpuratus), one tunicate (Ciona intestinalis), one cephalochordate (Branchiostoma floridae), three vertebrates (Danio rerio, Mus musculus and Homo sapiens), three fungi (Neurospora crassa, Saccharomyces cerevisiae and Ustilago maydis) and one plant (Arabidopsis thaliana). Information on the selected taxa is provided as Supplementary data.

This dataset is particularly rich in metazoans (76% of the selected species) that cover important evolutionary innovations, for example, the origin of bilateral symmetry, the third germ layer, the development of organs, systems, complex patterns of communication and the emergence of the adaptive immune system, which makes it especially suitable for addressing the evolutionary innovations in *S. mansoni* in comparison with other metazoan species (phylomedb.org).

The *S. mansoni* predicted proteome dataset was downloaded from SchistoDB version 2.0 (schistodb.net) (Zerlotini et al. 2009). Proteomes derived from the 16 fully sequenced genomes were downloaded from the Broad Institute *Ustilago maydis* Database, Ensembl, Intergr8, JGI Genome Projects, National Center for Biotechnology Information Genome Database and SilkDB, which can be collectively accessed through the Genomes OnLine Database (genomesonline.org).

Endopeptidase protein families - Peptidases are hydrolases that act on peptide bonds [Enzyme Commission (EC) 3.4]. Three endopeptidase families were selected and analysed in detail in the present work. They include the metallopeptidase M8 family (EC 3.4.24.-), serine peptidase S1 family (EC 3.4.21.-) and aspartic peptidase A1 family (EC 3.4.23.-) members and belong to three peptidase clans (MA, PA and AA, respectively), as described in the MEROPS database (Rawlings et al. 2010).

Information on enzymes was collected from the literature and database references and included in the Supplementary data. The EC numbers were collected from the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology database, which is available online (chem.qmul.ac.uk/iubmb/enzyme/).

Alignments and phylogenetic trees - Sequence alignments and phylogenetic trees of the endopeptidase families selected for analysis were retrieved from the *S. mansoni* phylome data, which were reconstructed through a comparative analysis among all proteins encoded by the parasite genome and their potential homologs in 16 other eukaryotic species (phylomedb.org) (Huerta-Cepas et al. 2011).

Briefly, the *S. mansoni* phylome was reconstructed using each protein encoded in the *S. mansoni* genome ("seed" proteins) and the potential homologs identified through similarity-based searches (Smith & Waterman 1981) against the dataset of selected proteome data described above. The groups of homologous sequences were aligned using MUSCLE v3.6 (Edgar 2004) and gap-rich columns were filtered using trimAl (Capella-Gutiérrez et al. 2009). Phylogenetic analyses were performed using the neighbour-joining and maximum likelihood (ML) methods, as implemented in PhyML (Guindon & Gascuel 2003).

For the phylogenetic reconstruction of each "seed protein", we tested four different evolutionary models (JTT, WAG, BLOSUM62 and VT). In all cases, a discrete gamma-distribution model with four rate categories plus invariant positions was assumed with the gamma parameter and the fraction of invariant positions estimated from the data. Tree support values were computed using the approximate likelihood ratio test as implemented in PhyML (Guindon & Gascuel 2003, Anisimova & Gascuel 2006). The evolutionary model best fitting the data was determined by comparing the likelihood of the used models according to the Akaike Information Criterion (Akaike 1973). The resulting alignments, phylogenies and homology prediction can be accessed at PhylomeDB (phylomedb.org) (Huerta-Cepas et al. 2011) through protein sequence identifiers (e.g., UniProt: C4PZH6; SchistoDB: Smp 127030; PhylomeDB: Phy000V7EC SCHMA).

To integrate information from SchistoDB (Zerlotini et al. 2009) and PhylomeDB (Huerta-Cepas et al. 2011), we built a local relational database, named SchistoPhylomeSQL, which allowed us to extract and interpret the large amount of data in this work (Fig. 1). Access to this local database was implemented using DbVisualizer version 7.0.5 (dbvis.com). The SchistoPhylomeSQL database was the main resource for data mining in this work. In-house Perl scripts and Structured Query Language queries were used to parse data files during the database building and searching processes.

Paralogy and orthology relationships - To derive a complete catalogue of the paralogy and orthology relationships between *S. mansoni* proteins and those from other eukaryotic proteomes, we applied a "species-overlap" algorithm, as previously described (Huerta-Cepas et al. 2007). This algorithm uses the level of species overlap between the two daughter partitions of a given node to define it as a duplication or speciation event, which give rise to paralogs and orthologs, respectively. Once all the nodes have been classified, the algorithm establishes the paralogy and orthology relationships between the "seed

protein" and other proteins included in the phylogenetic tree, according to the original definition of these terms (Fitch 1970, Gabaldón 2008).

Lineage-specific duplications - Using a python Environment for Tree Exploration (Huerta-Cepas et al. 2010a), we analysed the *S. mansoni* phylome data (phylomedb. org) to identify protein families that were specifically expanded in the *S. mansoni* lineage since its diversification from the other selected taxa (Supplementary data). The duplication events defined by the "species-overlap" algorithm that only comprised paralogs from *S. mansoni* were considered lineage-specific duplications. In cases where more than one phylogenetic tree contained the same paralogous proteins, by changing only the "seed" protein position, the data were filtered to obtain a non-redundant list of in-paralogs.

Protein architecture and critical residues - In this study, we used the Pfam database (Finn et al. 2010) to identify the presence and organisation of protein sequence domains as well as critical residues present in the three S. mansoni endopeptidase families. Pfam is a large and widely used database of protein domains families. This database contains multiple sequence alignments and profile hidden Markov models (profile HMMs) for each protein family. Pfam-A entries are derived from the underlying sequence database, which is termed Pfamseq. This database is built from the most recent release of UniProtKB at a given time point (Finn et al. 2010, Apweiler et al. 2011). To predict active sites in new sequences. Pfam uses the information available in UniProtKB for homologous proteins, whose catalytic residues have been experimentally characterized (Mistry et al. 2007). Based on Pfam information, the illustrations of the S. mansoni protein domain architectures were generated using DOG 2.0 (Ren et al. 2009).

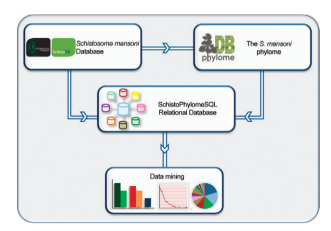


Fig. 1: flowchart of the applied methodology. The *Schistosoma mansoni* proteome data was retrieved from SchistoDB and each protein was used as "seed" to reconstruct the *S. mansoni* phylome. The resulting alignments, phylogenies, and homology predictions are available at PhylomeDB. To integrate information from SchistoDB and PhylomeDB, we built the SchistoPhylomeSQL, a local relational database as the main resource for data mining in this work.

#### **RESULTS**

Comparative genomics has revealed a great deal of sequence and/or functional diversity within and across organisms with respect to gene/protein family size, composition and the relatedness of their members (Huerta-Cepas et al. 2007, 2010b, Nahum et al. 2009, Andrade et al. 2011, Avelar et al. 2011). The rationale underlying the present work is that lineage-specific duplications may reflect molecular biodiversity and that the adaptation of organisms to different environments may ultimately help to identify potential therapeutic targets against parasitic diseases.

Our previous work identified lineage-specific gene duplications that led to the diversification of several families in *S. mansoni* (phylomedb.org). Furthermore, recent advances in proteomic analyses of schistosomes have revealed that peptidases are one of the main virulence factors involved in the pathogenesis of schistosomiasis (McKerrow et al. 2006, Kasný et al. 2009). In this work, we performed a phylogenomic analysis to address the two main questions of (i) whether peptidase families are expanded in the *S. mansoni* proteome and (ii) whether they share similar evolutionary histories.

Endopeptidase family members are duplicated in S. mansoni - To investigate which peptidase families are expanded in the S. mansoni genome, we explored the parasite phylome data available at PhylomeDB (Huerta-Cepas et al. 2011). Phylogenetic analyses were performed using an automated pipeline and a complete list of the paralogy relationships among the S. mansoni proteins was retrieved using a "species-overlap" algorithm that identifies family members originated by lineage-specific duplication events (Huerta-Cepas et al. 2007).

Based on the functional annotation available from the SchistoDB (Zerlotini et al. 2009) and UniProt (Apweiler et al. 2011) databases, the results revealed that the most significant peptidase expansions in the *S. mansoni* proteome corresponded to endopeptidases such as leishmanolysins, cercarial elastases and cathepsin D proteins. These enzymes belong to three distinct endopeptidase families, metallopeptidase M8 family (EC 3.4.24.-), serine peptidase S1 family (EC 3.4.21.-) and aspartic peptidase A1 family (EC 3.4.23.-), as described in the MEROPS database (Rawlings et al. 2010) and represent promising targets for vaccine and drug development.

In total, we identified 12 leishmanolysins, 13 cercarial elastases (Supplementary data) and 11 cathepsin D proteins (Supplementary data) in the predicted *S. mansoni* proteome. These proteins vary in length and sequence composition, but they are highly conserved with respect to the presence of a conserved sequence domain, which is distinct for each protein family as defined by the Pfam database (see details below). Currently, no crystal structure has been obtained for the *S. mansoni* peptidases described here.

Leishmanolysin (also called invadolysin) is a major surface peptidase member of the metallopeptidase M8 family. Leishmanolysins are believed to share the same mechanism used by the other zinc metalloproteinases, such as thermolysin. The conserved glutamate residue in the catalytic site acts in conjunction with a zinc ion

to deprotonate and activate a water molecule. In turn, the activated water molecule acts as a nucleophile to attack the carbonyl of the peptide bond of a variety of substrates (Macdonald et al. 1995, Schlagenhauf et al. 1998). In *Leishmania*, these proteins are involved in different types of processes, such as the inhibition or perturbations of host cell interactions and the degradation of the extracellular matrix (Fitzpatrick et al. 2009). These proteins may have similar activities in schistosomes. Indeed, the *S. mansoni* protein, SmPepM8 (Smp\_090100), is the second most abundant constituent in cercarial secretions, which provides insight on how it may contribute to tissue invasion by schistosomes and suggests this protein as a potential anti-parasitic target (Curwen et al. 2006, Fitzpatrick et al. 2009).

The catalytic triad of serine, histidine and aspartate residues is conserved in members of the serine protease family (Wilmouth et al. 2001, Hajjar et al. 2010). In elastases, this triad and an essential water molecule are involved in the catalysis. The peptide to be cleaved is bound noncovalently in the enzyme near the catalytic triad. In the first reaction step, the hydroxyl of the serine residue performs a nucleophilic attack on the substrate amide bond to form an ester. The amino terminus of the substrate is then covalently bound to the enzyme. The histidine residue abstracts a proton from a water molecule, which then attaches to the ester carbon to give rise to an oxyanion intermediate. Cercarial elastases play a key role in the penetration by the cercariae of mammalian skin to initiate infection and recent studies have revealed that these peptidases are also employed by the schistosomes to overcome or evade the host immune response (Salter et al. 2002, Aslam et al. 2008).

Cathepsin D is a member of the aspartic protease family. The active site of cathepsin D contains two aspartate residues, which perform an acid-base catalysis. This enzymatic mechanism involves the deprotonation of water by an ionised aspartate residue. This water molecule attacks the peptide carbonyl and there is a simultaneous protonation of the carbonyl oxygen by the other aspartate residue (e.g., Northrop 2001). Schistosome cathepsin D is involved in haemoglobin digestion, a process that provides the parasite with its main source of amino acid nutrients and that is essential for its development, growth and reproduction (Brindley et al. 2001, Caffrey et al. 2004, Delcroix et al. 2006). Given the essential function of cathepsin D in parasite nutrition and the ability of recombinant forms to cleave human immunoglobulin G, this protein is considered a potential target for novel anti-parasitic interventions (Verity et al. 2001, Morales et al. 2008).

The phylogenetic relationships of each endopeptidase family (Figs 2-4) are shown with protein sequences represented by identifiers in PhylomeDB (phylomedb. org) (Huerta-Cepas et al. 2011), UniProt (uniprot.org) (Apweiler et al. 2011) and/or SchistoDB (schistodb.net) (Zerlotini et al. 2009). In each phylogenetic tree, the *S. mansoni* endopeptidases form a well-supported clade of closely related proteins.

Together, the analysis of the *S. mansoni* proteome through an evolutionary approach identified endopeptidase family members that arose by gene duplication after

the divergence of this parasite from the other eukaryotic species studied in this work. These lineage-specific duplications are related to the parasite's biology and evolution.

Leishmanolysins (metallopeptidase M8 family) - Our pipeline identified 12 S. mansoni leishmanolysins (Supplementary data). Proteins Smp\_171330 and Smp\_171340 are located in the same genomic region of Smp\_090100 and Smp\_090110, respectively, and could not be retrieved from the UniProt (Apweiler et al. 2011) and GeneDB (genedb.org) databases, which suggests that these genes were incorrectly annotated and probably deleted from these databases. Similar findings were obtained in two previous studies (Berriman et al. 2009, Bos et al. 2009).

To reconstruct the evolutionary history of *S. man-soni* leishmanolysins and their homologs in selected taxa, we performed a sequence alignment of 32 protein sequences identified as potential homologs by our pipeline. The trimmed alignment contained 1,822 sites, which cover most of the conserved protein domain identified in these proteins.

By analysing the phylogenetic tree (Fig. 2), it is possible to demonstrate that *S. mansoni* leishmanolysins have homologs in most species analysed in the present work, with the exception of *C. intestinalis* (tunicata) and fungi. However, this result does not completely discard the presence of homologous proteins in other organisms because they may be very divergent from the others in the database and therefore be missed by the pipeline search. The same is true for the other protein families mentioned in this paper.

Based on the information available in the literature and curated databases, three leishmanolysin homologs have been experimentally confirmed in *D. melanogaster*, *M. musculus* and *H. sapiens* and their function is related to the coordination of mitotic progression and cell migration (for details see Supplementary data). Although predicted functions or experimental evidence are not yet available, the metallopeptidase M8 family is also expanded in the sea anemone (*N. vectensis*) and sea urchin (*S. purpuratus*). The metallopeptidase M8 family also has more paralogs in the schistosomes (12 proteins)

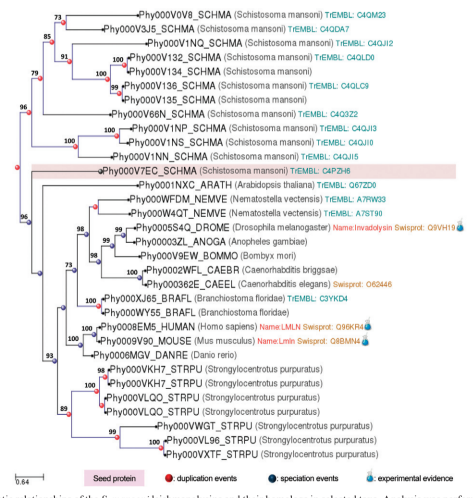


Fig. 2: phylogenetic relationships of the *S. mansoni* leishmanolysins and their homologs in selected taxa. Analysis was performed with trimmed sequence alignment by using the maximum likelihood method as implemented in PhyML. Best fit model (WAG) and support values for each node were estimated by the Akaike Likelihood Ratio Test (aLRT). Sequence labels follow the PhylomeDB internal identifier. For details, see Supplementary data.

compared to *H. sapiens* (4 proteins), which is in contrast with what is normally observed for the human peptidase families (Berriman et al. 2009).

A conserved protein domain (Pfam: PF01457), which characterises members of the metallopeptidase M8 family, was identified in all *S. mansoni* proteins analysed here (Fig. 5). Length variation and conservation of active sites were also observed. According to the Pfam profile HMMs, truncated domains were identified in all proteins, which possibly reflects the presence of different protein isoforms, as has been described elsewhere (Floris et al. 2008). The truncated domains could also indicate that parts of the sequences are missing at the N-terminal, C-terminal regions, or both due to annotation issues.

The data also reveals that the protein domain is duplicated in Smp\_167090, Smp\_167120 and Smp\_135530. Seven *S. mansoni* proteins (Smp\_090100, Smp\_090110, Smp\_127030, Smp\_135530, Smp\_153930, Smp\_167090 and Smp\_173070) were identified as active due to the presence of expected active site residues and metal ligand sites in the correct positions based on alignments with reference sequences, as previously described (Berriman et al. 2009).

Cercarial elastases (serine peptidase S1 family) - Our analysis identified a total of 13 cercarial elastases encoded in the *S. mansoni* genome (Supplementary data). Similar results were obtained by Berriman et al. (2009). However, with TreeFam (Ruan et al. 2008), Berriman et al. (2009) also

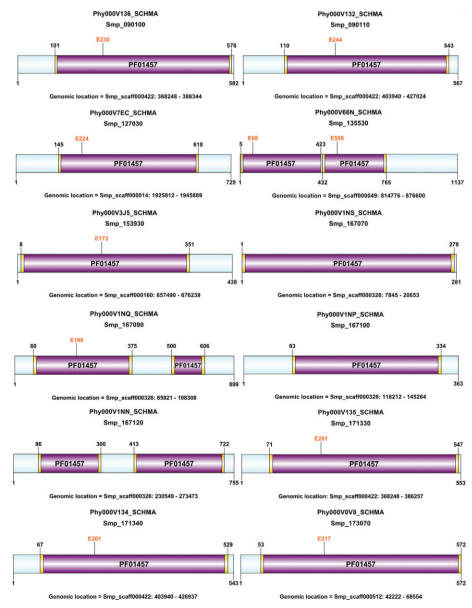


Fig. 3: conserved protein domain architecture of the *Schistosoma mansoni* leishmanolysins. Protein identifiers were assigned in SchistoDB. The conserved protein domain according to Pfam (PF01457) is present in all proteins. Truncated regions (yellow block) are indicated. Sequence length, domain limits, and active sites are also shown.

identified the Smp\_192850 protein, which is annotated as a hypothetical protein and only contains 69 amino acids.

Two proteins, Smp\_152560.2 and Smp\_056680.2, are encoded in the same genomic location and could not be recovered in UniProt (Apweiler et al. 2011). Searches for the former protein in GeneDB (genedb.org) retrieved only the latter (Smp\_056680), which indicated that the Smp\_152560.2 gene was improperly annotated and thus was eliminated from both databases. In the original version of the *S. mansoni* genome, some sequences were interpreted as isoforms and different gene models were constructed. However, further studies indicated that these were actually mistakes in the genome assembly/annotation due to low sequence coverage. In the new version of the parasite genome, which is to be released by the Wellcome Trust Sanger Institute (sanger.ac.uk), many of these sequences have been collapsed.

Whole amino acid sequences from 35 proteins were aligned and filtered to remove gap-rich columns as previously described. The trimmed alignment contains 583 sites, which cover the conserved protein domain.

The phylogenetic analysis of the *S. mansoni* elastases and their homologs in the other species included in this work was performed as already described. The parasite elastases form a well-supported monophyletic clade, which suggests that these proteins originated from a common ancestor by gene duplication events followed by divergence in the *Schistosoma* lineage.

In observing the resulting phylogeny (Fig. 3), it is possible to demonstrate that *S. mansoni* elastases have homologs in six of the 16 other species considered in this analysis (*N. vectensis*, *D. melanogaster*, *An. gambiae*, *B. floridae*, *M. musculus* and *H. sapiens*). The serine peptidase S1 family is also expanded in all of these species except for one, *D. melanogaster*. According to the information available in UniProt (Apweiler et al. 2011), seven homologs have been experimentally confirmed in *D. melanogaster*, *M. musculus* and *H. sapiens*, and their function is related to a digestive process and immune response (Supplementary data). It is believed that similar activities are performed by elastases in schistosomes (Salter et al. 2002, Aslam et al. 2008).

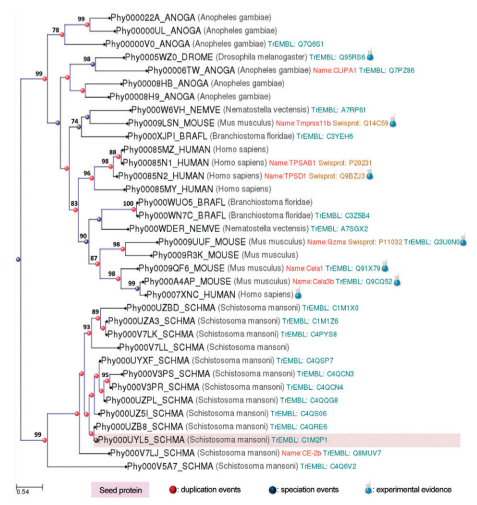


Fig. 4: phylogenetic relationships of the *Schistosoma mansoni* elastases and their homologs in selected taxa. Analysis was performed with trimmed sequence alignment by using the maximum likelihood method as implemented in PhyML. Best fit model (WAG) and support values for each node were estimated by the Akaike Likelihood Ratio Test (aLRT). Sequence labels follow the PhylomeDB internal identifier. For details, see Supplementary data.

A conserved protein sequence domain (Pfam: PF00089), which is found in all characterised members of the serine peptidase S1 family, was identified in the *S. mansoni* elastases and ranges in length from 141-265 amino acids (Fig. 6). The catalytic triad of histidine, aspartate and serine residues is present in most of these proteins. Based on profile HMMs available in Pfam, truncated regions were assigned to all 12 of these elastases, perhaps reflecting their degree of divergence in relation to other proteins in the database. Meanwhile, it is important to emphasise that protein databases do not cover all of the existing diversity in nature.

Together, these results indicate that the correct number of cercarial elastases encoded in the *S. mansoni* genome is 12 and not 13 as described before. However, only Smp\_006510, Smp\_006520 and Smp\_141450 were previously predicted as active proteins (Berriman et al. 2009). Smp\_194800 has a much shorter domain compared to others. This difference could reflect either the presence of an elastase pseudogene in the parasite genome or that the sequence was incorrectly annotated due to an error in the gene model. Considering that the first-pass annotation of the *S. mansoni* genome was produced by a combination of gene-finding algorithms (Augustus,

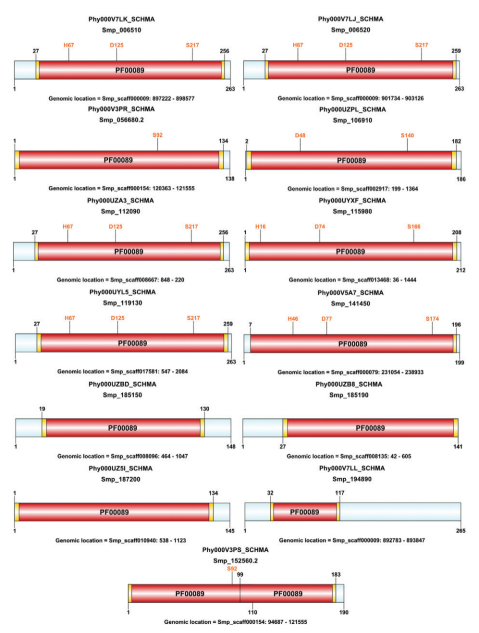


Fig. 5: conserved protein domain architecture of the *Schistosoma mansoni* elastases. Protein identifiers were assigned in SchistoDB. The conserved protein domain according to Pfam (PF00089) is present in all proteins. Truncated regions (yellow block) are indicated. Sequence length, domain limits, and the catalytic triad of histidine (H), aspartate (D), and serine (S) are also shown.

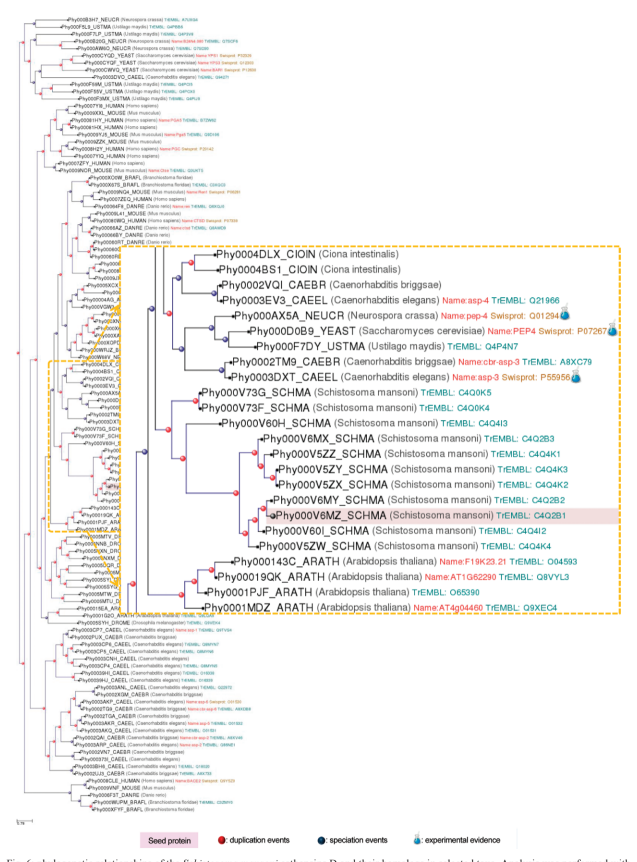


Fig. 6: phylogenetic relationships of the *Schistosoma mansoni* cathepsins D and their homologs in selected taxa. Analysis was performed with trimmed sequence alignment by using the maximum likelihood method as implemented in PhyML. Best fit model (WAG) and support values for each node were estimated by the Akaike Likelihood Ratio Test (aLRT). Sequence labels follow the PhylomeDB internal identifier. For details, see Supplementary data.

Twinscan and GlimmerHMM) (Berriman et al. 2009), this genome has not received extensive manual curation and therefore, many gene models will be refined in the future. Furthermore, EVidenceModeler (Haas et al. 2008) has also been used to incorporate expressed sequence tag (EST) evidence into the data.

Cathepsin D proteins (aspartic peptidase A1 family) - Our pipeline identified 11 S. mansoni cathepsin D proteins (Supplementary data) that were duplicated after the divergence of S. mansoni from the other metazoans analysed here. The evolutionary history of cathepsin D proteins was reconstructed from the sequence alignment of 111 protein sequences from S. mansoni and the selected taxa. The final trimmed alignment contained 1,676 sites, which covered most of the conserved protein domain (Pfam: PF00026). Two S. mansoni proteins corresponded to alternative splicing products (Smp\_136830.2 and Smp\_013040.2). Similar results were found by Berriman et al. (2009).

The phylogenetic tree indicates that the S. mansoni cathepsin D proteins have homologs in all but one species (S. purpuratus) analysed in this work (Fig. 4). The aspartic peptidase A1 family has also been expanded in 12 of the 15 species in which homologous proteins were identified (A. thaliana, U. maydis, S. cerevisiae, N. crassa, C. elegans, C. briggsae, D. melanogaster, C. intestinalis, B. floridae, D. rerio, M. musculus and H. sapiens). The number of paralogous proteins ranges from two-17 and includes different aspartic peptidases, such as pepsins, renins, gastricsin and cathepsin D proteins. Based on the information available in the literature and curated databases, these homologous proteins are involved in digestion and protein degradation (Supplementary data). In schistosomes, cathepsin D proteins play an integral role in haemoglobin proteolysis (Brindley et al. 2001, Caffrey et al. 2004, Delcroix et al. 2006).

To predict the protein domain architecture of S. mansoni cathepsin D proteins, we applied the same methodology as previously described. The conserved domain (Pfam: PF00026), which has been found in all characterised aspartic peptidase A1 family members, was also identified in the S. mansoni proteins with sequence lengths ranging from 94-430 amino acids (Fig. 7). Active sites are also indicated. Based on the profile HMMs available in Pfam, truncated regions were observed in the N-terminal, C-terminal or both regions. The data also indicate that an additional short sequence domain (Pfam: PF07966), which is known as the A1 propertide domain, is present at the N-terminal region of two S. mansoni proteins, Smp 013040.1 and Smp 013040.2. Smp 136840 has a much shorter domain compared to other proteins in the same family.

In a previous study, four *S. mansoni* cathepsin D proteins (Smp\_013040.1, Smp\_013040.2, Smp\_136730 and Smp\_136830.2) were identified as active proteins (Berriman et al. 2009), but the variation in the domain architecture and its implications in functional complexity were not investigated. One interesting study would be to analyse the functional properties of Smp\_013040.1 and Smp\_013040.2, which contain the A1 propeptide domain (PF07966).

#### **DISCUSSION**

We found that three endopeptidase families are expanded in the helminth parasite *S. mansoni*, which include members of the metallopeptidases (M8 family), serine peptidases (S1 family) and aspartic peptidases (A1 family). In this work, a comparative analysis of these three protein families in *S. mansoni* and 16 other eukaryotic proteomes revealed their distinct evolutionary histories and provided further information with respect to the sequence and functional features of the parasite family members.

Based on the *S. mansoni* genomic data, 335 peptidases were identified, which comprise 2.5% of the predicted proteome (Berriman et al. 2009). They include members of five major classes of peptidases (aspartic, cysteine, metallo, serine and threonine). Of the 61 peptidase families, 44 are expanded in this parasite and the number of paralogous proteins range from two-26.

Using a computational approach, Bos et al. (2009) analysed all putative peptidases encoded in the parasite's genome in addition to using EST data, which is similar to work by Berriman et al. (2009). After removing redundant sequences, inactive homologs, likely pseudogenes and sequences smaller than 100 amino acids from the dataset, they identified a total of 255 peptidase sequences from the five catalytic classes.

Our results are not fully comparable to those obtained by Bos et al. (2009) with respect to elastases and cathepsin D proteins. However, it is worth noting that the phylogenetic analysis of the serine peptidase S1 family performed by these authors also indicated a well-supported clade of four *S. mansoni* elastases, which are corroborated by our findings. The other homologs with high similarities to the cercarial elastases were likely pseudogenes and, for this reason, they were excluded from the analysis by Bos et al. (2009).

Our results suggest that *Schistosoma* members of these endopeptidase families originated from successive gene duplication events in the parasite lineage after its diversification from the other metazoans analysed here. These results were corroborated by previous proteomic and phylogenetic analyses on *Fasciola hepatica* peptidases, which showed that the repertoire of virulence-associated cathepsin L proteins was established by a series of gene duplication events (Irving et al. 2003, Robinson et al. 2008). These studies also indicate that the gene duplications were followed by active site residue refinements, which interfere with the substrate specificity of the *F. hepatica* cathepsin L proteins. Whether the *S. mansoni* proteins share a similar refinement remains to be established.

Gene duplication followed by divergence is known to be the most predominant mechanism of molecular evolution and represents the main source of raw material for the generation of new genes and proteins through the processes of neo and sub-functionalisation (Ohno 1970, Conant & Wolfe 2008, Nahum & Pereira 2008, Hamilton et al. 2009). Although in some cases sequences have diverged to the extent that it is impossible to recognise homologous relationships, different proteins that arose by gene duplication may be related at distinct levels, such as sequence, structure, function or a combination of these features and can be grouped into families and superfamilies (Nahum & Pereira 2008).

Gene fusion, gene fission and domain shuffling were not observed as mechanisms shaping the evolution of the *S. mansoni* endopeptidase families analysed in this work. Whether gene fusion/fission also plays a role in the evolution of the *S. mansoni* genome will be a subject of a future work. Our previous study indicated that domain shuffling is one of the main evolutionary forces driving the sequence and functional diversification of the protein kinases of this parasite (Andrade et al. 2011, Avelar et al. 2011).

Peptidases have been implicated in various processes that are crucial to the development and survival of parasites, including host invasion, degradation of haemoglobin in blood feeding, immune evasion and activation of inflammation (McKerrow et al. 2006, Kasný et al. 2009).

Experimental work suggests that the SmPepM8 metallopeptidase (leishmanolysin) may contribute to tissue invasion by schistosome cercariae. This peptidase was the second most abundant protein released during the transformation of *S. mansoni* cercariae into schistosomula (Curwen et al. 2006). Leishmanolysins are a major surface peptidase member of the metallopeptidase M8 family, which in leishmaniasis are involved in different types of processes, such as the inhibition or perturbation of host cell interactions and the degradation of the extracellular matrix (Fitzpatrick et al. 2009). It is speculated that these proteins could perform similar activities in schistosomes during host-parasite interactions (Curwen et al. 2006, Fitzpatrick et al. 2009).

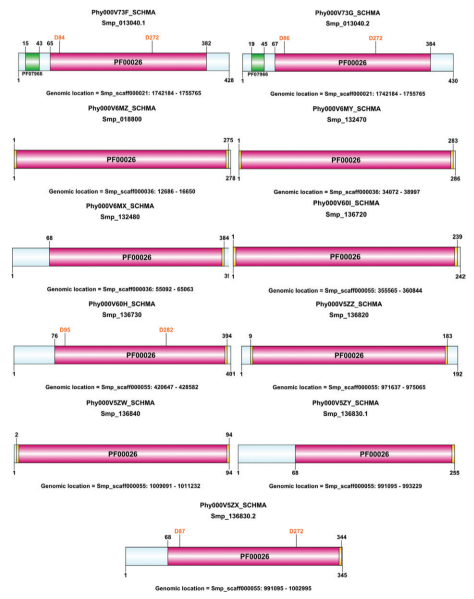


Fig. 7: conserved protein domain architecture of the *S. mansoni* cathepsin Ds. Protein identifiers were assigned in SchistoDB. The conserved protein domain according to Pfam (PF00026) is present in all proteins. The PF07966 additional N-terminal domain was identified in two proteins. Truncated regions (yellow block) are indicated. Sequence length, domain limits, and active sites are also shown.

Invasion of host skin is the initial event in establishing an infection in mammalian hosts. Considering the complexity of host skin barriers that the cercariae must go through during invasion, it has been suggested that multiple enzyme activities are required for this process (Salter et al. 2002). However, only one peptidase (cercarial elastase) has been identified as a major secretory product released during skin penetration (Knudsen et al. 2005, Hansell et al. 2008). These proteins may also be involved in eliminating the outer layer of the cercariae during transformation. Although cercarial elastases were named based on their ability to degrade insoluble elastin, numerous substrates for these enzymes have been identified, which include collagen, keratin and extracellular matrix proteins (Salter et al. 2002, McKerrow 2003, Knudsen et al. 2005).

Orthologous genes encoding elastase proteins were found in Schistosoma haematobium, Schistosoma japonicum and Schistosoma douthitti (Salter et al. 2002, Zhou et al. 2009). The expression of S. japonicum cercarial elastases was confirmed in both the sporocyst and cercarial stages and evidence that this peptidase is released by the parasite during the invasion of mammalian skin was obtained by anti-recombinant SiCE antibodies in infected mouse skin (Zhou et al. 2009). However, orthologous peptidases to S. mansoni cercarial elastases were not detected in the acetabular secretions of S. japonicum (Dvorák et al. 2008). Furthermore, the faster penetration by S. japonicum into the host skin may reflect the differential use of proteolytic enzymes in addition to those characterised in S. mansoni or even involve new peptidases not yet characterised (Chlichlia et al. 2005, He et al. 2005). Recent studies have also demonstrated that S. mansoni elastases are capable of cleaving IgE molecules from human, mouse and rat, indicating that the parasite may be able to overcome or evade the IgE response (Aslam et al. 2008). However, this subject remains controversial.

The biological complexity of *S. mansoni* is related to evolutionary innovations that took place before and after its diversification from other metazoans. Because duplicated genes are important substrates for improving an organism's adaptation to its environment, understanding how members of protein families evolved may link evolutionary studies to parasite biology. In turn, this knowledge will provide insights into host-parasite relationships and accelerate the identification of novel vaccine and drug targets aimed at the treatment and eradication of schistosomiasis.

In conclusion, this paper provides an evolutionary view of three *S. mansoni* peptidase families, thus allowing for a deeper understanding of the genomic complexity and lineage-specific adaptations potentially related to the parasitic lifestyle. In the future, our results obtained using a systemic approach (proteome-wide analyses) may accelerate the understanding of schistosomiasis, its etiologic agents and host-parasite interactions and optimise the discovery of therapeutic targets for the development of new drugs and vaccines.

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TABLE Selected taxa of leishmanolysins

Kingdom	Phylum	Subphylum	Code	Scientific name	NCBI_taxid	Proteome_data
Fungi	Ascomycota	Pezizomycotina	NEUCR	Neurospora crassa	5141	broadinstitute.org/annotation/genome/neurospora
Fungi	Ascomycota	Saccharomycotina	YEAST	Saccharomyces cerevisiae	4932	fungi.ensembl.org/Saccharomyces_cerevisiae
Fungi	Basidiomycota	Ustilaginomycotina	USTMA	Ustilago maydis	5270	broadinstitute.org/annotation/genome/ustilago_maydis
Metazoa	Arthropoda	-	DROME	Drosophila melanogaster	7227	metazoa.ensembl.org/Drosophila_melanogaster
Metazoa	Arthropoda	-	ANOGA	Anopheles gambiae	7165	metazoa.ensembl.org/Anopheles_gambiae
Metazoa	Arthropoda	-	BOMMO	Bombyx mori	7091	silkworm.genomics.org.cn
Metazoa	Chordata	Cephalochordata	BRAFL	Branchiostoma floridae	7739	genome.jgi-psf.org/Brafl1/Brafl1.home.html
Metazoa	Chordata	Craniata	DANRE	Danio rerio	7955	ensembl.org/Danio rerio
Metazoa	Chordata	Craniata	MOUSE	Mus musculus	10090	ensembl.org/Mus_musculus
Metazoa	Chordata	Craniata	HUMAN	Homo sapiens	9606	ensembl.org/Homo sapiens
Metazoa	Chordata	Tunicata	CIOIN	Ciona intestinalis	7719	ensembl.org/Ciona intestinalis
Metazoa	Cnidaria	-	NEMVE	Nematostella vectensis	45351	genome.jgi-psf.org/Nemve1/Nemve1.download.ftp.html
Metazoa	Echinodermata	-	STRPU	Strongylocentrotus purpuratus	7668	ncbi.nlm.nih.gov/projects/genome/guide/sea_urchin
Metazoa	Nematoda	-	CAEEL	Caenorhabditis elegans	6239	ensembl.org/Caenorhabditis elegans
Metazoa	Nematoda	-	CAEBR	Caenorhabditis briggsae	6238	metazoa.ensembl.org/Caenorhabditis_briggsae
Metazoa	Platyhelminthes	-	SCHMA	Schistosoma mansoni	6183	schistodb.net
Viridiplantae	Streptophyta	-	ARATH	Arabidopsis thaliana	3702	ebi.ac.uk/integr8

TABLE Functional annotation of leishmanolysins

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession	Original_ ID <sup>d</sup>	Lengthe	$Product^f$	Pfam_accession <sup>g</sup>	Function <sup>h</sup>	PubMed_ ID <sup>i</sup>	Organism <sup>j</sup>	Notes <sup>k</sup>
Phy000V136_SCHMA	Sch0003844	C4QLC9	Smp_090100	582 aa	SmPepM8; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V132_SCHMA	Sch0003840	C4QLD0	Smp_090110	567 aa	Metalloproteinase, putative; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V7EC_SCHMA	Sch0012282	C4PZH6	Smp_127030	729 aa	Protease family m8 leishmanolysin, putative; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V66N_SCHMA	Sch0010672	C4Q3Z2	Smp_135530	1,137 aa	Protease family m8 leishmanolysin, putative; Leishmanolysin-2 (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V3J5_SCHMA	Sch0007130	C4QDA7	Smp_153930	438 aa	Protease family m8 leishmanolysin, putative; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V1NS_SCHMA	Sch0004597	C4QJI0	Smp_167070	281 aa	Protease family m8 leishmanolysin, putative; Leishmanolysin-2 (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V1NQ_SCHMA	Sch0004595	C4QJI2	Smp_167090	699 aa	Protease family m8 leishmanolysin, putative; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V1NP_SCHMA	Sch0004594	C4QJI3	Smp_167100	363 aa	Expressed protein; Leishmanolysin-2 (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V1NN_SCHMA	Sch0004592	C4QJI5	Smp_167120	755 aa	Protease family m8 leishmanolysin, putative; Leishmanolysin-2 (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy000V135_SCHMA	Sch0003843	-	Smp_171330	553 aa	Protease family m8 leishmanolysin, putative	PF01457	-	-	Schistosoma mansoni	This sequence has genomic location overlap with Smp 090100.
Phy000V134_SCHMA	Sch0003842	-	Smp_171340	543 aa	Protease family m8 leishmanolysin, putative	PF01457	-	-	Schistosoma mansoni	This sequence has genomic location overlap with Smp_090110.
Phy000V0V8_SCHMA	Sch0003550	C4QM23	Smp_173070	572 aa	Protease family m8 leishmanolysin, putative; Invadolysin (M08 family)	PF01457	Predicted function	19606141	Schistosoma mansoni	-
Phy00003ZL_ANOGA	Aga0005169	Q7QD48	ENSANGP00000010959	621 aa	-	PF01457	Predicted function	12364791	Anopheles gambiae	-
Phy0001NXC_ARATH	Ath0029631	Q67ZD0	Q67ZD0	841 aa	Major surface like glycoprotein	PF07974; PF01457	Predicted function	11130714	Arabidopsis thaliana	-
Phy000V9EW_BOMMO	Bom0001650	-	BGIBMGA001650-PA	568 aa	-	-	-	-	Bombyx mori	-

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession <sup>c</sup>	$rac{ ext{Original}\_{ ext{ID}^d}$	Length <sup>e</sup>	Product <sup>f</sup>	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	PubMed_ ID <sup>i</sup>	Organism <sup>j</sup>	Notes <sup>k</sup>
Phy000WY55_BRAFL	Bra0014763	-	prot14763	1,029 aa	-	-	-	-	Branchiostoma floridae	-
Phy000XJ65_BRAFL	Bra0042485	C3YKD4	prot42485	585 aa	Putative uncharacterized protein	PF01457	Predicted function	18563158	Branchiostoma floridae	-
Phy0002WFLCAEBR	Cbr0010832	Q61YG1	Q61YG1	663 aa	Leishmanolysin-like peptidase	PF01457	Predicted function	14624247	Caenorhabditis briggsae	-
Phy000362E_CAEEL	Cel0010125	O62446	Y43F4A.1a	663 aa	Leishmanolysin-like peptidase	PF01457	Predicted function	9851916	Caenorhabditis elegans	-
Phy0005S4Q_DROME	Dme0007085	Q9VH19	CG3953-PA	683 aa	Leishmanolysin-like peptidase; Invadolysin	PF01457	Experimental evidence	15557119, 10731132, 12537572, 12537569	Drosophila melanogaster	-
Phy0006MGV_DANRE	Dre0029534	B0S5M4	ENSDARP00000059717	618 aa	Novel protein similar to human and mouse leishmanolysin-like (metallopeptidase M8 family)	-	-	-	Danio rerio	-
Phy0008EM5_HUMAN	Hsa0022469	Q96KR4	ENSP00000328829	665 aa	Leishmanolysin-like peptidase	PF01457	Experimental evidence	16641997, 15557119	Homo sapiens	-
Phy0009V90_MOUSE	Mms0018458	Q8BMN4	ENSMUSP00000023497	681 aa	Leishmanolysin-like peptidase	PF01457	Experimental evidence	16141072, 15489334	Mus musculus	-
Phy000W4QT_NEMVE	Nem0003341	A7ST90	prot3341	563 aa	Predicted protein	PF01457	Predicted function	17615350	Nematostella vectensis	-
Phy000WFDM_NEMVE	Nem0017544	A7RW33	prot17544	624 aa	Predicted protein	PF01457	Predicted function	17615350	Nematostella vectensis	-
Phy000VKH7_STRPU	Str0001413	-	XP_001186658.1	510 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VL96_STRPU	Str0002463	-	XP_001189142.1	1,096 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VLQO_STRPU	Str0003126	-	XP_001203887.1	529 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VLQO_STRPU	Str0008229	-	XP_785477.2	529 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VWGT_STRPU	Str0021529	-	XP_001193341.1	265 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VXTF_STRPU	Str0025031	-	XP_001192223.1	1,015 aa	-	-	-	-	Strongylocentrotus purpuratus	-
Phy000VKH7_STRPU	Str0041478	-	XP_001180406.1	510 aa	-	-	-	-	Strongylocentrotus purpuratus	-

a: new internal identifier in PhylomeDB; b: old internal identifier in PhylomeDB; c: UniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: functional annotation in SchistoDB and UniProt; g: protein sequence domain(s) identified in the Pfam database; h: functional information in the literature available in UniProt; i: identifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: functional annotation in SchistoDB and UniProt; i: identifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database; h: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database; h: scientifier in PhylomeDB; c: uniProt accession number; d: original identifier in the database; h: scientifier in PhylomeDB; d: original ident

TABLE Elastases

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession		Length <sup>e</sup>	Product	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	PubMed_ ID <sup>i</sup>	Organism <sup>i</sup>	Notes <sup>k</sup>
Phy000UYL5_SCHMA	Sch0000377	C1M2P1	Smp_119130	263 aa	Elastase 1a, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UYXF_SCHMA	Sch0000833	C4QSP7	Smp_115980	212 aa	Cercarial elastase HP1, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UZ5I_SCHMA	Sch0001129	C4QS06	Smp_187200	145 aa	Tryptase gamma precursor, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UZA3_SCHMA	Sch0001310	C1M1Z6	Smp_112090	263 aa	Elastase 2a, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UZB8_SCHMA	Sch0001364	C4QRE6	Smp_185190	141 aa	Elastase 1a, putative	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UZBD_SCHMA	Sch0001369	C1M1X0	Smp_185150	148 aa	Cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000UZPL_SCHMA	Sch0002010	C4QQG8	Smp_106910	186 aa	Elastase 1b, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000V3PR_SCHMA	Sch0007382	C4QCN4	Smp_056680.2	138 aa	Cercarial elastase (S01 family); tryptase precursor, putative	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000V3PS_SCHMA	Sch0007383	-	Smp_152560.2	190 aa	Tryptase precursor, putative	PF00089	Predicted function	19606141	Schistosoma mansoni	This sequence has genomic location overlap with Smp_056680.2.
Phy000V5A7_SCHMA	Sch0009474	C4Q6V2	Smp_141450	199 aa	Serine protease, putative; subfamily S1A unassigned peptidase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000V7LJ_SCHMA	Sch0012547	-	Smp_006520	263 aa	Elastase 2b, putative	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000V7LK_SCHMA	Sch0012548	C4PYS8	Smp_006510	263 aa	Serine protease, putative; cercarial elastase (S01 family)	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy000V7LL_SCHMA	Sch0012549	-	Smp_194890	265 aa	Elastase, truncated protein, possible pseudogene	PF00089	Predicted function	19606141	Schistosoma mansoni	-
Phy00000UL_ANOGA	Aga0001101	A7UTT9	ENSANGP00000021783	305 aa	- · ·	PF00089	Predicted function	12364791	Anopheles gambiae	-
Phy00000V0_ANOGA	Aga0001116	Q7Q6S1	ENSANGP00000015859	275 aa	-	-	Predicted function	12364791	Anopheles gambiae	-
-	Aga0002674	Q7QJ37	ENSANGP00000016585	240 aa	-	PF00089	Predicted function	12364791	Anopheles gambiae	-

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession <sup>c</sup>	Original_ID <sup>d</sup>	Lengthe	Product/	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	$\frac{\text{PubMed}\_}{\text{ID}^i}$	Organism <sup>/</sup>	Notes <sup>k</sup>
Phy00006TW_ANOGA	Aga0008852	Q7PZ86	ENSANGP00000020262	440 aa	CLIP-domain serine protease subfamily A	PF00089	Predicted function	12364791	Anopheles gambiae	-
Phy00008H9_ANOGA	Aga0010989	Q7PWE3	ENSANGP00000025294	248 aa	-	-	Predicted function	12364791	Anopheles gambiae	-
Phy00008HB_ANOGA	Aga0010991	-	ENSANGP00000006368	365 aa	-	-	-	-	Anopheles gambiae	-
hy000WN7C_BRAFL	Bra0000495	C3Z5B4	prot495	262 aa	Putative uncharacterized protein	PF00089	Predicted function	18563158	Branchiostoma floridae	-
hy000WUO5_BRAFL	Bra0010245	-	prot10245	218 aa	-	-	-	-	Branchiostoma floridae	-
hy000XJPI_BRAFL	Bra0043201	СЗҮЕН6	prot43201	236 aa	Putative uncharacterized protein	PF00089	Predicted function	18563158	Branchiostoma floridae	-
hy0005WZ0_DROME	Dme0013359	Q95RS6	CG40160-PA	421 aa	-	PF00089	Experimental evidence	10731132, 12537568, 12537574, 12537573, 12537572, 16110336, 17569856, 17569867	Drosophila melanogaster	-
ny0007XNC_HUMAN	Hsa0000480	P08861	ENSP00000338369	270 aa	Chymotrypsin-like elastase family member 3B; elastase IIIB; elastase-3B	PF00089	Experimental evidence	2826474, 14702039, 16710414, 15489334, 3477287, 2675835, 3178837, 2753124, 2737288	Homo sapiens	-
hy00085MY_HUMAN	Hsa0010834	-	ENSP00000234798	321 aa	-	-	-	-	Homo sapiens	-
ny00085MZ_HUMAN	Hsa0010835	-	ENSP00000344083	274 aa	-	-	-	-	Homo sapiens	-
hy00085N1_HUMAN	Hsa0010837	-	ENSP00000343577	275aa	-	-	-	-	Homo sapiens	-
ny00085N2_HUMAN	Hsa0010838	Q9BZJ3	ENSP00000211076	242 aa	Tryptase delta; mast cell mMCP-7-like; tryptase-3	PF00089	Experimental evidence	9920877, 11174199, 11157797, 15616553, 18854315, 12391231	Homo sapiens	-
hy0009LSN_MOUSE	Mms0006205	Q14C59	ENSMUSP00000042406	416 aa	Transmembrane protease serine 11B; airway trypsin-like protease 5	PF00089, PF01390	Experimental evidence	16141072, 15489334, 15328353	Mus musculus	-
· -	Mms0012200	Q91X79	ENSMUSP00000023775	266 aa	Elastase 1, pancreatic	PF00089	Experimental evidence	15489334, 12040188	Mus musculus	-
_	Mms0013078	-	ENSMUSP00000015576		-	-	-	-	Mus musculus	-
hy0009UUF_MOUSE	Mms0017933	P11032	ENSMUSP00000023897	260 aa	Granzyme A	PF00089	Experimental evidence	2976140, 1639378, 1460043, 15489334, 3292396, 2422755, 3555842, 3533635, 3260181	Mus musculus	-
hy000A4AP_MOUSE	Mms0030183	Q9CQ52	ENSMUSP00000024200	269 aa	Elastase 3, pancreatic	PF00089	Experimental evidence	10349636, 16141073, 11042159, 11076861, 15489334	Mus musculus	-
hy000W6H_NEMVE	Nem0006289	A7RP61	prot6289	252 aa	Predicted protein	PF00089	Predicted function	17615350	Nematostella vectensis	-
hy000WDER_NEMVE	Nem0014980	A7SGX2	prot14980	299 aa	Predicted protein	PF00089, PF00629	Predicted function	17615350	Nematostella vectensis	-

a: new internal identifier in PhylomeDB; b: old internal identifier in PhylomeDB; c: UniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: functional annotation in SchistoDB and UniProt; g: protein sequence domain(s) identified in the Pfam database; h: functional information in the literature available in UniProt; i: identifier in PhylomeDB; j: scientific name of source organism; k: notes from this work.

TABLE
Aspartic peptidases analyzed in this paper

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession		Length <sup>e</sup>	Product <sup>f</sup>	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	${\tt PubMed\_ID}^i$	Organism <sup>/</sup>	Notes <sup>k</sup>
Phy000V5ZW_SCHMA	Sch0010420	C4Q4K4	Smp_136840	94 aa	Cathepsin d, putative; subfamily A1A unassigned peptidase	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V5ZX_SCHMA	Sch0010421	C4Q4K2	Smp_136830.2	345 aa	Cathepsin d, putative; subfamily A1A unassigned peptidase (A01 family) EMBL CAZ30383.1	PF00026	Predicted function	19606141	Schistosoma mansoni	Alternative splicing product.
Phy000V5ZY_SCHMA	Sch0010422	C4Q4K3	Smp_136830.1	255 aa	Cathepsin d, putative; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V5ZZ_SCHMA	Sch0010423	C4Q4K1	Smp_136820	192 aa	Aspartic proteinase-related; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V60H_SCHMA	Sch0010441	C4Q4I3	Smp_136730	401 aa	Cathepsin D2-like protein; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V60I_SCHMA	Sch0010442	C4Q4I2	Smp_136720	242 aa	Aspartic proteinase-related; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V6MX_SCHMA	Sch0011276	C4Q2B3	Smp_132480	393 aa	Aspartyl proteases, putative; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V6MY_SCHMA	Sch0011277	C4Q2B2	Smp_132470	286 aa	Aspartyl proteases, putative; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V6MZ_SCHMA	Sch0011278	C4Q2B1	Smp_018800	278 aa	Aspartyl proteases, putative; subfamily A1A unassigned peptidase (A01 family)	PF00026	Predicted function	19606141	Schistosoma mansoni	-
Phy000V73F_SCHMA	Sch0011883	C4Q0K4	Smp_013040.1	428 aa	Aspartyl proteases, putative; cathepsin D (A01 family)		Predicted function	19606141	Schistosoma mansoni	-
Phy000V73G_SCHMA	Sch0011884	C4Q0K5	Smp_013040.2	430 aa	Aspartyl proteases, putative; cathepsin D (A01 family)	PF07966,	Predicted function	19606141	Schistosoma mansoni	Alternative splicing product.
Phy00004AG_ANOGA	Aga0005560	Q7PTQ9	ENSANGP00000013568	389 aa	-	PF00026	Predicted function	12364791	Anopheles gambiae	-
Phy000143C_ARATH	Ath0003927	O04593	O04593	433 aa	-	PF00026, PF05184, PF03489	-	-	Arabidopsis thaliana	-
Phy00015EA_ARATH	Ath0005617	O65453	O65453	336 aa	Aspartic proteinase like protein	PF00026	Predicted function	10617198	Arabidopsis thaliana	-

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession <sup>c</sup>	$ \begin{array}{c} \text{Original}_{-} \\ \text{ID}^{d} \end{array} $	Lengthe	Product/	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	PubMed_ID <sup>i</sup>	Organism <sup>/</sup>	Notes <sup>k</sup>
Phy00019QK_ARATH	Ath0011243	Q8VYL3	Q8VYL3	513 aa	-	PF00026, PF05184, PF03489	Predicted function	11130712	Arabidopsis thaliana	-
Phy0001G2O_ARATH	Ath0019455	Q9LQA9	Q9LQA9	375 aa	-	PF00026	-	-	Arabidopsis thaliana	-
Phy0001MDZ_ARATH	Ath0027638	Q9XEC4	Q9XEC4	508 aa	Putative aspartic proteinase	PF00026, PF05184, PF03489	Predicted function	10617198	Arabidopsis thaliana	-
Phy0001PJF_ARATH	Ath0031722	O65390	O65390	506 aa	Putative aspartic proteinase	PF00026, PF05184, PF03489	Experimental evidence	11130712; 12093376	Arabidopsis thaliana	-
Phy000VGW3_BOMMO	Bom0011344	-	BGIBMGA011344-PA	326 aa	-	-	-	-	Bombyx mori	-
Phy000WP82_BRAFL	Bra0003157	-	prot3157	423 aa	-	-	-	-	Branchiostoma floridae	-
Phy000WRJZ_BRAFL	Bra0006192	C3YUT2	prot6192	388 aa	Putative uncharacterized protein	PF07966, PF00026	Predicted function	18563158	Branchiostoma floridae	-
Phy000WUPM_BRAFL	Bra0010298	C3ZMY0	prot10298	493 aa	Putative uncharacterized protein	PF00026	Predicted function	18563158	Branchiostoma floridae	-
Phy000X67S_BRAFL	Bra0025329	C3XQC3	prot25329	392 aa	Putative uncharacterized protein	PF00026	Predicted function	18563158	Branchiostoma floridae	-
Phy000X6BF_BRAFL	Bra0025469	C3YBT8	prot25469	423 aa	Putative uncharacterized protein	PF07966, PF00026	Predicted function	18563158	Branchiostoma floridae	-
Phy000XA8D_BRAFL	Bra0030645	-	prot30645	488 aa	-	-	-	-	Branchiostoma floridae	-
Phy000XFYF_BRAFL	Bra0038243	-	prot38243	439 aa	-	-	-	-	Branchiostoma floridae	-
Phy000XNWK_BRAFL	Bra0048871	-	prot48871	398 aa	-	-	-	-	Branchiostoma floridae	-
Phy000XO0W_BRAFL	Bra0049039	-	prot49039	243 aa	-	-	-	-	Branchiostoma floridae	-
Phy000XOPD_BRAFL	Bra0050000	-	prot50000	387 aa	-	-	-	-	Branchiostoma floridae	-
Phy0002PUX_CAEBR	Cbr0002312	-	Q60TT2	393 aa	-	-	-	-	Caenorhabditis briggsae	-
Phy0002QAI_CAEBR	Cbr0002873	A8XV46	Q60W85	704 aa	CBR-ASP-2 protein; Cbr-asp-2	PF07966, PF00026	Predicted function	14624247	Caenorhabditis briggsae	-
Phy0002TG9_CAEBR	Cbr0006968	A8XDB8	Q61GA5	389 aa	CBR-ASP-6 protein; Cbr-asp-6	PF00026	Predicted function	14624247	Caenorhabditis briggsae	-
Phy0002TGA_CAEBR	Cbr0006969	-	Q61GA7	-	-	-	-	-	Caenorhabditis briggsae	-
Phy0002TM9_CAEBR	Cbr0007184	A8XC79	Q61H49	397 aa	CBR-ASP-3 protein	PF00026	Predicted function	14624247	Caenorhabditis briggsae	-

PhylomeDB_ new <sup>a</sup>	PhylomeDB_ old <sup>b</sup>	UniProt_ accession <sup>c</sup>	$\begin{array}{c} \text{Original}\_\\ \text{ID}^d \end{array}$	Length <sup>e</sup>	Product <sup>f</sup>	Pfam_ accession <sup>g</sup>	Function <sup>h</sup>	PubMed_ID <sup>/</sup>	Organism <sup>j</sup>	Notes <sup>k</sup>
Phy0002UJ3_CAEBR	Cbr0008366	A8X733	Q61MA1	428 aa	Putative uncharacterized protein	PF00026	Predicted function	14624247	Caenorhabditis briggsae	-
Phy0002VN7_CAEBR	Cbr0009810	-	Q61U20	393 aa	-	-	-	-	Caenorhabditis briggsae	-
Phy0002VQI_CAEBR	Cbr0009929	-	Q61UL2	446 aa	-	-	-	-	Caenorhabditis briggsae	-
Phy0002XGM_CAEBR	Cbr0012165	-	Q624R1	386 aa	-	-	-	-	Caenorhabditis briggsae	-
Phy000373I_CAEEL	Cel0011461	-	C11D2.2	394 aa	-	-	-	-	Caenorhabditis elegans	-
Phy00039HI_CAEEL	Cel0014557	O16338	F59D6.3	474 aa	Putative uncharacterized protein	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy00039HJ_CAEEL	Cel0014558	O16339	F59D6.2	380 aa	Putative uncharacterized protein	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003AKP_CAEEL	Cel0015968	O01530	F21F8.7.1	389 aa	Aspartic protease 6; asp-6	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003AKQ_CAEEL	Cel0015969	O01531	F21F8.4.1	395 aa	Putative uncharacterized protein	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003AKR_CAEEL	Cel0015970	O01532	F21F8.3.2	393 aa	Aspartyl protease protein 5; asp-5	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003ANL_CAEEL	Cel0016072	Q22972	F28A12.4.1	389 aa	Putative uncharacterized protein F28A12.4	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003ARP_CAEEL	Cel0016220	Q86NE1	T18H9.2a	635 aa	Aspartyl protease protein 2; asp-2	PF07966, PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003BH6_CAEEL	Cel0017137	Q18020	C15C8.3	428 aa	-	PF00026	Experimental evidence	17761667, 9851916	Caenorhabditis elegans	-
Phy0003CNH_CAEEL	Cel0018660	-	ZK384.3	638 aa	-	-	-	-	Caenorhabditis elegans	-
Phy0003CP4_CAEEL	Cel0018719	Q8MYN5	Y39B6A.24.1	391 aa	-	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003CP5_CAEEL	Cel0018720	Q8MYN6	Y39B6A.23	395 aa	-	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003CP6_CAEEL	Cel0018721	Q8MYN7	Y39B6A.22	390 aa	-	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003CP7_CAEEL	Cel0018722	Q9TVS4	Y39B6A.20.4	396 aa	Aspartic protease 1; aps-1	PF07966, PF00026	Experimental evidence	10854422, 9851916,	Caenorhabditis elegans	-
Phy0003DVO_CAEEL	Cel0020251	Q94271	K10C2.3	410 aa	Putative uncharacterized protein	PF00026	Predicted function	9851916	Caenorhabditis elegans	-
Phy0003DXT_CAEEL	Cel0020328	P55956	H22K11.1	398 aa	Aspartic protease 3	PF00026	Experimental evidence	9851916, 9150941, 17761667	Caenorhabditis elegans	-
Phy0003EV3_CAEEL	Cel0021526	Q21966	R12H7.2	444 aa	Protein R12H7.2, confirmed by transcript evidence EMBL CAA90633.1; asp-4	PF07966, PF00026	Predicted function	9851916	Caenorhabditis elegans	-

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Phy0004BS1_CIOIN	Cin0006471	-	ENSCINP00000010874	430 aa	-	-	-	-	Ciona intestinalis	-
Phy0004DLX_CIOIN	Cin0008843	-	ENSCINP00000014585	400 aa	-	-	-	-	Ciona intestinalis	-
Phy0005MTU_DROME	Dme0000213	Q9VQ11	CG31928-PA	418 aa	-	PF00026	Predicted function	10731132, 12537572	Drosophila melanogaster	-
Phy0005MTV_DROME	Dme0000214	Q9VQ12	CG33128-PA	405 aa	-	PF00026	Predicted function	10731132, 12537572	Drosophila melanogaster	-
Phy0005MTW_DROME	Dme0000215	Q9VQ13	CG31926-PA	410 aa	-	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005MTX_DROME	Dme0000216	Q9VQ14	CG31661-PA	393 aa	-	PF00026	Predicted function	10731132, 12537572	Drosophila melanogaster	-
Phy0005NNB_DROME	Dme0001274	Q9VLK3	CG13095-PA	372 aa	Beta-site APP-cleaving enzyme; bace	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005NXM_DROME	Dme0001645	Q9VKP7	CG6508-PA	423 aa	-	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005NXN_DROME	Dme0001646	Q9VKP6	CG17134-PA	391 aa	-	PF00026	Predicted function	10731132, 12537572	Drosophila melanogaster	-
Phy0005SYG_DROME	Dme0008155	Q9VEK5	CG17283-PA	465 aa	-	PF00026	Predicted protein	12537572, 10731132	Drosophila melanogaster	-
Phy0005SYH_DROME	Dme0008156	Q9VEK4	CG5860-PA	370 aa	-	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005SYI_DROME	Dme0008157	Q9VEK3	CG5863-PA	395 aa	-	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005UQR_DROME	Dme0010470	Q9W5G3	CG13374-PA	407 aa	Pepsinogen-like; pcl	PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005XCX_DROME	Dme0013860	Q7K485	CG1548-PA	392 aa	Cathepsin D	PF07966, PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy0005Y92_DROME	Dme0015017	A1Z9Q9	CG10104-PA	404 aa	-	PF07966, PF00026	Predicted function	12537572, 10731132	Drosophila melanogaster	-
Phy00060Q9_DANRE	Dre0001360	-	ENSDARP00000055495	395 aa	-	-	-	-	Danio rerio	-
Phy00060RL_DANRE	Dre0001408	-	ENSDARP00000012342	412 aa	-	-	-	-	Danio rerio	-
Phy00060RT_DANRE	Dre0001416	-	ENSDARP00000013587	200 aa	-	-	-	-	Danio rerio	-
Phy00064F8_DANRE	Dre0006147	Q6XQJ0	ENSDARP00000061335	395 aa	Renin	PF07966, PF00026	Predicted function	14645735	Danio rerio	-
Phy00066AZ_DANRE	Dre0008586	Q8AWD9	ENSDARP00000061373	398 aa	Ctsd protein	PF07966, PF00026	-	-	Danio rerio	-
Phy00066BY_DANRE	Dre0008621	-	ENSDARP00000061099	395 aa	-	-	-	-	Danio rerio	-
Phy0006F3T_DANRE	Dre0019992	-	ENSDARP00000003409	503 aa	-	-	-	-	Danio rerio	-
Phy0007YI8_HUMAN	Hsa0001592	-	ENSP00000304306	307 aa	-	-	-	-	Homo sapiens	-
Phy0007YIQ_HUMAN	Hsa0001610	-	ENSP00000329601	391 aa	-	-	-	-	Homo sapiens	-
Phy0007ZEQ_HUMAN	Hsa0002762	-	ENSP00000272190	406 aa	-	-	-	-	Homo sapiens	-

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Phy0007ZFY HUMAN	Hsa0002806	_	ENSP00000354337	401 aa	-	_	_	-	Homo sapiens	_
Phy00080WQ_HUMAN	Hsa0004706	P07339	ENSP00000236671	412 aa	Cathepsin D	PF07966, PF00026	Experimental evidence	3927292, 3588310, 2069717, 15489334, 8262386, 7935485, 12643545, 12754519, 16335952, 16670177, 17081065, 16263699, 19159218, 8467789, 8393577, 10716266, 16685649	Homo sapiens	-
Phy00081HX_HUMAN	Hsa0005469	-	ENSP00000322192	388 aa	-	-	-	-	Homo sapiens	-
Phy00081HY_HUMAN	Hsa0005470	B7ZW62	ENSP00000309542	388 aa	Pepsinogen 5, group I (Pepsinogen A)	PF07966, PF00026	Predicted function	15489334	Homo sapiens	-
Phy00089N3_HUMAN	Hsa0016023	-	ENSP00000253720	445 aa	-	-	-	-	Homo sapiens	-
Phy00089N4_HUMAN	Hsa0016024	O96009	ENSP00000253719	420 aa	Napsin-A	PF00026	Experimental evidence	9877162, 10580105, 10591213, 15489334	Homo sapiens	-
Phy0008CLE_HUMAN	Hsa0019850	Q9Y5Z0	ENSP00000332979	518 aa	Beta-secretase 2; aspartyl protease 1; memapsin-1	PF00026	Experimental evidence	10591213, 10838186, 10965118, 10683441, 10749877, 11083922, 10677483, 12975309, 14702039, 10830953, 15489334, 15857888, 11423558, 16816112, 16305800, 11316808, 16965550	Homo sapiens	-
Phy0008H2Y_HUMAN	Hsa0025666	P20142	ENSP00000211310	388 aa	Gastricsin	PF07966, PF00026	Experimental evidence	3335549, 2909526, 2567697, 14574404, 15489334, 2515193, 6816595, 7714902, 9406551	Homo sapiens	-
Phy0009JXB_MOUSE	Mms0003781	O09043	ENSMUSP00000002274	419 aa	Napsin-A	PF00026	Experimental evidence	9013890, 11082205, 15489334, 16944957	Mus musculus	-
Phy0009L41_MOUSE	Mms0005319	-	ENSMUSP00000008035	-	-	-	-	-	Mus musculus	-
Phy0009NOR_MOUSE	Mms0008657	Q3UKT5	ENSMUSP00000073072	397 aa	Cathepsin E	PF07966, PF00026	Experimental evidence	10349636, 16141073, 11042159, 11076861, 12040188	Mus musculus	-
Phy0009NQ4_MOUSE	Mms0008706	P06281	ENSMUSP00000000788	402 aa	Renin-1	PF07966, PF00026	Experimental evidence	6370686, 2685761, 2691339, 16141072, 15489334, 6089205, 6392850, 9030738, 6327270	Mus musculus	-
Phy0009VNF_MOUSE	Mms0018977	Q9JL18	ENSMUSP00000043918	514 aa	-	-	Experimental evidence (PubMed ID: 16141072, 15489334, 10683441)	-	Mus musculus	-
Phy0009XXL_MOUSE	Mms0021935	-	ENSMUSP00000077032	381 aa	-	-	-	-	Mus musculus	-
Phy0009YJ5_MOUSE	Mms0022711	Q9D106	ENSMUSP00000025647	387 aa	Pepsinogen 5, group I	PF07966, PF00026	Experimental evidence	10349636, 16141073, 11042159, 11076861, 15489334,	Mus musculus	-
Phy0009ZZK_MOUSE	Mms0024598	-	ENSMUSP00000024782	395 aa	-	-	-	-	Mus musculus	-
Phy000AW6O_NEUCR	Ncr0000966	Q7SD30	(NCU00994.2)	434 aa	Endothiapepsin	PF00026	Predicted function	12712197	Neurospora crassa	-
Phy000AX5A_NEUCR	Ncr0002212	Q01294	(NCU02273.2)	396 aa	Vacuolar protease A, pep-4	PF00026	Experimental evidence	8702999, 12655011, 12712197	Neurospora crassa	-

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Phy000B20G_NEUCR	Ncr0008518	Q7SCF6	(NCU08739.2)	439 aa	Endothiapepsin	PF00026	Predicted function	12712197	Neurospora crassa	-
Phy000B3H7_NEUCR	Ncr0010417	A7UXG4	(NCU10907.2)	529 aa	Predicted protein	PF00026	Predicted function	12712197	Neurospora crassa	-
Phy000W68V_NEMVE	Nem0005466	A7RH56	prot5466	370 aa	Predicted protein	PF07966, PF00026	Predicted function	17615350	Nematostella vectensis	-
Phy000CWVQ_YEAST	Sce0001780	P12630	YIL015W	587 aa	Barrierpepsin	PF00026	Predicted function	3124102, 9169870, 14562106	Saccharomyces cerevisiae	-
Phy000CYQD_YEAST	Sce0004179	P32329	YLR120C	569 aa	Aspartic proteinase 3	PF00026	Experimental evidence	2183521, 9090053, 9169871, 8389368, 7779785, 7657670, 9417119, 11737827, 14617149, 14562106, 16087741, 18591427, 18573178, 9485427	Saccharomyces cerevisiae	-
Phy000CYQF_YEAST	Sce0004181	Q12303	YLR121C	508 aa	Aspartic proteinase yapsin-3	PF00026	Experimental evidence	9090053, 9169871, 10191273, 11016834, 16087741	Saccharomyces cerevisiae	-
Phy000D0B9_YEAST	Sce0006227	P07267	YPL154C	405 aa	Saccharopepsin	PF00026	Experimental evidence	3537721, 3023936, 8948103, 9169875, 1618910, 8840499, 1959673, 9135120	Saccharomyces cerevisiae	-
Phy000F3MX_USTMA	Uma0000064	Q4PIJ9	UM00064.1	397 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-
Phy000F55V_USTMA	Uma0002043	Q4PCX0	UM02043.1	481 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-
Phy000F59M_USTMA	Uma0002178	Q4PCI5	UM02178.1	452 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-
Phy000F5L9_USTMA	Uma0002597	Q4PBB6	UM02597.1	603 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-
Phy000F7DY_USTMA	Uma0004926	Q4P4N7	UM04926.1	418 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-
Phy000F7LP_USTMA	Uma0005205	Q4P3V8	UM05205.1	768 aa	Putative uncharacterized protein	PF00026	Predicted function	17080091	Ustilago maydis	-

a: new internal identifier in PhylomeDB; b: old internal identifier in PhylomeDB; c: UniProt accession number; d: original identifier in the database from which the proteome data was downloaded; e: amino acid (aa) sequence length; f: functional annotation in SchistoDB and UniProt; g: protein sequence domain(s) identified in the Pfam database; h: functional information in the literature available in UniProt; i: identifier in PubMed (PMID); j: scientific name of source organism; k: notes from this work.