Original article

Identification and prioritization of novel uncharacterized peptidases for biochemical characterization

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Introduction

Enzymes that act on biopolymers are poorly classified in terms of the reactions they catalyse. It is frequently difficult, or even impossible, to fully describe the signals, which may be sequence- or structure-based, by which these enzyme recognize their target substrates. Classification by reaction catalysed also masks evolutionary relationships because convergent evolution can give rise to enzymes with similar specificities that are otherwise unrelated, and divergent evolution may give rise to homologous enzymes with different specificities. These factors apply to enzymes, such as nucleases, which act on nucleic acids; glycosyl hydrolases, which act on carbohydrates; lipases, which act on lipids; and protein kinases and peptidases, which act on proteins and peptides. For each of these types of enzymes, because the method of classification by reaction catalysed in Enzyme Nomenclature (1) has been found to be...
Peptidases are enzymes that hydrolyse the carbon–nitrogen bonds between amino acids (peptide bonds). Peptidases are not only important for the complete degradation of proteins and peptides, so that the component amino acids can be re-used, but also for post-translational processing of proteins, so that activation is delayed until the protein has reached the correct cellular or extracellular compartment, or the necessary developmental, physiological or environmental conditions prevail. Examples of the former include digestion of proteins in food and the turnover of cytoplasmic and phagocytosed proteins in the lysosome. Examples of the latter include activation of enzymes and the processing of prohormone precursors. Proteolytic cascades exist to amplify the original biological signal, such as in blood coagulation and apoptosis. The latter process highlights a function of peptidases in the destruction of proteins and peptidases in switching off biological signals, including the degradation of peptide hormones and neurotransmitters.

A peptidase is targeted to a specific substrate by recognition of one or more binding sites. A substrate may bind directly at the peptidase active site, where the peptidase recognizes a sequence or structural motif in the substrate, or to a site known as an exosite, some distance from the enzyme active site. For a peptidase, such as the lysosomal cysteine peptidase cathepsin B, no clear sequence motif for substrate binding has emerged, despite the identification of hundreds of cleavage sites in substrates (7). This makes it difficult to classify peptidases by reaction catalysed.

The MEROPS database classifies peptidases along evolutionary principles, so that homologous sequences are included in the same family, and homologous tertiary structures are included in the same clan, according to the methods of Rawlings and Barrett (8). A family consists of sequences that can be shown to be directly or indirectly homologous to a single sequence designated the family-type example. Sequences that represent the same peptidase from different organisms are given a unique identifier, composed of the family name, a dot and a sequential number. For example, the well-characterized lysosomal cysteine peptidase cathepsin B is C01.060, being a member of family C1 (9). A MEROPS identifier is initially applied to a single sequence, known as the holotype, representing a peptidase that has been biochemically characterized. Other sequences given the same MEROPS identifier may or may not have been characterized, but all are assumed to represent an enzyme with similar biochemical properties (10). Not all sequences can be assigned to a particular MEROPS identifier, because some sequences are divergent and not closely related to any existing holotype.

A peptidase is assumed to be biochemically characterized when it can be measured quantitatively, which usually means that an assay of some kind is possible. In practical terms, this means that an article has been published, which describes characterization of the peptidase. Ideally, this would include information on substrates cleaved (even if the cleavage positions are not known), inhibitor profiles and the amino acid sequence of the peptidase. MEROPS maintains an extensive collection of references (11), known cleavage sites in substrates [including synthetic peptides, as well as proteins (12)] and a collection of interactions between peptidases and inhibitors, both naturally occurring and synthetic (13). An identifier is only established if the peptidase in question can be shown to differ from an existing peptidase because there is a significant difference in the action on one or more substrates or in the interaction with one or more inhibitors, or, more likely, because the sequence is too divergent to be included within an existing MEROPS identifier.

In the current release of the MEROPS database (release 9.7, 1 August 2012), for peptidases there are 60 clans, 236 families and 3944 identifiers, derived from 264 093 sequences. Of the sequences included in the database, 104 568 sequences (39.6%) cannot be assigned to a MEROPS identifier and have been assigned to miscellaneous family codes. Almost all of these sequences are derived from genome sequencing projects, often from unusual microbes, and are unlikely to be characterized. Besides their physiological roles, many peptidases have biomedical, scientific and industrial uses (14), and several peptidases from unfamiliar organisms have been used, in cheese-making and biological washing powders, for example. It is thus possible that many of the uncharacterized peptidases will prove to have unique specificities and be equally useful, or at least physiologically interesting. The problem of genome sequencing projects yielding large amounts of data relating to uncharacterized proteins has previously been addressed by the curators of the Pfam database because the database contains a considerable number of domain families of unknown function (15).

Figure 1 shows the number of peptidase homologue sequences in the MEROPS database at the end of each year from 1998 to 2012, and the number of these sequences assigned to a MEROPS identifier. Also shown is the number of MEROPS identifiers, which equates to the number of different characterized peptidases. In 1998, there were 7381 sequences and 117 identifiers, with 3786 sequences assigned to identifiers (51.3%). The growth in sequences is exponential, yet the increase in identifiers is linear, which implies that the gap between the number of
sequences and the proportion that can be assigned to identifiers will only widen. Clearly, characterization of peptidases is lagging behind genome sequencing, and a high-throughput mechanism for characterization is required. An important initial aspect of such a high-throughput system would be to recognize those peptidases most worthy of characterization. In this article, I will identify the uncharacterized peptidases that are most widely distributed among different species because widely distributed peptidases are most likely to have key physiological roles and to be worthy of biochemical characterization.

**Materials and Methods**

**Sequence sources**

Sequences homologous to peptidase family-type examples were identified from BLASTP searches of the NCBI non-redundant protein sequence database (17). A Perl program was written to submit that part of the peptidase protein sequence that bears the active site residues (known as the peptidase unit) from each family-type example, to the BLASTP search, with low complexity sequence masked. Searches were performed so that the maximum number of hits in which the expect value (e) was 0.001 or lower was returned. If the same hit sequence was returned in more than one results file, only the match with the lowest E value was retained. A Perl program was written to check that no residues had been filed more than once in the MEROPS MySQL database.

**Classification of sequences**

A pipeline consisting of a series of Perl programs was assembled to enter sequence data returned by the BLASTP searches into the MEROPS MySQL database. For each BLASTP search, all hits were entered as either a miscellaneous peptidase homologue or a miscellaneous non-peptidase homologue. A non-peptidase homologue was identified as a sequence having one or more active site residues replaced with an amino acid not known to occupy this position in an active peptidase. A list of peptidase active site residues was published by Rawlings and Morton (18). For most peptidases, the active site consists of a catalytic dyad or triad, but in the case of metallopeptidases, the ligands that bind the catalytic metal ion or ions are also considered active site residues. Non-peptidase homologues are not considered further in this article.

An alignment of peptidase units from all homologues in a family considered likely to be active peptidases was prepared using the Muscle software (19). Sequences in which active site residues were misaligned because the sequences were too divergent to align correctly, or missing, because the known sequence is only a fragment, were removed from the alignment. This ensured that false-positive results were removed from the analyses. A difference matrix was calculated from the alignment and converted to accept point mutations using the PAM250 matrix (20). A phylogenetic tree was then computed using the UPGMA algorithm (21) as implemented in the Quicktree program (22).

A bespoke Perl program was written to traverse the tree, to identify the tip that corresponded to each holotype and then to walk down the tree so that all sequences derived from the same node as the holotype were assigned the same MEROPS identifier, provided that the following three conditions were met:

1. No other holotype was included in the cluster.
2. No tip in which the sequence length differed from the holotype by a hundred or more residues was included in the cluster.
3. The node from which all tips in the cluster were derived did not correspond to an average sequence difference $>50\%$ of aligned residues.

In practice, the first two of these criteria had exceptions, as discussed later in the text.
Results

Establishment of MEROPS identifiers for all peptidases from model organisms

MEROPS identifiers were set-up for all homologues assumed to be proteolytically active from a range of model organisms (Table 1). The organisms chosen represent the major groups of cellular life: animals, a plant, fungi, a protist, a Gram-negative bacterium, a Gram-positive bacterium, and an archaean. For each of the bacteria, a single reference strain was selected: *Escherichia coli* strain K12 substrain MG1655 and *Bacillus subtilis* strain 168, which are also the strains chosen to be representative by the UniProt database (23). The number of peptidases in a model organism varies greatly. It may seem surprising that a mouse has more peptidase genes than any other model organism, or that the plant *Arabidopsis thaliana* has more peptidase genes than the fruit fly *Drosophila melanogaster* or the nematode *Caenorhabditis elegans*. The slime mould *Dictyostelium discoideum* has more peptidase genes than either of the yeasts *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*. The Gram-positive bacterium *B. subtilis* has nearly twice as many peptidases as the Gram-negative bacterium *E. coli*, and more than any of the three unicellular eukaryotes has. The model organism with fewest peptidase genes is the archaean *Pyrococcus furiosus*, although it is entirely possible that with so many uncharacterized proteins in this proteome, many of these will subsequently be discovered to be peptidases from novel families. Of the 2045 proteins from the *P. furiosus* genome, only 1095 have domains included in families in the Pfam database (24), of which 198 are included in domains of unknown function. In other words, 63.2% of proteins from *P. furiosus* have unknown functions.

Table 1 shows the number of holotypes established in each model organism, and the numbers that have been biochemically characterized, i.e. those that have been the subject of a published study and those with known substrates and inhibitor interactions. The holotypes that are biochemically uncharacterized are the putative peptidases that are the subject of this study.

Table 2 shows the number of peptidases common to all model organisms. Equivalent peptidases are tentatively assumed to perform the same physiological roles and to have similar protein architectures and substrate specificities. It is clear from the table that only human and mouse have a significant proportion of equivalent peptidases (77.4% of human peptidases and 72.7% of mouse peptidases). These two organisms are from the same taxonomic class, Mammalia, whereas the other model organisms are from different phyla. This indicates that there are very few sequences from organisms in different phyla that can be considered to be representatives of the same peptidase.

The new identifiers were set-up to be similar to, but easily distinguishable from, existing MEROPS identifiers. For the new identifiers, the first character after the dot was replaced by the letter ‘A’, and then the letter ‘B’ if >26 new identifiers were created for the family, and then the letter ‘C’ if more than 52 identifiers were created.

Assignment of the new identifiers to other sequences

Once the new peptidase identifiers had been established, the program to assign identifiers using the phylogenetic
tree was re-run so that the new identifiers could be assigned to sequences from species other than that of the holotype. Figure 2 shows the consequence of running the program on the phylogenetic tree for subfamily M3A. Table 3 shows the new identifiers that were assigned to ≥50 different species.

Table 4 shows the peptidase families for which identifiers for either 20 characterized or 20 putative peptidases have been assigned. For many of these families, the numbers for both are approximately equal (for example: A1, M1, M14, M16 and M20). Family A2 includes peptidases from viruses and retrotransposons, neither of which has been included in this study; hence, there are no identifiers for putative peptidases. The number of identifiers for uncharacterized peptidases in families S63 and T3 is also disproportionally small. There are families for which the number of identifiers for uncharacterized peptidases now exceeds the number for those that have been characterized, especially C26, M13, S9, S10, S12, S28 and S33. The numbers of identifiers for uncharacterized peptidases in S9 and S33 (both 100) are exceeded only by the number in S1 (180).

In total, the new identifiers were applied to 15 149 sequences, which had previously been unassigned, which is ~4.6% of all the peptidase homologues currently in the MEROPS database.

Discussion

Criteria for assigning sequences to a MEROPS identifier

The criteria for limiting the extension of a MEROPS identifier to other members of a family, as described in ‘Materials and Methods’ section, had to be relaxed in some cases. Peptidase holotypes are assumed to represent gene duplications that precede speciation events, but there are known instances where speciation precedes gene duplication. An example is the renin gene in mouse. Most mammals have only one renin gene, expressed in the kidney, and the enzyme is important for controlling blood pressure by processing the precursor of the peptide hormone angiotensin. But mice have two genes, one expressed in the kidney and one expressed only in the salivary gland, where the enzyme is probably more important as a chemical signal (25). A holotype exists for both renin 1 (the kidney enzyme) and renin 2 (the mouse salivary gland enzyme), and that the classification software should work correctly, the renin 2 holotype is considered a subset of the renin 1 cluster.

The condition that a sequence is not included in a cluster if the sequence length differs from that of the holotype by a hundred residues or more has to be relaxed in the few situations where the holotype sequence has been derived from sequencing of the mature protein and is being compared with the sequences of precursors (and vice versa). It was also discovered that many sequences derived from eukaryote genome sequencing projects were considerably longer than many holotype sequences, preventing the MEROPS identifier from being applied fully to other sequences in the tree. Gene building in eukaryote sequencing projects is notoriously difficult because of the problem of correctly identifying introns. Misidentifying introns can lead to incorrect chaining together of proteins derived from adjacent genes, or mistranslation of intron sequence as protein coding, both of which inadvertently increase the length of the protein sequence. To avoid this problem, the length criterion was not applied to sequences derived entirely from eukaryote genome sequencing projects. The
Figure 2. Use of a phylogenetic tree to assign MEROPS identifiers. Part of the tree for family M3 subfamily A is shown. The tips correspond to individual sequences. An arrow indicates the tip corresponding to the sequence that is the holotype for a particular MEROPS identifier. Tips assigned to the same MEROPS identifier are bracketed together, and the MEROPS identifier is shown. Sequences that are not included within a bracket are unassigned. Tips 63–70 are derived from a node ancestral to M03.003 and M03.009; hence, they cannot be assigned to either identifier; similar situations apply to the unassigned tips 135–157, 158–166 and 184–246. The tree shows two identifiers for putative peptidases. M03.A03 was initially assigned to the At1g67690 gene product from *A. thaliana* and has been extended to include sequences from nine other plants. On the other hand, M03.A07, which was originally assigned to the DDB_G0292362 gene product from the slime mould *D. discoideum*, but cannot be extended to sequences from other species because no others are derived from the same node on the tree.
<table>
<thead>
<tr>
<th>MEROPS identifier</th>
<th>Peptidase name</th>
<th>Number of species</th>
<th>Phyla</th>
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</table>
| C26.A05           | \(\gamma\)-Glutamyl peptidase 1 (\textit{A. thaliana}) | 78 | Bacteria (Bacteroidetes, Proteobacteria, Xenobacteria)  
Archaea (Euryarchaeota)  
Fungi (Ascomycota, Basidiomycota, Chytridiomycota)  
Plantae (Tracheophyta)  
Animalia (Annelida) |
| C26.A28           | SPBP8282.05 (\textit{S pombe}) | 297 | Bacteria (Acidobacteria, Bacteroidetes, Chlamydiae, Chloroflexi, Chrysiogenetes, Cyanobacteria, Firmicutes, Fusobacteria, Gemmatimonadetes, Nitrospira, Planctomycetes, Proteobacteria, Spirochaetes, Thermotogae, Xenobacteria)  
Fungi (Ascomycota)  
Plantae (Tracheophyta) |
| C26.A31           | GMP synthase (\textit{P. furiosus}) | 72 | Bacteria (Chlamydiae)  
Archaea (Crenarchaeota, Euryarchaeota) |
| C26.A32           | Imidazoleglycerol-phosphate synthase (\textit{P. furiosus}) | 176 | Bacteria (Acidobacteria, Bacteroidetes, Chloroflexi, Dictyoglomi, Fibrobacteres, Firmicutes, Fusobacteria, Proteobacteria, Synergistetes, Thermotogae, Xenobacteria)  
Archaea (Crenarchaeota, Euryarchaeota, Korarchaeota)  
Fungi (Ascomycota)  
Plantae (Tracheophyta)  
Animalia (Chordata) |
| C40.A01           | NlpC protein (\textit{E. coli}) | 60 | Bacteria (Proteobacteria)  
Animalia (Porifera) |
| C44.A08           | Glutamine-fructose-6-phosphate transaminase precursor (\textit{D. discoideum}) | 74 | Bacteria (Bacteroidetes, Chlorobi, Chloroflexi, Cyanobacteria, Firmicutes, Proteobacteria, Verrucomicrobia, Xenobacteria)  
Archaea (Crenarchaeota, Euryarchaeota)  
Protozoa (Alveolata, Apicomplexa, Microspora, Sarcomastigophora)  
Plantae (Heterokontophyta, Ochrophyta)  
Animalia (Arthropoda, Chordata) |
| C56.A06           | DDB_G0276405 (\textit{D. discoideum}) | 350 | Bacteria (Acidobacteria, Bacteroidetes, Chloroflexi, Cyanobacteria, Deferribacteres, Firmicutes, Ignavibacteria, Lentisphaerae, Planctomycetes, Proteobacteria, Spirochaetes, Thermodesulfobacteria, Verrucomicrobia, Xenobacteria)  
Archaea (Euryarchaeota)  
Protozoa (Sarcomastigophora)  
Fungi (Ascomycota)  
Plantae (Chlorophyta)  
Animalia (Arthropoda, Nematoda) |
| C82.A01           | YafK protein (\textit{E. coli}) | 56 | Bacteria (Bacteroidetes, Proteobacteria) |
| C82.A07           | Murein transglycosylase (\textit{B. subtilis}) | 74 | Bacteria (Cyanobacteria, Firmicutes, Proteobacteria) |
| M03.A08           | BSSC8_09230 protein (\textit{B. subtilis}) | 234 | Bacteria (Chlamydiae, Chloroflexi, Cyanobacteria, Firmicutes, Planctomycetes, Proteobacteria, Spirochaetes, Thermotogae, Xenobacteria)  
Plantae (Tracheophyta) |

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<th>MEROPS identifier</th>
<th>Peptidase name</th>
<th>Number of species</th>
<th>Phyla</th>
</tr>
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</table>
| M13.A32           | T25B6.2 protein (C. elegans) | 140 | Bacteria (Bacteroidetes, Firmicutes, Proteobacteria)  
Fungi (Ascomycota, Basidiomycota)  
Plantae (Chlorophyta, Rhodophyta, Tracheophyta)  
Animalia (Arthropoda, Chordata, Echinodermata, Hemichordata, Nematoda) |
| M15.A04           | YokZ protein (B. subtilis) | 274 | Bacteria (Bacteroidetes, Chloroflexi, Cyanobacteria, Firmicutes, Spirochaetaes, Xenobacteria) |
| M16.A04           | Insulysin homologue (S. cerevisiae) | 73 | Fungi (Ascomycota, Basidiomycota) |
| M16.A05           | PqqL protein (E. coli) | 103 | Bacteria (Bacteroidetes, Chlorobi, Deferribacteres, Fusobacteria, Gemmatimonadetes, Proteobacteria, Spirochaetes, Xenobacteria) |
| M20.A08           | YgeY protein (E. coli) | 61 | Bacteria (Bacteroidetes, Caldiserica, Chloroflexi, Firmicutes, Fusobacteria, Proteobacteria, Spirochaetes, Synergistetes, Thermotogae) |
| M20.A18           | DD8_G0279291 protein (D. discoideum) | 72 | Bacteria (Firmicutes, Proteobacteria)  
Protozoa (Alveolata, Parabasalidea, Sarcomastigophora)  
Plantae (Tracheophyta) |
| M20.A21           | YodQ protein (B. subtilis) | 81 | Bacteria (Chloroflexi, Firmicutes, Fusobacteria, Proteobacteria, Spirochaetes)  
Archaea (Euryarchaeota) |
| M20.A23           | Bsub1_01010013116 protein (B. subtilis) | 52 | Bacteria (Firmicutes, Fusobacteria, Planctomycetes, Proteobacteria, Synergistetes) |
| M20.A27           | YkuR protein (B. subtilis) | 82 | Bacteria (Firmicutes, Fusobacteria, Thermotogae) |
| M24.A09           | YFR006W protein (S. cerevisiae) | 57 | Fungi (Ascomycota, Basidiomycota)  
Plantae (Chlorophyta) |
Fungi (Ascomycota, Basidiomycota)  
Plantae (Tracheophyta) |
| M48.A02           | At3g27110 (A. thaliana) | 50 | Bacteria (Cyanobacteria, Firmicutes, Proteobacteria, Spirochaetes)  
Archaea (Euryarchaeota)  
Plantae (Tracheophyta) |
| M50.A04           | PF0392 protein (P. furiosus) | 79 | Bacteria (Acidobacteria, Bacteroidetes, Chlorobi, Chloroflexi, Cyanobacteria, Nitrospirae, Planctomycetes, Proteobacteria)  
Archaea (Crenarchaeota, Euryarchaeota) |
| M50.A05           | YwhC protein (B. subtilis) | 203 | Bacteria (Acidobacteria, Aquificae, Chloroflexi, Chrysogentes, Deferrribacteres, Dictyoglomi, Firmicutes, Fusobacteria,  
Nitrospirae, Proteobacteria, Spirochaetes, Synergistetes, Thermodesulfovibacteria, Verrucomicrobia, Xenobacteria)  
Archaea (Nanoarchaeota) |

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<th>MEROPS identifier</th>
<th>Peptidase name</th>
<th>Number of species</th>
<th>Phyla</th>
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<tbody>
<tr>
<td>M50.A07</td>
<td>PF0457 protein (<em>P. furiosus</em>)</td>
<td>76</td>
<td>Bacteria (Bacteroidetes, Chloroflexi, Cyanobacteria, Firmicutes, Gemmatimonadetes, Proteobacteria, Xenobacteria) Archaea (Crenarchaeota, Euryarchaeota) Plantae (Chlorophyta)</td>
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<tr>
<td>M79.A04</td>
<td>At3g26085 (<em>A. thaliana</em>)</td>
<td>58</td>
<td>Bacteria (Chloroflexi, Cyanobacteria, Firmicutes, Proteobacteria, Xenobacteria)</td>
</tr>
<tr>
<td>M79.A09</td>
<td>YdlL protein (<em>B. subtilis</em>)</td>
<td>140</td>
<td>Bacteria (Bacteroidetes, Chlamydiae, Chloroflexi, Cyanobacteria, Deferrribacteres, Firmicutes, Fusobacteria, Proteobacteria, Spirochaetes, Verrucomicrobia) Archaea (Crenarchaeota, Euryarchaeota) Protozoa (Sarcomastigophora)</td>
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<tr>
<td>N06.A01</td>
<td>FlhB protein (<em>E. coli</em>)</td>
<td>450</td>
<td>Bacteria (Acidobacteria, Aquificae, Chloroflexi, Deferrribacteres, Firmicutes, Proteobacteria, Spirochaetes, Synergistetes, Thermotogae)</td>
</tr>
<tr>
<td>S01.A08</td>
<td>At5g27660 (<em>A. thaliana</em>)</td>
<td>58</td>
<td>Bacteria (Acidobacteria, Bacteroidetes, Caldiseria, Chlamydiae, Chloroflexi, Firmicutes, Fusobacteria, Ignavibacteria, Planctomycetes, Proteobacteria, Spirochaetes, Thermotogae, Xenobacteria) Archaea (Crenarchaeota) Plantae (Heterokontophyta, Tracheophyta) Animalia (Chordata)</td>
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<tr>
<td>S09.A43</td>
<td>YpfH protein (<em>E. coli</em>)</td>
<td>101</td>
<td>Bacteria (Acidobacteria, Chlamydiae, Chlorobi, Chloroflexi, Cyanobacteria, Firmicutes, Proteobacteria) Fungi (Ascomycota)</td>
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<tr>
<td>S09.A77</td>
<td>dpf-6 protein (<em>C. elegans</em>)</td>
<td>334</td>
<td>Bacteria (Acidobacteria, Bacteroidetes, Chlamydiae, Chloroflexi, Cyanobacteria, Firmicutes, Gemmatimonadetes, Nitrospirae, Planctomycetes, Proteobacteria, Synergistetes, Thermotogae, Verrucomicrobia, Xenobacteria) Archaea (Crenarchaeota, Euryarchaeota) Protozoa (Sarcomastigophora) Fungi (Ascomycota) Plantae (Ochrophyta, Tracheophyta) Animalia (Chordata, Nematoda)</td>
</tr>
<tr>
<td>S09.B04</td>
<td>BSU23640 protein (<em>B. subtilis</em>)</td>
<td>56</td>
<td>Bacteria (Acidobacteria, Bacteroidetes, Chloroflexi, Firmicutes, Planctomycetes, Proteobacteria, Xenobacteria) Archaea (Crenarchaeota, Euryarchaeota)</td>
</tr>
<tr>
<td>S10.A47</td>
<td>At2g22960 (<em>A. thaliana</em>)</td>
<td>122</td>
<td>Protozoa (Alveolata, Apicomplexa, Sarcomastigophora) Fungi (Ascomycota, Basidiomycota) Plantae (Chlorophyta, Ochrophyta, Oomycota, Rhodophyta, Streptophyta, Tracheophyta) Animalia (Arthropoda, Chordata, Echinodermata, Hemichordata, Nematoda)</td>
</tr>
<tr>
<td>S12.A03</td>
<td>EcHS_A2566 protein (<em>E. coli</em>)</td>
<td>180</td>
<td>Bacteria (Bacteroidetes, Chlorobi, Chloroflexi, Cyanobacteria, Firmicutes, Fusobacteria, Nitrospirae, Planctomycetes, Proteobacteria, Spirochaetes, Thermotogae, Xenobacteria) Archaea (Euryarchaeota) Fungi (Ascomycota)</td>
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<th>MEROPS identifier</th>
<th>Peptidase name</th>
<th>Number of species</th>
<th>Phyla</th>
</tr>
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</table>
| S12.A23           | GYO_0385 protein (B. subtilis) | 51 | Bacteria (Bacteroidetes, Firmicutes, Proteobacteria, Spirochaetes, Xenobacteria) 
Archaea (Euryarchaeota) |
| S16.A10           | YcbZ (E. coli) | 110 | Bacteria (Firmicutes, Proteobacteria, Spirochaetes) 
Animalia (Chordata) |
| S16.A12           | PF1438 (P. furiosus) | 270 | Bacteria (Firmicutes, Proteobacteria) 
Archaea (Crenarchaeota, Euryarchaeota, Korarchaeota) 
Protozoa (Apicomplexa) |
| S49.A08           | BSn5_05605 protein (B. subtilis) | 189 | Bacteria (Chrysiogenetes, Cyanobacteria, Deferrribateres, Firmicutes, Proteobacteria, Spirochaetes, Thermodesulfobacteria, Xenobacteria) 
Archaea (Crenarchaeota, Euryarchaeota) |
| S49.A09           | PF0240 protein (P. furiosus) | 67 | Bacteria (Aquificae, Bacteroidetes, Chloroflexi, Cyanobacteria, Deferrribateres, Dictyoglomi, Firmicutes, Nitrospirae, Proteobacteria, Synergistetes, Thermodesulfobacteria, Thermotogae, Xenobacteria) 
Archaea (Crenarchaeota, Euryarchaeota) |
| S54.A18           | YdcA protein (B. subtilis) | 54 | Bacteria (Firmicutes, Proteobacteria) |
| T03.A09           | GYO_3966 protein (B. subtilis) | 761 | Bacteria (Acidobacteria, Bacteroidetes, ChloroFlexi, Cyanobacteria, Firmicutes, Fusobacteria, Lentisphaerae, Planctomycetes, Proteobacteria, Xenobacteria) 
Archaae (Euryarchaeota) 
Protozoa (Sarcomastigophora) 
Fungi (Ascomycota, Basidiomycota, Oomycota) 
Plantae (Chlorophyta, Heterokontophyta, Ochrophyta, Tracheophyta) 
Animalia (Annelida, Arthropoda, Chordata, Cnidaria, Placozoa, Porifera) |
| U32.A01           | YhbV protein (E. coli) | 131 | Bacteria (Proteobacteria) |

Columns are the MEROPS identifier (linked to the peptidase summary page in the MEROPS database); the name used in the MEROPS database, which is often derived from the gene name; the number of species containing a sequence to which the identifier has been assigned; and the phyla that includes these species, grouped by kingdom. When one of these putative peptidases is characterized, the MEROPS identifier will be replaced with an identifier that follows the normal naming convention used in the MEROPS database. The obsolete identifier will not be re-used, and so that the links below will remain useful the user will be automatically redirected to the summary page for the replacement identifier.
length criterion was introduced so that the domain architecture would be more or less the same for all sequences assigned to the same identifier because sequences with similar domain architectures are unlikely to have different exosites. Relaxing the length criterion for eukaryotes until automated gene assembly improves does carry the risk of assigning sequences with genuinely different domain architectures, and possibly different exosites, to the same identifier.

Species-distribution of new peptidase identifiers

The species-distribution of a peptidase varies greatly, from just one known species, such as renin-2, which is known only from the mouse, to many species from all major groups of organic life. The most widely distributed peptidases are peptidase Clp (present in organisms from 51 of the 100 known phyla), methionyl aminopeptidase 1 (48 phyla) and the self-processing ornithine acetyltransferase precursor (42 phyla). T03.A09 is the new identifier assigned to the sequences from the most species. This is a homologue of \(\gamma\)-glutamyltransferase, which degrades glutathione (\(\gamma\)-Glu-Cys-Gly) by cleavage of the \(\gamma\)-glutamyl bond. T03.A09 is present in species from all six major kingdoms of organisms (bacteria, archaea, protista, fungi, plants and animals), and it is widely distributed in bacteria, plants and animals. Despite having not been characterized, crystal structures have been deposited for T03.A09 from \(B.\) halodurans (PDB: 2NLZ) and \(T.\) acidophilum (PDB: 2I3O), but neither has been published. T03.A09 from \(B.\) halodurans is described as ‘cephalosporin acylase’ (cephalosporin is a \(\beta\)-lactam–based antibiotic and a bacteriocide), whereas in \(T.\) acidophilum, it is described as a ‘\(\gamma\)-glutamyltransferase–related protein’.

More genomes from bacteria have been sequenced than for any other kingdom of organisms; thus, it is no surprise that most of the identifiers listed in Table 3 are from bacteria. The identifiers with the widest distribution among bacterial phyla are C26.A28 (15), C56.A06 (14), S09.A77 (42), and S33.100.

<table>
<thead>
<tr>
<th>Family</th>
<th>Type example</th>
<th>Characterized peptidases</th>
<th>Putative peptidases</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>Pepsin</td>
<td>84</td>
<td>92</td>
</tr>
<tr>
<td>A2</td>
<td>Retropepsin</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>C1</td>
<td>Papain</td>
<td>148</td>
<td>59</td>
</tr>
<tr>
<td>C2</td>
<td>Calpain</td>
<td>25</td>
<td>10</td>
</tr>
<tr>
<td>C19</td>
<td>Ubiquitin-specific peptidase 14</td>
<td>94</td>
<td>66</td>
</tr>
<tr>
<td>C26</td>
<td>(\gamma)-Glutamyl hydrolase</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>C48</td>
<td>Ulp1 peptidase</td>
<td>32</td>
<td>19</td>
</tr>
<tr>
<td>M1</td>
<td>Aminopeptidase N</td>
<td>31</td>
<td>31</td>
</tr>
<tr>
<td>M12</td>
<td>Astacin</td>
<td>180</td>
<td>51</td>
</tr>
<tr>
<td>M13</td>
<td>Neprilysin</td>
<td>17</td>
<td>32</td>
</tr>
<tr>
<td>M14</td>
<td>Carboxypeptidase A1</td>
<td>34</td>
<td>34</td>
</tr>
<tr>
<td>M16</td>
<td>Pitrilysin</td>
<td>17</td>
<td>20</td>
</tr>
<tr>
<td>M20</td>
<td>Glutamate carboxypeptidase</td>
<td>20</td>
<td>21</td>
</tr>
<tr>
<td>M41</td>
<td>FtsH peptidase</td>
<td>22</td>
<td>14</td>
</tr>
<tr>
<td>S1</td>
<td>Chymotrypsin</td>
<td>462</td>
<td>180</td>
</tr>
<tr>
<td>S8</td>
<td>Subtilisin</td>
<td>147</td>
<td>58</td>
</tr>
<tr>
<td>S9</td>
<td>Prolyl oligopeptidase</td>
<td>45</td>
<td>100</td>
</tr>
<tr>
<td>S10</td>
<td>Carboxypeptidase Y</td>
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<td>69</td>
</tr>
<tr>
<td>S12</td>
<td>(\alpha)-Ala-(\alpha)-Ala carboxypeptidase B</td>
<td>10</td>
<td>21</td>
</tr>
<tr>
<td>S26</td>
<td>Signal peptidase</td>
<td>27</td>
<td>10</td>
</tr>
<tr>
<td>S28</td>
<td>Lysosomal Pro-Xaa carboxypeptidase</td>
<td>5</td>
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</tr>
<tr>
<td>S33</td>
<td>Prolyl aminopeptidase</td>
<td>18</td>
<td>100</td>
</tr>
<tr>
<td>S54</td>
<td>Rhomboid-1</td>
<td>28</td>
<td>18</td>
</tr>
<tr>
<td>S63</td>
<td>EGF-like module containing mucin-like hormone receptor-like 2</td>
<td>35</td>
<td>2</td>
</tr>
<tr>
<td>T3</td>
<td>(\gamma)-Glutamyltransferase 1</td>
<td>21</td>
<td>9</td>
</tr>
</tbody>
</table>

The total number of identifiers for characterized and uncharacterized peptidases is shown for all families where there are \(\geq 20\) examples in either category. The name of the type example peptidase is given for each family.
(14), S01.A08 (13), S12.A03 (12), C26.A32 (11), M79.A09 (10) and T03.A09 (10). The few identifiers that are exclusively bacterial are found only in species from a few phyla. An exception is N06.A01, which is found in species from nine bacterial phyla. Members of family N6 are not peptidases, but self-processing proteins that use asparagine as a nucleophile and have been described as ‘asparagine peptide lyases’ (26). Members of family N6 are known to be important for type III secretion by which virulence factors are injected into host cells through a flagellum-like filament (27).

There are three other identifiers found only in bacteria for which examples are known from over a hundred species. These are M15.A04, M16.A05 and U32.A01. Family M15 includes predominantly D-Ala-D-Ala carboxypeptidases, important for processing of the precursor of the cross-linking peptide of a bacterial cell wall (28). The cross-linking peptide precursor in both B. subtilis (from which the holotype is derived) and E. coli is L-Ala-L-Glu-L-meso-diaminopimelic acid-D-Ala-D-Ala (29), so presumably the M15.A04 peptidase is not involved in processing of this precursor because this peptidase is not present in E. coli. Peptidase family M16 includes the inverzincins, metallopeptidases in which the zinc-binding residues are the histidines in the motif HXXEH, the reverse of the motif found in thermolysin (30), and includes the oligopeptidases pitrilysin, which is periplasmic in E. coli, and insulysin, which is cytoplasmic in vertebrates. Insulysin consists of two homologous domains, of which only the N-terminal one bears the active site (31). The PqQL protein (M16.A05), which was sequenced independently of the genome sequencing project (32) but never characterized, is similarly composed of two homologous domains. Unlike pitrilysin, the PqQL protein does not have an N-terminal signal peptide and is presumably cytosolic. Peptidases in family U32 are of unknown catalytic type, and the only characterized peptidases have been shown to degrade soluble collagen, such as the PrtC protein from Porphyromonas gingivalis (33). U32.A01 is restricted entirely to Proteobacteria. However, not only is U32.A01 the only identifier in Table 3 for a peptidase of unknown catalytic type, it is also the only one for which active site residues are unknown. Therefore, it is impossible to tell whether this protein represents an active peptidase or a non-peptidase homologue.

Table 3 lists three identifiers that are not found in bacteria: M16.A04 (found in 73 species), M24.A09 (found in 57) and S10.A47 (found in 122). M16.A04, which is restricted to fungi, presumably represents a novel oligopeptidase and M24.A09, found only in fungi and green algae, may be a novel exopeptidase because family M24 contains aminopeptidases (such as methionyl aminopeptidase), dipeptidases (such as Xaa-Pro dipeptidase) and aminopeptidases (such as aminopeptidase P) (34, 35). Family S10 contains carboxypeptidases (35), and S10.A47 is presumably also a carboxypeptidase. An example of S10.A47 was partially sequenced, but never characterized, from barley and was shown to be expressed in the aleurone layer of the seed (36).

Examination of the distribution of source organisms among the sequences assigned to a particular MEROPS identifier can highlight the problem of contamination when the genome was being sequenced, or reveal possible examples of horizontal gene transfer if the unusual distribution is consistent. Examples of possible contamination include M03.A08 from the moss Physcomitrella patens (the identifier has otherwise only been assigned to sequences from bacteria); M79.A04 from the plants A. thaliana and poplar (otherwise the identifier has been assigned only to sequences from bacteria and archaea); M79.A09 from the protozoan Naegleria gruberi (otherwise the identifier has been assigned exclusively to prokaryote sequences); S09.A43 from the yeast Gibberella zeae (otherwise the identifier has been assigned only to sequences from bacteria); and S16.A10 from the armadillo (otherwise the identifier has been assigned only to sequences from bacteria). The presence of M48.A02 in A. thaliana and rice also probably represents bacterial contamination, but there is also a sequence from the uncultured methanogenic archaeon RC-I, which could be an example of a horizontal gene transfer. M50.A07 is again mostly prokaryotic apart from three sequences from three different green algae species, which may represent a single horizontal gene transfer to an ancestral alga, because the algae sequences cluster together on the phylogenetic tree, from a bacterium.

Non-peptidase homologues

It should be stressed that not all the identifiers listed in Table 3 may relate to peptidases. Almost all families have non-peptidase homologues where one or more active site residues are missing or have been replaced. However, there are members of some families in which the active site residues are conserved but are known to perform reactions other than proteolysis. Examples include aminoacylase and acetylornithine deacytelyase (from family M20), ureases (family M38), esterases (family S9) and lipases (family S33). Succinyl-diaminopimelate desuccinylase (family M20) was initially characterized as a non-peptidase homologue, but it was then shown to possess proteolytic activity (37). Leukotriene A4 hydrolase is an example of a protein with dual enzymatic activity, including that of a peptidase (38).

Table 3 lists five members of family C26, which includes not only the mammalian peptidase γ-glutamyl hydrolase but is also known to include several activities other than peptidases, including guanine 5’-monophosphate (GMP), carbamyl phosphate and aminodeoxychorismate synthases. The holotypes for C26.A31 and C26.A32, both from P. furiosus, are described as a GMP synthase and an imidazolglycerol-phosphate synthase, respectively, even though neither has...
been biochemically characterized. It remains likely that some or even all of the C26 homologues listed in Table 3 are not peptidases.

Characterization

Although none of the putative peptidases listed in Table 3 has been biochemically characterized, some have been subjects of study, such as PqqL (M16.A05, see earlier in the text). Additionally, C26.A05 has been implicated, along with other members of the γ-glutamyl hydrolase family, in the production of glucosinolates, which are defence-related plant metabolites using glutathione as a source of sulphur (39, 40), although the pathway has not been established. Tertiary structures have been solved for some of the putative peptidases listed in Table 3 by consortia who are trying to solve structures for all protein folds without necessarily knowing the functions of the proteins being solved. Structures have been solved for M32.A01 from B. subtilis [PDB: 3HQ2; (41)] and M13.A32 (see earlier in the text).

The identifiers for these putative peptidases will be established in the next release of the MEROPS database (release 9.8, due December 2012). When one of these putative peptidases becomes characterized, the identifier will be replaced with a standard MEROPS identifier. The obsolete identifier will not be re-used, and an automatic redirection to the new identifier will be set up in the MEROPS database.

Conclusions

Methods have been established for distinguishing a cluster of sequences within a peptidase family that represents species variants of a common peptidase. The methods involve detection of homologues by a sequence similarity search using that part of the sequence bearing the catalytic residues from a designated type example, assembly of an alignment, generation of a phylogenetic tree from the alignment, followed by computerized assessment of the tree to identify clusters of related sequences. By using these methods and the extensive collections of references, substrate cleavages and inhibitor interactions in the MEROPS database, it has been possible to distinguish a cluster of sequences representing a biochemically characterized peptidase from a cluster that represents a putative peptidase that has not been characterized. By assigning identifiers to every peptidase homologue from the complete genome sequences of 11 model organisms, both those that are characterized and those that are not, the methods have been applied to detect a cluster of sequences that includes each of the putative peptidases from these model organisms. The clusters of putative peptidases with the widest organism distribution are presented in this article as ideal targets for biochemical characterization. Data derived from experimentation should then be transferable to all members of the cluster.

Versions of these methods should be applicable to any collection of sequence information where uncharacterized proteins or genes need to be distinguished from those that are characterized and should be especially applicable to enzymes that act on biological polymers.

By combining these methods with an analysis of the taxonomic distribution within the cluster, it is possible to identify potential contaminants in genome sequencing projects and examples of possible horizontal gene transfer.

The new identifiers for putative peptidases will be included in release 9.8 of the MEROPS database.

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Conflict of interest

None declared.

References