

# BAR-PLUS: the Bologna Annotation Resource Plus for functional and structural annotation of protein sequences

Damiano Piovesan<sup>1</sup>, Pier Luigi Martelli<sup>1</sup>, Piero Fariselli<sup>2</sup>, Andrea Zauli<sup>3</sup>,  
Ivan Rossi<sup>3</sup> and Rita Casadio<sup>1,\*</sup>

<sup>1</sup>Department of Biology, Bologna Biocomputing Group, Bologna Computational Biology Network, <sup>2</sup>Department of Computer Science, University of Bologna, Bologna and <sup>3</sup>BioDec srl, Bologna, Italy

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## ABSTRACT

We introduce **BAR-PLUS (BAR<sup>+</sup>)**, a web server for functional and structural annotation of protein sequences. **BAR<sup>+</sup>** is based on a large-scale genome cross comparison and a non-hierarchical clustering procedure characterized by a metric that ensures a reliable transfer of features within clusters. In this version, the method takes advantage of a large-scale pairwise sequence comparison of 13 495 736 protein chains also including 988 complete proteomes. Available sequence annotation is derived from UniProtKB, GO, Pfam and PDB. When PDB templates are present within a cluster (with or without their SCOP classification), profile Hidden Markov Models (HMMs) are computed on the basis of sequence to structure alignment and are cluster-associated (Cluster-HMM). Therefrom, a library of 10858 HMMs is made available for aligning even distantly related sequences for structural modelling. The server also provides pairwise query sequence–structural target alignments computed from the correspondent Cluster-HMM. **BAR<sup>+</sup>** in its present version allows three main categories of annotation: PDB [with or without SCOP (\*)] and GO and/or Pfam; PDB (\*) without GO and/or Pfam; GO and/or Pfam without PDB (\*) and no annotation. Each category can further comprise clusters where GO and Pfam functional annotations are or are not statistically significant. **BAR<sup>+</sup>** is available at <http://bar.biocomp.unibo.it/bar2.0>.

## INTRODUCTION

In the post-genomic era, with the advent of rapid sequencing techniques, reliable and efficient functional annotation methods are needed. Routinely, a translated

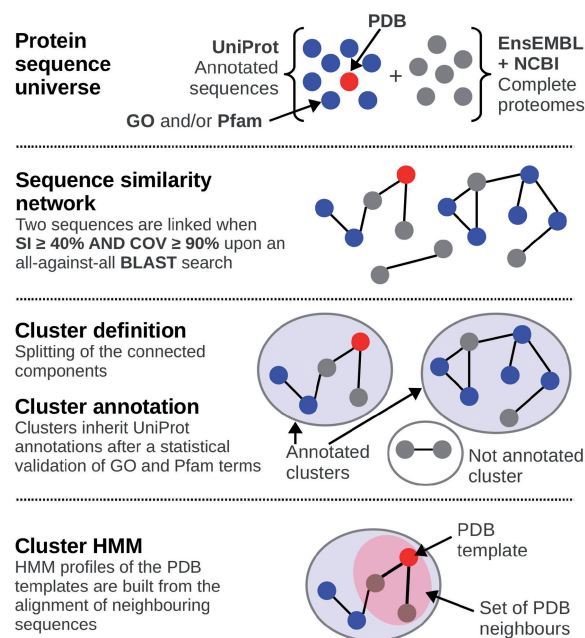
protein sequence is aligned towards a data base of already annotated sequences and by this it is endowed with different features depending on the level of sequence identity (SI). This similarity search is the basis for transfer of annotation by homology. The UniProt Knowledgebase (UniProtKB; <http://www.UniProtKB.org/>) is presently our major resource of information of protein sequences and of corresponding functions and structures, when available. It provides links also to other resources/data bases, allowing a comprehensive knowledge of experimental and computational characteristics of known/putative proteins and genes. However, only 4.4% of the all protein universe that presently (UniProtKB release 2011\_03; 8 March 2011) includes some 14 million of sequences has evidence at the protein and at the transcript level. With this scenario, inference of function and structure among related sequences requires the definition of rules to increase the reliability of annotation. This is routinely obtained with clustering methods by which sequences are included into sets of similarity. Clustering can be hierarchical and non-hierarchical. Hierarchical clustering categorizes sequences into a tree-structure. Examples of hierarchical clustering include SYSTERS (1), Picasso (2) and iProClass (3). CluSTr (4,5) and ProtoNet (6,7) are the only web servers that comprise the large number of sequences made available by fully sequenced genomes and the entire UniProtKB. Both CluSTr and ProtoNet cluster sequences according to different levels of SI, as set by different *E*-value thresholds, and with different hierarchical algorithms. Alternatively, non-hierarchical clustering partitions a sequence data set into disjoint clusters (8,9). However, neither hierarchical nor non-hierarchical methods consider explicitly proteins containing multiple domains or proteins that sharing common domains do not necessarily have the same function. Proteins with different combinations of shared domains can have different molecular and biological functions, as recently re-discussed (10). In order to address these problems, we

\*To whom correspondence should be addressed. Tel: +39 0512094005; Fax: +39 0512094005; Email: [casadio@biocomp.unibo.it](mailto:casadio@biocomp.unibo.it)

developed BAR (11), an annotation procedure that relies on a non-hierarchical clustering method and a large-scale genome comparison where pairs of sequences are selected with very strict criteria of similarity and overlapping of the alignment as described in the next section. We provided statistical validation that BAR allows reliable functional and structural annotation in addition to that given by commonly used databases (11). Here, we introduce BAR<sup>+</sup>, an updated and extended version of BAR that includes: (i) a 5-fold increase in sequences; (ii) GO terms from the three main roots (molecular function, biological process and cellular localization; <http://www.geneontology.org/>); (iii) Pfam domains (<http://pfam.sanger.ac.uk/>); (iv) known ligands and (v) for clusters containing PDB structure/s, a Cluster HMM model and the corresponding alignment of the target sequence to the optimal template in the cluster for computing its 3D structure.

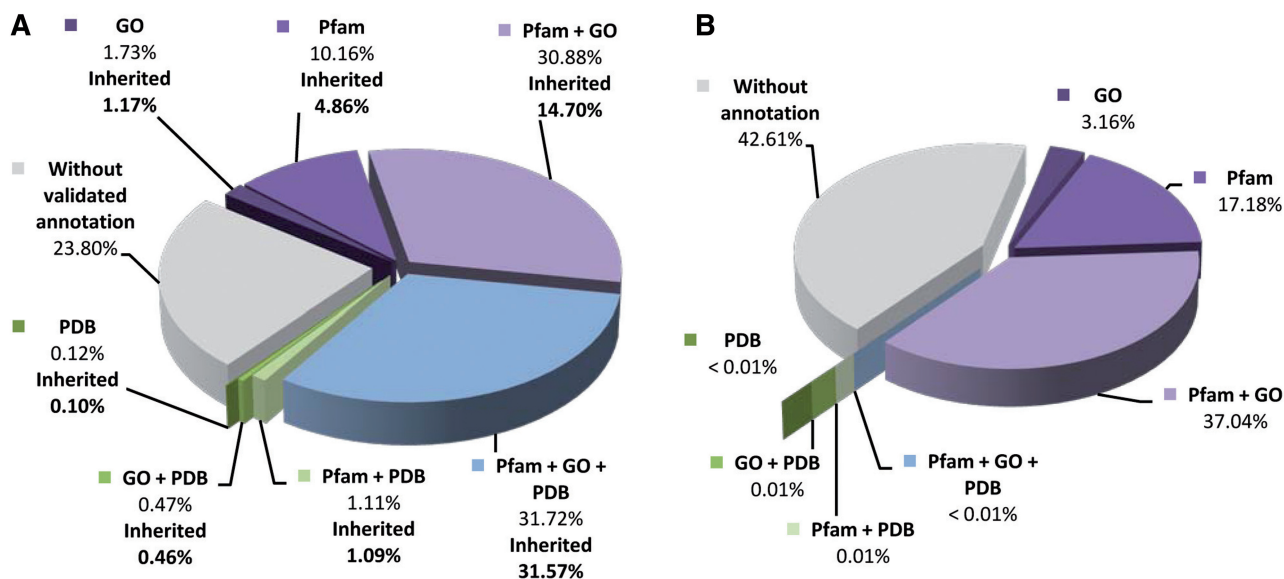
## BAR<sup>+</sup> IMPLEMENTATION

BAR<sup>+</sup> is constructed by performing an all-against-all pairwise alignment of all protein sequences (collected from the entire UniProtKB 05\_2010, with the exclusion of fragments (9 399 063 sequences), and from the proteome of complete sequenced genomes available on the same date at the National Center for Biotechnology Information (NCBI) [[www.ncbi.nlm.nih.gov/genomes/lproks.cgi](http://www.ncbi.nlm.nih.gov/genomes/lproks.cgi) (Prokaryotes); [www.ncbi.nlm.nih.gov/genomes/leuks.cgi](http://www.ncbi.nlm.nih.gov/genomes/leuks.cgi) (Eukaryotes)] and at Ensembl (<http://www.ensembl.org/info/data/ftp/index.html>) for a total of 988 complete proteomes (the list of the species is available at BAR<sup>+</sup> web site). For the sake of comparison, we also used the entire SwissProt 03\_2011 (8 March). Similarly to BAR (11), BAR<sup>+</sup> is also a non-hierarchical clustering method relying on a comparative large-scale genome analysis. The method relies on a non-hierarchical clustering procedure characterized by a stringent metric that ensures a reliable transfer of features within clusters. In this new version, the method takes advantage of a larger scale pairwise sequence comparison than BAR, including 13 495 736 protein sequences. Alignment is performed with BLAST (12) in a GRID environment (11). From this we compute for each pair both the SI and the Coverage (COV) defined as the ratio of the length of the intersection of the aligned regions on the two sequences and the overall length of the alignment (namely the sum of the lengths of the two sequences minus the intersection length). Each protein is then taken as a node and a graph is built allowing links among nodes only when the following similarity constraints are found among two proteins: their SI is  $\geq 40\%$  and COV is  $\geq 90\%$ . By this, clusters are simply the connected components of the graph (11). A workflow of the method is shown in Figure 1. Seventy percent of the whole data set (9 401 223 sequences) falls into 913 962 clusters. Noticeably, 55% of the clusters include 84% of the cluster-included sequences. The number of sequence in the clusters ranges from two up to 87 893 in the most populated (Molecular Function: ABC transporter). Given our stringent criteria, 87% of the clusters contain



**Figure 1.** BAR<sup>+</sup> implementation. Our method collects sequences from the protein universe (UniProtKB) including also some 988 genomes. By this, all the features [PDB ( $\pm$  SCOP classification) (red circles), GO terms (including Molecular Function, Biological Process and Cellular Localization) and Pfam models (blue circles)] are also included. An extensive BLAST alignment is performed of all the 13 495 736 sequences in a GRID environment. The sequence similarity network is built by connecting two sequences only if their SI is  $\geq 40\%$  with an overlapping COV  $\geq 90\%$ . About 913 762 clusters are obtained by splitting of the connected components. By this, any cluster may contain from 2 up to 87 893 sequences (one cluster containing ABC transporters from Prokaryotes, Eukaryotes and Archaea). Stand alone sequences are called Singletons (30.4% of the total protein universe). Sequences inherit the annotations within a cluster. When clusters are endowed with PDB template/s, a Cluster-HMM is generated by considering all the sequences that have an identity  $\geq 40\%$  and a COV  $\geq 90\%$  with the structure/s (pink subset). The Cluster-HMM can be used to align all the other sequences in the cluster to template/s.

sequences whose standard deviation (SD) of the protein length is  $\leq 5$  residues. The remaining sequences (30% of the total) originate singletons (containing just one sequence). Well annotated sequences are characterized by functional and structural annotations derived from UniProtKB entries (Figure 1). These include GO, Pfam, PDB and SCOP (<http://scop.mrc-lmb.cam.ac.uk/scop/>) (when available). To assess whether GO and Pfam terms are significant in a cluster, we compute *P*-values and given the multiplicity of the terms, we applied the Bonferroni correction (11). We evaluated the cumulative distribution of Bonferroni corrected *P*-values by adopting a bootstrapping procedure. From this we set the threshold *P*-value at 0.01 in order to discriminate among random and significant (cluster associated) features (11). Validated features (significant for the cluster) are those endowed with  $P \leq 0.01$ . According to our procedure when hypothetical and or putative proteins fall into an annotated and validated cluster, they can safely inherit GO terms and Pfam domain/s even in the case of very low SI with the most annotated proteins. These sequences can



**Figure 2.** Different types of annotations are possible with BAR<sup>+</sup>. After clustering and depending on the features (structure, domains and function) annotated in the cluster, sequences within a cluster can inherit different types of annotation. The percentage of sequences endowed with a given annotation type and inheriting validated annotation ( $P < 0.01$ ) is indicated. (A) Sequences within clusters. Percentage is computed with respect to 9401 223 comprised in 913 762 clusters. Inherited: sequences that inherit annotations by falling into a cluster. Without validated annotation: the slice comprises sequences with no annotation and not validated annotations. (B) Singletons (stand alone sequences). Percentage is computed with respect to 4 091 908 singleton sequences.

therefore be labelled as distantly related homologues and inherit function and structure (when available) in a validated manner. We previously discussed that this procedure can increase the level of annotation of UniProtKB (11). Here we increase the level of structural and functional annotations of cluster-included sequences by 54% (Figure 2A). When sequences are standing alone (according to our criteria) they are singletons. They can anyway carry along information (Figure 2B), provided that each singleton is endowed with PDB and/or Pfam and/or GO annotation.

### CLUSTER-HMMs

In BAR<sup>+</sup>, when PDB templates are present within a cluster (with or without their SCOP classification), profile HMMs are computed on the basis of sequence to structure alignment and are cluster associated (Cluster-HMM) (Figure 1). When different templates are present in a cluster the structural alignment among them is computed with MUSTANG (13). Multiple alignments comprising all the overlapping templates and the sequences similar to them (with  $SI \geq 40\%$  and  $COV \geq 90\%$ ) are computed with MUSCLE (14) and fed to HMMER 2.3 (15) in order to train the profile-HMM. By this, a library of 10 858 HMMs is made available for aligning even distantly related sequences to a given PDB template/s. The server also provides the pairwise query sequence–structural target alignment computed with the Viterbi decoding implemented in HMMER from the correspondent Cluster-HMM and useful for further processing and/or computing the corresponding 3D structure.

### DIFFERENT ANNOTATIONS with BAR<sup>+</sup>

BAR<sup>+</sup> allows 35 possible fine grain types of annotations (plus no annotation) (Table 1). The most complete type of annotation is the one with PDB (with and without SCOP annotation) and GO terms and Pfam domains with  $P \leq 0.01$  (validated) (first row in Table 1). Interestingly, enough 0.11% of the total sequences in our database are sufficient to annotate in a validated manner and with the most complete annotation another 21.99% sharing common clusters (8251; 0.90% of the total), with an annotation gain factor higher than 200. Summing up (along the first row of Table 1), we can conclude that validated functional annotation is possible within 10% of the clusters. Eleven percent of the sequences remains without annotation and are included in 45% of the clusters. About 57% of singletons (corresponding to 17% of the total set) are annotated with different features (Figure 2B and Table 1).

### SUBMITTING A PROTEIN SEQUENCE TO BAR<sup>+</sup>

When a query sequence is submitted, there are three possible outcomes (Figure 3). The sequence can match a sequence already present in the cluster (or in a singleton). By this, non-annotated proteins can inherit functional and structural annotation from other proteins within the same cluster. Validated annotations are inherited when clusters are endowed with validated GO and Pfam ( $P < 0.01$ ). Alternatively a BLAST alignment starts. The query sequence may then align with any other sequence in BAR<sup>+</sup> with the stringent criteria of our procedure and, therefore, find a cluster from where it can safely inherit all the corresponding structural and functional features.

**Table 1.** The fine grain types of annotation with BAR<sup>+</sup>

	PDB (%)	SCOP Mono	SCOP Multi	Without PDB
GO validated				
Pfam validated				
Clusters	8251 (0.90)	3613 (0.40)	1461 (0.16)	83 266 (9.11)
Sequences	2 982 449 (22.10)	1 408 542 (10.44)	1 028 565 (7.62)	2 903 431 (21.51)
<b>Inherited</b>	<b>2 967 743 (21.99)</b>	<b>1 404 011 (10.40)</b>	<b>1 026 154 (7.60)</b>	<b>1 382 310 (10.24)</b>
Pfam				
Clusters	8334 (0.91)	3647 (0.40)	1463 (0.16)	85 886 (9.40)
Sequences	2 984 057 (22.11)	1 409 647 (10.45)	1 028 569 (7.62)	2 922 876 (21.66)
<b>Inherited</b>	<b>2 969 285 (22.00)</b>	<b>1 405 095 (10.41)</b>	<b>1 026 156 (7.60)</b>	<b>1 398 603 (10.36)</b>
Without Pfam				
Clusters	320 (0.04)	123 (0.01)	25 <sup>a</sup>	6251 (0.68)
Sequences	42 202 (0.31)	15 415 (0.11)	7363 (0.05)	143 533 (1.06)
<b>Inherited</b>	<b>41 825 (0.31)</b>	<b>15 303 (0.11)</b>	<b>7331 (0.05)</b>	<b>93 568 (0.69)</b>
GO				
Pfam validated				
Clusters	8938 (0.98)	3887 (0.43)	1504 (0.16)	133 895 (14.65)
Sequences	3 042 649 (22.55)	1 450 437 (10.75)	1 029 707 (7.63)	3 311 421 (24.54)
<b>Inherited</b>	<b>3 026 916 (22.43)</b>	<b>1 445 521 (10.71)</b>	<b>1 027 219 (7.61)</b>	<b>1 617 763 (11.99)</b>
Pfam				
Clusters	9357 (1.02)	4033 (0.44)	1526 (0.17)	322 937 (35.34)
Sequences	3 045 465 (22.57)	1 451 928 (10.76)	1 029 755 (7.63)	3 739 076 (27.71)
<b>Inherited</b>	<b>3 029 337 (22.45)</b>	<b>1 446 890 (10.72)</b>	<b>1 027 247 (7.61)</b>	<b>1 852 223 (13.72)</b>
Singletons	2608 (0.02)	10 <sup>a</sup>	5 <sup>a</sup>	1 515 720 (11.23)
Without Pfam				
Clusters	452 (0.05)	176 (0.02)	30 <sup>a</sup>	45 539 (4.98)
Sequences	46 311 (0.34)	17 020 (0.13)	7400 (0.05)	330 354 (2.45)
<b>Inherited</b>	<b>45 803 (0.34)</b>	<b>16 862 (0.12)</b>	<b>7362 (0.05)</b>	<b>226 500 (1.68)</b>
Singletons	279 <sup>a</sup>	2 <sup>a</sup>	2 <sup>a</sup>	129 212 (0.96)
Without GO				
Pfam validated				
Clusters	679 (0.07)	345 (0.04)	15 <sup>a</sup>	54 314 (5.94)
Sequences	44 172 (0.33)	27 775 (0.21)	654 <sup>a</sup>	547 459 (4.06)
<b>Inherited</b>	<b>43 416 (0.32)</b>	<b>27 410 (0.20)</b>	<b>633<sup>a</sup></b>	<b>221 585 (1.64)</b>
Pfam				
Clusters	779 (0.09)	377 (0.04)	16 <sup>a</sup>	122 236 (13.38)
Sequences	44 582 (0.33)	27 983 (0.21)	656 <sup>a</sup>	695 684 (5.15)
<b>Inherited</b>	<b>43 735 (0.32)</b>	<b>27 592 (0.20)</b>	<b>634<sup>a</sup></b>	<b>301 792 (2.24)</b>
Singletons	205 <sup>a</sup>	1 <sup>a</sup>	0 <sup>a</sup>	702 834 (5.21)
Without Pfam				
Clusters	270 (0.03)	83 (0.01)	5 <sup>a</sup>	412 192 (45.11)
Sequences	5308 (0.04)	1771 (0.01)	154 <sup>a</sup>	1 494 443 (11.07)
<b>Inherited</b>	<b>5023 (0.04)</b>	<b>1689 (0.01)</b>	<b>149<sup>a</sup></b>	
Singletons	129 <sup>a</sup>	1 <sup>a</sup>	0 <sup>a</sup>	1 743 526 (12.92)

Percentage is evaluated with respect to the total number of sequences in the data base (13 495 736 sequences). Bold character: sequences that inherit the annotation type

<sup>a</sup>Values are negligible. Validated:  $P \leq 0.01$  (See text for details, 11). Within BAR<sup>+</sup> clusters, 35 different types of annotations are possible: (i) +GO+Pfam+PDB [with or without SCOP (Monodomain, Multidomain)\*]; GO and Pfam are or not validated (no. of levels = 12). (ii) +Pfam+PDB (with or without SCOP)\* (no. of levels = 6). (iii) +GO+PDB (with or without SCOP)\* (number of levels = 6). (iv) +Pfam+GO (no. of levels = 4). (v) +PDB (with or without SCOP)\* (number of levels = 3). (vi) +GO (no. of levels = 2). (vii) +Pfam (no. of levels = 2). Seventy percent of the initial set fall into clusters (913 962) and 53% in validated clusters. Some 6% of the sequences are annotated without validation and the remaining 11% are not annotated (rightmost bottom cell). About 17 and 13% of the sequences are singletons with and without annotations, respectively.

Alternatively, when the criteria are not met, all the BLAST matches are returned. This allows anyway locating the sequence within a cluster. However, in this case, annotation through inheritance should be manually curated. Singletons may be or not source of information depending on their annotation.

## BAR<sup>+</sup> UPDATE

BAR<sup>+</sup> collects sequences and their features from UniProtKB and genome repositories. Our re-clustering is programmed on a yearly base. BAR<sup>+</sup> cluster annotation

will be updated every 6 months. This is based on the notion that indeed the BAR<sup>+</sup> annotation system increases its capacity only when we add information. This is achieved when proteins with evidence at the transcript and protein level (e.g.: PDB new files and/or proteins with GO/Pfam terms) are included in the system. For example, by comparing UniprotKB 05\_2010 with SwissProt 03\_2011, we collected some 2445 sequences carrying information according to our criteria (evidence at protein/transcript level). By aligning this set towards BAR<sup>+</sup> clusters, we find that 62% of the sequences fall into already validated clusters. About 8% aligns with singletons and only 0.03% of the total number of BAR<sup>+</sup>

**BAR+ matching table (SI ≥ 40% and Coverage ≥ 90%)**

**Cluster:** Cluster identification, Annotations (GO,PDB,Pfam), Cluster HMM (when possible)  
**Query:** Your input, Alignment to template/s (when possible)  
**SI:** Pairwise Sequence Identity  
**COV:** Overlapping of the pairwise alignment as defined in the Help  
**BLAST Match:** Sequences that match your query, UniProt code (if available) or other description  
**cov:** Overlapping of the PDB template over the query (if available)  
**id:** PDB template/query sequence identity (if available)

**GREEN:** Identical sequences  
**RED:** SI ≥ 40% and COV ≥ 90%  
**GREY:** SI < 40% and COV < 90%

BLAST matching sequences. Sequence Identity (SI) and Coverage (COV) are displayed. A brief description is also provided

Cluster	Query	SI	COV	BLAST Match	
<p><b>Cluster-ID: 5516</b></p> <p>No. sequences: 195 Average length: 712.0 (6.9% SD)</p> <ul style="list-style-type: none"> <li>Eukaryota 195</li> </ul> <p><b>Gene Ontology function (Total: 203)</b></p> <p><b>Molecular Function:</b> (Validated:6/18)</p> <ul style="list-style-type: none"> <li>heparin binding</li> <li>serine-type endopeptidase inhibitor activity</li> <li>endopeptidase inhibitor activity</li> <li>peptidase inhibitor activity</li> <li>carbohydrate binding</li> <li>enzyme regulator activity</li> </ul> <p><b>Biological Process:</b> (Validated:107/164)</p> <ul style="list-style-type: none"> <li>smooth endoplasmic reticulum calcium ion homeostasis</li> <li>calcium ion homeostasis</li> <li>positive regulation of transcription from RNA polymerase II promoter</li> <li>mRNA polyadenylation</li> <li>copper ion homeostasis</li> </ul> <p>● developmental process ● multi-organism process</p> <p><b>Cellular Component:</b> (Validated:1/1)</p> <ul style="list-style-type: none"> <li>integral to membrane</li> <li>Golgi apparatus</li> <li>insoluble fraction</li> <li>cell projection</li> <li>membrane-bounded organelle</li> </ul> <p><b>Structural annotation (No. PDB: 8)</b></p> <ul style="list-style-type: none"> <li>2yszA 1oqnC 2wk3C 1nmjA 3dxeB 1rw6A 1aapA 3ktmA</li> <li>Ligands: BU4</li> <li>SCOP: j.42.1.1 # a.47.4.1 # g.8.1.1</li> </ul> <p><b>Pfam annotation (Total: 1)</b></p> <ul style="list-style-type: none"> <li>beta-amyloid precursor protein C-terminus(APP_amyloid)</li> </ul> <p><b>Cluster HMM</b></p> <p>Download HMM</p> <p>Download <a href="#">cluster annotation data</a></p>	<p><b>D2GUX3</b></p> <p><b>2wk3C</b> 696-728 (start-end) 4.5% cov   70.0% id</p> <ul style="list-style-type: none"> <li>PIR alignment</li> </ul> <p><b>1rw6A</b> 369-573 (start-end) 28.2% cov   60.0% id</p> <ul style="list-style-type: none"> <li>PIR alignment</li> </ul> <p><b>3ktmA</b> 19-190 (start-end) 23.6% cov   60.0% id</p> <ul style="list-style-type: none"> <li>PIR alignment</li> </ul> <p><b>1aapA</b> 291-347 (start-end) 7.8% cov   60.0% id</p> <ul style="list-style-type: none"> <li>PIR alignment</li> </ul>	<p><b>97.4</b></p>	<p><b>100.0</b></p>	<p><b>613_GeneScaffold.466_3620</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Felis catus <b>Description:</b> AMYLOID BETA A4 PRECURSOR APP ABPP ALZHEIMER DISEASE AMYLOID [CONTAINS: SOLUBLE APP ALPHA S APP ALPHA ; SOLUBLE APP BETA S APP BETA ; BETA AMYLOID 42 BETA APP42 ; BETA AMYLOID 40 BETA APP40 ; C83 ; P3 42 ; P3 40 ; GAMMA CTF 59 GAMMA SECRETASE C TERMINAL FR</p> <hr/> <p><b>717_7_14778</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Equus caballus <b>Description:</b> ENSECAP00000019671 pep:novel chromosome:EquCab2:7:38599732:38634456:1 gene:ENSECAG00000021265 transcript:ENSECAT00000023727</p> <hr/> <p><b>616_11_16870</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Homo sapiens <b>Description:</b> AMYLOID BETA A4 PRECURSOR APP ABPP ALZHEIMER DISEASE AMYLOID [CONTAINS: SOLUBLE APP ALPHA S APP ALPHA ; SOLUBLE APP BETA S APP BETA ; BETA AMYLOID 42 BETA APP42 ; BETA AMYLOID 40 BETA APP40 ; C83 ; P3 42 ; P3 40 ; GAMMA CTF 59 GAMMA SECRETASE C TERMINAL FR</p> <hr/> <p><b>Q60709</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Mus musculus (Mouse) <b>Description:</b> Amyloid-like protein 2, isoform 751</p> <hr/> <p><b>Q3UDL6</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Mus musculus (Mouse) <b>Description:</b> Putative uncharacterized protein</p> <hr/> <p><b>621_GeneScaffold.695_6960</b></p> <p><b>Taxonomy:</b> Eukaryota <b>Organism:</b> Myotis lucifugus <b>Description:</b> AMYLOID BETA A4 PRECURSOR APP ABPP ALZHEIMER DISEASE AMYLOID [CONTAINS: SOLUBLE APP ALPHA S APP ALPHA ; SOLUBLE APP BETA S APP BETA ; BETA AMYLOID 42 BETA APP42 ; BETA AMYLOID 40 BETA APP40 ; C83 ; P3 42 ; P3 40 ; GAMMA CTF 59 GAMMA SECRETASE C TERMINAL FR</p>	
			95.5	100.0	
			95.5	95.4	
			95.3		
		90.5	95.3		
		89.4	100.0		

Query column with the list of query/templates alignments (when available)

Cluster summary and list of cluster-specific annotations (GO,Pfam,PDB,Ligands, SCOP,HMM)

**Figure 3.** BAR+ at work. A query sequence has been submitted. Provided that the sequence after running BLAST has a level of SI ≥ 40% with a COV ≥ 90% to any sequence of BAR+, it is included into a cluster. In the above example, the cluster is well annotated and the sequence inherits all the possible annotations from the cluster including GO terms (203), PDB/s, ligands, SCOP and Pfam annotations and the Cluster-HMM. Furthermore in PIR format alignment/alignments of the query sequence to the cluster template/s with Cluster HMM is/are also provided. All the sequences that align with the query are returned. (●●●) Only the top and bottom portions of the page are shown.

singletons become new clusters (with two protein sequences). Another 7% fall into non-validated clusters without affecting the statistical significance of the cluster-specific annotation. The remaining 23% originate new singletons. We are currently planning to include other annotation resources in order to extend our annotation process with more protein domains and their interactions.

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*Conflict of interest statement.* None declared.

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