CDD/SPARCLE: the conserved domain database in 2020

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ABSTRACT

As NLM's Conserved Domain Database (CDD) enters its 20th year of operations as a publicly available resource, CDD curation staff continues to develop hierarchical classifications of widely distributed protein domain families, and to record conserved sites associated with molecular function, so that they can be mapped onto user queries in support of hypothesisdriven biomolecular research. CDD offers both an archive of pre-computed domain annotations as well as live search services for both single protein or nucleotide queries and larger sets of protein query sequences. CDD staff has continued to characterize protein families via conserved domain architectures and has built up a significant corpus of curated domain architectures in support of naming bacterial proteins in RefSeq. These architecture definitions are available via SPARCLE, the Subfamily Protein Architecture Labeling Engine. CDD can be accessed at https://www.ncbi.nlm.nih.gov/Structure/ cdd/cdd.shtml.

CDD CONTENT

At the time of writing, the CDD version v3.17 is the live production version with 52,910 protein- and protein domainmodels obtained from Pfam (1), SMART (2), the COGs collection (3), TIGRFAMS (4), the NCBI Protein Clusters collection (5), NCBIfam (6) and CDD's in-house data curation effort (7). CDD version v3.18 will be released in the winter 2019/2020 and will include Pfam version 32 and a total of 55 434 protein and protein-domain models. For CDD v3.18, the fixed assumed size of the domain model database has again been increased to match the current size of the model collection, resulting in marginally higher E-values reported by RPS-BLAST (8). The NCBIfam collection in CDD is a set of models derived from HMMs that have been developed for improving the annotation of bacterial genomes. Currently, CDD excludes NCBIfam models that were built to identify proteins involved in antimicrobial resistance, due to their narrow scope.

Table 1 shows the 20 largest classifications for common and functionally diverse domain families that have recently been updated or added to CDD. In total, over 4,700 models curated by the CDD group have been newly published or updated since CDD release v3.16. At the time of the CDD v3.17 release, CDD annotated about 85% of the sequences in the Entrez/protein database (excluding sequences from environmental sampling). CDD also covered about 94% of the protein sequences (longer than 30 residues) derived from protein 3-dimensional structures as provided via MMDB. CDD curation staff monitors structure-derived sequences that do not yet have coverage in CDD for novel protein domain families with wide taxonomic distribution and generates corresponding domain family models de novo.

For CDD v3.18, a total of 33 980 site annotations are available on 12 418 out of 16 069 CDD staff-curated domain models. Sequence patterns have been recorded for 3250 of these site annotations, so that pattern matches determine whether a site annotation is being mapped onto a query sequence.

SPARCLE

Protein domain architectures can be defined as a sequential (N- to C-terminal) list of one or more domain footprints annotated on a protein sequence. In CDD, we distinguish between superfamily architectures (where hits to several different models that are redundant or related to each other are treated as the same superfamily hit) and specific or subfamily domain architectures (SDAs), where high-confidence (specific) domain annotation is taken into consideration. The CDART (Conserved Domain Architec-

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This work is written by (a) US Government employee(s) and is in the public domain in the US.

Table 1. The largest domain family hierarchies created or updated since CDD release v3.16
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Root	models	Name	
cd14964	589	Seven-transmembrane G protein-coupled receptor superfamily	
cd00196	315	Beta-grasp ubiquitin-like fold	
cd01165	242	BTB/POZ domain superfamily	
cd00083	202	basic Helix Loop Helix (bHLH) domain superfamily	
cd01391	192	Type 1 periplasmic binding fold superfamily	
cd06174	187	Major Facilitator Superfamily	
cd14494	172	Cys-based protein tyrosine phosphatase and dual-specificity phosphatase superfamily	
cd17912	169	N-terminal helicase domain of the DEAD-box helicase superfamily	
cd09852	137	PIN (PilT N terminus) domain superfamily	
cd02208	113	RmlC-like cupin superfamily	
cd00021	97	B-box-type zinc finger superfamily	
cd06660	96	Aldo-keto reductase (AKR) superfamily	
cd14733	93	BACK (BTB and C-terminal Kelch) domain	
cd08161	86	SET (Su(var)3–9, Enhancer-of-zeste, Trithorax) domain superfamily	
cd04433	85	Adenylate forming domain, Class I superfamily	
cd03873	82	Zinc peptidases M18, M20, M28, and M42	
cd00156	82	phosphoacceptor receiver (REC) domain of response regulators/pseudo response regulators	
cd16961	77	Type I restriction-modification system specificity (S) subunit Target Recognition Domain	
cd07346	75	Six-transmembrane helical domain of the ATP-binding cassette transporters	
cd00172	74	SERine Proteinase INhibitors (serpin) family	
cd00301	71	lipocalin/cytosolic fatty acid-binding protein family	
cd00048	69	double-stranded RNA binding motif (DSRM) superfamily	

The table lists the root node of each hierarchy, the number of models in the hierarchy (including the root node and intermediate nodes if present), and the name of the protein domain (super)family.

sigma-54-dependent Fis family transcriptional regulator [Pseudomonas indica]

Sigina-04-0	rependent i is fanning transcriptional regulator	I li seudoi	nonus ma	licaj				
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Identical Proteins FASTA Graphics			bi.nlm.nih.gov/Structure/sp	< < 				
<u>Go to:</u> 🕑			ional Library of Me	Logi				
FINITION sigm [Pse cESSION WP_0 RSION WP_0 YWORDS Ref5 URCE Pseu ORGANISM Pseu MMENT REFS sequ from	udomonas indica udomonas indica teria; Proteobacteria; Gammaproteobacteria; Pseudomonadales; udomonadaceae; Pseudomonas. SEQ: This record represents a single, non-redundant, protein uence which naw by non-third on many different RefSeq genomes one same, or different, species.	sigma-54 regulator sigma-54-deper which may be a caulinodans nit	Architecture View 4 – depende 4 – depende 5 1 – depende 1 – depen	Table of Contents Description Sequences with this architecture Name, label and taxonomic scope Conserved domains				
<pre>##Evidence-For-Name-Assignment-START## Evidence Category :: Conserved Domain (CDO) Evidence Accession :: Domain architecture ID 11454220 Evidence Source :: ICBI SPARCLE ##Evidence-For-Name-Assignment-END## COMPLETENESS: full length.</pre>		Arch. ID 🔞	Version 🕲	Date Published 🗭	Review Level 🗈	Actions Download JSON data for this architecture		
source	Location/Qualifiers 1305 "Dimenism="Pseudomonas indice"			AtoC		Similar superfamily architectures		
Protein	<pre>/db_xref="taxon:<u>117658</u>" 1305 /product="sigma-54-dependent Fis family transcriptional regulator" /calculated mol_wt=33022</pre>	Sequences w	rith this architectu	ure		New search by architecture ID or sequence		
Region	<pre>/calculated_moi_wt=>3022 (27.298 /region_name="AtoC" /note="DNA-binding transcriptional response regulator, NtrC family, contains REC, AAA-type ATPase, and a Fis-type DNA-binding domains [Signal transduction mechanisms]; CC062204" /db_xref="CDD:225114"</pre>	Swiss-Pro	Otein w/ PubMed Rafereno t Annotated (73)	SPARCLE downloads Search SPARCLE				
	spall tlsdadrspl siraralvfa dptsqrlrae verlaastlp vliqgetgsg rhihr ssdrqglfva vncgalspkh aeaelfghaa garagtagra gwfgsanggs	Identifier WP_097572	Descriptio sigma-54-dependent Fis ranscriptional regulator, p	family Mesorhizobiur				

Figure 1. 'flatfile' (GenPept format) view of a bacterial protein from the RefSeq collection. Bacterial of proteins with the accession prefix 'WP' are now being equipped with evidence for the name assignment (highlighted with a red oval). Evidence accessions are hot-linked to provide more information about the specific annotation rule, in this case a conserved domain architecture curated in SPARCLE. Other evidence types with hotlinks to an annotation rule viewer are Hidden Markov Models (HMMs) and BLAST rules, which have higher precedence than domain architectures and will overrule the name suggested by SPARCLE.

Table 2. URLs and o	other resources	associated	with the	CDD	project
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URL	Description			
https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi	CD-Search interface utilizing the RPS-BLAST algorithm and the model database, and to the CDART database of pre-computed domain annotation			
https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi	BATCH CD-Search interface utilizing the RPS-BLAST algorithm and the model database, and to the CDART database of pre-computed domain annotation. Up to 4000 protein queries may be submitted per request			
https://www.ncbi.nlm.nih.gov/cdd	Entrez interface to CDD			
https://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml	CDD project home page			
https://www.ncbi.nlm.nih.gov/Structure/lexington/lexington.cgi	CDART domain architecture viewer			
https://ftp.ncbi.nih.gov/pub/mmdb/cdd	CDD FTP site, see README file for content			
https://www.ncbi.nlm.nih.gov/Structure/cdtree/cdtree.shtml	Domain hierarchy editor/viewer and protein structure/alignment viewer			
https://ftp.ncbi.nlm.nih.gov/toolbox executables can be obtained	RPS-BLAST stand-alone tool for searching databases of			
from: https://www.ncbi.nlm.nih.gov/BLAST/download.shtml	profile models, part of the NCBI toolkit distribution			
https://www.ncbi.nlm.nih.gov/sparcle	Entrez interface to SPARCLE (Subfamily Protein Architecture Labeling Engine)			
https://ftp.ncbi.nlm.nih.gov/pub/mmdb/cdd/rpsbproc	Standalone utility for enriching and formatting RPS-BLAST results			
https://ftp.ncbi.nlm.nih.gov/pub/mmdb/cdd/SparcleLabel/	Standalone utility for naming/labeling proteins using SPARCLE			

ture Retrieval Tool) service (9) groups proteins in the Entrez database by common domain superfamily architecture. SPARCLE for 'Subfamily Protein Architecture Labeling Engine', on the other hand, groups proteins by SDA, and we have engaged in a curation effort that reviews SDAs that are well-represented in the protein sequence collection and associates them with protein name suggestions and short functional descriptions. To date, CDD curators have assigned names and functional labels to ~25 000 SDAs, with a focus on SDAs common in bacterial genomes. A publicly accessible Entrez database supports text queries and points to summary information for SDAs as well as links to other databases, most importantly the NCBI protein collection.

CD-Search displays not only domain and feature annotation, but also the name and functional characterization assigned to the corresponding SDA, if available via SPAR-CLE.

The SPARCLE curation effort is focused on architectures common in bacteria, and supports the automated, evidence-based assignment of names to proteins in Ref-Seq and the Prokaryotic Genome Annotation Pipeline (PGAP)6. Protein names provided by the curated subset of SPARCLE have relatively low preference in the hierarchy of naming evidence sources but cover a lot of ground and often provide the only suggestion available for a gene product. At this time, about 42 million bacterial RefSeq proteins are named via SPARCLE (out of 126 million total bacterial proteins and 92 million proteins with naming evidence). Figure 1 shows an example of how naming evidence is currently being displayed by the sequence 'flatfile' (GenPept format) viewer.

DATA AVAILABILITY

Table 2 lists URLs for services, tools, and data collections provided by CDD. RPS-BLAST is part of NCBI's BLAST software distribution. Pre-formatted RPS-BLAST search databases are available so that conserved domain searches can be run locally, and the results can be formatted with the rpsbproc utility so that they correspond to reports generated by CD-Search (10) and BATCH CD-Search, including site annotations. A new utility, sparclbl (SparcleLabel) is available via FTP; sparclbl processes results from local RPS-BLAST searches and provides suggestions for protein names based on domain architecture. An in-house version of sparclbl is part of NCBI's prokaryotic genome annotation pipeline (PGAP) (6).

CDD shares domain models with the InterPro at the European Bioinformatics Institute to supplement sequence annotation with data that are uniquely provided by the CDD curation effort, including protein domain models for very specific subfamilies and the annotation of functional sites. To date, >3100 domain signatures provided by CDD have been integrated by InterPro (11).

FUTURE WORK

The CDD group is investigating whether model-specific word-score thresholds can be applied when building RPS-BLAST search databases and help speed searching while keeping the loss of annotation at a minimum. Instructions for how to use such a search set will be announced via the CDD news page at https://www.ncbi.nlm.nih.gov/Structure/cdd/docs/cdd_news.html, once available.

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Comments, suggestions, and questions are welcome and should be directed to: info@ncbi.nlm.nih.gov.

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REFERENCES

- El-Gebali, S., Mistry, J., Bateman, A., Eddy, S.R., Luciani, A., Potter, S.C., Qureshi, M., Richardson, L.J., Salazar, G.A., Smart, A. *et al.* (2019) The Pfam protein families database in 2019. *Nucleic Acids Res.*, 47, D427–D432.
- 2. Letunic, I. and Bork, P. (2018) 20 years of the SMART protein domain annotation resource. *Nucleic Acids Res.*, **46**, D493–D496.
- Tatusov,R.L., Natale,D.A., Garkavtsev,I.V., Tatusova,T.A., Shankavaram,U.T., Rao,B.S., Kiryutin,B., Galperin,M.Y., Fedorova,N.D. and Koonin,E.V. (2001) The COG database: new developments in phylogenetic classification of proteins from complete genomes. *Nucleic Acids Res.*, 29, 22–28.
- 4. Haft,D.H., Selengut,J.D., Richter,A.R., Harkins,D., Basu,M.K. and Beck,E. (2013) TIGRFAMs and genome properties in 2013. *Nucleic Acids Res.*, **41**, D387–D395.

- Klimke, W., Agarwala, R., Badretdin, A., Chetvernin, S., Ciufo, S., Fedorov, B., Kiryutin, B., O'Neill, K., Resch, W., Resenchuk, S. *et al.* (2009) The National center for biotechnology information's protein clusters database. *Nucleic Acids Res.*, 37, D216–D223.
- Haft,D.H., DiCuccio,M., Badretdin,A., Brover,V., Chetvernin,V., O'Neill,K., Li,W., Chitsaz,F., Derbyshire,M.K., Gonzales,N.R. *et al.* (2018) RefSeq: an update on prokaryotic genome annotation and curation. *Nucleic Acids Res.*, 46, D851–D860.
- Marchler-Bauer, A., Bo, Y., Han, L., He, J., Lanczycki, C.J., Lu, S., Chitsaz, F., Derbyshire, M.K., Geer, R.C., Gonzales, N.R. *et al.* (2017) CDD/SPARCLE: functional classification of proteins via subfamily domain architecture. *Nucleic Acids Res.*, 45, D200–D203.
- Marchler-Bauer, A., Panchenko, A.R., Shoemaker, B.A., Thiessen, P.A., Geer, L.Y. and Bryant, S.H. (2002) CDD: a database of conserved domain alignments with links to domain three-dimensional structure. *Nucleic Acids Res.*, **30**, 281–283.
- Geer, L.Y., Domrachev, M., Lipman, D.J. and Bryant, S.H. (2002) CDART: protein homology by domain architecture. *Genome Res.*, 12, 1619–1623.
- Marchler-Bauer, A. and Bryant, S.H. (2005) CD-Search: protein domain annotations on the fly. *Nucleic Acids Res.*, 32, W327–W331.
- Mitchell,A.L., Attwood,T.K., Babbitt,P.C., Blum,M., Bork,P., Bridge,A., Brown,S.D., Chang,H.Y., El-Gebali,S., Fraser,M. *et al.* (2019) InterPro in 2019: improving coverage, classification and access to protein sequence annotations. *Nucleic Acids Res.*, 47, D351–D360.