

# Genome Databases

## The CATH database

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### Abstract

The CATH database provides hierarchical classification of protein domains based on their folding patterns. Domains are obtained from protein structures deposited in the Protein Data Bank and both domain identification and subsequent classification use manual as well as automated procedures. The accompanying website ([www.cathdb.info](http://www.cathdb.info)) provides an easy-to-use entry to the classification, allowing for both browsing and downloading of data. Here, we give a brief review of the database, its corresponding website and some related tools.

**Keywords:** CATH, protein domains, classification, protein structure, database

### Introduction

The number of solved protein structures is increasing at an exceptional rate. At the time of writing, the Protein Data Bank<sup>1,2</sup> (PDB) contains more than 61,000 structures. The CATH database<sup>3,4</sup> is a classification of protein domains (sub-sequences of proteins that may fold, evolve and function independently of the rest of the protein), based not only on sequence information, but also on structural and functional properties. CATH offers an important tool to researchers, as proteins with even very little sequence similarity often are both structurally and functionally related.<sup>5</sup>

The most recent version of CATH (version 3.2.0, released July 2008<sup>6</sup>) contains 114,215 domains, classified in a hierarchical scheme with four main levels (listed from the top and down) called class (C), architecture (A), topology (T) and homologous superfamily (H) — hence the name CATH. More than 20,000 domains have been added since the previous release (version 3.1.0, January 2007), and the rate of new additions is

expected to increase. (The first CATH release<sup>3</sup> from 1997 contained only 8,078 domains.)

At the C-level, domains are grouped according to their secondary structure content into four categories: mainly alpha, mainly beta, mixed alpha-beta; and a fourth category which contains domains with only few secondary structures. The A-level groups domains according to the general orientations of their secondary structures. At the T-level, the connectivity (ie the order) of the secondary structures is taken into account. The grouping of domains at the H-level is based on a combination of both sequence similarity and a measure of structural similarity obtained from the dynamic programming algorithm SSAP.<sup>7</sup> To supplement the traditional alignment of the  $\alpha$ -carbon atoms of the protein backbone, SSAP gains additional strength by also aligning  $\beta$ -carbon atoms of the amino acid side chains and thus also takes into account the rotational conformation of the protein chains.

In addition to the four main levels, CATH comprises five more layers, called S, O, L, I and D. The first four layers group domains according to

increasing sequence overlap and similarity (eg two domains with the same CATHSOLI classification must have 80 per cent overlap, with 100 per cent sequence identity), whereas the D-level assigns a unique identifier to every domain, thus ensuring that no two domains have exactly the same CATHSOLID classification.

A combination of automated procedures and manual inspections are used in the CATH classification. In particular, at the A-level, similarity is difficult to detect using automated methods only.

Other similar databases are available online.<sup>8</sup> Among these, SCOP<sup>9</sup> is the most widely used, and by being a hierarchical classification too, it provides a supplement to CATH. Despite their hierarchical architectures, the two databases are not entirely comparable. For example, at the class level, SCOP contains two mixed alpha-beta classes; the  $\alpha + \beta$  class comprises domains with mostly antiparallel  $\beta$ -sheets and segregated  $\alpha$ - and  $\beta$ -regions, while the  $\alpha/\beta$  class comprises domains with many parallel  $\beta$ -sheets and  $\beta$ - $\alpha$ - $\beta$  units. It is still possible to compare CATH and SCOP, however — for example, in a recent study,<sup>10</sup> where a consensus set on which the hierarchical structures of both databases agree was extracted. The consensus set contained 64,016 domains, which amounts to 56 per cent of the domains in CATH.

Various other databases exist that are non-hierarchical and use more standard clustering methods. Among these, the most widely cited are DALI,<sup>11</sup> HOMSTRAD<sup>12</sup> and COMPASS.<sup>13,14</sup>

## Organisation of the CATH homepage

The CATH homepage (<http://www.cathdb.info/>) provides easy access to the CATH classification. The first site element contains a quick description of CATH, with a link to a more thorough introduction. The language is very non-technical and the reader can quickly grasp the overall structure of CATH; more details are provided by Greene *et al.*,<sup>15</sup> for example. Links to more details are provided in the Documentation section in the main menu. A useful glossary of terms and definitions used in CATH is available, alongside a thorough

tutorial on how to use CATH and the related Gene3D server,<sup>16–19</sup> which, by scanning sequences in CATH predicts the domain compositions of proteins from sequences alone. At the time of writing, the Gene3D database comprises more than 10 million protein sequences, from over 1,100 fully sequenced species genomes, from all three kingdoms of life.<sup>19</sup>

## Data accessibility

Besides a Quick Search box, which facilitates easy searching, links are provided to various other ways of accessing the data: (1) search by keyword or domain ID; (2) search using a sequence in FASTA format; (3) browse the database from the top of the hierarchy; and (4) download datasets. The ability to browse the database provides a way to get acquainted with the structure of CATH and is also a convenient way to locate and compare similar structures.

Figure 1 shows an example of what a domain looks like in the CATH browser. The domain 3cx5B01 (chain B, domain 1 of the PDB entry 3cx5) is classified as 3.30.830.10, making it a *Mixed Alpha-Beta* domain (C = 3) in the *2-Layer Sandwich* architecture (A = 30). Besides a picture of the domain's three-dimensional structure, a schematic depiction of the arrangement of secondary structures is shown in the Structure pane, also present in Figure 1. The Sequence pane contains the amino acid sequence of the domain, and the History pane describes the history of the domain in the CATH database, with information about when the domain was added and if the classification has changed over time.

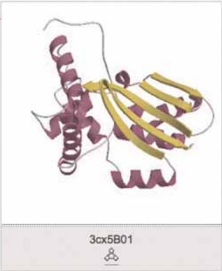
Browsing is not only possible at the domain level. Figure 2 shows the entry corresponding to the *Alpha/alpha barrel* architecture (with CATH classification 1.50). A summary of the lower levels is provided, alongside links to the adjacent sub-levels in the hierarchy — in this case, the two topologies 1.50.10 and 1.50.30. By clicking on a link to a representative domain, an output as in Figure 1 is obtained.

The Download section provides access to various kinds of data. Large compressed archives of chopped PDB files corresponding to representative CATH domains are available. These sets are the

**CATH Domain: 3cx5B01** [XML](#)

PDB 3cx5, Chain B, Domain 1


CATH Code	Level Description	Links
3	Alpha Beta	
3.30	2-Layer Sandwich	
3.30.830	Cytochrome Bc1 Complex; Chain A, domain 1	
3.30.830.10	Cytochrome Bc1 Complex; Chain A, domain 1	[Gene3D]
3.30.830.10.1		
3.30.830.10.1.1		
3.30.830.10.1.1.1		
3.30.830.10.1.1.1.1		
3.30.830.10.1.1.1.1.1		[Gene3D]



3cx5B01

**Structure** | **Sequence** | **History**

Segment boundaries for domain 3cx5B01



Domain ID	Start Res	Stop Res	Name	Length
3cx5B01	17	227		211

**Figure 1.** Screenshot of the domain 3cx5B01 in the CATH browser. The domain is classified as 3.30.830.10, which means that it belongs to the *Mixed Alpha-Beta* class ( $C = 3$ ), the *2-Layer Sandwich* architecture ( $A = 30$ ) and so forth. The CATH Code column allows for easy browsing both up and down levels in the hierarchy, and the Links column provides links to relevant entries in the Gene3D database. An XML file containing all information on the page can be downloaded by clicking on the XML link next to the domain name. The icon below the image links to a structure file in the Rasmol format. The panes in the bottom of the screen provide additional information about the domain. The content of the Structure pane, which contains secondary structure information, is shown in the figure. The Sequence pane contains the amino acid sequence of the domain and the History pane contains the history of the domain in CATH, with information about when the domain was added to the database and whether its classification has changed over time.

**Architecture: 1.50**

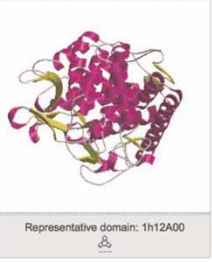
Alpha/alpha barrel

Classification Lineage (1.50)

CATH Code	Level Description	Links
1	Mainly Alpha	
1.50	Alpha/alpha barrel	

Summary of Child Nodes



-	-	2	6	34	43	45	80	390	



Representative domain: 1h12A00

**Summary** | **History**

Topology Entries in Architecture 1.50 (2)

CATH Level	CATH code	Name	Domains	Representative Domain	Representative Thumbnail
1	1.50.10	Glycosyltransferase	370	1h12A00	
1	1.50.30	Isoprenoid Synthase; domain 1	20	1n1bA01	

**Figure 2.** View of the *Alpha/alpha barrel* architecture (CATH classification 1.50) on the CATH website. The Classification Lineage shows the selected architecture is placed in the CATH hierarchy, and the Summary of Child Nodes gives the number of nodes further down. The selected architecture comprises two topologies, 1.50.10 and 1.50.30, shown in the Summary pane below. Direct links to the CATH pages corresponding to the topologies, as well as links to representative domains, are available alongside the topology names. By clicking a link to a representative domain, an output as in Figure 1 is obtained. Navigation on all levels of the CATH hierarchy is facilitated by an analogous page layout.

so-called S100, S95, S60 and S35 sets containing representatives from domain clusters obtained from clusterings based on sequence overlaps and similarities. For example, in the S95 set, two domains must have at least 80 per cent sequence overlap, with 95 per cent sequence identity. Furthermore, files describing how to chop the PDB files of complete proteins, to obtain the domains, can be downloaded; since all PDB files are available at the PDB homepage (<http://www.pdb.org/>), it is possible to construct PDB files of all 114,215 CATH domains by applying the chopping instructions provided in the files. The ability to download complete datasets is of paramount importance for establishing tools like the Gene3D server, discussed above, and, hence, CATH may be seen as more than a resource for acquiring information about single domains only. Furthermore, as CATH is often viewed as a gold standard for automated classification procedures,<sup>20–22</sup> the availability of complete datasets is crucial.

A list containing the names of all domains in CATH — together with their respective classifications — is also available, and the amino acid sequences of all domains classified in CATH are accessible for download in the FASTA format. Finally, the Download section provides a list of 14,652 putative domains that have not yet been assigned a classification, let alone been verified as genuine domains. This dataset may be a valuable ingredient in any development of new, automated classification methods.

## Tools

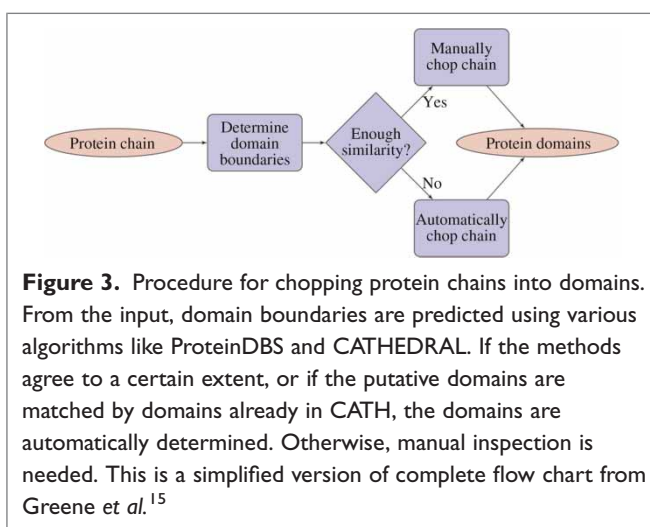
The main menu located in the upper right corner of the homepage links to various tools for use in combination with the CATH database.

1) The **sequential structure alignment program (SSAP)** server<sup>7</sup> takes as input two domains, either provided as PDB/CATH identifiers or as uploaded files, and performs a structural alignment. This allows the user also to **compare domains by structural similarity, rather than sequence homology only**. The SSAP algorithm is computationally feasible; it is a dynamic programming algorithm, like the familiar algorithms for sequence alignment. In

this way, SSAP is able to align not only the  $\alpha$ -carbon atoms of the protein backbones, but also the  $\beta$ -carbon atoms of the amino acid side chains. The output shows the alignment, together with **SSAP score, root mean square deviation (RMSD), overlap and sequence identity**. It is also possible to download a PDB file with the two structures superposed to facilitate additional visual inspection.

2) The **CATHEDRAL server**<sup>23</sup> is used for **discovering known domains in new multi-domain structures**. By either entering a CATH/PDB identifier or by uploading a PDB file, an automated assignment of domain boundaries is performed by querying the structure against a set of representative domains from CATH. This task is accomplished using a modified version of the SSAP algorithm, and the output is a **list of candidate domains ordered according to increasing E-value**. Furthermore, **CATHEDRAL score, SSAP score and RMSD are reported for each candidate**.

3) When a structure has been selected in the CATH browser (see Figure 1), links to the **Gene3D server**<sup>16–19</sup> are also available. For example, clicking the **Gene3D link next to the D-level 3.30.830.10.1.1.1.1.1** presents the Gene3D entry corresponding to the domain 3cx5B01 (recall that any full CATHSOLID classification uniquely defines a domain). From there, several links are available to lists of, for example, **complexes, pathways and functional categories (GO)** in which the domain is involved.



Both CATHEDRAL and SSAP allow the user to sign up for optional e-mail notifications regarding the progress of queries.

## Database construction

The data in CATH are obtained from PDB files deposited in the Protein Data Bank.<sup>1–2</sup> Only structures determined with a resolution of 4Å or better are included. Furthermore, CATH requires the domains to be of minimum 40 residues in length, with 70 per cent or more of the side chains resolved.<sup>15</sup> As mentioned in the introduction, the most recent version of CATH contains 114,215 domains, processed from the proteins in PDB.

Two main steps are involved in adding new structures to CATH: 1) submitted protein chains are chopped to obtain the domains; and 2) classifications are assigned to the resulting domains.

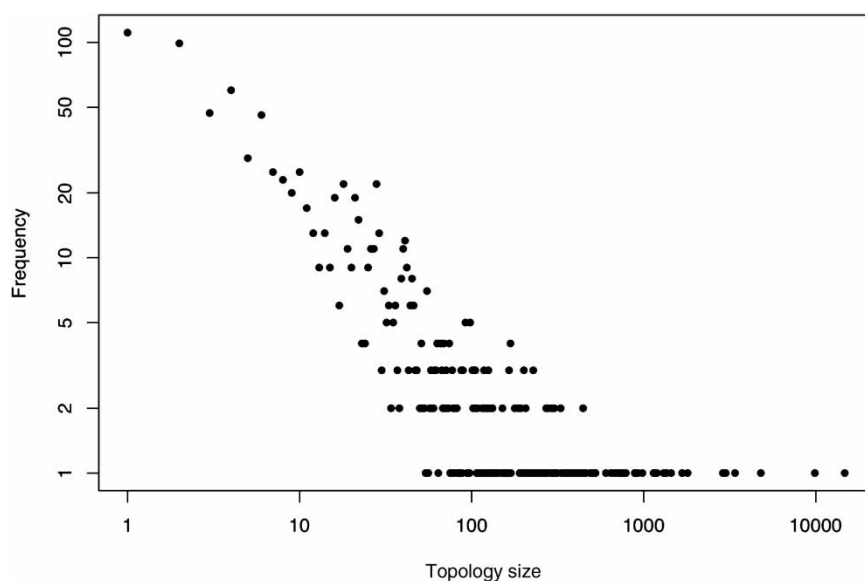
The chopping of protein chains is far from an easy task, and several different measures obtained from, for example, CATHEDRAL<sup>23</sup> and ProteinDBS<sup>24</sup> are taken into account to reduce the need for human intervention. The procedure is illustrated in Figure 3 in a simplified version of the complete flow chart in Greene *et al.*<sup>15</sup> A very

similar flow chart applies to the classification assignment; the domains obtained in the previous step are compared with already known domains using CATHEDRAL and hidden Markov models, and, based on the output, it is decided whether to do an auxiliary manual inspection.

## Future directions

It has long been a matter of debate whether the hierarchical organisation of CATH (and of other domain databases like SCOP) is appropriate,<sup>25</sup> and whether the space of protein structures is better viewed as a continuum. The evolutionary relationships between sequences, however, should allow for discretising the structure space to some extent.

As noted already by the CATH group,<sup>5</sup> a few topologies — often referred to as **superfolds** — contain a disproportionate number of structures (see Figure 4). This was further discussed in the first description of CATH,<sup>3</sup> where the **Russian doll effect** was also considered: a series of small structural changes in a domain's embellishments (ie parts of the structure not belonging to the highly conserved core) could mediate a walk from one topology to another. Furthermore, large structural



**Figure 4.** The distribution of topology sizes in the most recent version of CATH (version 3.2.0) resembles a power law. A few topologies, so-called **superfolds**, contain a disproportionate number of structures. The largest topology, the *Rossmann fold* (3.40.50), comprises 14,720 structures, whereas 111 topologies have one member only.



divergences are observed within several topologies. In Cuff *et al.*,<sup>26</sup> **structurally similar groups (SSGs) are defined as clusters originating from a clustering procedure where two domains are regarded as similar if their normalised RMSD is less than 5 Å.** The study revealed that while the majority of topologies comprise only one or two SSGs, a few contain more than ten (see also Reeves *et al.*<sup>27</sup>). Moreover, these topologies represent a large proportion of the domains in CATH.

Despite the complications caused by the structural overlaps between topologies and the vast structural divergence within some topologies, the CATH database is still a valuable tool if one focuses on **domains that share a common structure in their topological cores and neglects features of the less constrained outer layers of the domains.**

A planned update of CATH (version 3.3.0) will, besides the current hierarchical structure, also contain horizontal links between related topologies.<sup>26</sup>

## Conclusion

The CATH database is valuable for biologists and bioinformaticians alike. For biologists with very specific tasks, browsing for individual domains is made easy by the user-friendly web interface, while bioinformaticians with a focus on large-scale analyses can find complete datasets available for downloading. Thus, working with CATH is remarkably uncomplicated. Updates are frequent, and, given the significant upcoming extension<sup>26</sup> with horizontal layers complementary to the hierarchical structure, CATH is likely to become an even more valuable resource in the future.

## References

- Berman, H.M., Westbrook, J., Feng, Z., Gilliland, G. *et al.* (2000), 'The Protein Data Bank', *Nucl. Acids Res.* Vol. 28, pp. 235–242.
- Berman, H.M., Battistuz, T., Bhat, T.N., Bluhm, W.F. *et al.* (2002), 'The Protein Data Bank', *Acta Cryst.* Vol. D58, pp. 899–907.
- Orengo, C.A., Michie, A.D., Jones, D.T., Swindells, M.B. *et al.* (1997), 'CATH: A hierarchic classification of protein domain structures', *Structure* Vol. 5, pp. 1093–1108.
- Orengo, C.A., Martin, A.M., Hutchinson, G., Jones, S. *et al.* (1998), 'Classifying a protein in the CATH database of domain structures', *Acta Cryst.* Vol. D54, pp. 1155–1167.
- Orengo, C.A., Jones, D.T., Taylor, W. and Thornton, J.M. (1994), 'Protein superfamilies and domain superfolds', *Nature* Vol. 372, pp. 631–634.
- Cuff, A.L., Sillitoe, I., Lewis, T., Redfern, O.C. *et al.* (2009), 'The CATH classification revisited — Architectures reviewed and new ways to characterize structural divergence in superfamilies', *Nucl. Acids Res.* Vol. 37, pp. D310–D314.
- Taylor, W.R. and Orengo, C.A. (1989), 'Protein structure alignment', *J. Mol. Biol.* Vol. 208, pp. 1–22.
- Redfern, O., Grant, A., Maibaum, M. and Orengo, C. (2005), 'Survey of current protein family databases and their applications in comparative, structural and functional genomics', *J. Chromatogr. B* Vol. 815, pp. 97–107.
- Murzin, A.G., Brenner, S.E., Hubbard, T. and Chothia, C. (1995), 'SCOP: A structural classification of proteins database for the investigation of sequences and structures', *J. Mol. Biol.* Vol. 247, pp. 536–540.
- Csaba, G., Birzele, F. and Zimmer, R. (2009), 'Systematic comparison of SCOP and CATH: A new gold standard for protein structure analysis', *BMC Struct. Biol.* Vol. 9, p. 23.
- Dietmann, S., Park, J., Notredame, C., Heger, A. *et al.* (2001), 'A fully automatic evolutionary classification of protein folds: Dali Domain Directory version 3', *Nucl. Acids Res.* Vol. 29, pp. 55–57.
- Mizuguchi, K., Deane, C.M., Blundell, T.L. and Overington, J.P. (1997), 'HOMSTRAD: A database of protein structure alignments for homologous families', *Protein Science* Vol. 7, pp. 2469–2471.
- Sadreyev, R. and Grishin, N. (2003), 'COMPASS: A tool for comparison of multiple protein alignments with assessment of statistical significance', *J. Mol. Biol.* Vol. 326, pp. 317–336.
- Sadreyev, R., Tang, M., Kim, B.-H. and Grishin, N.V. (2007), 'COMPASS server for remote homology inference', *Nucl. Acids Res.* Vol. 35, pp. W653–W658.
- Greene, L.H., Lewis, T.E., Addou, S., Cuff, A. *et al.* (2007), 'The CATH domain structure database: New protocols and classification levels give a more comprehensive resource for exploring evolution', *Nucl. Acids Res.* Vol. 35, pp. D291–D297.
- Buchan, D.W., Shepard, A.J., Lee, D., Pearl, F.M. *et al.* (2002), 'Gene3D: Structural assignment for whole genes and genomes using the CATH domain structure database', *Genome Res.* Vol. 12, pp. 503–514.
- Buchan, D.W., Rison, S.C., Bray, J.E., Lee, D. *et al.* (2003), 'Structural assignments for the biologist and bioinformaticist alike', *Nucl. Acids Res.* Vol. 31, pp. 469–473.
- Yeats, C., Maibaum, M., Marsden, R., Dibley, M. *et al.* (2006), 'Gene3D: Modelling protein structure, function and evolution', *Nucl. Acids Res.* Vol. 34, D281–D284.
- Lees, J., Yeats, C., Redfern, O., Clegg, A. *et al.* (2010), 'Gene3D: merging structure and function for a thousand genomes', *Nucl. Acids Res.* Vol. 38, pp. D296–D300.
- Røgen, P. and Fain, B. (2003), 'Automatic classification of protein structure by using Gauss integrals', *Proc. Natl. Acad. Sci. USA* Vol. 100, pp. 119–124.
- Choi, I.-G., Kwon, J. and Kim, S.-H. (2004), 'Local feature frequency profile: A method to measure structural similarity in proteins', *Proc. Natl. Acad. Sci. USA* Vol. 101, pp. 3797–3802.
- Getz, G., Vendruscolo, M., Sachs, D. and Domany, E. (2002), 'Automated assignment of SCOP and CATH protein structure classification from FSSP scores', *Proteins* Vol. 46, pp. 405–411.
- Redfern, O.C., Harrison, A., Dallman, T., Pearl, F.M. *et al.* (2007), 'CATHEDRAL: A fast and effective algorithm to predict folds and domain boundaries from multidomain protein structures', *PLoS Comp. Biol.* Vol. 3, p. e232.
- Shyu, C.-R., Chi, P.-H., Scott, G. and Xu, D. (2004), 'ProteinDBS: A real-time retrieval system for protein structure comparison', *Nucl. Acids Res.* Vol. 32, pp. W572–W575.
- Dietmann, S. and Holm, L. (2001), 'Identification of homology in protein structure classification', *Nature Struct. Biol.* Vol. 8, pp. 953–957.
- Cuff, A., Redfern, O.C., Greene, L., Sillitoe, I. *et al.* (2009), 'The CATH hierarchy revisited — Structural divergence in domain superfamilies and the continuity of fold space', *Structure* Vol. 17, pp. 1051–1062.
- Reeves, G.A., Dallmann, T.J., Redfern, O.C., Akpor, A. *et al.* (2006), 'Structural diversity of domain superfamilies in the CATH database', *J. Mol. Biol.* Vol. 360, pp. 725–741.