

Protein Three-Dimensional Structural Databases: Domains, Structurally Aligned Homologues and Superfamilies

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Abstract

This paper reports the availability of a database of protein structural domains (DDBASE), an alignment database of homologous proteins (HOMSTRAD) and a database of structurally aligned superfamilies (CAMPASS) on the World Wide Web (WWW). DDBASE contains information on the organization of structural domains and their boundaries; it includes only one representative domain from each of the homologous families. This database has been derived by identifying the presence of structural domains in proteins on the basis of inter-secondary structural distances using the program *DIAL* [Sowdhamini & Blundell (1995), *Protein Sci.* **4**, 506–520]. The alignment of proteins in superfamilies has been performed on the basis of the structural features and relationships of individual residues using the program *COMPARER* [Sali & Blundell (1990), *J. Mol. Biol.* **212**, 403–428]. The alignment databases contain information on the conserved structural features in homologous proteins and those belonging to superfamilies. Available data include the sequence alignments in structure-annotated formats and the provision for viewing superposed structures of proteins using a graphical interface. Such information, which is freely accessible on the WWW, should be of value to crystallographers in the comparison of newly determined protein structures with previously identified protein domains or existing families.

1. Introduction

The Brookhaven Protein Data Bank (PDB) (Bernstein *et al.*, 1977) currently contains over 7000 entries; after removing the repeated entries of identical proteins (such

as the same protein in different complexes or at different resolutions), there remain 1729 proteins (Brenner *et al.*, 1997), including many homologues (see Fig. 1). If only representative structures from the homologous protein ‘family’ are retained such that no two proteins have more than 25% sequence identity (Hobohm *et al.*, 1992; May 1997 release), the resultant data set still includes 687 proteins. This corresponds to 463 superfamilies of protein domains with 96 superfamilies arising from more than one family (Brenner *et al.*, 1997).

Proteins that have diverged but retain high sequence identity fold into similar three-dimensional structures and usually perform similar functions – these clearly belong to a homologous family (Richardson, 1981; Rossmann & Argos, 1977; Chothia, 1984; Overington *et al.*, 1990, 1993). Proteins or domains of proteins that adopt the same three-dimensional fold despite poor sequence identity and perform remotely similar functions (Blundell & Humbel, 1980; Murzin & Chothia, 1992; Murzin *et al.*, 1995; Murzin, 1996) are termed superfamilies. The identification of new members belonging to pre-existing families and superfamilies is straightforward only when contiguous residues forming a functional motif are conserved, where *PROSITE* searches may be appropriate (Bairoch, 1991). Furthermore these should be distinguished from proteins with no sequence identity and no similarity of functions that nevertheless have the same fold or superfolds (Orengo *et al.*, 1994).

An analysis of protein sequence and structure entries indicates that about 50% of the ‘new’ sequences could be attributed a previously known function and roughly 20% of the sequences have homologues of known structure (Bork *et al.*, 1992, 1994; Koonin *et al.*, 1994). When the crystal structure of a ‘new’ protein is determined, it is important to compare its structure with the previously determined structures. This is facilitated by the existence of databases of aligned protein structures and sequences (Overington *et al.*, 1990, 1993; Johnson *et al.*, 1993).

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Often homology or structural similarity exists between parts of two different proteins; one or two domains only may be conserved (Wetlauber, 1973; Richardson, 1981; Wodak & Janin, 1981; Go, 1981). Although algorithms to identify such compact sub-structures have been developed (Schulz, 1977; Crippen, 1978; Rose, 1979; Zehfus & Rose, 1986), it is convenient to use automatic methods so that the information of domain organization can be compiled for the large number of protein structures now available (Islam *et al.*, 1995; Siddiqui & Barton, 1995; Swindells, 1995; Nichols *et al.*, 1995). We have constructed a database of protein structural domains (DDBASE) (Sowdhamini *et al.*, 1996) using the procedure *DIAL* (Sowdhamini & Blundell, 1995).

Structure-based alignment of sequences of related protein domains provides a basis for understanding evolutionary relationships as well as diversity in function and specificity. Such alignments can be used to derive information on amino-acid replacements which are of value also

in comparative modelling and fold recognition (Overington *et al.*, 1990). Databases of structural alignments of homologous proteins (HOMSTRAD: HOMologous STRucture Alignment Database) (Overington *et al.*, 1990, 1993; Mizuguchi *et al.*, 1998) and protein superfamilies (CAMPASS: CAMbridge database of Protein Alignments organized as Structural Superfamilies) (*RS*, Sowdhamini *et al.*, 1998) will be described in this paper. Because of the low percentage of sequence identities amongst distantly related proteins, it is difficult, on the basis of sequence alone, to obtain reliable alignments where secondary structures and functionally important residues are aligned correctly. Alignment of proteins in superfamilies, therefore, is based on the conservation of structural features and relationships using the program *COMPARER* (Sali & Blundell, 1990; Zhu *et al.*, 1992). The three databases, described here, are available on the WWW (<http://www-cryst.bioc.cam.ac.uk/~ddbbase> for DDBASE, <http://www-cryst.bioc.cam.ac.uk/~homstrad> for HOMSTRAD and <http://www-cryst.bioc.cam.ac.uk/~campass> for CAMPASS).

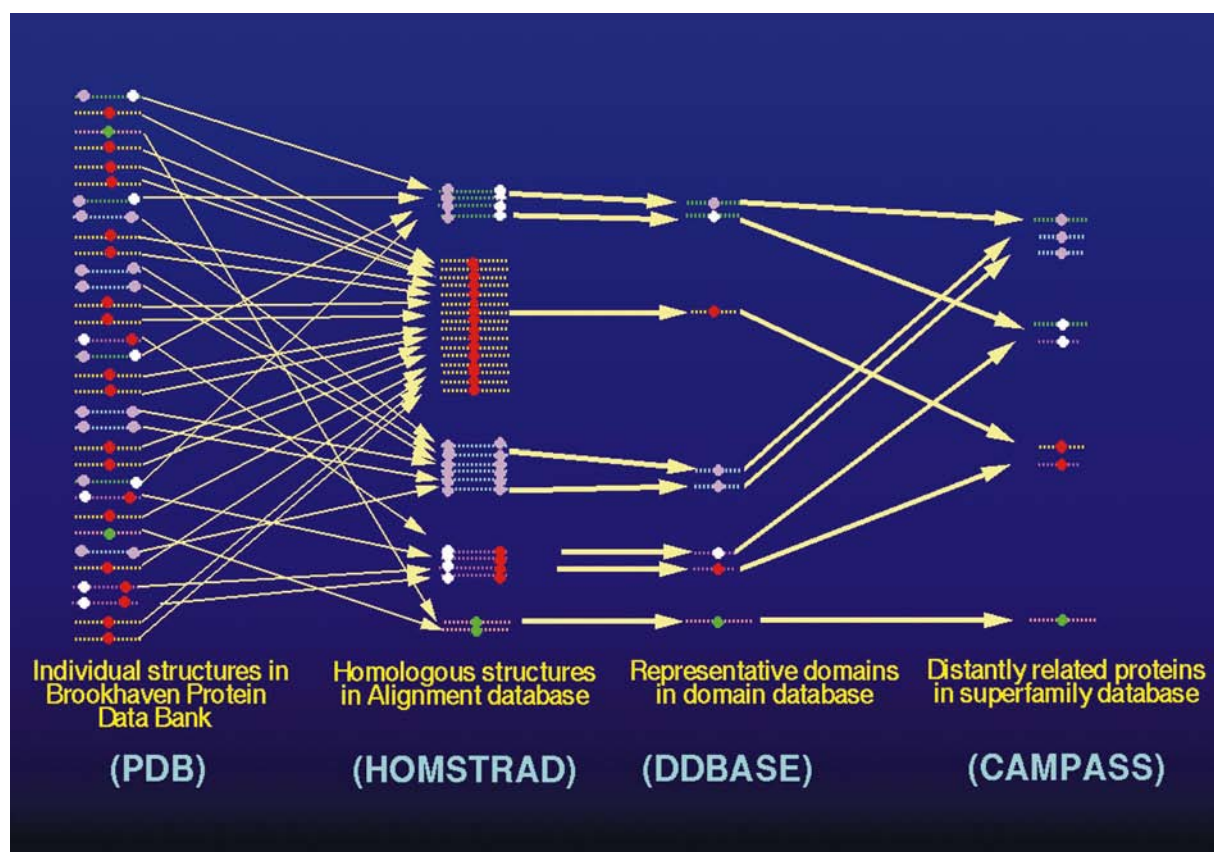


Fig. 1. A cartoon representation of the classification and alignment of proteins at various structural hierarchies. HOMSTRAD database contains alignments of homologous sequences. Some of them exist as multi-domain proteins (denoted by different coloured spheres). DDBASE is a compilation of structural domains found in representatives of homologous proteins. CAMPASS is a database of aligned protein domains belonging to superfamilies.

2. DDBASE

2.1. Description and availability

DDBASE is a compilation of the information on structural domains that are present in a representative set of 436 protein chains (Sowdhamini *et al.*, 1996). The identification of structural domains in a protein chain was performed using the program *DIAL* (Sowdhamini & Blundell, 1995), where elements of secondary structure are clustered on the basis of the proximity to each other. This gave rise to 695 structural domains, of which 206 are α -rich, 191 are β -rich and 294 fall under the α - and- β class. 63% of the domains are from multi-domain proteins and 73% of the identified domains have less than 150 residues.

The organization of structural domains in individual protein chains is described on the WWW page assigned to that protein chain; an example is shown in Fig. 2. Secondary-structural dendrograms are provided that correspond to the clustering based on distances between all possible pairs of secondary structures. All possible combinations of nodes in the secondary-structural dendrogram are automatically examined for compactness of putative domains corresponding to clusters and listed with their disjoint-factor values (see Sowdhamini & Blundell, 1995, for details). It is possible for the user to extract the domain boundary corresponding to any situation by clicking on that entry. However, the 'best' domain boundaries, defined by the program, have been identified and the domain organization may be viewed

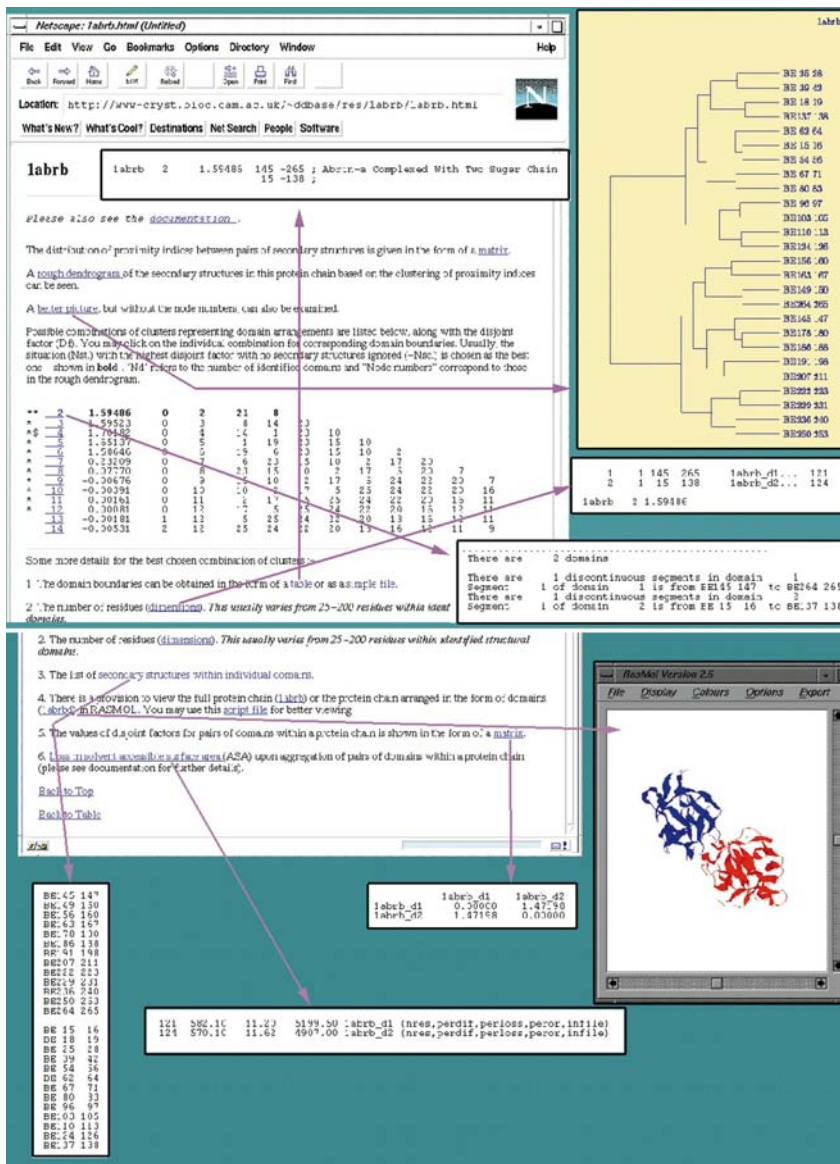


Fig. 2. Domain database (DDBASE) WWW page for the B chain of abrin (PDB code, 1abr) as an example. Domains have been identified using the program *DIAL* (Sowdhamini & Blundell, 1995). The organization of structural domains can be viewed as secondary structural dendrograms where helices and extended strands have been clustered on the basis of intersecondary structural inter-C α distances. Various combinations of nodes, corresponding to secondary-structural clusters, have been examined for structural compactness and listed along with their disjoint factor (see Sowdhamini & Blundell, 1995, for details). Domain boundaries for all these possibilities can be accessed by clicking on that entry. Further, detailed outputs can be accessed for the 'best' combination. The 'best' combination is usually the one with the highest disjoint factor (Df) without any secondary structures being ignored (-Nst. column shows the number of secondary structures that are ignored while examining various nodes in the dendrogram). The protein chain can be viewed using *RASMOL* (Sayle & Milner-White, 1995) where domains are coloured differently in the case of multi-domain proteins.

Table 1. *Proteins in superfamily and homologous databases*

N_{mem} is the number of members in the superfamily. The first four characters of the member codes correspond to the PDB code, the fifth to the chain identifier and the last character to the domain number. Superfamily name is as defined in SCOP (Murzin *et al.*, 1995). In a few cases where there is considerable functional similarity, we have considered a broader class of proteins under one superfamily (marked as fold). In a few other cases, we have restricted our choice of superfamily members to a group of proteins, defined as a family in SCOP (marked as family), to permit reliable structural superposition and structure-based sequence alignment. N_{hom} is the number of homologous proteins in this family. Many of them are single member families.

| Superfamily code (N_{mem}) | Member codes | Superfamily name | Homologous family name | N_{hom} |
|--|---|---|---|-----------------------------------|
| 4helud (3) | 256ba0 11bbha0, 2ccya0 | Cytochromes | Cytochrome <i>b562</i> Cytochrome <i>c'</i> | 1 2 |
| FAD-binding-like (13) | 1gal-1, 1pbe-2†, 3cox-1 1gnd-2 1npx-2, 1fcda2†, 1fcda1† 1trb-1†, 1trb-2†, 3grs-1 3grs-2, 3lada2 2tmda2 | FAD/NAD(P)-binding domain | Cholesterol oxidase (full protein) Guanine nucleotide dissociation inhibitor Disulfide oxidoreductase As above As above Trimethylamine dehydrogenase | 3 1 10 1 |
| FMN_typeI (2) | 2tmda1†, 1oyb-0† | FMN-linked oxidoreductases | Flavin-binding beta-barrel | 2 |
| PH (3) | btn-0, 1dyna0, 1mai-0 | PH domain-like | Pleckstrin-homology domain | 7‡ |
| SH3(2) | 1lck-2, 1pht-0 | SH3 domain | SH3 domain | 7 |
| ab5_toxins (5) | 1bova0, 1chbd0, 1ptob2, 1ptod0, 1ptof0 | Bacterial enterotoxins | Bacterial AB5 toxins | 8‡ |
| ab_hydrolases (8) | 1broa0 2had-0 1thta0 1gpl-0 1tca-0, 2ace-0 1din-0 1whta0 | Alpha/beta-hydrolases | Bromoperoxidase A2 Haloalkane dehalogenase Thioesterases Lipase alpha beta-hydrolase Dienelactone hydrolase Serine carboxypeptidase | 1 1 1 2 3 1 3 |
| actinIA (3) | 1atna3, 3hsc-2 1glcg1 | Actin-like ATPase domain | Actin Glycerate kinase | 2 1 |
| actinIIA (3) | 1atna1, 3hsc-3 1glcg2 | Actin-like ATPase domain | See actinIA See actinIA | |
| actin_binding (2) | 1vil-0, 1svq-0 | Actin depolymerizing proteins | Gelsolin-like | 3‡ |
| adk (2) | 2ak3a1, 1gky-1 | Nucleotide and nucleoside kinases | Nucleotide kinase | 5 |
| adp (4) | 1ddt-3, 1dmaa0, 1ltaa0 1ptoa0 | ADP-ribosylation | ADP-ribosylating toxins As above | 6‡ |
| animal_viral (5) | 1bbt30, 2rhn3m, 1cov1m 61bbt10, 1bbt2m | Animal virus proteins (family) | Picornavirus coat proteins As above | 6 |
| anticodon_binding (2) | 1asya2, 1lyla2 | An anticodon-binding domain (family) | An anticodon-binding domain | |
| asp_hiv (3) | 1hiva0 45pep-2, 5pep-1 | Acid proteases | Retroviral proteinase Aspartic proteinase | 4‡ 11 |
| bacteriophage (2) | 1gpc-0, 2gva-0 | Bacteriophage ssDNA- binding proteins | Bacteriophage ssDNA- binding proteins | 3‡ |
| beta-gamma- crystallin_like (3) | 4gcr-1, 1prs-1 | Crystallins/protein S/killer toxin | Crystallin | 5 |
| bgt-gpb (2) | 1wkt-0 2bgu-0 | Beta-glucosyltransferase & glycosyltransferase | Yeast killer toxin Beta-glucosyltransferase | 1 1 |
| cbp (7) | 1gpb-0 3cln-2, 2scpa2, 2scpa1 2sas-1, 2sas-2, 1rec-1† 1rro-m† | EF-hand | Oligosaccharide phosphorylase Calcium binding protein – calmodulin-like As above Parvalbumin | 3a 6 5 5 |
| ccperoxy (3) | 1lgaa0, 1scha0†, 2cyp-0 | Heme-dependent peroxidases | Peroxidase | 4 |
| creatinase (2) | 1chma2, 1mat-0 | Creatinase/methionine aminopeptidase | Creatinase/methionine aminopeptidase | 3‡ |
| ctt (2) | 1ctt-1, 1ctt-2 | Cytidine deaminase | Cytidine deaminase | 1 |
| cys (2) | 2act-0, 1gcb-1† | Papain-like | Cysteine proteinase | 5 |
| cystineknot (6) | 1bet-0 a1aoca2 1pdga0 1hcna0, 1hcnb0 | Cystine-knot cytokines | Neurotrophin Coagulogen Platelet-derived growth factor Gonadotropin | 3‡ 1 1 1 |

Table 1 (cont.)

| Superfamily code (N_{mem}) | Member codes | Superfamily name | Homologous family name | N_{hom} |
|--|--|--|---|------------------|
| cytc (3) | 2tgi-0 | Monodomain cytochrome c (family) | Transforming growth factor β | 4‡ |
| | 351c-0, 1cyi-0† | | Cytochrome-c5 | 5 |
| cytokine (2) | 1ycc-0 | Cytokine | Cytochrome-c | 9 |
| | 1ilb-0, 4fgf-0 (2fgf) | | Interleukin 1- β -like growth factor | 5 |
| exopeptidase (3) | 1amp-0 | Zn-dependent exopeptidases | Bacterial aminopeptidases | 2‡ |
| | 1lcpa1 | | Leucine aminopeptidase, C-domain | 1 |
| ferredoxin_reductases (3) | 2ctb-0 | Ferredoxin reductase-like C-terminal domain | Pancreatic carboxypeptidases | 3‡ |
| | 2pia-3 | | Phthalate dioxygenase reductase | 1 |
| flav (7) | 1ndh-2, 1fnc-2 | Flavodoxin-like(fold) | Reductases | 5‡ |
| | 1bmta1 | | Methionine synthase C- Ornithine decarboxylase N-domain | 1 |
| | 1orda4 | | Ornithine decarboxylase N-domain | 1 |
| | 1cus-m | | Cutinase | 1 |
| | 3chy-0 | | CHEY-like | 5‡ |
| | 1scua2 | | Succinyl-CoA synthetase- α -chain C-domain | 1 |
| | 4fxn-0 | | Flavodoxin | 6 |
| 1qora1 | Alcohol/glucose dehydrogen- ase, C-domain | 2 | | |
| globins (7) | 1flp-0, 1ithb0, 3sdha0, 2gdm-0, 1mbc-0, 2hbg-0, 1ash-0 | Globin-like | Globin | 23 |
| glucoamylase_like (3) | 1gai-0 | Glycosyltransferases of the superhelical fold | Glucoamylase | 1 |
| glycosyltransferases (18) | 1clc-1, 1cem-0 | Glycosyltransferases | Cellulase catalytic domain | 3‡ |
| | 1bgl-2, 1ecea0, 1edg-0 | | beta-glycanases | 11‡ |
| | 1ghsa0, 1xyza0, 1cec-0 | | As above | |
| | 1byb-0 | | beta-amylase | 1 |
| | 1cbg-0 | | Family 1 of glycosyl hydrolase | 4‡ |
| | 1cgt-1, 1bpla1†, 1ppi-1† | | Amylase (full protein) | 6 |
| | 2amg-1† | | As above | |
| | 1ctn-1, 2ebn-0, 2hvm-0 | | Type II chitinase | 6‡ |
| | 1nar-0 | | As above | |
| | 1qba-1 | | Bacterial chitobiase ca. domain | 1 |
| gshase_2 (4) | 4xiaa1 | Glutathione synthetase ATP-binding-like | Xylose isomerase | 5 |
| | 1gsh-3, 2dln-2 | | Peptide synthetases C-domain | 2‡ |
| | 1scub3 | | Succinyl-CoA synthetase beta- N- | 1 |
| | 1dik-2 | | Pyruvate phosphate dikinase N- | 1 |
| gshase_3 (5) | 1gsh-2 | Glutathione synthetase ATP-binding like | See gshase_2 | |
| | 2dln-1 | | See gshase_2 | |
| | 1scub2 | | See gshase_2 | |
| | 1bnca3 | | Biotin carboxylase | 1 |
| | 1dik-3 | | See gshase_2 | |
| | 1cid-2, 1vcaa2, 3 cd4-1 | | Immunoglobulin - C2 set | 2 |
| ig (12) | 1hsaa2, 1vabb0 | Immunoglobulin | Histocompatibility antigen- binding domain | 5 |
| | 1nct-0, 1tit-0, 1tlk-0 | | I set domains | 7‡ |
| | 1vcaa1, 1wit-0 | | As above | |
| | 2fbjl2, 3hflh1 | | Immunoglobulin domain C1 set - constant immunoglobulin | 17 |
| | | | | |
| il8_like (2) | 1huma, 1ikl- (1il8) | Interleukin 8-like chemokines | Interleukin 8-like protein | 5 |
| kinases (3) | 1atpe0, 1csn-0, 1irk-0 | Protein kinases (PK) ca. core | kinase(1apm) | 7 |
| lectins (6) | 1saca0 | ConA-like lectins/glucanases | Pentraxin | 2‡ |

Table 1 (*cont.*)

| Superfamily code (N_{mem}) | Member codes | Superfamily name | Homologous family name | N_{hom} |
|--|---|--|---|------------------------|
| | 1ayh-m | | Bacillus 1-3,1-4- β -glucanase (2ayh) | 3‡ |
| | 2ltm-m | | Plant lectin | 7 |
| | 1slt-0 | | S-lectin | 2 |
| lipocalin (5) | 1kit-2, 1kit-3 1icm-0 (1ifb), 1mup-0 1epba0 (1bbp), 1bbpa0 1fel-0 (1rbp) | Lipocalins | Vibrio cholerae sialidase, N- Lipocalin As above | 1 12 |
| methyltransferases (5) | 1vpt-1 | S-adenosyl-L-methionine -dependent methyltransferases | Polymerase regulatory subunit VP39 | 1 |
| | 2adma2, 1hmy-1 1vid-1 | | DNA methylases Catechol O-methyltransferase COMT | 3‡ 1 |
| muconate_lactonizing (3) | 1xvaa1 1muca1, 2mnr-1 | Enolase & muconate- lactonizing C-domain | Glycine N-methyltransferase Muconate lactonizing enzyme-like | 1 3‡ |
| nip (3) | 4enl-1 1dts-0, 1adea1, 1nipb0 | P-loop containing nucleotide triphosphate hydrolases | Enolase Nitrogenase iron protein-like | 2‡ 3‡ |
| p450 (4) | 2cpp-0, 2hpd0, 1cpt-0 1oxa-0 | Cytochrome P450 | Cytochrome p450 As above | 3 |
| pbgd1 (4) | 1pda-1, 1sbp-2, 1omp-1 1lfg-1 | Periplasmic binding II | Phosphate binding protein-like Transferrin | 12‡ 5‡ |
| pbgd2 (4) | 1pda-2, 1omp-2, 1sbp-1 1lfg-3 | Periplasmic binding II | See pbgd1 See pbgd1 | |
| phospholipase (2) | 1bp2-0 1poc-m | Phospholipase A2 | Phospholipase A2 Insect phospholipase A2 | 7 1 |
| plant_viral (5) | 1bmv21, 1cwpam, 1bmv10 1bmv22, 2stv-m | Plant virus proteins (family) | Plant virus coat protein (4sbv) As above | 2 |
| plp1 (4) | 1ars-2 1dge-2 | PLP-dependent transferases | Aspartate aminotransferase (3aat) omega-Amino acid pyruvate aminotransferase-like | 2 2‡ |
| | 1orda2 | | Ornithine decarboxylase major domain | 1 |
| plq (2) | 1tpla1 1plq-1, 1plq-2 | DNA-clamp | Tyrosine phenol-lyase DNA polymerase processivity factor | 1 1 |
| porins (3) | 2omf-0, 2por-0† 1mal-0 | Porins | Porin Maltoporin | 3 2‡ |
| ppase1 (3) | 1spia2 2hhma1 1inp-1 | Sugar phosphatases | Fructose-1,6-bisphosphatase Inositol monophosphatase Inositol polyphosphate 1-phosphatase | 3‡ 1 1 |
| ppase2 (3) | 1spia1 2hhma2 1inp-2 | Sugar phosphatases | See ppase1 See ppase1 See ppase1 | |
| ras (4) | 5p21-0, 1eft-1 (1etu) 1tada1†, 1hura0† | G proteins(family) | GTP-binding protein As above | 4 |
| repressor_like (4) | 1copd0, 1r69-0, 1neq-0† | Lambda repressor-like DNA-binding domains | DNA-binding repressor (2cro) | 5 |
| ribonucleaseh_like (5) | 1octc0 1bco-1 1kfd-1 | Ribonuclease H-like | Oct-1 POU-specific domain Mu transposase core domain Exonuclease domain of DNA polymerase KF | 1 1 2‡ |
| rubredoxins (3) | 1hjra0 2rn2-0 1itg-0 8rxna0 4at1b2 | Rubredoxin-like(fold) | RuvC resolvase Ribonuclease H (1rnh) Retroviral integrase Rubredoxin (7rxn) Aspartate carbamoyl transferase_RC | 1 3 2‡ 5 1 |
| | 1tfi-0 | | A transcriptional factor domain | 2‡ |

Table 1 (cont.)

| Superfamily code (N_{mem}) | Member codes | Superfamily name | Homologous family name | N_{hom} |
|--|------------------------|--|--|------------------|
| serineproteases1 (5) | 1sgt-1 | Trypsin-like serine proteases | Serine proteinase, mammalian | 16 |
| | 1hava1 | | picornain | 2‡ |
| | 2alp-2, 1arb-1† | | Serine proteinase, bacterial | 4 |
| | 1svpa1 | | Viral proteases | 2‡ |
| serineproteases2 (4) | 2alp-1, 1arb-2 | Trypsin-like serine proteases | See serineproteases1 | |
| | 1hava | | See serineproteases1 | |
| | 1svpa2 | | See serineproteases1 | |
| sial_neur (3) | 1eus-0 (1nsb), 1dim-0 | Sialidases (neuraminidases) | Neuraminidase | 4 |
| sslipid (2) | 1nsca0 | As above | | |
| | 1hyp-0 | Bifunctional inhibitor/lipid-transfer Seed storage 2S albumin | Plant lipid-transfer and hydrophobic proteins | 4‡ |
| strep (2) | 1bip-0 | Avidin/streptavidin | Bifunctional proteinase | 1 |
| | 1sria0 | | Avidin (1pts) | 2 |
| | 1smpi0 | | Metalloprotease inhibitor | 1 |
| superantigen_toxins (2) | 1tssa1, 1se2-1 | Superantigen toxins N-domain (family) | Superantigen toxins N-domain | 4‡ |
| thiamin_binding (6) | 1pyda1, 1pyda2, 1powa1 | Thiamin-binding | Pyruvate oxidase and decarboxylase | 3‡ |
| | 1powa2 | | As above | |
| thioredoxin (6) | 1trka1, 1trka2 | Thioredoxin-like | Transketolase | 1 |
| | 1erv-0, 1thx-0, 1aba-0 | | Thioredoxin (3trx) | 4 |
| | 1dsba1 | | Disulfide-bond formation facilitator | 2 |
| | 2gsta1 | | Glutathione S-transferase (5gst) | 7 |
| trp-biosynthesis (3) | 1gp1a0 | Tryptophan biosynthesis enzymes | Glutathione peroxidase | 1 |
| | 1igs-0, 1pii-2, 1wsya0 | | Tryptophan biosynthesis enzyme | 2 |
| tyrosine_phosphatases (3) | 2hnq-0, 1ypta0 | Phosphotyrosine protein phosphatases I | Higher molecular-weight phosphotyrosine | 3‡ |
| | 1vhra0 | | Dual-specificity phosphatase | 1 |
| viral_coat (3) | 2bbva0 | Viral coat and capsid proteins | Insect virus proteins | 1 |
| | 2tbva2 | | Plant virus coat protein | 2 |
| | 2cas1m† | | Picornavirus coat proteins | 7 |

† This entry is yet to be added in one of the existing families in the homologous alignment database. ‡ This family is yet to be added in the homologous alignment database.

on graphics using *RasMol* (Sayle & Milner-White, 1995). Each domain can be identified by its unique six-character code (the first four characters correspond to the PDB code of the protein, the fifth to the chain identifier and the sixth, as a subscript, corresponds to the domain numbering as in the individual domain pages).

2.2. Application

DDBASE can be used to trace similarities where particular domains are shared between proteins. It is especially useful where there are discontinuous domains. 400 large (with seven or more secondary structures) domains can be grouped into 30 classes on the basis of the structural similarity estimated from structural environments of individual secondary structures (Rufino & Blundell, 1994; Sowdhamini *et al.*, 1996). The clustering of individual protein domains into structurally similar classes can also be examined on the DDBASE WWW page.

3. HOMSTRAD and CAMPASS

3.1. Description and availability

HOMSTRAD and CAMPASS are databases of structure-based alignments of protein sequences, grouped into homologous families and superfamilies, respectively. Aligned sequences of families of homologous protein structures are available in HOMSTRAD (Overington *et al.*, 1990, 1993) and categorized according to the secondary-structural classes. There are 130 homologous protein families with at least two members in the March 1998 version. The sequences of homologous proteins within a family are initially aligned using the rigid-body superposition program *MNYFIT* (Sutcliffe *et al.*, 1987) or *COMPARER* (Sali & Blundell, 1990; Zhu *et al.*, 1992) and later subjected to a careful manual examination. Similar types of information are available for CAMPASS, the database of protein (domain)s belonging to superfamilies (*RS*, Sowdhamini *et al.*,

cytochrome-c

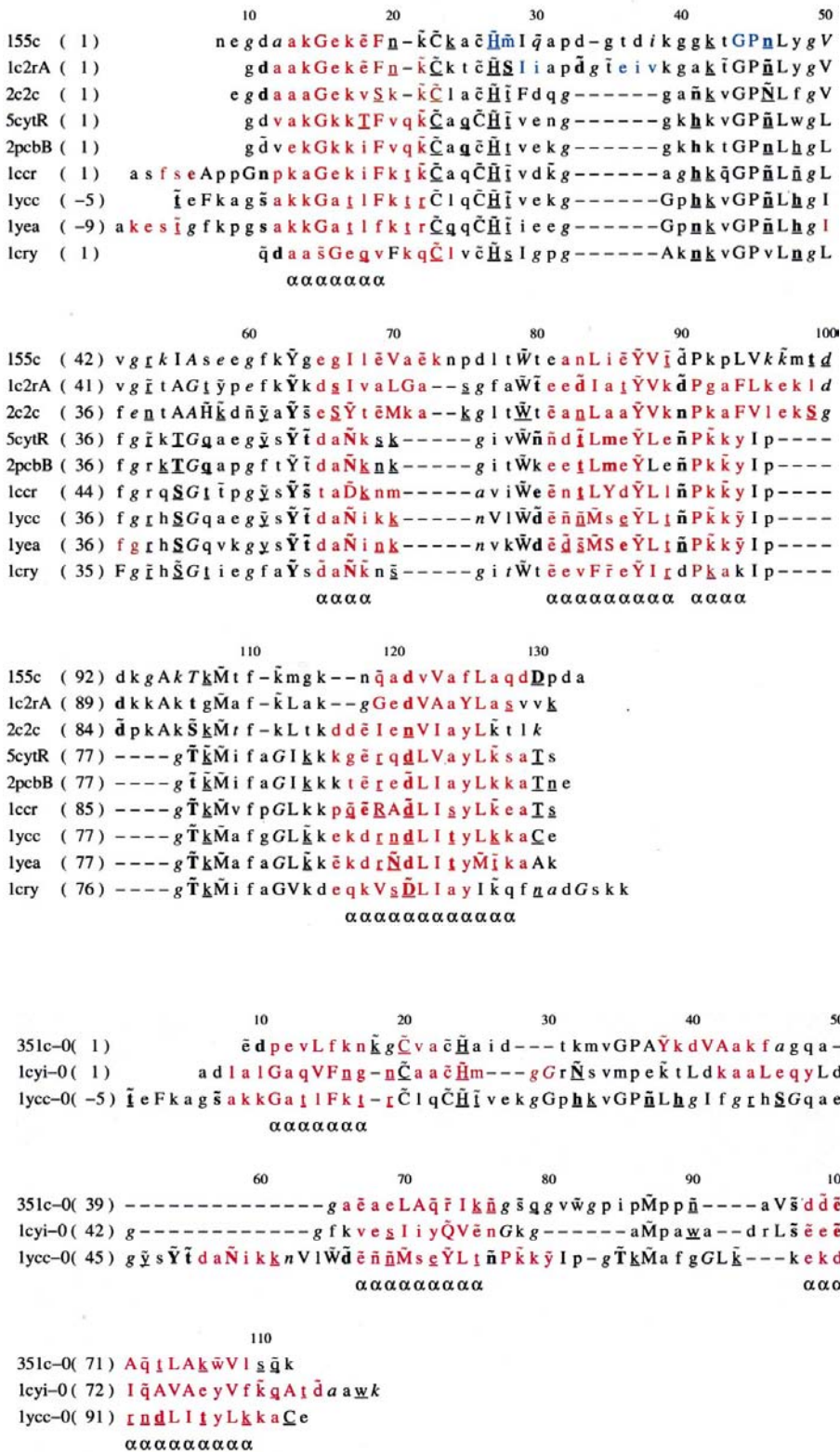


Fig. 3. HOMSTRAD database. Structure-based alignment of proteins in the family of cytochrome c. The first four characters of the code of the protein corresponds to the PDB code. Numbers in brackets correspond to residue numbers and residues are shown in single letter code. The alignment has been formatted using JOY (Overington *et al.*, 1990). The conserved helices are important to the structural integrity of the proteins; functionally important residues (for example CXXCH, residue number 13 of 1ycc) are conserved. Residues are classified into two categories: those which are in the interior and those which are solvent-exposed (with solvent accessibility (ASA) values more than 7% (Hubbard & Blundell, 1987). In the sequence alignment, the solvent-exposed and solvent-buried residues are shown in lower case and upper case, respectively. Residues which have a positive ϕ value and a *cis*-peptide bond in their backbone conformation are shown in italics and with a breve accent on top, respectively. Disulfide-bonded cystine residues are shown by a cedilla symbol. Hydrogen bonding to other side chains, main-chain amides and main-chain carbonyl groups are shown by a tilde (indicated in non-HTML files), in bold and underlined, respectively. Residues in β -strands, α -helices and 3(10)-helices are shown in blue, red and maroon, respectively.

Fig. 4. CAMPASS database. Structure-based alignment of the cytochrome superfamily including distantly related proteins such as c550. Helix 2 of 1ycc, conserved within the homologues (see Fig. 3), occurs as an insertion in this alignment. Despite poor sequence identity, the functionally important residues (CXXCH) are conserved amongst the members in this superfamily.

1998). Superfamilies of structural domains were selected initially on the basis of structural environment at secondary structural units (Rufino & Blundell, 1994; Sowdhamini *et al.*, 1996). The selection of superfamilies has been extended by referring to SCOP (Murzin *et al.*, 1995) and by including smaller domains like the cystine-knots, not considered earlier in the clustering analysis since they were not easy to compare using automatic structure-based procedures. 367 of 451 superfamilies annotated in SCOP have single families (Brenner *et al.*, 1997; the more recent February 1998 release of SCOP has 419 of the 571 superfamilies with single families). Superfamily members were chosen such that no two domains within a superfamily share more than 25% sequence identity (alignments of closely related proteins are available in HOMSTRAD). This cut-off is consistent with the DDBASE definition in choosing representative protein chains. A rigorous sequence-alignment program, *COMPARER* (Sali & Blundell, 1990; Zhu *et al.*, 1992), was used to align the members of a superfamily on the basis of structural features and relationships, which are equivalenced using simulated annealing. Table 1 lists protein superfamilies, with at least two members within the above-defined cut-off of sequence identity, whose alignments have been compiled in the March 1998 version. This includes 67 multi-member superfamilies which involves 293 domains representing 464 homologous proteins. There are a further 357 superfamilies, annotated in SCOP, which have single members (Murzin *et al.*, 1995; Brenner *et al.*, 1997). A few other multi-member superfamilies included in SCOP, such as the DNA-binding HMG box, pheromones, annexins and insulin-superfamily, were excluded from CAMPASS as members exhibited more than 25% sequence identity.

3.2. Availability

The WWW site of HOMSTRAD (Mizuguchi *et al.*, 1998) provides a page for each of the families. The name of the protein, source, resolution and *R* factor are given for each family member corresponding to a PDB entry. The alignment of sequences is formatted in *JOY* (Overington *et al.*, 1990) which highlights the conservation of local-residue structural features such as secondary structure, solvent accessibility and hydrogen bonding. Fig. 3 shows the alignment of cytochrome *c* from different sources and its homologues (cytochrome *c2* and cytochrome *c550*), as an example.

CAMPASS, on the WWW, provides information on the superfamilies: for each superfamily member, the name, source, resolution and domain boundaries are given. The beginning and end residue numbers for each segment of discontinuous domains are recorded. The pairwise percentage identity matrix of the members is provided. The structure-based alignment in the *JOY*-

annotated form (Overington *et al.*, 1990), similar to that described in HOMSTRAD, is shown and also available for extraction in the form of PostScript files, or as LATEX or HTML files or as a plain text file. Fig. 4 shows the alignment of the cytochrome superfamily as an example. A single representative (1yc) of the nine cytochrome homologues (see above and Fig. 3) has been aligned with rather distantly related cytochromes such as cytochrome *c6* and *c551*. The structures of the proteins within a family/superfamily have been superposed using *MNYFIT* (Sutcliffe *et al.*, 1987), where the equivalent residues correspond to the final alignment. These superposed structures can be viewed on the WWW using the *RASMOL* graphics interface (Sayle & Milner-White, 1995).

Fig. 5 shows the distribution of pairwise percentage identities in the two alignment databases. Protein pairs in HOMSTRAD have a broad range of pairwise sequence identities with a slightly bimodal distribution (237 pairs have sequence identities between 25 and 30% and 121 pairs have sequence identities between 60 and 65% out of a total of 1962 pairs). However, the majority of homologous proteins in the database have sequence identities between 15 and 65%. The distribution of pairwise sequence identity of members within superfamilies (CAMPASS) is restricted to a maximum of 25%. A vast majority of protein pairs (449 out of 665) have pairwise percentage identities between 5 and 15%.

4. Conclusions

HOMSTRAD and CAMPASS are distinct from but complementary to other databases. SCOP (Murzin *et al.*, 1995) has classified the entire Protein Data Bank at different levels of structural hierarchy and structural domains are defined. There is emphasis on functionality in the clustering of folds. SCOP does not attempt to perform or present sequence or structural alignments. CATH (Orengo *et al.*, 1993, 1994) was originally designed and developed for whole proteins where the

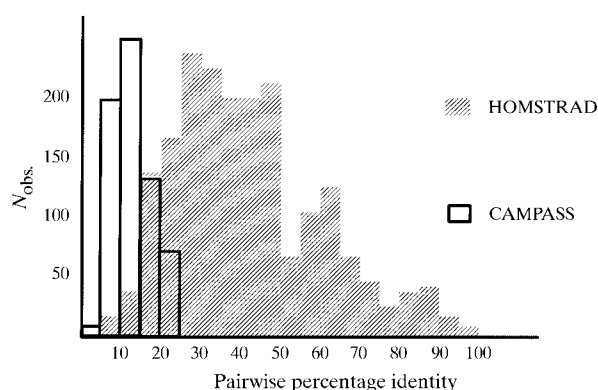


Fig. 5. Distribution of pairwise percentage sequence identities amongst members in the homologue alignment database (HOMSTRAD) and superfamily alignment database (CAMPASS).

authors had taken particular caution to exclude multi-domain proteins. Subsequently, the structures have been systematically classified at the level of domains (Orengo *et al.*, 1997). CATH does not include structure-based alignments of sequences. *FSSP* (Holm & Sander, 1994) is most similar to HOMSTRAD and CAMPASS due to the fact that *FSSP* also provides structure-based sequence alignments, even incorporating remote homologues. However, the alignments do not distinguish homologues and superfamilies from those which only share a similar fold. The databases described in this paper contain structure-based alignments that have been specially annotated to describe the structural environment at residue positions. This should provide extra information useful in the comparison of protein structures.

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