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Six Rossmannoid folds, including the Class I aminoacyl-tRNA synthetases, share a partial core with the anti-codon-binding domain of a Class II aminoacyl-tRNA synthetase

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ABSTRACT

Motivation: Similarities in core residue packing provide evidence for divergence or convergence not reported using other methods.

Results: We apply a new method for rapid structure comparison based on Simplicial Neighborhood Analysis of Protein Packing (SNAPP) to the diverse structural classification of proteins (SCOP) α/β -class of protein folds. The procedure identifies inter-residue packing motifs shared by protein pairs from different folds. A threshold of 0.67 Å RMSD for all atoms of corresponding residues ensures inclusion of only highly significant similarities comparable with those observed for identical catalytic residues in homologues. Many tertiary packing motifs are shared among the three classical Rossmannoid folds, as well as thousands of other motifs that occur in at least two distinct folds. Merging of neighboring packing motifs facilitated recognition of larger, recurrent substructures or cores. The anti-codon-binding domain of an archeal aminoacyl-tRNA synthetase (aaRS) was discovered to possess a packed core in which eight identical amino acid residues are within 0.55 Å RMSD of the comparable structure in the FixJ receiver, a member of the Rossmannoid family that also includes the CheY signaling protein and flavodoxin-like proteins. Further investigation identified close variants of this core in five other Rossmannoid folds, including a functionally relevant core in Class Ia aminoacyl-tRNA synthetases. Although it is possible that the two essentially identical cores in the ProRS anti-codon-binding domain and the FixJ receiver converged to the same structure, the consensus core obtained from the structural and sequence alignments suggests that all the implicated protein folds descended from a simpler ancestral protein in which this core provided nucleotide binding and proto-allosteric functions. Availability: Programs are available at http://staff.vbi.vt.edu/ cammer/snapp/download/

Implementation: Programs were written in Perl and c and run under Linux.

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1 INTRODUCTION

Demonstrating structural similarities between proteins has long provided the initial evidence for common structural motifs

representing evolutionary divergence or convergence. Most approaches rely on comparing the arrangement of secondary structures or other local substructures in 3D (Dror et al., 2003; Holm and Sander, 1993; Madej et al., 1995; Orengo and Taylor, 1990; Shindyalov and Bourne, 1998). Considerably less development has been devoted to identifying more precise structural details.

Comparison methods that focus on precise arrangements of amino acid residues are usually used to identify catalytic, ion binding and small molecule binding sites (Barker and Thornton, 2003; Fetrow and Skolnik, 1998; Fischer et al., 1994; Jambon et al., 2003). Often, the aim is to recognize similarities across distinct protein folds for function prediction or active-site comparisons among homologous protein structures (Fetrow et al., 1999). Such atomic level comparisons have generally not been extended to features outside active sites. Yet, as distantly related proteins with different folds usually have substantial active-site differences, comparing structural features outside the active sites may yield more definitive evidence for divergence from a common ancestor for proteins with specific sub-structural motifs. Central to the present work is identification of tertiary contact motifs found in pairs of proteins from at least two different folds.

The method identifies recurrent tertiary contacts between protein secondary structure elements. Similarities are observed at the atomic level of detail where conformations of individual amino acid residues are repeated in similarly packed arrangements of side chains on the secondary structure faces. In conventional analyses, the most regularly repeated features of tertiary contacts observed are residue pairs separated as (i, i+2) in β -strands, and as (i, i+3) and (i, i+4)on the surface of α -helices (Chothia and Janin, 1982; Chothia *et al.*, 1977, 1981; Janin and Chothia, 1980). In this work, these residue pairs will be referred to as secondary structure-coupled residues (SCRs). SCRs yield 'knobs-and-holes' (Crick, 1953) that form the close-packed interfaces between secondary structure elements. The regular spacing in sequence and in 3D of SCRs leads to preferred orientations for the secondary structures in 3D that have been observed in similar protein structures. Results presented here demonstrate that tertiary, i.e. non-sequence local arrangements of residues interacting at the interfaces of secondary structure elements yield evolutionary clues that cannot be obtained by comparing one-dimensional motifs.

We analyze the structural classification of protein (SCOP) α/β class of proteins, in which $\beta - \alpha - \beta$ motifs are repeated to form layered sandwich structures with helices on both sides of a central, parallel β -sheet. The largest related superfamily of proteins in this

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class is the Rossmannoids, which comprises 12 different folds presumed to be present in the last common ancestor (LCA) of all organisms (Aravind *et al.*, 2002; Shakhnovich *et al.*, 2003). Many motifs have been identified here that connect the three classical Rossmannoid folds, as well as the others, and many more motifs have been discovered that are shared by at least two different folds in the α/β -class.

A large, interesting and widely shared motif is a core of eight residues linking an α -helix to three parallel β -strands in flavodoxinlike (Rossmannoid) fold members, FixJ and CheY, and in the anti-codon-binding domain of Class II prolyl- and glycyl-tRNA synthetases and the accessory β -subunit of mitochondrial DNA polymerase γ . These two fold families have not been previously identified as possessing such detailed similarity. Interestingly, this core is partially represented in five other Rossmannoid folds, including Class I aminoacyl-tRNA synthetases, where it appears also to serve an allosteric function (Kapustina et al., 2007; Weinreb et al., 2009). The prevalence of this motif in proteins that bind to nucleotides, and the presumptive functional importance in Class I aminoacyl-tRNA synthetases, suggest that a single ancestral protein may have diverged to form the Rossmannoids and the anti-codonbinding domain. As the Rossmannoids diverged before the LCA of all organisms, the detected core similarity would represent one of the most ancient protein structural modules yet identified.

2 METHODS

2.1 Database

Five classes from the SCOP 1.73 database guided assembly of a near-exhaustive collection of representative protein structures from the Protein Data Bank (Murzin *et al.*, 1995). One protein structure was selected for each species represented and for every domain in the SCOP α , β , α / β , $\alpha + \beta$ and multi-domain proteins classes. In all, 11 610 fold representatives were compiled from 9197 structures in SCOP.

2.2 Structure representation and packing decomposition

All structures were represented by a single-point per residue based on sidechain centroids. Tetrahedral packing simplices were identified using SNAPP (Cammer *et al.*, 2002; Tropsha *et al.*, 2003), which uses Delaunay tessellation to decompose the set of side-chain centroids into non-overlapping tetrahedra where four residues form nearest neighbors.

Each four-body contact was represented by combining the single-letter codes in sequence order with the three distances of sequence separation of the residues along the chain. Distances used were 2, 3 and 4 for SCRs, and L for distances greater than four residues of separation. For example, IILV_2_L_L refers to a Delaunay simplex composed of residues 151I, 153I, 165L and 176V. A four-body pattern that includes at least one SCR is referred to here as Secondary structure-Coupled Residues And Packing Environment (SCRAPE). SCRAPEs constitute elemental links between secondary structures and are therefore elementary tertiary packing motifs. SCRAPEs common to at least two different folds were screened to identify packing arrangements within 0.67 Å RMSD for all atoms. This degree of similarity is comparable with similarities in ser-asp-his catalytic triads among homologous trypsin-like serine proteases (Stark *et al.*, 2003).

2.3 Motif finding

SCRAPEs repeated in protein structures from two different folds initially were compared geometrically using tetrahedra defined by connecting

side-chain centroids. Motifs were identified and analyzed using the following workflow:

- SCRAPEs, i.e. residue side chain centroid tetrahedra, are identified in all protein structures in the database using SNAPP based on Delaunay tetrahedralization.
- (2) Tetrahedra with identical residue types and sequence spacing found in pairs of structures from different folds are superimposed and compared.
- (3) When all-atom RMSD for the four residues in a tetrahedron is <0.67 Å, the tetrahedron is considered to be a motif shared by both structures.
- (4) When neighboring motifs are identified in the same structure pair, these motifs are combined to form a composite motif linking the two structures.
- (5) All motifs linking structures in different fold pairs are tabulated and used for generating similarity networks that graph motif distribution of among different folds in the database.

Network edges were drawn based on various cut-offs for number of residues shared. Many of these structure similarity networks were generated and visualized for subsets of the data. Superimpositions of hundreds of structure pairs were visualized directly.

3 RESULTS

Protein structures from the SCOP α/β -class were analyzed for SCRAPEs in recurring packing motifs that might help discriminate between evolutionary divergence and convergence. There were 32 180 instances of protein structure pairs from different folds possessing SCRAPE pairs that were within 0.67 Å RMSD for all atoms. SCRAPES were combined to form composite motifs, revealing a total of 18 107 similarities between protein pairs from distinct folds in the α/β -class.

The overall motif distribution can be visualized by drawing a graph where edges represent the highly significant structural similarities shared by two protein structures. Figure 1A illustrates the protein structure similarity network obtained for the SCOP α/β -class. The network reveals many precise similarities between proteins from different folds, suggesting many evolutionary connections between members of the α/β -class.

The most prominent feature of the α/β similarity network is a sub-structure formed from edges connecting the three classical Rossmannoid folds defined in SCOP: Class 2—NAD(P)-binding Rossmann; Class 3—FAD/NAD(P)-binding domains; Class 4—nucleotide-binding domains (Fig. 1B). Figure 1B shows 1841 connections between c.2 and c.3, 266 connections between c.2 and c.4 and 285 connections between c.3 and c.4. Each edge represents similarity of at least four residues between fold representatives.

The most significant similarities can be observed in graphs that impose a minimum threshold of similar residues. Figure 1C shows a graph representing the α/β -class connections when at least seven residues are structurally similar. This network reveals that the c.23 flavodoxin-like fold (a Rossmannoid) and the c.26 adenine nucleotide hydrolase-like fold [a Rossmannoid that includes the Class I archeal aminoacyl-tRNA synthetase (aaRS) catalytic core] are connected to the classical Rossmannoids.

The network also shows that the c.51 anti-codon-binding domain of Class II aaRS shares a common motif of eight residues with the FixJ receiver (c.23). This network illustrates how the method

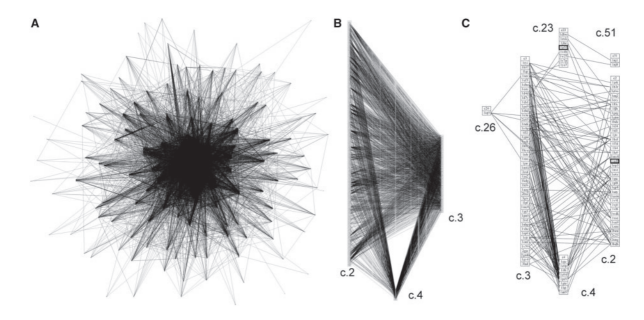


Fig. 1. Protein structure similarity networks for α/β proteins, classical Rossmannoids and a subset of motifs. (**A**) Edges were drawn by connecting representatives of each fold when the two motif structures were within 0.67 Å RMSD of each other. Each edge represents a similarity of at least four residues. (**B**) The structure similarity network is shown for the three classical Rossmannoid fold subsets; a fold is a column of PDB representatives in boxes: c.2, NAD(P)-binding Rossmann-fold domains; c.3, the FAD/NAD(P)-binding domain; and c.4, the nucleotide-binding domain. These classical Rossmannoids are the most densely connected fold clique of the similarity network in (A). (**C**) The structure similarity network is shown for the cases when at least seven residues are shared between the two proteins. The classical Rossmannoids are well-connected, and the remaining connections are to two different Rossmannoid folds (c.23 and c.26), as well as the connection between the anti-codon-binding domain of a Class II aminoacyl-tRNA synthetase (c.51) and c.23, the flavodoxin-like fold (FixJ structure). c.26 is the adenine nucleotide α-hydrolase-like fold, which includes the Class I aminoacyl-tRNA synthetase catalytic core; the representative in the c.26 box (1iq0) is the Class I argininyl-tRNA synthetase. All structures in 1C except the two highlighted by bold boxes possess at least partial representation of the core identified in the Rossmannoids and described further in this work.

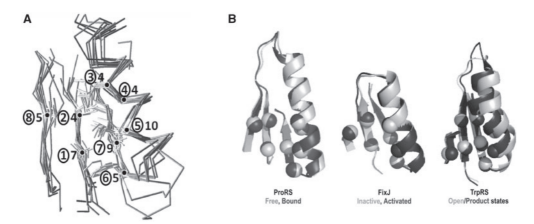


Fig. 2. Structurally similar cores in the ProRS anti-codon-binding domain, the FixJ receiver and other Rossmannoid representatives. (**A**) Cores from Figure 1C and Table 1 are shown as superimposed Cα traces and stick figures for residue side chains. The sequential β-α-β motif is formed from the central and rightmost strands and the α-helix. The number of consensus amino acids is given for each residue position (circled). (**B**) Functionally relevant allosteric changes in ProRS, FixJ and TrpRS core motifs. These three motifs exhibit conformational changes in response to ligand binding [ProRS anti-codon-binding domain, 1H4S, 1H4T; TrpRS catalytic domain, 1MAW(F), 1I6L] or phosphorylation [FixJ, 1D5W(A), 1DBW(A)]. Activated conformations are darker. The eight positions indicated in Table 1 are indicated by Cα positions that rearrange similarly in the three proteins. These changes have been implicated experimentally in long-range communication with the active site catalytic Mg²⁺ ion in TrpRS (Kapustina *et al.*, 2007; Weinreb *et al.*, 2009).

of comparison used here can reveal relationships among proteins that have diverged to the point of having distinct folds, as well as possibly identifying cases of convergence to a common motif. Further analyses identified five additional Rossmannoid folds that possess partial representation of this core. Superimposed representatives are shown in Figure 2, and statistics for core residues for each representative are collected in Table 1, together with crystal

 Table 1 Residues comprising cores shown in Figure 2

SCOP fold	NCBI/pfam Domains	PDB	ď	Core Pos	2	3	4	5	9	7	8
			RMSD	MSA							
c51—ClassII aaRS anti-codon-binding domain	ProRS cd00862:	1nj1A 2.55 Å 1h4sB		V297	1299	L318	R319	L322	F327	V329	V354
c51—mitochondrial polymerase gamma		2.63 A	0.4 Å	V357	14/221 L359	U.22E L376	U.22.A L.377	13/2/2L L380	1385	14/22 v V387	V415
(β)		1.95 Å		2/8V	4/8L 2I	78/L	1/8L	78/8	4/8L	A8/9	4/8V
c23—flavodoxin-like	FixJ Receiver cd00156:	1d5wA 2.3 Å 1dbwA	0.26 Å	9/	81	L19	A20	L23	F28	V30	V52
		1.68 Å			97/307I 95V	144/307L	41/307A	237/307L	58/307F 79Y	184/307V	54/307V 80I 120L
c2—NAD(P)-binding Rossmann-fold	PRK06179: PRK06179	ljtvA	0.44 Å	V5	17	L18	A19	L22	F30	V32	V88
domains		1.54 Å		12/24V 12A	9/24I 15V	1/24L	24/24A	14/24L 10F	8/24 F 7Y	24/24V	24/24V
c3—FAD/NAD(P)- binding	pfam07992:	1m6i	0.84 Å	1303	1305	L315	A316	L319	A323	V330	V395
domain	Pyr_redox_2	1.80 Å		46/146 V 28I 24L	53/146I 84V	28/146L 22F 20V 109/146A	109/146A	62/146L 32F 28A Gapped	Gapped	117/146V	31/146V 46L 26I
c66—SAM-dependent methyltransferases	pfam08241: Methyltransf_11	1r74A 2.55 Å	0.43 Å	09A	D62 113/177D	D70 26/177L	S71 30/177A	L74 85/177L	F79 4/177F Gapped	V81 81/177V	1136 24/177V 221 29L
c26—adenine nucleotide PRK00509: α-hydrolase-like PRK0050	PRK00509: PRK00509	1v12A 2.10 Å	0.65 Å	V5 134/143V	L7 138/143L	I18 83/1431 43A	L19 51/143L 88I	L22 119/143L	F27 V29 1/143F 18Y Gapped 125/143V	V29 125/143V	A113 95/143A
c4—nucleotide-binding domain	pfam00070: Pyr_redox	1094A 2.0 Å	0.46 Å	V392 37/131V	I394 46/131I	A404 25/131L 21F 7A	A405 102/131A	L408 55/131L 33F	Y413 4/131Y 13L Gapped 111/131V	V415 111/131V	1484 25/1311 46L 22V
c26—arginyl-tRNA synthetase	Class Ia aaRS Pfam00750	1iq0A 2.3 Å	0.42Å	V105 20/43V	V107 16/43V, 16I	1132 28/431	A133 23/43A	L136 25/43L	R141 19/43Y 4F	V143 37/43V	1350 31/431
c26—Tryptophanyl- tRNA	Class Ic aaRS	1161	1.83 Å	14	F37	(Missing)	F26	L29	Y33	C35	1140
synthetase	Pfam00579	1.72 Å	(0.91Å)		(Out of order)						
		3.0Å		19/99I 50V	42/99F		16/99F 47W 46/99L	46/99L	45/99Y	47/99C 11V 16/99I 35L	16/99I 35L
	Structure			>	I	Г	A	L	ц	>	>
	MSA's			I/Λ	I/V	I/L	A	Г	F/Y	>	I/L/V

Core residues are listed with MSA results for each domain. Structure resolutions (\mathring{a}) and $C\alpha$ RMSDs (\mathring{a}) for core residues compared with the ProRS anti-codon-binding domain also are shown. In boldface are consensus homologs from the Class I aminoacyl-tRNA synthetase family. ArgRS is identified in Figure 1C. TrpRS has a larger RMSD, as its C-terminal β -strand is displaced to form the tryptophan-binding site. The double line separates the variant TrpRS core from those identified in Figure 1C.

structure resolutions and RMSDs. Despite the stringent cut-off, only two structures in Figure 1C failed to exhibit this motif.

Pre-computed multiple sequence alignments (MSAs) were obtained from the conserved domain representatives at NCBI using the structure sequences as queries to determine the consensus core positions. Consensus sequences are shown in Table 1. Most positions followed the structures in Figure 1C closely, although some positions were represented by nearly equal numbers of similar amino acid types in the MSAs.

4 DISCUSSION AND CONCLUSIONS

(i) Four-body simplices afford a valuable extension of graphtheoretic applications from active site configurations to a far broader class of evolutionary problems: SNAPP detects precisely repeated structural details in many proteins from a nearly exhaustive representation of protein structures in the SCOP α/β class. The similarities indicate either evolutionary divergence or convergence. Although convergence cannot be ruled out, the N-terminal location and the coincidence of primary, secondary and tertiary structural similarities (Table 1; Fig. 2A) highlighted here—between classical Rossmannoids in the protein structure similarity network—are most naturally interpreted in terms of an early adaptive radiation and hence of divergent evolution. This interpretation is reinforced (Shakhnovich et al., 2003) by the fact that divergence could have been driven by the adaptive radiation of the same primordial functions of the core motif-nucleotide (phosphate) binding and conformational switching (Fig. 2B) shared by descendant Rossmannoid folds.

(ii) Construction of composite packing motifs involving elemental links between secondary structures affords a new metric for ancient protein phylogeny: any core motif identified by our procedure implicitly comprises many *bona fide* examples of divergent evolution represented in the MSAs belonging to each fold family. Importantly, there is less variation between the motifs in different fold families in Figure 1C than is evident within single fold families (i.e. the ArgRS and TrpRS core motifs in Table 1). Comparison of the core motif in ArgRS and TrpRS illustrates that using a strict threshold for RMSD and residue identity for identification helps assure significance of the motifs identified, especially in the face of considerable variation evident in motif instances within consensus superfamilies.

(iii)One eight-residue motif, in particular, links the N-terminal β - α - β cross-over to a downstream β -strand especially frequently in the Rossmannoid family: common amino acids in all seven topologies are consistently reinforced by the associated MSAs, raising the possibility that at some time before the LCA, a small ancestral protein with this N-terminal β - α - β cross-over connection of 30-40 amino acids, tied to a downstream β -strand, diverged to eventually form all Rossmannoids including the Class I aminoacyltRNA synthetases, as well as the anti-codon-binding domain of Class II aminoacyl-tRNA synthetases for proline and glycine, and the accessory β -subunit of the mitochondrial DNA polymerase γ (Kaguni, 2004). This example, therefore, shows that examining precisely repeated inter-residue packing arrangements using an SNAPP-based bioinformatics approach can lead to consistent and novel inferences about evolutionarily divergent and/or convergent motifs shared by proteins with similar but distinct folds.

(iv) The motif at the N-terminus of the Rossmann fold has independently emerged as having important nucleotide-binding and proto-allosteric functionality. The widespread occurrence of this motif is especially interesting in light of its consensus association with nucleotide-binding functions and its likely allosteric behavior (Fig. 2B). Anti-codon binding by the ProRS domain leads to structural rearrangement that is likely communicated to the active site (Yaremchuk et al., 2000). Structural changes in the FixJ motif upon phosphorylation of D54 complement the rearrangement of F106 associated with its activation for transcriptional regulation (Birck et al., 1999), and a similar change occurs in the bacterial chemotaxis response regulator, CheY (Schuster et al., 2001). The homologous motif in Class I tryptophanyl-tRNA synthetase functions in molecular switching that accompanies both induced fit and catalysis (Kapustina et al., 2007). Mutation (i.e. of F37I) exerts long-range effects on the relative stability of different TrpRS conformational states and reduces the catalytic contribution of the Mg^{2+} ion (Weinreb et al., 2009).

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REFERENCES

Aravind, L. et al. (2002) Trends in protein evolution inferred from sequence and structure analysis. Curr. Opin. Struct. Biol., 12, 392–399.

Barker, J.A. and Thornton, J.W. (2003) An algorithm for constraint-based structural template matching: application to 3D templastes with statistical analysis. *Bioinformatics*, 19, 1644–1649.

Birck, C. et al. (1999) Conformational changes induced by phosphorylation of the FixJ receiver domain. Struct. Fold. Design, 7, 1505–1515.

Cammer,S.A. et al. (2002) Identification of sequence-specific tertiary packing motifs in protein structures using Delaunay tessellation. In Schlick,T. and Gan,H.H. (eds) Lecture Notes in Computational Science and Engineering, Vol. 24. Springer, New York, pp. 477–494.

Chothia, C. and Janin, J. (1982) Orthogonal packing of b-pleated sheets in proteins. Biochemistry, 21, 3955–3965.

Chothia, C. et al. (1977) Structure of proteins: packing of a-helices and pleated sheets. Proc. Natl Acad. Sci. USA, 74, 4130–4134.

Chothia, C. et al. (1981) Helix to helix packing in proteins. J. Mol. Biol., 145, 215–250.
Crick, F.H.C. (1953) The packing of a-helices: simple coiled-coils. Acta Cryst., 6, 689–697.

Dror, O. et al. (2003) Multiple structural alignment by secondary structures: algorithm and applications. Prot. Sci., 12, 2492–2507.

Fetrow, J.S. et al. (1999) Structure-based functional motif identifies a potential disulfide oxidoreductase active site in the serine/threonine protein phosphatase-I subfamily. FASEB J., 13, 1866–1874.

Fetrow,J.S. and Skolnik,J. (1998) Method for prediction of protein function from sequence using the sequence-to-structure-to-function paradigm with application to glutaredoxins/thioredoxins and t1 ribonucleases. J. Mol. Biol., 281, 949–968.

Fischer, D. et al. (1994) Three-dimensional, sequence order-independent structural comparison of a serine protease against the crystallographic database reveals active site similarities: potential implications to evolution and to protein folding. Prot. Sci., 3, 769–778.

Holm,L. and Sander,C. (1993) Protein structure comparison by alignment of distance matrices. J. Mol. Biol., 233, 123–138.

Janin, J. and Chothia, C. (1980) Packing of alpha-helices onto beta-pleated sheets and the anatomy of alpha/beta proteins. J. Mol. Biol., 143, 95–128.

- Jambon, M. et al. (2003) A new bioinformatic approach to detect common 3D sites in protein structures. Proteins, Structure, Function and Bioinformatics, 52, 137–145.
- Kaguni,L.S. (2004) DNA polymerase g, the mitochondrial replicase. Ann. Rev. Biochem., 73, 293–320.
- Kapustina, M. et al. (2007) A conformational transition state accompanies amino acid activation by B. stearothermphilus tryptophanyl-trna synthetase. Structure, 15, 1272–1284.
- Madej,T. et al. (1995) Threading a database of protein cores. Proteins, Structure, Function and Genetics, 23, 536–540.
- Murzin, A.G. et al. (1995) SCOP: a structural classification of proteins database for the investigation of sequences and structures. J. Mol. Biol., 247, 536–540.
- Orengo, C.A. and Taylor, W.R. (1990) A rapid method for protein structure alignment. *J. Theor. Biol.*, **147**, 517–551.
- Schuster, M. et al. (2001) Conformational coupling in the chemotaxis response regulator chey. Proc. Natl Acad. Sci. USA, 98, 6003–6008.

- Shakhnovich, B.E. et al. (2003) Functional fingerprints of folds: evidence for correlated structure-function evolution. J. Mol. Biol., 326, 1–9.
- Shindyalov,I.N. and Bourne,P.E. (1998) Protein structure alignment by incremental combinatorial extension of the optimal path. *Prot. Eng.*, 11, 739–747.
- Stark, A. et al. (2003) A model for the statistical significance of local similarities in structure. J. Mol. Biol., 326, 1307–1316.
- Tropsha, A. et al. (2003) Simplicial Neighborhood Analysis of Protein Packing (SNAPP): a computational geometry approach to studying proteins. *Meth. Enz.*, **374**, 509–544.
- Weinreb, V. et al. (2009) Mg²⁺-assisted catalysis by B. stearothermophilus trprs is promoted by allosteric effects. Structure, 17, 952–964.
- Yaremchuk, A. et al. (2000) Crystal structure of a eukaryote/archaeon-like protyltRNA synthetase and its complex with tRNApro(cgg). EMBO J., 19, 4745–4758.