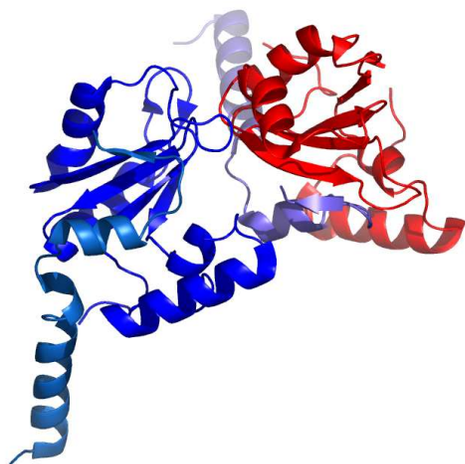


2f9d

Evolutionary trace report by `report_maker`

March 12, 2010



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1 INTRODUCTION

- 1 From the original Protein Data Bank entry (PDB id 2f9d):
- 1 **Title:** 2.5 angstrom resolution structure of the spliceosomal protein p14 bound to region of sf3b155
- 1 **Compound:** Mol id: 1; molecule: pre-mrna branch site protein p14; chain: a, b; synonym: sf3b 14 kda subunit; engineered: yes; mol id: 2; molecule: splicing factor 3b subunit 1; chain: p, q; fragment: residues: 373-415; synonym: spliceosome associated protein 155, sap 155, sf3b155, pre-mrna splicing factor sf3b 155 kda subunit; engineered: yes
- 2 **Organism, scientific name:** Homo Sapiens;
- 2 2f9d contains unique chains 2f9dA (114 residues) and 2f9dP (39 residues) 2f9dB is a homologue of chain 2f9dA. 2f9dQ is a homologue of chain 2f9dP.
- 4

5 2 CHAIN 2F9DA

5 2.1 Q6IQM0 overview

- 5 From SwissProt, id Q6IQM0, 88% identical to 2f9dA:
- 5 **Description:** Hypothetical protein zgc:86708 (Novel protein similar to vertebrate pre-mRNA branch site protein p14) (P14).
- 5 **Organism, scientific name:** Brachydanio rerio (Zebrafish) (Danio rerio).
- 6 **Taxonomy:** Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Actinopterygii; Neopterygii; Teleostei; Ostariophysi; Cypriniformes; Cyprinidae; Danio.
- 7 **Subcellular location:** Nuclear (By similarity).
- 7

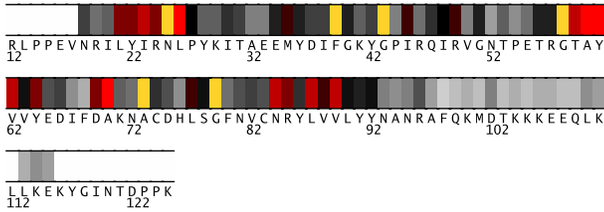


Fig. 1. Residues 12-125 in 2f9dA colored by their relative importance. (See Appendix, Fig.12, for the coloring scheme.)

2.2 Multiple sequence alignment for 2f9dA

For the chain 2f9dA, the alignment 2f9dA.msf (attached) with 142 sequences was used. The alignment was downloaded from the HSSP database, and fragments shorter than 75% of the query as well as duplicate sequences were removed. It can be found in the attachment to this report, under the name of 2f9dA.msf. Its statistics, from the *alistat* program are the following:

```

Format:                MSF
Number of sequences:   142
Total number of residues: 14399
Smallest:              86
Largest:               114
Average length:        101.4
Alignment length:      114
Average identity:       32%
Most related pair:     99%
Most unrelated pair:   10%
Most distant seq:      30%

```

Furthermore, <1% of residues show as conserved in this alignment.

The alignment consists of 45% eukaryotic (15% vertebrata, 6% arthropoda, 9% fungi, 3% plantae) sequences. (Descriptions of some sequences were not readily available.) The file containing the sequence descriptions can be found in the attachment, under the name 2f9dA.descr.

2.3 Residue ranking in 2f9dA

The 2f9dA sequence is shown in Fig. 1, with each residue colored according to its estimated importance. The full listing of residues in 2f9dA can be found in the file called 2f9dA.ranks.sorted in the attachment.

2.4 Top ranking residues in 2f9dA and their position on the structure

In the following we consider residues ranking among top 25% of residues in the protein. Figure 2 shows residues in 2f9dA colored by their importance: bright red and yellow indicate more conserved/important residues (see Appendix for the coloring scheme). A Pymol script for producing this figure can be found in the attachment.

2.4.1 Clustering of residues at 25% coverage. Fig. 3 shows the top 25% of all residues, this time colored according to clusters they belong to. The clusters in Fig.3 are composed of the residues listed in Table 1.

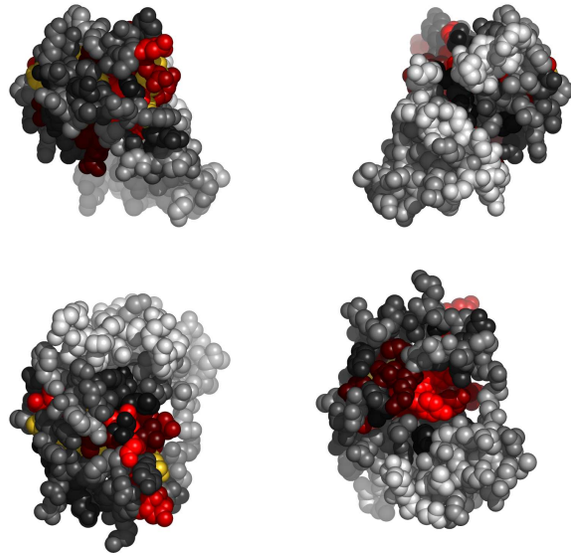


Fig. 2. Residues in 2f9dA, colored by their relative importance. Clockwise: front, back, top and bottom views.

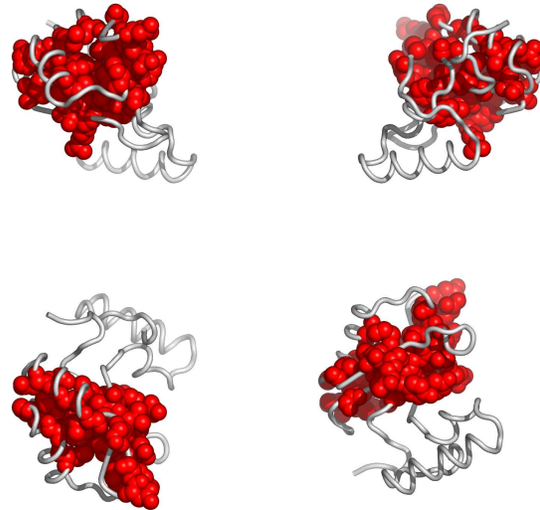


Fig. 3. Residues in 2f9dA, colored according to the cluster they belong to: red, followed by blue and yellow are the largest clusters (see Appendix for the coloring scheme). Clockwise: front, back, top and bottom views. The corresponding Pymol script is attached.

Table 1.		
cluster color	size	member residues
red	29	21, 22, 23, 24, 25, 26, 27, 35, 39 43, 45, 48, 49, 58, 59, 60, 61, 62 64, 69, 70, 73, 77, 79, 84, 85, 87 88, 89

Table 1. Clusters of top ranking residues in 2f9dA.

2.4.2 Overlap with known functional surfaces at 25% coverage.

The name of the ligand is composed of the source PDB identifier and the heteroatom name used in that file.

Interface with 2f9dP. By analogy with 2f9dB – 2f9dP interface. Table 2 lists the top 25% of residues at the interface with 2f9dP. The following table (Table 3) suggests possible disruptive replacements for these residues (see Section 4.6).

Table 2.					
res	type	subst's (%)	cvg	noc/ bb	dist (Å)
24	R	F (9)	0.18	1/0	4.58
		G (19)			
		N (11)			
		K (21)			
		T (4)			
		R (29)			
88	V	D (2) YS.	0.24	1/0	4.46
		L			
		R (33)			
		K (9)			
		V (31)			
		H (2)			
		I (9)			
		E (4) N			
		Y (2)			
		T (1)			
		Q (2)			
		L (1)			
F (1)					

Table 2. The top 25% of residues in 2f9dA at the interface with 2f9dP. (Field names: res: residue number in the PDB entry; type: amino acid type; substs: substitutions seen in the alignment; with the percentage of each type in the bracket; noc/bb: number of contacts with the ligand, with the number of contacts realized through backbone atoms given in the bracket; dist: distance of closest approach to the ligand.)

Table 3.		
res	type	disruptive mutations
24	R	(D) (T) (E) (Y)
88	V	(Y) (E) (R) (K)

Table 3. List of disruptive mutations for the top 25% of residues in 2f9dA, that are at the interface with 2f9dP.

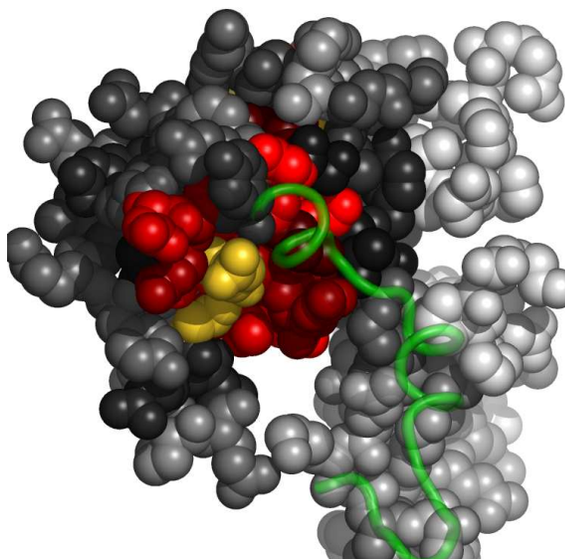


Fig. 4. Residues in 2f9dA, at the interface with 2f9dP, colored by their relative importance. 2f9dP is shown in backbone representation (See Appendix for the coloring scheme for the protein chain 2f9dA.)

Figure 4 shows residues in 2f9dA colored by their importance, at the interface with 2f9dP.

Interface with 2f9dB. Table 4 lists the top 25% of residues at the interface with 2f9dB. The following table (Table 5) suggests possible disruptive replacements for these residues (see Section 4.6).

Table 4.					
res	type	subst's (%)	cvg	noc/ bb	dist (Å)
43	G	G (99) A	0.02	17/17	3.03
69	D	D (68)	0.15	10/0	2.54
		S (12)			
		A (7)			
		T (4)			
		E (1) G			
45	I	N (1) QHC	0.21	1/1	4.40
		P			
		L (11)			
		I (64)			
		V (22) AG			
		P			

Table 4. The top 25% of residues in 2f9dA at the interface with 2f9dB. (Field names: res: residue number in the PDB entry; type: amino acid type; substs: substitutions seen in the alignment; with the percentage of each type in the bracket; noc/bb: number of contacts with the ligand, with the number of contacts realized through backbone atoms given in the bracket; dist: distance of closest approach to the ligand.)

Table 5.		
res	type	disruptive mutations
43	G	(KER) (QHD) (FYMW) (N)
69	D	(R) (H) (FW) (Y)
45	I	(R) (Y) (H) (KE)

Table 5. List of disruptive mutations for the top 25% of residues in 2f9dA, that are at the interface with 2f9dB.

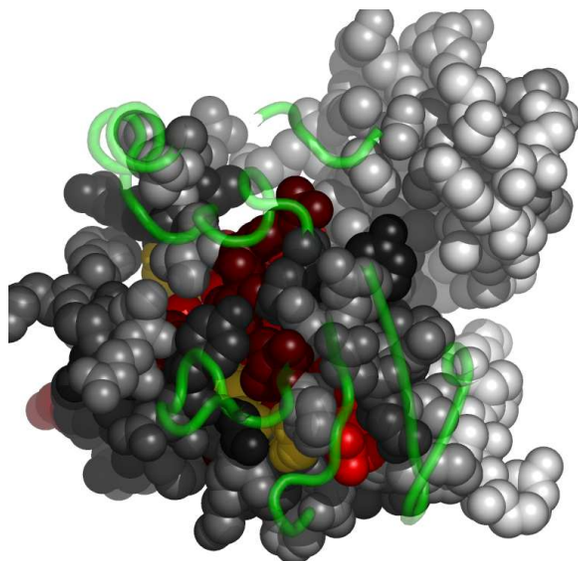


Fig. 5. Residues in 2f9dA, at the interface with 2f9dB, colored by their relative importance. 2f9dB is shown in backbone representation (See Appendix for the coloring scheme for the protein chain 2f9dA.)

Figure 5 shows residues in 2f9dA colored by their importance, at the interface with 2f9dB.

2.4.3 Possible novel functional surfaces at 25% coverage. One group of residues is conserved on the 2f9dA surface, away from (or substantially larger than) other functional sites and interfaces recognizable in PDB entry 2f9d. It is shown in Fig. 6. The right panel shows (in blue) the rest of the larger cluster this surface belongs to. The residues belonging to this surface "patch" are listed in Table 6, while Table 7 suggests possible disruptive replacements for these residues (see Section 4.6).

Table 6.			
res	type	substitutions(%)	cvg
43	G	G(99)A	0.02
58	G	G(96)PS(2)T	0.03
25	N	G(14)N(78)EY(4) PH.D	0.04
79	G	G(86)N(2)MYD(4)	0.05

continued in next column

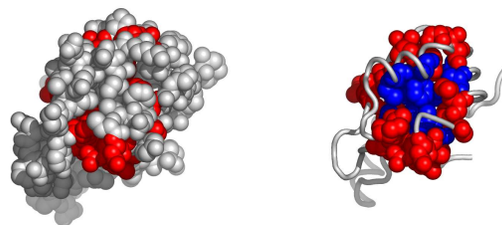


Fig. 6. A possible active surface on the chain 2f9dA. The larger cluster it belongs to is shown in blue.

Table 6. <i>continued</i>			
res	type	substitutions(%)	cvg
26	L	R(3)KE(1) L(82)I(10)V(4) F(1).	0.07
61	Y	F(77)W(1)H(1) Y(14)I(3)L(1)	0.09
89	V	V(74)I(18)AL(2) T(1)SYED	0.10
84	N	G(47)D(16)N(31) E(2).P	0.13
59	T	F(31)Y(11)Q(11) T(40)C(2)I(1)S	0.14
69	D	D(68)S(12)A(7) T(4)E(1)GN(1)QH CP	0.15
64	Y	F(38)Y(60)M(1)	0.16
22	Y	G(8)Y(47)F(27) K(2)L(6)I(4) S(1).(1)	0.17
85	R	R(71)G(1)K(16)Q VMFC(1)E(2).SL H(1)T	0.17
24	R	F(9)G(19)N(11) K(21)T(4)R(29) D(2)YS.L	0.18
77	L	A(6)R(4)M(14) L(67)KTV(1)I(1) C(2)	0.20
45	I	L(11)I(64)V(22) AGP	0.21
49	R	N(5)H(2)L(3) V(7)R(56)FE(1) Y(6)D(4)W(3) Q(4)TKS(1)I(1)	0.22
88	V	R(33)K(9)V(31) H(2)I(9)E(4)N Y(2)T(1)Q(2) L(1)F(1)	0.24

continued in next column

res	type	substitutions(%)	cvg
27	P	S(16)P(52)N(14) FD(2)T(2)A(3) G(4)HR.Y	0.25
48	I	V(24)I(62)F(2) L(3)C(4)MAH	0.25

Table 6. Residues forming surface "patch" in 2f9dA.

res	type	disruptive mutations
43	G	(KER)(QHD)(FYMW)(N)
58	G	(R)(K)(E)(H)
25	N	(Y)(FWH)(TR)(VA)
79	G	(R)(FW)(H)(KE)
26	L	(R)(Y)(T)(H)
61	Y	(K)(Q)(E)(R)
89	V	(R)(K)(Y)(H)
84	N	(Y)(H)(FW)(R)
59	T	(KR)(Q)(H)(M)
69	D	(R)(H)(FW)(Y)
64	Y	(K)(Q)(R)(E)
22	Y	(K)(Q)(R)(EM)
85	R	(D)(T)(Y)(E)
24	R	(D)(T)(E)(Y)
77	L	(Y)(R)(H)(T)
45	I	(R)(Y)(H)(KE)
49	R	(T)(D)(Y)(E)
88	V	(Y)(E)(R)(K)
27	P	(R)(Y)(K)(H)
48	I	(R)(Y)(TE)(KH)

Table 7. Disruptive mutations for the surface patch in 2f9dA.

3 CHAIN 2F9DP

3.1 Q4QR12 overview

From SwissProt, id Q4QR12, 94% identical to 2f9dP:

Description: Hypothetical protein.

Organism, scientific name: *Xenopus laevis* (African clawed frog).

Taxonomy: Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Amphibia; Batrachia; Anura; Mesobatrachia; Pipoidae; Pipidae; Xenopodinae; Xenopus; Xenopus.

3.2 Multiple sequence alignment for 2f9dP

For the chain 2f9dP, the alignment 2f9dP.msf (attached) with 26 sequences was used. The alignment was downloaded from the HSSP database, and fragments shorter than 75% of the query as well as duplicate sequences were removed. It can be found in the attachment to this report, under the name of 2f9dP.msf. Its statistics, from the *alistat* program are the following:



Fig. 7. Residues 377-415 in 2f9dP colored by their relative importance. (See Appendix, Fig.12, for the coloring scheme.)

```

Format:                               MSF
Number of sequences:                   26
Total number of residues:              984
Smallest:                              32
Largest:                               39
Average length:                        37.8
Alignment length:                      39
Average identity:                       67%
Most related pair:                      97%
Most unrelated pair:                   35%
Most distant seq:                      56%

```

Furthermore, 7% of residues show as conserved in this alignment.

The alignment consists of 30% eukaryotic (7% vertebrata, 7% arthropoda, 3% plantae) sequences. (Descriptions of some sequences were not readily available.) The file containing the sequence descriptions can be found in the attachment, under the name 2f9dP.descr.

3.3 Residue ranking in 2f9dP

The 2f9dP sequence is shown in Fig. 7, with each residue colored according to its estimated importance. The full listing of residues in 2f9dP can be found in the file called 2f9dP.ranks.sorted in the attachment.

3.4 Top ranking residues in 2f9dP and their position on the structure

In the following we consider residues ranking among top 26% of residues in the protein (the closest this analysis allows us to get to 25%). Figure 8 shows residues in 2f9dP colored by their importance: bright red and yellow indicate more conserved/important residues (see Appendix for the coloring scheme). A Pymol script for producing this figure can be found in the attachment.

3.4.1 Clustering of residues at 26% coverage. Fig. 9 shows the top 26% of all residues, this time colored according to clusters they belong to. The clusters in Fig.9 are composed of the residues listed in Table 8.

cluster color	size	member residues
red	3	401, 403, 404
blue	3	395, 396, 398
yellow	3	379, 380, 382

Table 8. Clusters of top ranking residues in 2f9dP.

3.4.2 Overlap with known functional surfaces at 26% coverage. The name of the ligand is composed of the source PDB identifier and the heteroatom name used in that file.

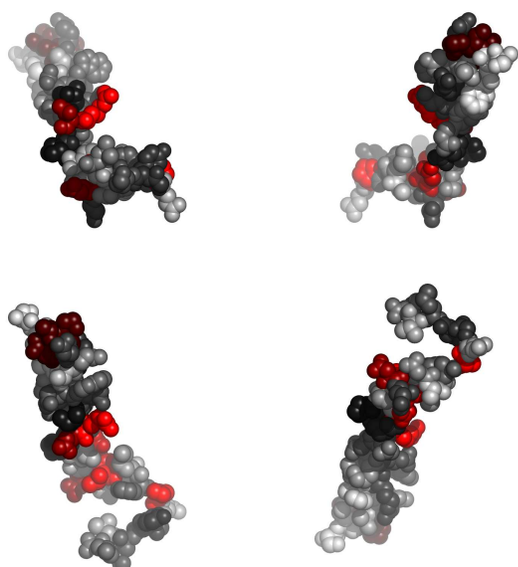


Fig. 8. Residues in 2f9dP, colored by their relative importance. Clockwise: front, back, top and bottom views.

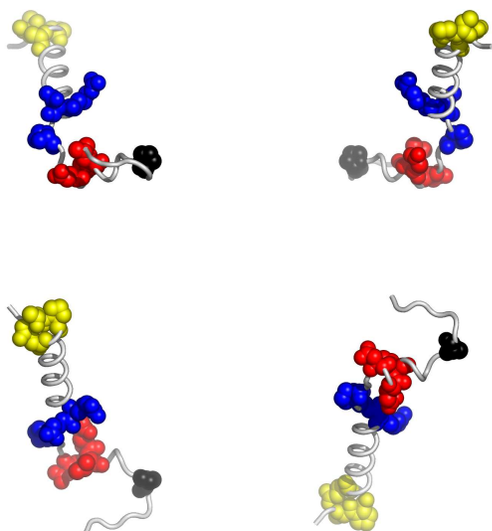


Fig. 9. Residues in 2f9dP, colored according to the cluster they belong to: red, followed by blue and yellow are the largest clusters (see Appendix for the coloring scheme). Clockwise: front, back, top and bottom views. The corresponding Pymol script is attached.

Interface with 2f9dA. Table 9 lists the top 26% of residues at the interface with 2f9dA. The following table (Table 10) suggests possible disruptive replacements for these residues (see Section 4.6).

res	type	subst's (%)	cvg	noc/bb	dist (Å)
395	R	R (100)	0.08	70/18	2.66
404	L	L (100)	0.08	29/4	3.61
409	P	P (100)	0.08	33/9	3.17
403	E	E (96) D (3)	0.10	42/16	3.42
396	N	N (96) X (3)	0.15	68/20	2.83
401	D	D (96) X (3)	0.15	12/1	3.12

Table 9. The top 26% of residues in 2f9dP at the interface with 2f9dA. (Field names: res: residue number in the PDB entry; type: amino acid type; substs: substitutions seen in the alignment; with the percentage of each type in the bracket; noc/bb: number of contacts with the ligand, with the number of contacts realized through backbone atoms given in the bracket; dist: distance of closest approach to the ligand.)

res	type	disruptive mutations
395	R	(TD) (SYEVCLAPIG) (FMW) (N)
404	L	(YR) (TH) (SKECG) (FQWD)
409	P	(YR) (TH) (SKECG) (FQWD)
403	E	(FWH) (R) (YVCAG) (T)
396	N	(Y) (FTWH) (ER) (SVCAG)
401	D	(R) (FWH) (KY) (VCAG)

Table 10. List of disruptive mutations for the top 26% of residues in 2f9dP, that are at the interface with 2f9dA.

Figure 10 shows residues in 2f9dP colored by their importance, at the interface with 2f9dA.

Interface with 2f9dB. Table 11 lists the top 26% of residues at the interface with 2f9dB. The following table (Table 12) suggests possible disruptive replacements for these residues (see Section 4.6).

res	type	subst's (%)	cvg	noc/bb	dist (Å)
396	N	N (96) X (3)	0.15	2/0	4.50

Table 11. The top 26% of residues in 2f9dP at the interface with 2f9dB. (Field names: res: residue number in the PDB entry; type: amino acid type; substs: substitutions seen in the alignment; with the percentage of each type in the bracket; noc/bb: number of contacts with the ligand, with the number of contacts realized through backbone atoms given in the bracket; dist: distance of closest approach to the ligand.)

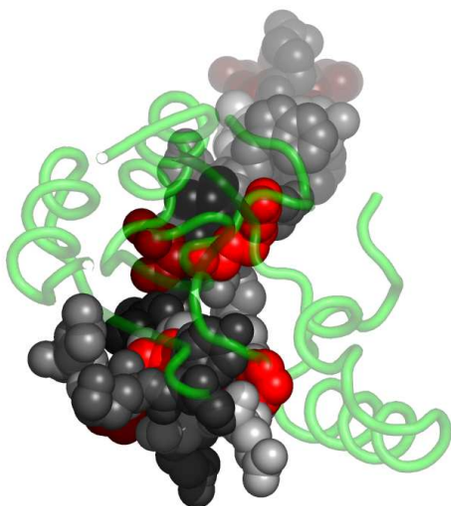


Fig. 10. Residues in 2f9dP, at the interface with 2f9dA, colored by their relative importance. 2f9dA is shown in backbone representation (See Appendix for the coloring scheme for the protein chain 2f9dP.)

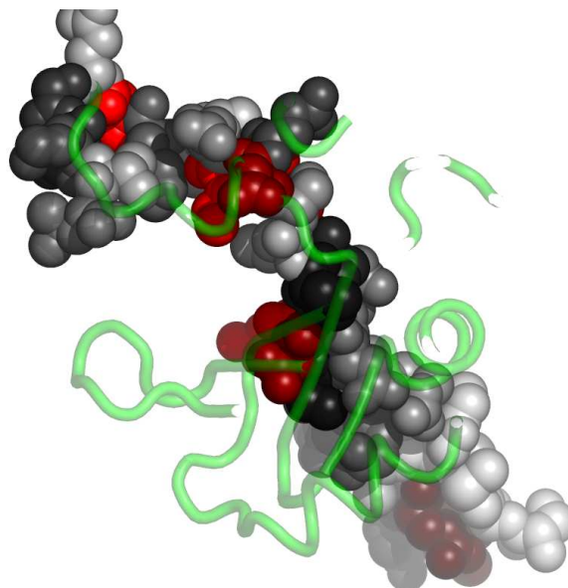


Fig. 11. Residues in 2f9dP, at the interface with 2f9dB, colored by their relative importance. 2f9dB is shown in backbone representation (See Appendix for the coloring scheme for the protein chain 2f9dP.)

Table 12.		
res	type	disruptive mutations
396	N	(Y) (FTWH) (ER) (SVCAG)

Table 12. List of disruptive mutations for the top 26% of residues in 2f9dP, that are at the interface with 2f9dB.

Figure 11 shows residues in 2f9dP colored by their importance, at the interface with 2f9dB.

4 NOTES ON USING TRACE RESULTS

4.1 Coverage

Trace results are commonly expressed in terms of coverage: the residue is important if its “coverage” is small - that is if it belongs to some small top percentage of residues [100% is all of the residues in a chain], according to trace. The ET results are presented in the form of a table, usually limited to top 25% percent of residues (or to some nearby percentage), sorted by the strength of the presumed evolutionary pressure. (I.e., the smaller the coverage, the stronger the pressure on the residue.) Starting from the top of that list, mutating a couple of residues should affect the protein somehow, with the exact effects to be determined experimentally.

4.2 Known substitutions

One of the table columns is “substitutions” - other amino acid types seen at the same position in the alignment. These amino acid types may be interchangeable at that position in the protein, so if one wants to affect the protein by a point mutation, they should be avoided. For example if the substitutions are “RVK” and the original protein has

an R at that position, it is advisable to try anything, but RVK. Conversely, when looking for substitutions which will *not* affect the protein, one may try replacing, R with K, or (perhaps more surprisingly), with V. The percentage of times the substitution appears in the alignment is given in the immediately following bracket. No percentage is given in the cases when it is smaller than 1%. This is meant to be a rough guide - due to rounding errors these percentages often do not add up to 100%.

4.3 Surface

To detect candidates for novel functional interfaces, first we look for residues that are solvent accessible (according to DSSP program) by at least 10\AA^2 , which is roughly the area needed for one water molecule to come in the contact with the residue. Furthermore, we require that these residues form a “cluster” of residues which have neighbor within 5\AA from any of their heavy atoms.

Note, however, that, if our picture of protein evolution is correct, the neighboring residues which *are not* surface accessible might be equally important in maintaining the interaction specificity - they should not be automatically dropped from consideration when choosing the set for mutagenesis. (Especially if they form a cluster with the surface residues.)

4.4 Number of contacts

Another column worth noting is denoted “noc/bb”; it tells the number of contacts heavy atoms of the residue in question make across the interface, as well as how many of them are realized through the backbone atoms (if all or most contacts are through the backbone, mutation presumably won’t have strong impact). Two heavy atoms are considered to be “in contact” if their centers are closer than 5\AA .

4.5 Annotation

If the residue annotation is available (either from the pdb file or from other sources), another column, with the header “annotation” appears. Annotations carried over from PDB are the following: site (indicating existence of related site record in PDB), S-S (disulfide bond forming residue), hb (hydrogen bond forming residue, jb (james bond forming residue), and sb (for salt bridge forming residue).

4.6 Mutation suggestions

Mutation suggestions are completely heuristic and based on complementarity with the substitutions found in the alignment. Note that they are meant to be **disruptive** to the interaction of the protein with its ligand. The attempt is made to complement the following properties: small [AVGSTC], medium [LPNQDEMIK], large [WFYHR], hydrophobic [LPVAMWFI], polar [GTCY]; positively [KHR], or negatively [DE] charged, aromatic [WFYH], long aliphatic chain [EK RQM], OH-group possession [SDETY], and NH2 group possession [NQRK]. The suggestions are listed according to how different they appear to be from the original amino acid, and they are grouped in round brackets if they appear equally disruptive. From left to right, each bracketed group of amino acid types resembles more strongly the original (i.e. is, presumably, less disruptive) These suggestions are tentative - they might prove disruptive to the fold rather than to the interaction. Many researcher will choose, however, the straightforward alanine mutations, especially in the beginning stages of their investigation.

5 APPENDIX

5.1 File formats

Files with extension “ranks_sorted” are the actual trace results. The fields in the table in this file:

- alignment# number of the position in the alignment
- residue# residue number in the PDB file
- type amino acid type
- rank rank of the position according to older version of ET
- variability has two subfields:
 1. number of different amino acids appearing in in this column of the alignment
 2. their type
- rho ET score - the smaller this value, the lesser variability of this position across the branches of the tree (and, presumably, the greater the importance for the protein)
- cvg coverage - percentage of the residues on the structure which have this rho or smaller
- gaps percentage of gaps in this column

5.2 Color schemes used

The following color scheme is used in figures with residues colored by cluster size: black is a single-residue cluster; clusters composed of more than one residue colored according to this hierarchy (ordered by descending size): red, blue, yellow, green, purple, azure, turquoise, brown, coral, magenta, LightSalmon, SkyBlue, violet, gold, bisque, LightSlateBlue, orchid, RosyBrown, MediumAquamarine, DarkOliveGreen, CornflowerBlue, grey55, burlywood, LimeGreen, tan, DarkOrange, DeepPink, maroon, BlanchedAlmond.

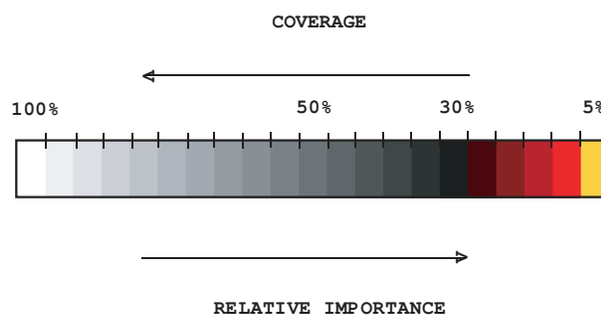


Fig. 12. Coloring scheme used to color residues by their relative importance.

The colors used to distinguish the residues by the estimated evolutionary pressure they experience can be seen in Fig. 12.

5.3 Credits

5.3.1 **Alistat** *alistat* reads a multiple sequence alignment from the file and shows a number of simple statistics about it. These statistics include the format, the number of sequences, the total number of residues, the average and range of the sequence lengths, and the alignment length (e.g. including gap characters). Also shown are some percent identities. A percent pairwise alignment identity is defined as $(\text{idents} / \text{MIN}(\text{len1}, \text{len2}))$ where *idents* is the number of exact identities and *len1*, *len2* are the unaligned lengths of the two sequences. The “average percent identity”, “most related pair”, and “most unrelated pair” of the alignment are the average, maximum, and minimum of all $(N)(N-1)/2$ pairs, respectively. The “most distant seq” is calculated by finding the maximum pairwise identity (best relative) for all *N* sequences, then finding the minimum of these *N* numbers (hence, the most outlying sequence). *alistat* is copyrighted by HHMI/Washington University School of Medicine, 1992-2001, and freely distributed under the GNU General Public License.

5.3.2 **CE** To map ligand binding sites from different source structures, *report_maker* uses the CE program: <http://cl.sdsc.edu/>. Shindyalov IN, Bourne PE (1998) “Protein structure alignment by incremental combinatorial extension (CE) of the optimal path”. *Protein Engineering* 11(9) 739-747.

5.3.3 **DSSP** In this work a residue is considered solvent accessible if the DSSP program finds it exposed to water by at least 10\AA^2 , which is roughly the area needed for one water molecule to come in the contact with the residue. DSSP is copyrighted by W. Kabsch, C. Sander and MPI-MF, 1983, 1985, 1988, 1994 1995, CMBI version by Elmar.Krieger@cmbi.kun.nl November 18,2002,

<http://www.cmbi.kun.nl/gv/dssp/descrip.html>.

5.3.4 **HSSP** Whenever available, *report_maker* uses HSSP alignment as a starting point for the analysis (sequences shorter than 75% of the query are taken out, however); R. Schneider, A. de Daruvar, and C. Sander. “The HSSP database of protein structure-sequence alignments.” *Nucleic Acids Res.*, 25:226–230, 1997.

<http://swift.cmbi.kun.nl/swift/hssp/>

5.3.5 LaTeX The text for this report was processed using L^AT_EX; Leslie Lamport, "LaTeX: A Document Preparation System Addison-Wesley," Reading, Mass. (1986).

5.3.6 Muscle When making alignments "from scratch", report maker uses Muscle alignment program: Edgar, Robert C. (2004), "MUSCLE: multiple sequence alignment with high accuracy and high throughput." Nucleic Acids Research 32(5), 1792-97.

<http://www.drive5.com/muscle/>

5.3.7 Pymol The figures in this report were produced using Pymol. The scripts can be found in the attachment. Pymol is an open-source application copyrighted by DeLano Scientific LLC (2005). For more information about Pymol see <http://pymol.sourceforge.net/>. (Note for Windows users: the attached package needs to be unzipped for Pymol to read the scripts and launch the viewer.)

5.4 Note about ET Viewer

Dan Morgan from the Lichtarge lab has developed a visualization tool specifically for viewing trace results. If you are interested, please visit:

<http://mammoth.bcm.tmc.edu/traceview/>

The viewer is self-unpacking and self-installing. Input files to be used with ETV (extension .etvx) can be found in the attachment to the main report.

5.5 Citing this work

The method used to rank residues and make predictions in this report can be found in Mihalek, I., I. Reš, O. Lichtarge. (2004). "A Family of Evolution-Entropy Hybrid Methods for Ranking of Protein Residues by Importance" J. Mol. Bio. **336**: 1265-82. For the original version of ET see O. Lichtarge, H.Bourne and F. Cohen (1996). "An Evolutionary Trace Method Defines Binding Surfaces Common to Protein Families" J. Mol. Bio. **257**: 342-358.

report_maker itself is described in Mihalek I., I. Res and O. Lichtarge (2006). "Evolutionary Trace Report Maker: a new type of service for comparative analysis of proteins." Bioinformatics **22**:1656-7.

5.6 About report_maker

report_maker was written in 2006 by Ivana Mihalek. The 1D ranking visualization program was written by Ivica Reš. **report_maker**

is copyrighted by Lichtarge Lab, Baylor College of Medicine, Houston.

5.7 Attachments

The following files should accompany this report:

- 2f9dA.complex.pdb - coordinates of 2f9dA with all of its interacting partners
- 2f9dA.etvx - ET viewer input file for 2f9dA
- 2f9dA.cluster_report.summary - Cluster report summary for 2f9dA
- 2f9dA.ranks - Ranks file in sequence order for 2f9dA
- 2f9dA.clusters - Cluster descriptions for 2f9dA
- 2f9dA.msf - the multiple sequence alignment used for the chain 2f9dA
- 2f9dA.descr - description of sequences used in 2f9dA msf
- 2f9dA.ranks_sorted - full listing of residues and their ranking for 2f9dA
- 2f9dA.2f9dP.if.pml - Pymol script for Figure 4
- 2f9dA.cbcbvg - used by other 2f9dA – related pymol scripts
- 2f9dA.2f9dB.if.pml - Pymol script for Figure 5
- 2f9dP.complex.pdb - coordinates of 2f9dP with all of its interacting partners
- 2f9dP.etvx - ET viewer input file for 2f9dP
- 2f9dP.cluster_report.summary - Cluster report summary for 2f9dP
- 2f9dP.ranks - Ranks file in sequence order for 2f9dP
- 2f9dP.clusters - Cluster descriptions for 2f9dP
- 2f9dP.msf - the multiple sequence alignment used for the chain 2f9dP
- 2f9dP.descr - description of sequences used in 2f9dP msf
- 2f9dP.ranks_sorted - full listing of residues and their ranking for 2f9dP
- 2f9dP.2f9dA.if.pml - Pymol script for Figure 10
- 2f9dP.cbcbvg - used by other 2f9dP – related pymol scripts
- 2f9dP.2f9dB.if.pml - Pymol script for Figure 11