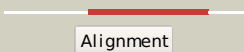
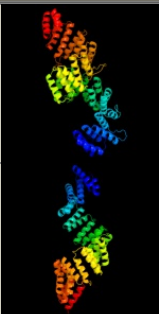
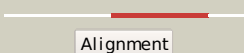



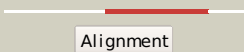

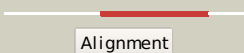














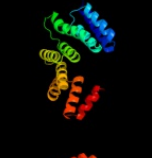
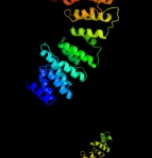





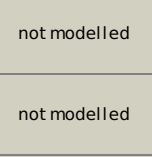


# Phyre2

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Date	Thu Jan 5 12:24:52 GMT 2012
Unique Job ID	3e24117744b4e7d2


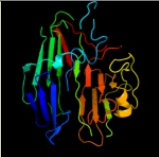
Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	<a href="#">c2xpiA_</a>	 Alignment		100.0	12	<b>PDB header:</b> cell cycle <b>Chain:</b> A: <b>PDB Molecule:</b> anaphase-promoting complex subunit cut9; <b>PDBTitle:</b> crystal structure of apc/c hetero-tetramer cut9-hcn1
2	<a href="#">d1w3ba_</a>	 Alignment		100.0	18	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
3	<a href="#">c3fp4A_</a>	 Alignment		100.0	11	<b>PDB header:</b> transport protein <b>Chain:</b> A: <b>PDB Molecule:</b> tpr repeat-containing protein yhr117w; <b>PDBTitle:</b> crystal structure of tom71 complexed with ssa1 c-terminal2 fragment
4	<a href="#">c2gw1A_</a>	 Alignment		100.0	10	<b>PDB header:</b> protein transport <b>Chain:</b> A: <b>PDB Molecule:</b> mitochondrial precursor proteins import receptor; <b>PDBTitle:</b> crystal structure of the yeast tom70
5	<a href="#">d2o0ea1</a>	 Alignment		100.0	8	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> HAT/Suf repeat
6	<a href="#">c2y4tA_</a>	 Alignment		99.9	18	<b>PDB header:</b> chaperone <b>Chain:</b> A: <b>PDB Molecule:</b> dnaj homolog subfamily c member 3; <b>PDBTitle:</b> crystal structure of the human co-chaperone p58(ipk)
7	<a href="#">c3iegB_</a>	 Alignment		99.9	17	<b>PDB header:</b> chaperone <b>Chain:</b> B: <b>PDB Molecule:</b> dnaj homolog subfamily c member 3; <b>PDBTitle:</b> crystal structure of p58(ipk) tpr domain at 2.5 a
8	<a href="#">c3hymB_</a>	 Alignment		99.9	9	<b>PDB header:</b> cell cycle, ligase <b>Chain:</b> B: <b>PDB Molecule:</b> cell division cycle protein 16 homolog; <b>PDBTitle:</b> insights into anaphase promoting complex tpr subdomain2 assembly from a cdc26-apc6 structure
9	<a href="#">d1qsaa1</a>	 Alignment		99.9	9	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> Bacterial muramidases <b>Family:</b> Bacterial muramidases
10	<a href="#">c1fchB_</a>	 Alignment		99.9	16	<b>PDB header:</b> signaling protein <b>Chain:</b> B: <b>PDB Molecule:</b> peroxisomal targeting signal 1 receptor; <b>PDBTitle:</b> crystal structure of the pts1 complexed to the tpr region2 of human pex5
11	<a href="#">d1fcha_</a>	 Alignment		99.8	14	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)

12	<a href="#">c2uy1A_</a>	Alignment		99.8	11	<b>PDB header:</b> rna-binding protein <b>Chain:</b> A: <b>PDB Molecule:</b> cleavage stimulation factor 77; <b>PDBTitle:</b> crystal structure of cstf-77
13	<a href="#">c3pe3D_</a>	Alignment		99.8	13	<b>PDB header:</b> transferase <b>Chain:</b> D: <b>PDB Molecule:</b> udp-n-acetylglucosamine--peptide n- <b>PDBTitle:</b> structure of human o-glcna transferase and its complex with a peptide2 substrate
14	<a href="#">d1dcea1</a>	Alignment		99.8	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> Protein prenyltransferase <b>Family:</b> Protein prenyltransferase
15	<a href="#">c1xi4D_</a>	Alignment		99.8	13	<b>PDB header:</b> endocytosis/exocytosis <b>Chain:</b> D: <b>PDB Molecule:</b> clathrin heavy chain; <b>PDBTitle:</b> clathrin d6 coat
16	<a href="#">c1tnol_</a>	Alignment		99.8	13	<b>PDB header:</b> transferase <b>Chain:</b> I: <b>PDB Molecule:</b> geranylgeranyltransferase type i alpha subunit; <b>PDBTitle:</b> rat protein geranylgeranyltransferase type-i complexed with2 a ggpp analog and a kkskktkvim peptide derived from k-3 ras4b
17	<a href="#">d2h6fa1</a>	Alignment		99.8	11	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> Protein prenyltransferase <b>Family:</b> Protein prenyltransferase
18	<a href="#">d1d8da_</a>	Alignment		99.8	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> Protein prenyltransferase <b>Family:</b> Protein prenyltransferase
19	<a href="#">d1hz4a_</a>	Alignment		99.8	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Transcription factor MalT domain III
20	<a href="#">c3cvpA_</a>	Alignment		99.8	11	<b>PDB header:</b> transport protein <b>Chain:</b> A: <b>PDB Molecule:</b> peroxisome targeting signal 1 receptor pex5; <b>PDBTitle:</b> structure of peroxisomal targeting signal 1 (pts1) binding2 domain of trypanosoma brucei peroxin 5 (tbpex5)complexed3 to pts1 peptide (10-skl)
21	<a href="#">c3q75A_</a>	Alignment	not modelled	99.8	15	<b>PDB header:</b> transferase <b>Chain:</b> A: <b>PDB Molecule:</b> farnesyltransferase alpha subunit; <b>PDBTitle:</b> cryptococcus neoformans protein farnesyltransferase in complex with2 fpt-ii and tkcvvm peptide
22	<a href="#">c2q7fA_</a>	Alignment	not modelled	99.8	23	<b>PDB header:</b> protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> yrrb protein; <b>PDBTitle:</b> crystal structure of yrrb: a tpr protein with an unusual peptide-2 binding site
23	<a href="#">c3draA_</a>	Alignment	not modelled	99.8	10	<b>PDB header:</b> transferase <b>Chain:</b> A: <b>PDB Molecule:</b> protein <b>PDBTitle:</b> candida albicans protein geranylgeranyltransferase-i2 complexed with ggpp
24	<a href="#">d1xnfa_</a>	Alignment	not modelled	99.7	15	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
25	<a href="#">c3mkrA_</a>	Alignment	not modelled	99.7	15	<b>PDB header:</b> transport protein <b>Chain:</b> A: <b>PDB Molecule:</b> coatomer subunit epsilon; <b>PDBTitle:</b> crystal structure of yeast alpha/epsilon-cop subcomplex of the cop12 vesicular coat
26	<a href="#">c2ho1B_</a>	Alignment	not modelled	99.7	16	<b>PDB header:</b> protein binding <b>Chain:</b> B: <b>PDB Molecule:</b> type 4 fimbrial biogenesis protein pilf; <b>PDBTitle:</b> functional characterization of pseudomonas aeruginosa pilf
27	<a href="#">c3q15A_</a>	Alignment	not modelled	99.7	10	<b>PDB header:</b> hydrolase/kinase <b>Chain:</b> A: <b>PDB Molecule:</b> response regulator aspartate phosphatase h; <b>PDBTitle:</b> crystal structure of raph complexed with spo0f
28	<a href="#">c3mv3B_</a>	Alignment	not modelled	99.7	11	<b>PDB header:</b> protein transport <b>Chain:</b> B: <b>PDB Molecule:</b> coatomer subunit epsilon; <b>PDBTitle:</b> crystal structure of a-cop in complex with e-cop
						<b>PDB header:</b> structural protein

29	<a href="#">c2vq2A_</a>	Alignment	not modelled	99.6	19	<b>Chain:</b> A: <b>PDB Molecule:</b> putative fimbrial biogenesis and twitching <b>PDBTitle:</b> crystal structure of pilw, widely conserved type iv pilus2 biogenesis factor
30	<a href="#">c3sf4B_</a>	Alignment	not modelled	99.6	13	<b>PDB header:</b> signaling protein/protein binding <b>Chain:</b> B: <b>PDB Molecule:</b> g-protein-signaling modulator 2; <b>PDBTitle:</b> crystal structure of the complex between the conserved cell polarity2 proteins inscuteable and lgn
31	<a href="#">c2r5sB_</a>	Alignment	not modelled	99.6	13	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> B: <b>PDB Molecule:</b> uncharacterized protein vp0806; <b>PDBTitle:</b> the crystal structure of a domain of protein vp0806 (unknown function)2 from vibrio parahaemolyticus rimd 2210633
32	<a href="#">c3as5A_</a>	Alignment	not modelled	99.6	13	<b>PDB header:</b> protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> mama; <b>PDBTitle:</b> mama amb-1 p212121
33	<a href="#">c3ly8A_</a>	Alignment	not modelled	99.6	10	<b>PDB header:</b> signaling protein <b>Chain:</b> A: <b>PDB Molecule:</b> transcriptional activator cadc; <b>PDBTitle:</b> crystal structure of mutant d471e of the periplasmic domain of cadc
34	<a href="#">c2hyzA_</a>	Alignment	not modelled	99.6	24	<b>PDB header:</b> de novo protein <b>Chain:</b> A: <b>PDB Molecule:</b> synthetic consensus tpr protein; <b>PDBTitle:</b> crystal structure of an 8 repeat consensus tpr superhelix2 (orthorhombic crystal form)
35	<a href="#">c2e2eA_</a>	Alignment	not modelled	99.6	13	<b>PDB header:</b> lyase <b>Chain:</b> A: <b>PDB Molecule:</b> formate-dependent nitrite reductase complex nrfg subunit; <b>PDBTitle:</b> tpr domain of nrfg mediates the complex formation between heme lyase2 and formate-dependent nitrite reductase in escherichia coli o157:h7
36	<a href="#">c1wao4_</a>	Alignment	not modelled	99.5	21	<b>PDB header:</b> hydrolase <b>Chain:</b> 4: <b>PDB Molecule:</b> serine/threonine protein phosphatase 5; <b>PDBTitle:</b> pp5 structure
37	<a href="#">d2c2la1</a>	Alignment	not modelled	99.5	14	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
38	<a href="#">d1hh8a_</a>	Alignment	not modelled	99.5	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
39	<a href="#">c3urzB_</a>	Alignment	not modelled	99.5	15	<b>PDB header:</b> protein binding <b>Chain:</b> B: <b>PDB Molecule:</b> uncharacterized protein; <b>PDBTitle:</b> crystal structure of a hypothetical protein (bacova_03105) from2 bacteroides ovatus atcc 8483 at 2.19 a resolution
40	<a href="#">c2vyiA_</a>	Alignment	not modelled	99.5	18	<b>PDB header:</b> chaperone <b>Chain:</b> A: <b>PDB Molecule:</b> sgta protein; <b>PDBTitle:</b> crystal structure of the tpr domain of human sgt
41	<a href="#">c1slyA_</a>	Alignment	not modelled	99.5	10	<b>PDB header:</b> glycosyltransferase <b>Chain:</b> A: <b>PDB Molecule:</b> 70-kda soluble lytic transglycosylase; <b>PDBTitle:</b> complex of the 70-kda soluble lytic transglycosylase with2 bulgecin a
42	<a href="#">c2pl2A_</a>	Alignment	not modelled	99.5	19	<b>PDB header:</b> protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> hypothetical conserved protein ttc0263; <b>PDBTitle:</b> crystal structure of ttc0263: a thermophilic tpr protein in thermus2 thermophilus hb27
43	<a href="#">c3ulqA_</a>	Alignment	not modelled	99.5	10	<b>PDB header:</b> gene regulation/transcription activator <b>Chain:</b> A: <b>PDB Molecule:</b> response regulator aspartate phosphatase f; <b>PDBTitle:</b> crystal structure of the anti-activator rapf complexed with the2 response regulator coma dna binding domain
44	<a href="#">c3gyzB_</a>	Alignment	not modelled	99.5	12	<b>PDB header:</b> chaperone <b>Chain:</b> B: <b>PDB Molecule:</b> chaperone protein ipgc; <b>PDBTitle:</b> crystal structure of ipgc from shigella flexneri
45	<a href="#">c3gw4B_</a>	Alignment	not modelled	99.5	14	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> B: <b>PDB Molecule:</b> uncharacterized protein; <b>PDBTitle:</b> crystal structure of uncharacterized protein from deinococcus2 radiodurans. northeast structural genomics consortium target drr162b.
46	<a href="#">c3qdnA_</a>	Alignment	not modelled	99.5	13	<b>PDB header:</b> oxidoreductase <b>Chain:</b> A: <b>PDB Molecule:</b> putative thioredoxin protein; <b>PDBTitle:</b> putative thioredoxin protein from salmonella typhimurium
47	<a href="#">c2xcbA_</a>	Alignment	not modelled	99.5	11	<b>PDB header:</b> protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> regulatory protein pcrh; <b>PDBTitle:</b> crystal structure of pcrh in complex with the chaperone2 binding region of popd
48	<a href="#">d1a17a_</a>	Alignment	not modelled	99.4	21	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
49	<a href="#">c3sz7A_</a>	Alignment	not modelled	99.4	16	<b>PDB header:</b> chaperone regulator <b>Chain:</b> A: <b>PDB Molecule:</b> hsc70 cochaperone (sgt); <b>PDBTitle:</b> crystal structure of the sgt2 tpr domain from aspergillus fumigatus
50	<a href="#">c2c2ID_</a>	Alignment	not modelled	99.4	14	<b>PDB header:</b> chaperone <b>Chain:</b> D: <b>PDB Molecule:</b> carboxy terminus of hsp70-interacting protein; <b>PDBTitle:</b> crystal structure of the chip u-box e3 ubiquitin ligase
51	<a href="#">d1elwa_</a>	Alignment	not modelled	99.4	17	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
52	<a href="#">c1ihgA_</a>	Alignment	not modelled	99.4	13	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> cyclophilin 40; <b>PDBTitle:</b> bovine cyclophilin 40, monoclinic form
53	<a href="#">d1kt1a1</a>	Alignment	not modelled	99.4	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)

54	<a href="#">c3nreB_</a>	Alignment		99.4	14	<b>PDB header:</b> isomerase <b>Chain:</b> B: <b>PDB Molecule:</b> aldose 1-epimerase; <b>PDBTitle:</b> crystal structure of a putative aldose 1-epimerase (b2544) from2 escherichia coli k12 at 1.59 a resolution
55	<a href="#">c2vsnB_</a>	Alignment	not modelled	99.4	16	<b>PDB header:</b> transferase <b>Chain:</b> B: <b>PDB Molecule:</b> xcogt; <b>PDBTitle:</b> structure and topological arrangement of an o-glcna2 transferase homolog: insight into molecular control of3 intracellular glycosylation
56	<a href="#">dlihga1</a>	Alignment	not modelled	99.4	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
57	<a href="#">c2ifuA_</a>	Alignment	not modelled	99.3	13	<b>PDB header:</b> endocytosis/exocytosis <b>Chain:</b> A: <b>PDB Molecule:</b> gamma-snap; <b>PDBTitle:</b> crystal structure of a gamma-snap from danio rerio
58	<a href="#">d1p5qa1</a>	Alignment	not modelled	99.3	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
59	<a href="#">c2dbaA_</a>	Alignment	not modelled	99.3	16	<b>PDB header:</b> structural protein <b>Chain:</b> A: <b>PDB Molecule:</b> smooth muscle cell associated protein-1, isoform <b>PDBTitle:</b> the solution structure of the tetratricopeptide repeat of2 human smooth muscle cell associated protein-1, isoform 2
60	<a href="#">c3rkva_</a>	Alignment	not modelled	99.3	13	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> putative peptidylprolyl isomerase; <b>PDBTitle:</b> c-terminal domain of protein c56c10.10, a putative peptidylprolyl2 isomerase, from caenorhabditis elegans
61	<a href="#">d2buga1</a>	Alignment	not modelled	99.3	22	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
62	<a href="#">d2onda1</a>	Alignment	not modelled	99.3	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> HAT/Suf repeat
63	<a href="#">c1kt0A_</a>	Alignment	not modelled	99.3	12	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> 51 kda fk506-binding protein; <b>PDBTitle:</b> structure of the large fkbp-like protein, fkbp51, involved in steroid2 receptor complexes
64	<a href="#">c1qz2B_</a>	Alignment	not modelled	99.3	11	<b>PDB header:</b> isomerase/chaperone <b>Chain:</b> B: <b>PDB Molecule:</b> fk506-binding protein 4; <b>PDBTitle:</b> crystal structure of fkbp52 c-terminal domain complex with2 the c-terminal peptide meevd of hsp90
65	<a href="#">d1zu2a1</a>	Alignment	not modelled	99.3	20	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
66	<a href="#">d2fba1</a>	Alignment	not modelled	99.3	11	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
67	<a href="#">c2fbaA_</a>	Alignment	not modelled	99.3	11	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> 70 kda peptidylprolyl isomerase, putative; <b>PDBTitle:</b> plasmodium falciparum putative fk506-binding protein2 pfl2275c, c-terminal tpr-containing domain
68	<a href="#">c3q49B_</a>	Alignment	not modelled	99.3	16	<b>PDB header:</b> ligase/chaperone <b>Chain:</b> B: <b>PDB Molecule:</b> stip1 homology and u box-containing protein 1; <b>PDBTitle:</b> crystal structure of the tpr domain of chip complexed with hsp70-c2 peptide
69	<a href="#">c2kckA_</a>	Alignment	not modelled	99.3	10	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> tpr repeat; <b>PDBTitle:</b> nmr solution structure of the northeast structural genomics2 consortium (nesg) target mrr121a
70	<a href="#">c3mwxA_</a>	Alignment		99.3	18	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> aldose 1-epimerase; <b>PDBTitle:</b> crystal structure of a putative galactose mutarotase (bsu18360) from2 bacillus subtilis at 1.45 a resolution
71	<a href="#">d1hxia_</a>	Alignment	not modelled	99.3	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
72	<a href="#">c1ltxA_</a>	Alignment	not modelled	99.2	11	<b>PDB header:</b> transferase/protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> rab geranylgeranyltransferase alpha subunit; <b>PDBTitle:</b> structure of rab escort protein-1 in complex with rab2 geranylgeranyl transferase and isoprenoid
73	<a href="#">c3os7B_</a>	Alignment		99.2	13	<b>PDB header:</b> isomerase <b>Chain:</b> B: <b>PDB Molecule:</b> galactose mutarotase-like protein; <b>PDBTitle:</b> crystal structure of a galactose mutarotase-like protein (ca_c0697)2 from clostridium acetobutylicum at 1.80 a resolution
74	<a href="#">c2yhca_</a>	Alignment	not modelled	99.2	19	<b>PDB header:</b> membrane protein <b>Chain:</b> A: <b>PDB Molecule:</b> upf0169 lipoprotein yfio; <b>PDBTitle:</b> structure of bamd from e. coli
75	<a href="#">c3nf1A_</a>	Alignment	not modelled	99.2	14	<b>PDB header:</b> motor protein, transport protein <b>Chain:</b> A: <b>PDB Molecule:</b> kinesin light chain 1; <b>PDBTitle:</b> crystal structure of the tpr domain of kinesin light chain 1
						<b>PDB header:</b> chaperone

76	<a href="#">c2vgxA</a>	Alignment	not modelled	99.2	9	<b>Chain:</b> A: <b>PDB Molecule:</b> chaperone sycd; <b>PDBTitle:</b> structure of the yersinia enterocolitica type iii secretion2 translocator chaperone sycd
77	<a href="#">c3os7D</a>	Alignment		99.2	15	<b>PDB header:</b> isomerase <b>Chain:</b> D: <b>PDB Molecule:</b> galactose mutarotase-like protein; <b>PDBTitle:</b> crystal structure of a galactose mutarotase-like protein (ca_c0697)2 from clostridium acetobutylicum at 1.80 a resolution
78	<a href="#">dliygA</a>	Alignment	not modelled	99.2	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
79	<a href="#">c3k9iA</a>	Alignment	not modelled	99.2	17	<b>PDB header:</b> protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> bh0479 protein; <b>PDBTitle:</b> crystal structure of putative protein binding protein (np_241345.1)2 from bacillus halodurans at 2.71 a resolution
80	<a href="#">c2katA</a>	Alignment	not modelled	99.2	13	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> uncharacterized protein; <b>PDBTitle:</b> solution structure of protein bpp2914 from bordetella2 parapertussis. northeast structural genomics consortium3 target bpr206
81	<a href="#">c3q1nA</a>	Alignment		99.1	17	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> galactose mutarotase related enzyme; <b>PDBTitle:</b> crystal structure of a galactose mutarotase-like protein (lsei_2598)2 from lactobacillus casei atcc 334 at 1.61 a resolution
82	<a href="#">d1elra</a>	Alignment	not modelled	99.1	18	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
83	<a href="#">c2xeVB</a>	Alignment	not modelled	99.1	17	<b>PDB header:</b> metal binding <b>Chain:</b> B: <b>PDB Molecule:</b> ybgf; <b>PDBTitle:</b> crystal structure of the tpr domain of xanthomonas2 campestris ybgf
84	<a href="#">c3cegB</a>	Alignment	not modelled	99.1	17	<b>PDB header:</b> motor protein, transport protein <b>Chain:</b> B: <b>PDB Molecule:</b> kinesin light chain 2; <b>PDBTitle:</b> the tpr domain of human kinesin light chain 2 (hklc2)
85	<a href="#">d1nzna</a>	Alignment	not modelled	99.1	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
86	<a href="#">d1qqea</a>	Alignment	not modelled	99.1	9	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
87	<a href="#">c3qkYA</a>	Alignment	not modelled	99.1	17	<b>PDB header:</b> membrane protein <b>Chain:</b> A: <b>PDB Molecule:</b> outer membrane assembly lipoprotein yfio; <b>PDBTitle:</b> crystal structure of rhodothermus marinus bamd
88	<a href="#">c1na3A</a>	Alignment	not modelled	99.1	24	<b>PDB header:</b> de novo protein <b>Chain:</b> A: <b>PDB Molecule:</b> designed protein ctrp2; <b>PDBTitle:</b> design of stable alpha-helical arrays from an idealized tpr2 motif
89	<a href="#">d1pc2a</a>	Alignment	not modelled	99.1	7	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
90	<a href="#">d2hr2a1</a>	Alignment	not modelled	99.0	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> CT2138-like
91	<a href="#">c2if4A</a>	Alignment	not modelled	99.0	10	<b>PDB header:</b> signaling protein <b>Chain:</b> A: <b>PDB Molecule:</b> atfkbp42; <b>PDBTitle:</b> crystal structure of a multi-domain immunophilin from2 arabidopsis thaliana
92	<a href="#">c2xm6A</a>	Alignment	not modelled	99.0	10	<b>PDB header:</b> unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> protein corresponding to locus c5321 from cft073 e.coli <b>PDBTitle:</b> crystal structure of the protein corresponding to locus c5321 from2 cft073 e.coli strain
93	<a href="#">c2kcvA</a>	Alignment	not modelled	99.0	19	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> tetratricopeptide repeat domain protein; <b>PDBTitle:</b> solution nmr structure of tetratricopeptide repeat domain2 protein sru_0103 from salinibacter ruber, northeast3 structural genomics consortium (nesg) target srr115c
94	<a href="#">c3k25B</a>	Alignment	not modelled	99.0	15	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> B: <b>PDB Molecule:</b> slr1438 protein; <b>PDBTitle:</b> crystal structure of slr1438 protein from synechocystis sp. pcc 6803,2 northeast structural genomics consortium target sgr112
95	<a href="#">c3fflC</a>	Alignment	not modelled	99.0	12	<b>PDB header:</b> cell cycle <b>Chain:</b> C: <b>PDB Molecule:</b> anaphase-promoting complex subunit 7; <b>PDBTitle:</b> crystal structure of the n-terminal domain of anaphase-2 promoting complex subunit 7
96	<a href="#">d1zbpa1</a>	Alignment	not modelled	98.9	11	<b>Fold:</b> ImpE-like <b>Superfamily:</b> ImpE-like <b>Family:</b> ImpE-like
97	<a href="#">c3dcdA</a>	Alignment	not modelled	98.9	17	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> galactose mutarotase related enzyme; <b>PDBTitle:</b> x-ray structure of the galactose mutarotase related enzyme q5fkd7 from2 lactobacillus acidophilus at the resolution 1.9a. northeast3 structural genomics consortium target iar33.
98	<a href="#">c3ty1B</a>	Alignment	not modelled	98.9	14	<b>PDB header:</b> isomerase <b>Chain:</b> B: <b>PDB Molecule:</b> hypothetical aldose 1-epimerase; <b>PDBTitle:</b> crystal structure of a hypothetical aldose 1-epimerase (kpn_04629)2 from klebsiella pneumoniae subsp. pneumoniae mgh 78578 at 1.90 a3 resolution <b>PDB header:</b> protein binding



99	<a href="#">c2l6jA</a>	Alignment	not modelled	98.8	15	<b>Chain:</b> A: <b>PDB Molecule:</b> tpr repeat-containing protein associated with hsp90; <b>PDBTitle:</b> tah1 complexed by meevd
100	<a href="#">c3n71A</a>	Alignment	not modelled	98.8	9	<b>PDB header:</b> transcription <b>Chain:</b> A: <b>PDB Molecule:</b> histone lysine methyltransferase smyd1; <b>PDBTitle:</b> crystal structure of cardiac specific histone methyltransferase smyd1
101	<a href="#">c2avpA</a>	Alignment	not modelled	98.8	25	<b>PDB header:</b> de novo protein <b>Chain:</b> A: <b>PDB Molecule:</b> synthetic consensus tpr protein; <b>PDBTitle:</b> crystal structure of an 8 repeat consensus tpr superhelix
102	<a href="#">d1tjca</a>	Alignment	not modelled	98.8	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
103	<a href="#">d1y8ma1</a>	Alignment	not modelled	98.7	10	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
104	<a href="#">d1z45a1</a>	Alignment	not modelled	98.6	12	<b>Fold:</b> Supersandwich <b>Superfamily:</b> Galactose mutarotase-like <b>Family:</b> Aldose 1-epimerase (mutarotase)
105	<a href="#">c2kc7A</a>	Alignment	not modelled	98.5	14	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> A: <b>PDB Molecule:</b> bfr218_protein; <b>PDBTitle:</b> solution nmr structure of bacteroides fragilis protein2 bf1650. northeast structural genomics consortium target3 bfr218
106	<a href="#">d2ff4a2</a>	Alignment	not modelled	98.5	13	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> BTAD-like
107	<a href="#">d1ouva</a>	Alignment	not modelled	98.5	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> HCP-like <b>Family:</b> HCP-like
108	<a href="#">c2qfcB</a>	Alignment	not modelled	98.4	8	<b>PDB header:</b> transcription regulation <b>Chain:</b> B: <b>PDB Molecule:</b> plcr protein; <b>PDBTitle:</b> crystal structure of bacillus thuringiensis plcr complexed with papr
109	<a href="#">c1ygaA</a>	Alignment	not modelled	98.4	12	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> hypothetical 37.9 kda protein in bio3-hxt17 <b>PDBTitle:</b> crystal structure of saccharomyces cerevisiae yn9a protein,2 new york structural genomics consortium
110	<a href="#">d2pqrbl</a>	Alignment	not modelled	98.3	12	<b>Fold:</b> alpha-alpha superhelix <b>Superfamily:</b> TPR-like <b>Family:</b> Tetratricopeptide repeat (TPR)
111	<a href="#">c3beeB</a>	Alignment	not modelled	98.3	10	<b>PDB header:</b> structural genomics, unknown function <b>Chain:</b> B: <b>PDB Molecule:</b> putative yfre protein; <b>PDBTitle:</b> crystal structure of putative yfre protein from vibrio2 parahaemolyticus
112	<a href="#">c3qwvA</a>	Alignment	not modelled	98.3	10	<b>PDB header:</b> transferase <b>Chain:</b> A: <b>PDB Molecule:</b> set and mynd domain-containing protein 2; <b>PDBTitle:</b> crystal structure of histone lysine methyltransferase smyd2 in complex2 with the cofactor product adohcy
113	<a href="#">c3e4bD</a>	Alignment	not modelled	98.3	10	<b>PDB header:</b> protein binding <b>Chain:</b> D: <b>PDB Molecule:</b> algk; <b>PDBTitle:</b> crystal structure of algk from pseudomonas fluorescens wcs374r
114	<a href="#">c3q3hA</a>	Alignment	not modelled	98.3	6	<b>PDB header:</b> transferase <b>Chain:</b> A: <b>PDB Molecule:</b> hmw1c-like glycosyltransferase; <b>PDBTitle:</b> crystal structure of the actinobacillus pleuropneumoniae hmw1c2 glycosyltransferase in complex with udp-glc
115	<a href="#">d1nsza</a>	Alignment	not modelled	98.3	12	<b>Fold:</b> Supersandwich <b>Superfamily:</b> Galactose mutarotase-like <b>Family:</b> Aldose 1-epimerase (mutarotase)
116	<a href="#">d1jova</a>	Alignment	not modelled	98.2	8	<b>Fold:</b> Supersandwich <b>Superfamily:</b> Galactose mutarotase-like <b>Family:</b> Hypothetical protein HI1317
117	<a href="#">c2cisA</a>	Alignment	not modelled	98.2	17	<b>PDB header:</b> isomerase <b>Chain:</b> A: <b>PDB Molecule:</b> hexose-6-phosphate mutarotase; <b>PDBTitle:</b> structure-based functional annotation: yeast ymr099c codes2 for a d-hexose-6-phosphate mutarotase. complex with3 tagatose-6-phosphate
118	<a href="#">c3mekA</a>	Alignment	not modelled	98.2	13	<b>PDB header:</b> transferase <b>Chain:</b> A: <b>PDB Molecule:</b> set and mynd domain-containing protein 3; <b>PDBTitle:</b> crystal structure of human histone-lysine n-2 methyltransferase smyd3 in complex with s-adenosyl-l-3 methionine
119	<a href="#">c3bxaA</a>	Alignment	not modelled	98.2	12	<b>PDB header:</b> hydrolase, protein binding <b>Chain:</b> A: <b>PDB Molecule:</b> 26s proteasome regulatory complex subunit p42b; <b>PDBTitle:</b> crystal structure of rpn6 from drosophila melanogaster, gd(3+) complex
120	<a href="#">c3imhB</a>	Alignment	not modelled	98.1	15	<b>PDB header:</b> isomerase <b>Chain:</b> B: <b>PDB Molecule:</b> galactose-1-epimerase; <b>PDBTitle:</b> crystal structure of galactose 1-epimerase from lactobacillus2 acidophilus ncfm