






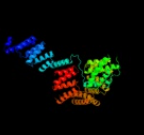














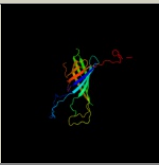





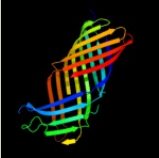
#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c2xpiA_	 Alignment		100.0	16	PDB header: cell cycle Chain: A: PDB Molecule: anaphase-promoting complex subunit cut9; PDBTitle: crystal structure of apc/c hetero-tetramer cut9-hcn1
2	dlw3ba_	 Alignment		100.0	17	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
3	d2ooea1	 Alignment		99.9	9	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: HAT/Suf repeat
4	c3fp4A_	 Alignment		99.8	13	PDB header: transport protein Chain: A: PDB Molecule: tpr repeat-containing protein yhr117w; PDBTitle: crystal structure of tom71 complexed with ssa1 c-terminal2 fragment
5	c2gw1A_	 Alignment		99.8	11	PDB header: protein transport Chain: A: PDB Molecule: mitochondrial precursor proteins import receptor; PDBTitle: crystal structure of the yeast tom70
6	c2y4tA_	 Alignment		99.8	17	PDB header: chaperone Chain: A: PDB Molecule: dnaj homolog subfamily c member 3; PDBTitle: crystal structure of the human co-chaperone p58(ipk)
7	c3iegB_	 Alignment		99.8	19	PDB header: chaperone Chain: B: PDB Molecule: dnaj homolog subfamily c member 3; PDBTitle: crystal structure of p58(ipk) tpr domain at 2.5 a
8	dlqsaal	 Alignment		99.7	13	Fold: alpha-alpha superhelix Superfamily: Bacterial muramidases Family: Bacterial muramidases
9	clfchB_	 Alignment		99.7	12	PDB header: signaling protein Chain: B: PDB Molecule: peroxisomal targeting signal 1 receptor; PDBTitle: crystal structure of the pts1 complexed to the tpr region2 of human pex5
10	c3pe3D_	 Alignment		99.7	14	PDB header: transferase Chain: D: PDB Molecule: udp-n-acetylglucosamine--peptide n- PDBTitle: structure of human o-glcna transferase and its complex with a peptide2 substrate
11	dlfcha_	 Alignment		99.7	11	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)

12	c2ho1B_	Alignment		99.7	13	PDB header: protein binding Chain: B: PDB Molecule: type 4 fimbrial biogenesis protein pilf; PDBTitle: functional characterization of pseudomonas aeruginosa pilf
13	c3hymB_	Alignment		99.7	12	PDB header: cell cycle, ligase Chain: B: PDB Molecule: cell division cycle protein 16 homolog; PDBTitle: insights into anaphase promoting complex tpr subdomain2 assembly from a cdc26-apc6 structure
14	c3cvpA_	Alignment		99.7	13	PDB header: transport protein Chain: A: PDB Molecule: peroxisome targeting signal 1 receptor pex5; PDBTitle: structure of peroxisomal targeting signal 1 (pts1) binding2 domain of trypanosoma brucei peroxin 5 (tbpex5)complexed3 to pts1 peptide (10-skl)
15	c2uy1A_	Alignment		99.6	12	PDB header: rna-binding protein Chain: A: PDB Molecule: cleavage stimulation factor 77; PDBTitle: crystal structure of cstf-77
16	c2vq2A_	Alignment		99.6	14	PDB header: structural protein Chain: A: PDB Molecule: putative fimbrial biogenesis and twitching PDBTitle: crystal structure of pilw, widely conserved type iv pilus2 biogenesis factor
17	d1hz4a_	Alignment		99.6	17	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Transcription factor MalT domain III
18	c3draA_	Alignment		99.6	11	PDB header: transferase Chain: A: PDB Molecule: protein PDBTitle: candida albicans protein geranylgeranyltransferase-i2 complexed with ggpp
19	c1xi4D_	Alignment		99.5	12	PDB header: endocytosis/exocytosis Chain: D: PDB Molecule: clathrin heavy chain; PDBTitle: clathrin d6 coat
20	c2q7fA_	Alignment		99.5	13	PDB header: protein binding Chain: A: PDB Molecule: yrrb protein; PDBTitle: crystal structure of yrrb: a tpr protein with an unusual peptide-2 binding site
21	d2h6fa1	Alignment	not modelled	99.5	11	Fold: alpha-alpha superhelix Superfamily: Protein prenyltransferase Family: Protein prenyltransferase
22	d1d8da_	Alignment	not modelled	99.5	11	Fold: alpha-alpha superhelix Superfamily: Protein prenyltransferase Family: Protein prenyltransferase
23	c3q75A_	Alignment	not modelled	99.5	11	PDB header: transferase Chain: A: PDB Molecule: farnesyltransferase alpha subunit; PDBTitle: cryptococcus neoformans protein farnesyltransferase in complex with2 fpt-ii and tkcvvm peptide
24	d1dcea1	Alignment	not modelled	99.5	10	Fold: alpha-alpha superhelix Superfamily: Protein prenyltransferase Family: Protein prenyltransferase
25	d1xnfa_	Alignment	not modelled	99.5	13	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
26	c3mkrA_	Alignment	not modelled	99.5	14	PDB header: transport protein Chain: A: PDB Molecule: coatmer subunit epsilon; PDBTitle: crystal structure of yeast alpha/epsilon-cop subcomplex of the cop12 vesicular coat
27	c1tnol_	Alignment	not modelled	99.5	11	PDB header: transferase Chain: I: PDB Molecule: geranylgeranyltransferase type i alpha subunit; PDBTitle: rat protein geranylgeranyltransferase type-i complexed with2 a ggpp analog and a kkskktkcvim peptide derived from k-3 ras4b
28	c3urzB_	Alignment	not modelled	99.5	11	PDB header: protein binding Chain: B: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a hypothetical protein (bacova 03105) from2 bacteroides ovatus atcc 8483 at 2.19 a

						resolution
29	c2r5sB_	Alignment	not modelled	99.4	16	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: uncharacterized protein vp0806; PDBTitle: the crystal structure of a domain of protein vp0806 (unknown function)2 from vibrio parahaemolyticus rimd 2210633
30	c3mv3B_	Alignment	not modelled	99.4	15	PDB header: protein transport Chain: B: PDB Molecule: coatomer subunit epsilon; PDBTitle: crystal structure of a-cop in complex with e-cop
31	c2pl2A_	Alignment	not modelled	99.4	18	PDB header: protein binding Chain: A: PDB Molecule: hypothetical conserved protein ttc0263; PDBTitle: crystal structure of ttc0263: a thermophilic tpr protein in thermus2 thermophilus hb27
32	c3as5A_	Alignment	not modelled	99.4	15	PDB header: protein binding Chain: A: PDB Molecule: mama; PDBTitle: mama amb-1 p212121
33	c1slvA_	Alignment	not modelled	99.3	13	PDB header: glycosyltransferase Chain: A: PDB Molecule: 70-kda soluble lytic transglycosylase; PDBTitle: complex of the 70-kda soluble lytic transglycosylase with2 bulgecin a
34	c3qdnA_	Alignment	not modelled	99.3	9	PDB header: oxidoreductase Chain: A: PDB Molecule: putative thioredoxin protein; PDBTitle: putative thioredoxin protein from salmonella typhimurium
35	d2onda1	Alignment	not modelled	99.3	9	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: HAT/Suf repeat
36	c2e2eA_	Alignment	not modelled	99.1	17	PDB header: lyase Chain: A: PDB Molecule: formate-dependent nitrite reductase complex nrfg subunit; PDBTitle: tpr domain of nrfg mediates the complex formation between heme lyase2 and formate-dependent nitrite reductase in escherichia coli o157:h7
37	c2hyzA_	Alignment	not modelled	99.1	22	PDB header: de novo protein Chain: A: PDB Molecule: synthetic consensus tpr protein; PDBTitle: crystal structure of an 8 repeat consensus tpr superhelix2 (orthorhombic crystal form)
38	c3ulqA_	Alignment	not modelled	99.1	10	PDB header: gene regulation/transcription activator Chain: A: PDB Molecule: response regulator aspartate phosphatase f; PDBTitle: crystal structure of the anti-activator rapf complexed with the2 response regulator coma dna binding domain
39	c3q15A_	Alignment	not modelled	99.1	9	PDB header: hydrolase/kinase Chain: A: PDB Molecule: response regulator aspartate phosphatase h; PDBTitle: crystal structure of raph complexed with spo0f
40	c3ly8A_	Alignment	not modelled	99.1	13	PDB header: signaling protein Chain: A: PDB Molecule: transcriptional activator cadc; PDBTitle: crystal structure of mutant d471e of the periplasmic domain of cadc
41	c2yhCA_	Alignment	not modelled	99.1	11	PDB header: membrane protein Chain: A: PDB Molecule: upf0169 lipoprotein yfio; PDBTitle: structure of bamd from e. coli
42	d1hh8a_	Alignment	not modelled	99.1	12	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
43	c3gw4B_	Alignment	not modelled	99.0	19	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of uncharacterized protein from deinococcus2 radiodurans. northeast structural genomics consortium target drr162b.
44	c2xcBA_	Alignment	not modelled	99.0	18	PDB header: protein binding Chain: A: PDB Molecule: regulatory protein pcrh; PDBTitle: crystal structure of pcrh in complex with the chaperone2 binding region of popd
45	c1wao4_	Alignment	not modelled	99.0	16	PDB header: hydrolase Chain: 4: PDB Molecule: serine/threonine protein phosphatase 5; PDBTitle: pp5 structure
46	c3sf4B_	Alignment	not modelled	99.0	13	PDB header: signaling protein/protein binding Chain: B: PDB Molecule: g-protein-signaling modulator 2; PDBTitle: crystal structure of the complex between the conserved cell polarity2 proteins inscuteable and lgn
47	c3gyzB_	Alignment	not modelled	99.0	13	PDB header: chaperone Chain: B: PDB Molecule: chaperone protein ipgc; PDBTitle: crystal structure of ipgc from shigella flexneri
48	d2c2la1	Alignment	not modelled	99.0	16	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
49	c3qkyA_	Alignment	not modelled	99.0	13	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane assembly lipoprotein yfio; PDBTitle: crystal structure of rhodothermus marinus bamd
50	c2c2lD_	Alignment	not modelled	99.0	18	PDB header: chaperone Chain: D: PDB Molecule: carboxy terminus of hsp70-interacting protein; PDBTitle: crystal structure of the chip u-box e3 ubiquitin ligase
51	c2vsnB_	Alignment	not modelled	98.9	13	PDB header: transferase Chain: B: PDB Molecule: xcogt; PDBTitle: structure and topological arrangement of an o-glcnac2 transferase homolog: insight into molecular control of3 intracellular glycosylation
52	d1elwa_	Alignment	not modelled	98.9	17	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
53	d1hxia_	Alignment	not modelled	98.9	14	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)

54	c2ifuA	Alignment	not modelled	98.9	14	PDB header: endocytosis/exocytosis Chain: A: PDB Molecule: gamma-snap; PDBTitle: crystal structure of a gamma-snap from danio rerio
55	d1a17a	Alignment	not modelled	98.9	15	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
56	c2vviA	Alignment	not modelled	98.9	17	PDB header: chaperone Chain: A: PDB Molecule: sgta protein; PDBTitle: crystal structure of the tpr domain of human sgt
57	d1zbpa1	Alignment	not modelled	98.8	12	Fold: ImpE-like Superfamily: ImpE-like Family: ImpE-like
58	c3q49B	Alignment	not modelled	98.8	18	PDB header: ligase/chaperone Chain: B: PDB Molecule: stip1 homology and u box-containing protein 1; PDBTitle: crystal structure of the tpr domain of chip complexed with hsp70-c2 peptide
59	d1zu2a1	Alignment	not modelled	98.8	15	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
60	c3sz7A	Alignment	not modelled	98.8	19	PDB header: chaperone regulator Chain: A: PDB Molecule: hsc70 cochaperone (sgt); PDBTitle: crystal structure of the sg2 tpr domain from aspergillus fumigatus
61	c2kckA	Alignment	not modelled	98.8	8	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: tpr repeat; PDBTitle: nmr solution structure of the northeast structural genomics2 consortium (nesg) target mrr121a
62	c2xevB	Alignment	not modelled	98.7	14	PDB header: metal binding Chain: B: PDB Molecule: ybgf; PDBTitle: crystal structure of the tpr domain of xanthomonas2 campestris ybgf
63	c2katA	Alignment	not modelled	98.7	15	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: solution structure of protein bpp2914 from bordetella2 parapertussis. northeast structural genomics consortium3 target bpr206
64	d2buga1	Alignment	not modelled	98.7	16	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
65	c2dbaA	Alignment	not modelled	98.7	15	PDB header: structural protein Chain: A: PDB Molecule: smooth muscle cell associated protein-1, isoform PDBTitle: the solution structure of the tetratrico peptide repeat of2 human smooth muscle cell associated protein-1, isoform 2
66	d1qqea	Alignment	not modelled	98.7	10	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
67	c1lbaA	Alignment	not modelled	98.6	9	PDB header: transferase/protein binding Chain: A: PDB Molecule: rab geranylgeranyltransferase alpha subunit; PDBTitle: structure of rab escort protein-1 in complex with rab2 geranylgeranyl transferase and isoprenoid
68	c1na3A	Alignment	not modelled	98.6	24	PDB header: de novo protein Chain: A: PDB Molecule: designed protein ctr2; PDBTitle: design of stable alpha-helical arrays from an idealized tpr2 motif
69	d1kt1a1	Alignment	not modelled	98.6	16	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
70	c2vgxA	Alignment	not modelled	98.6	11	PDB header: chaperone Chain: A: PDB Molecule: chaperone sycd; PDBTitle: structure of the yersinia enterocolitica type iii secretion2 translocator chaperone sycd
71	d1ihga1	Alignment	not modelled	98.5	14	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
72	d1p5qa1	Alignment	not modelled	98.5	15	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
73	d1elra	Alignment	not modelled	98.5	12	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
74	c3k9iA	Alignment	not modelled	98.5	17	PDB header: protein binding Chain: A: PDB Molecule: bh0479 protein; PDBTitle: crystal structure of putative protein binding protein (np_241345.1)2 from bacillus halodurans at 2.71 a resolution
75	c3rkva	Alignment	not modelled	98.5	18	PDB header: isomerase Chain: A: PDB Molecule: putative peptidylprolyl isomerase; PDBTitle: c-terminal domain of protein c56c10.10, a putative peptidylprolyl2 isomerase, from caenorhabditis elegans
76	c1ihgA	Alignment	not modelled	98.5	13	PDB header: isomerase Chain: A: PDB Molecule: cyclophilin 40; PDBTitle: bovine cyclophilin 40, monoclinic form
77	d2fba1	Alignment	not modelled	98.5	10	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
78	c2fbaA	Alignment	not modelled	98.5	10	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: 70 kda peptidylprolyl isomerase, putative; PDBTitle: plasmodium falciparum putative fk506-binding protein2 pfl2275c, c-terminal tpr-containing domain
79	c2k7A	Alignment	not modelled	98.5	16	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: bfr218 protein;

79	c2k07A	Alignment	not modelled	98.3	10	PDBTitle: solution nmr structure of bacteroides fragilis protein2 bf1650. northeast structural genomics consortium target3 bfr218 PDB header: isomerase
80	c1kt0A	Alignment	not modelled	98.5	17	Chain: A: PDB Molecule: 51 kda fk506-binding protein; PDBTitle: structure of the large fkbp-like protein, fkbp51, involved in steroid2 receptor complexes
81	d2hr2a1	Alignment	not modelled	98.4	20	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: CT2138-like
82	c2kcvA	Alignment	not modelled	98.4	11	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: tetratricopeptide repeat domain protein; PDBTitle: solution nmr structure of tetratricopeptide repeat domain2 protein sru_0103 from salinibacter ruber, northeast3 structural genomics consortium (nesg) target srr115c
83	d1iyga	Alignment	not modelled	98.4	7	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
84	c2k0lA	Alignment		98.4	11	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane protein a; PDBTitle: nmr structure of the transmembrane domain of the outer2 membrane protein a from klebsiella pneumoniae in dhpc3 micelles.
85	c1qz2B	Alignment	not modelled	98.4	17	PDB header: isomerase/chaperone Chain: B: PDB Molecule: fk506-binding protein 4; PDBTitle: crystal structure of fkbp52 c-terminal domain complex with2 the c-terminal peptide meevd of hsp90
86	d1nzna	Alignment	not modelled	98.4	9	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
87	c2l6jA	Alignment	not modelled	98.4	16	PDB header: protein binding Chain: A: PDB Molecule: tpr repeat-containing protein associated with hsp90; PDBTitle: tah1 complexed by meevd
88	c2avpA	Alignment	not modelled	98.4	16	PDB header: de novo protein Chain: A: PDB Molecule: synthetic consensus tpr protein; PDBTitle: crystal structure of an 8 repeat consensus tpr superhelix
89	d1pc2a	Alignment	not modelled	98.3	13	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
90	c3ceqB	Alignment	not modelled	98.3	12	PDB header: motor protein, transport protein Chain: B: PDB Molecule: kinesin light chain 2; PDBTitle: the tpr domain of human kinesin light chain 2 (hklc2)
91	c3nf1A	Alignment	not modelled	98.3	10	PDB header: motor protein, transport protein Chain: A: PDB Molecule: kinesin light chain 1; PDBTitle: crystal structure of the tpr domain of kinesin light chain 1
92	c3beeB	Alignment	not modelled	98.3	11	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: putative yfpe protein; PDBTitle: crystal structure of putative yfpe protein from vibrio2 parahaemolyticus
93	c2if4A	Alignment	not modelled	98.3	13	PDB header: signaling protein Chain: A: PDB Molecule: atfkbp42; PDBTitle: crystal structure of a multi-domain immunophilin from2 arabidopsis thaliana
94	d1tjca	Alignment	not modelled	98.2	14	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
95	d2ff4a2	Alignment	not modelled	98.2	14	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: BTAD-like
96	c3n71A	Alignment	not modelled	98.1	12	PDB header: transcription Chain: A: PDB Molecule: histone lysine methyltransferase smyd1; PDBTitle: crystal structure of cardiac specific histone methyltransferase smyd1
97	c3fflC	Alignment	not modelled	98.0	20	PDB header: cell cycle Chain: C: PDB Molecule: anaphase-promoting complex subunit 7; PDBTitle: crystal structure of the n-terminal domain of anaphase-2 promoting complex subunit 7
98	d1y8ma1	Alignment	not modelled	98.0	14	Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratricopeptide repeat (TPR)
99	c2f1tB	Alignment		97.9	12	PDB header: membrane protein Chain: B: PDB Molecule: outer membrane protein w; PDBTitle: outer membrane protein ompw
100	d1p4ta	Alignment	not modelled	97.8	16	Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
101	c2x27X	Alignment		97.8	11	PDB header: membrane protein Chain: X: PDB Molecule: outer membrane protein oprg; PDBTitle: crystal structure of the outer membrane protein oprg from2 pseudomonas aeruginosa
102	c2lhfa	Alignment	not modelled	97.7	14	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane protein h1; PDBTitle: solution structure of outer membrane protein h (oprh)

					from p.2 aeruginosa in dhpc micelles
103	d2pqrbl	Alignment	not modelled	97.6	7 Fold: alpha-alpha superhelix Superfamily: TPR-like Family: Tetratrico peptide repeat (TPR)
104	c2x4mD	Alignment		97.3	17 PDB header: hydrolase Chain: D: PDB Molecule: coagulase/fibrinolysin; PDBTitle: yersinia pestis plasminogen activator pla
105	c3sljA	Alignment	not modelled	97.3	13 PDB header: protein transport Chain: A: PDB Molecule: serine protease espp; PDBTitle: pre-cleavage structure of the autotransporter espp - n1023a mutant
106	dli78a	Alignment		97.3	16 Fold: Transmembrane beta-barrels Superfamily: OMPT-like Family: Outer membrane protease OMPT
107	c3kvnA	Alignment	not modelled	97.3	12 PDB header: hydrolase Chain: A: PDB Molecule: esterase esta; PDBTitle: crystal structure of the full-length autotransporter esta from2 pseudomonas aeruginosa
108	c3qraA	Alignment	not modelled	97.2	10 PDB header: cell invasion Chain: A: PDB Molecule: attachment invasion locus protein; PDBTitle: the crystal structure of ail, the attachment invasion locus protein of2 yersinia pestis
109	d1g90a	Alignment	not modelled	97.1	15 Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
110	c2ff4B	Alignment	not modelled	97.1	14 PDB header: transcription Chain: B: PDB Molecule: probable regulatory protein embr; PDBTitle: mycobacterium tuberculosis embr in complex with low affinity2 phosphopeptide
111	d1qj8a	Alignment	not modelled	97.1	15 Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
112	c3q3hA	Alignment	not modelled	97.1	7 PDB header: transferase Chain: A: PDB Molecule: hmw1c-like glycosyltransferase; PDBTitle: crystal structure of the actinobacillus pleuropneumoniae hmw1c2 glycosyltransferase in complex with udp-glc
113	d1qjpa	Alignment	not modelled	97.0	11 Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
114	c3nb3C	Alignment	not modelled	96.9	16 PDB header: virus Chain: C: PDB Molecule: outer membrane protein a; PDBTitle: the host outer membrane proteins ompa and ompc are packed at specific2 sites in the shigella phage sf6 virion as structural components
115	c3mekA	Alignment	not modelled	96.9	10 PDB header: transferase Chain: A: PDB Molecule: set and mynd domain-containing protein 3; PDBTitle: crystal structure of human histone-lysine n-2 methyltransferase smyd3 in complex with s-adenosyl-l-3 methionine
116	c3mkqA	Alignment	not modelled	96.8	15 PDB header: transport protein Chain: A: PDB Molecule: coatomer beta'-subunit; PDBTitle: crystal structure of yeast alpha/betaprime-cop subcomplex of the copi2 vesicular coat
117	c2qomB	Alignment	not modelled	96.7	11 PDB header: hydrolase Chain: B: PDB Molecule: serine protease espp; PDBTitle: the crystal structure of the e.coli espp autotransporter beta-domain.
118	c3qwvA	Alignment	not modelled	96.7	11 PDB header: transferase Chain: A: PDB Molecule: set and mynd domain-containing protein 2; PDBTitle: crystal structure of histone lysine methyltransferase smyd2 in complex2 with the cofactor product adohcy
119	c3aehB	Alignment	not modelled	96.6	14 PDB header: hydrolase Chain: B: PDB Molecule: hemoglobin-binding protease hbp autotransporter; PDBTitle: integral membrane domain of autotransporter hbp
120	c3oxgA	Alignment	not modelled	96.5	13 PDB header: transferase Chain: A: PDB Molecule: set and mynd domain-containing protein 3; PDBTitle: human lysine methyltransferase smyd3 in complex with adohcy (form iii)