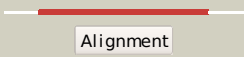

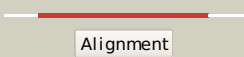

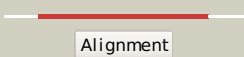
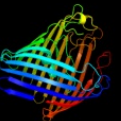
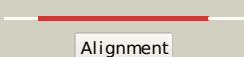

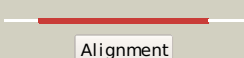

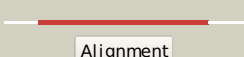

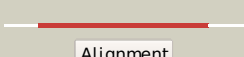

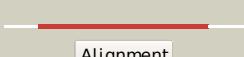

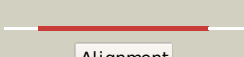
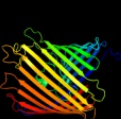






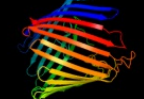




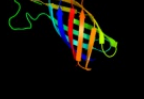



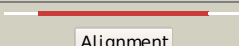

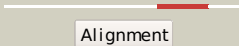

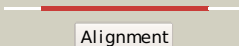


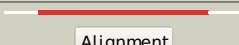



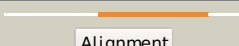
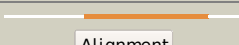
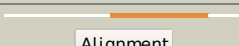



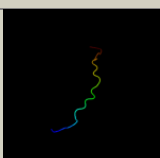



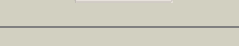
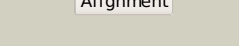



#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	d1phoa_	 Alignment		100.0	100	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
2	d2zfga1	 Alignment		100.0	63	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
3	d1osma_	 Alignment		100.0	59	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
4	c3nsgA_	 Alignment		100.0	52	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane protein f; PDBTitle: crystal structure of ompf, an outer membrane protein from salmonella2 typhi
5	d2fgqx1	 Alignment		100.0	18	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
6	c3a2rX_	 Alignment		100.0	18	PDB header: membrane protein Chain: X: PDB Molecule: outer membrane protein ii; PDBTitle: crystal structure of outer membrane protein porb from neisseria2 meningitidis
7	d2pora_	 Alignment		100.0	13	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
8	d3prna_	 Alignment		100.0	14	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Porin
9	c2o4vA_	 Alignment		99.9	12	PDB header: membrane protein Chain: A: PDB Molecule: porin p; PDBTitle: an arginine ladder in oprp mediates phosphate specific transfer across2 the outer membrane
10	c3qraA_	 Alignment		99.1	15	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
11	c2y0kA_	 Alignment		99.0	11	PDB header: transport protein Chain: A: PDB Molecule: pyroglutamate porin opdo; PDBTitle: crystal structure of pseudomonas aeruginosa opdo

12	d1p4ta_	Alignment		99.0	14	Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
13	d1qj8a_	Alignment		98.9	14	Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
14	c3jtyB_	Alignment		98.8	10	PDB header: transport protein Chain: B: PDB Molecule: benf-like porin; PDBTitle: crystal structure of a benf-like porin from pseudomonas fluorescens2 pf-5
15	c2odjA_	Alignment		98.8	13	PDB header: membrane protein Chain: A: PDB Molecule: porin d; PDBTitle: crystal structure of the outer membrane protein opod from pseudomonas2 aeruginosa
16	c2y0hA_	Alignment		98.7	11	PDB header: transport protein Chain: A: PDB Molecule: probable porin; PDBTitle: crystal structure of pseudomonas aeruginosa opd1
17	c2y0lA_	Alignment		98.6	13	PDB header: transport protein Chain: A: PDB Molecule: cis-aconitate porin opdh; PDBTitle: crystal structure of pseudomonas aeruginosa opdo
18	d1g90a_	Alignment		98.5	13	Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
19	d1qjpa_	Alignment		98.5	14	Fold: Transmembrane beta-barrels Superfamily: OMPA-like Family: Outer membrane protein
20	c2k0lA_	Alignment		98.4	12	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.
21	c3nb3C_	Alignment	not modelled	98.3	15	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
22	d1t16a_	Alignment	not modelled	98.2	12	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Outer membrane protein transport protein
23	c3brzA_	Alignment	not modelled	98.2	13	PDB header: transport protein Chain: A: PDB Molecule: todx; PDBTitle: crystal structure of the pseudomonas putida toluene2 transporter todx
24	c2qtkB_	Alignment	not modelled	98.2	12	PDB header: membrane protein Chain: B: PDB Molecule: probable porin; PDBTitle: crystal structure of the outer membrane protein opd2 from2 pseudomonas aeruginosa
25	c2f1tB_	Alignment	not modelled	98.1	15	PDB header: membrane protein Chain: B: PDB Molecule: outer membrane protein w; PDBTitle: outer membrane protein ompw
26	c2lhfA_	Alignment	not modelled	98.1	13	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
27	c2x27X_	Alignment	not modelled	98.0	14	PDB header: membrane protein Chain: X: PDB Molecule: outer membrane protein oprg; PDBTitle: crystal structure of the outer membrane protein oprg from2 pseudomonas aeruginosa
28	c3dwoX_	Alignment	not modelled	97.3	11	PDB header: membrane protein Chain: X: PDB Molecule: probable outer membrane protein; PDBTitle: crystal structure of a pseudomonas aeruginosa fadl homologue

29	c2jmmA		Alignment	not modelled	96.7	14	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane protein a; PDBTitle: nmr solution structure of a minimal transmembrane beta-2 barrel platform protein
30	c2wjgA		Alignment	not modelled	96.0	9	PDB header: transport protein Chain: A: PDB Molecule: probable n-acetylneuraminic acid outer membrane channel PDBTitle: nanc porin structure in hexagonal crystal form.
31	d2mpa		Alignment	not modelled	95.9	13	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Maltoporin-like
32	d1uynx		Alignment	not modelled	95.7	11	Fold: Transmembrane beta-barrels Superfamily: Autotransporter Family: Autotransporter
33	c2x4mD		Alignment	not modelled	95.5	19	PDB header: hydrolase Chain: D: PDB Molecule: coagulase/fibrinolysin; PDBTitle: yersinia pestis plasminogen activator pla
34	c3aehB		Alignment	not modelled	95.5	12	PDB header: hydrolase Chain: B: PDB Molecule: hemoglobin-binding protease hbp autotransporter; PDBTitle: integral membrane domain of autotransporter hbp
35	c3bryB		Alignment	not modelled	94.9	11	PDB header: transport protein Chain: B: PDB Molecule: tbux; PDBTitle: crystal structure of the ralstonia pickettii toluene2 transporter tbux
36	c3rbhC		Alignment	not modelled	94.6	8	PDB header: transport protein Chain: C: PDB Molecule: alginate production protein alge; PDBTitle: structure of alginate export protein alge from pseudomonas aeruginosa
37	c2iwwD		Alignment	not modelled	94.3	13	PDB header: ion channel Chain: D: PDB Molecule: outer membrane protein g; PDBTitle: structure of the monomeric outer membrane porin ompg in the2 open and closed conformation
38	d1af6a		Alignment	not modelled	93.3	11	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Maltoporin-like
39	d1a0tp		Alignment	not modelled	93.2	12	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Maltoporin-like
40	c3qq2C		Alignment	not modelled	93.1	14	PDB header: membrane protein/protein transport Chain: C: PDB Molecule: brka autotransporter; PDBTitle: crystal structure of the beta domain of the bordetella autotransporter2 brka
41	c2qomB		Alignment	not modelled	90.7	14	PDB header: hydrolase Chain: B: PDB Molecule: serine protease espp; PDBTitle: the crystal structure of the e.coli espp autotransporter beta-domain.
42	c3sljA		Alignment	not modelled	89.2	12	PDB header: protein transport Chain: A: PDB Molecule: serine protease espp; PDBTitle: pre-cleavage structure of the autotransporter espp - n1023a mutant
43	c3kvnA		Alignment	not modelled	86.2	14	PDB header: hydrolase Chain: A: PDB Molecule: esterase esta; PDBTitle: crystal structure of the full-length autotransporter esta from2 pseudomonas aeruginosa
44	d1i78a		Alignment	not modelled	83.3	16	Fold: Transmembrane beta-barrels Superfamily: OMPT-like Family: Outer membrane protease OMPT
45	d2gufa1		Alignment	not modelled	69.4	14	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Ligand-gated protein channel
46	c2iahA		Alignment	not modelled	65.3	16	PDB header: membrane protein Chain: A: PDB Molecule: ferripyoverdine receptor; PDBTitle: crystal structure of the ferripyoverdine receptor of the outer2 membrane of pseudomonas aeruginosa bound to ferripyoverdine.
47	d2jnaa1		Alignment		54.1	48	Fold: Dodecin subunit-like Superfamily: YdgH-like Family: YdgH-like
48	c2grxB		Alignment	not modelled	43.3	11	PDB header: metal transport Chain: B: PDB Molecule: ferrichrome-iron receptor; PDBTitle: crystal structure of tonb in complex with fhua, e. coli2 outer membrane receptor for ferrichrome
49	c3fhhA		Alignment	not modelled	39.0	13	PDB header: membrane protein Chain: A: PDB Molecule: outer membrane heme receptor shua; PDBTitle: crystal structure of the heme/hemoglobin outer membrane2 transporter shua from shigella dysenteriae
50	c3qlbA		Alignment	not modelled	35.6	12	PDB header: metal transport Chain: A: PDB Molecule: enantio-pyochelin receptor; PDBTitle: enantiopyochelin outer membrane tonb-dependent transporter from2 pseudomonas fluorescens bound to the ferri-enantiopyochelin
51	c1xkW		Alignment	not modelled	29.3	13	PDB header: membrane protein Chain: A: PDB Molecule: fe(iii)-pyochelin receptor; PDBTitle: pyochelin outer membrane receptor fpta from pseudomonas2 aeruginosa
52	c2hdiA		Alignment	not modelled	26.4	13	PDB header: protein transport, antimicrobial protein Chain: A: PDB Molecule: colicin i receptor; PDBTitle: crystal structure of the colicin i receptor cir from e.coli in complex2 with receptor binding domain of colicin ia.
53	c2k4tA		Alignment	not modelled	23.2	13	PDB header: membrane protein, apoptosis Chain: A: PDB Molecule: voltage-dependent anion-selective channel

						PDBTitle: solution structure of human vdac-1 in ldao micelles
54	c3cslB_	Alignment	not modelled	12.3	16	PDB header: membrane protein/heme binding protein Chain: B: PDB Molecule: hasr protein; PDBTitle: structure of the serratia marcescens hemophore receptor hasr in2 complex with its hemophore hasa and heme
55	c1qd6C_	Alignment	not modelled	10.8	12	PDB header: membrane protein Chain: C: PDB Molecule: protein (outer membrane phospholipase (ompla)); PDBTitle: outer membrane phospholipase a from escherichia coli
56	c2k3aA_	Alignment	not modelled	9.7	32	PDB header: hydrolase Chain: A: PDB Molecule: chap domain protein; PDBTitle: nmr solution structure of staphylococcus saprophyticus chap2 (cysteine, histidine-dependent amidohydrolases/peptidases)3 domain protein. northeast structural genomics consortium4 target syr11
57	d1by5a_	Alignment	not modelled	7.1	10	Fold: Transmembrane beta-barrels Superfamily: Porins Family: Ligand-gated protein channel
58	c2vdaB_	Alignment	not modelled	6.9	40	PDB header: protein transport Chain: B: PDB Molecule: maltoporin; PDBTitle: solution structure of the seca-signal peptide complex