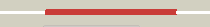
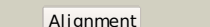
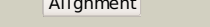
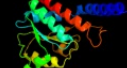
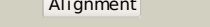
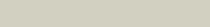
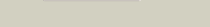
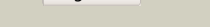
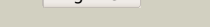


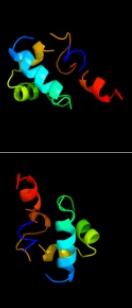
Phyre²

Email	i.a.kelley@imperial.ac.uk
Description	P0ACS7
Date	Thu Jan 5 11:18:55 GMT 2012
Unique Job ID	a49418611e7b07bd

Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c3shoA_			100.0	17	PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator, rpir family; PDBTitle: crystal structure of rpir transcription factor from sphaerobacter2 thermophilus (sugar isomerase domain)
2	c3fxaA_			99.9	17	PDB header: sugar binding protein Chain: A: PDB Molecule: sis domain protein; PDBTitle: crystal structure of a putative sugar-phosphate isomerase2 (lmof2365_0531) from listeria monocytogenes str. 4b2365 at 1.60 a3 resolution
3	c3etnD_			99.9	16	PDB header: isomerase Chain: D: PDB Molecule: putative phosphosugar isomerase involved in capsule PDBTitle: crystal structure of putative phosphosugar isomerase involved in2 capsule formation (yp_209877.1) from bacteroides fragilis nctc 93433 at 1.70 a resolution
4	c2xhzC_			99.9	22	PDB header: isomerase Chain: C: PDB Molecule: arabinose 5-phosphate isomerase; PDBTitle: probing the active site of the sugar isomerase domain from e. coli2 arabinose-5-phosphate isomerase via x-ray crystallography
5	d1vima_			99.9	16	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
6	d1m3sa_			99.9	19	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
7	d1jeoa_			99.9	15	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
8	c2zj3A_			99.8	17	PDB header: transferase Chain: A: PDB Molecule: glucosamine--fructose-6-phosphate PDBTitle: isomerase domain of human glucose:fructose-6-phosphate2 amidotransferase
9	c2puwA_			99.8	18	PDB header: transferase Chain: A: PDB Molecule: isomerase domain of glutamine-fructose-6-phosphate PDBTitle: the crystal structure of isomerase domain of glucosamine-6-phosphate2 synthase from candida albicans
10	c2yvaB_			99.8	19	PDB header: dna binding protein Chain: B: PDB Molecule: dnaa initiator-associating protein diaa; PDBTitle: crystal structure of escherichia coli diaa
11	d1x92a_			99.8	20	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain

12	d1tk9a_			99.8	16	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
13	c3trjC_			99.8	17	PDB header: isomerase Chain: C: PDB Molecule: phosphoheptose isomerase; PDBTitle: structure of a phosphoheptose isomerase from francisella tularensis
14	c2x3yA_			99.8	20	PDB header: isomerase Chain: A: PDB Molecule: phosphoheptose isomerase; PDBTitle: crystal structure of gmha from burkholderia pseudomallei
15	c3cvjB_			99.8	21	PDB header: isomerase Chain: B: PDB Molecule: putative phosphoheptose isomerase; PDBTitle: crystal structure of a putative phosphoheptose isomerase (bh3325) from2 bacillus halodurans c-125 at 2.00 a resolution
16	d1j5xa_			99.8	19	Fold: SIS domain Superfamily: SIS domain Family: double-SIS domain
17	d1x94a_			99.8	21	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
18	c2a3nA_			99.8	18	PDB header: sugar binding protein Chain: A: PDB Molecule: putative glucosamine-fructose-6-phosphate aminotransferase; PDBTitle: crystal structure of a putative glucosamine-fructose-6-phosphate2 aminotransferase (stm4540.s) from salmonella typhimurium lt2 at 1.353 a resolution
19	c1nria_			99.8	19	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: hypothetical protein hi0754; PDBTitle: crystal structure of putative phosphosugar isomerase hi0754 from2 haemophilus influenzae
20	d1nria_			99.8	19	Fold: SIS domain Superfamily: SIS domain Family: mono-SIS domain
21	c3iwfA_			99.8	18	PDB header: transcription regulator Chain: A: PDB Molecule: transcription regulator rpr family; PDBTitle: the crystal structure of the n-terminal domain of a rpr2 transcriptional regulator from staphylococcus epidermidis to 1.4a
22	c3euad_		not modelled	99.8	22	PDB header: isomerase Chain: D: PDB Molecule: putative fructose-aminoacid-6-phosphate deglycase; PDBTitle: crystal structure of a putative phosphosugar isomerase (bsu32610) from2 bacillus subtilis at 1.90 a resolution
23	c3fj1A_		not modelled	99.8	14	PDB header: isomerase Chain: A: PDB Molecule: putative phosphosugar isomerase; PDBTitle: crystal structure of putative phosphosugar isomerase (yp_167080.1)2 from silicibacter pomeroyi dss-3 at 1.75 a resolution
24	c3g68A_		not modelled	99.8	18	PDB header: isomerase Chain: A: PDB Molecule: putative phosphosugar isomerase; PDBTitle: crystal structure of a putative phosphosugar isomerase (cd3275) from2 clostridium difficile 630 at 1.80 a resolution
25	d1moqa_		not modelled	99.8	15	Fold: SIS domain Superfamily: SIS domain Family: double-SIS domain
26	c3hbaA_		not modelled	99.8	18	PDB header: isomerase Chain: A: PDB Molecule: putative phosphosugar isomerase; PDBTitle: crystal structure of a putative phosphosugar isomerase (sden_2705)2 from shewanella denitrificans os217 at 2.00 a resolution
						PDB header: transferase Chain: A: PDB Molecule: glucosamine--fructose-6-phosphate

27	c3tbfa	Alignment	not modelled	99.8	17	<p>PDB header:isomerase</p> <p>PDBTitle: c-terminal domain of glucosamine-fructose-6-phosphate aminotransferase2 from francisella tularensis.</p>
28	c2amlB	Alignment	not modelled	99.8	12	<p>PDB header:transferase</p> <p>Chain: B: PDB Molecule:sis domain protein;</p> <p>PDBTitle: crystal structure of lmo0035 protein (46906266) from listeria2 monocytogenes 4b f2365 at 1.50 a resolution</p>
29	c3knzA	Alignment	not modelled	99.7	17	<p>PDB header:sugar binding protein</p> <p>Chain: A: PDB Molecule:putative sugar binding protein;</p> <p>PDBTitle: crystal structure of putative sugar binding protein (np_459565.1) from2 salmonella typhimurium lt2 at 2.50 a resolution</p>
30	c3fkjA	Alignment	not modelled	99.7	16	<p>PDB header:isomerase</p> <p>Chain: A: PDB Molecule:putative phosphosugar isomerases;</p> <p>PDBTitle: crystal structure of a putative phosphosugar isomerase (stm_0572) from2 salmonella typhimurium lt2 at 2.12 a resolution</p>
31	c2decA	Alignment	not modelled	99.6	16	<p>PDB header:structural genomics, unknown function</p> <p>Chain: A: PDB Molecule:325aa long hypothetical protein;</p> <p>PDBTitle: crystal structure of the pho510 protein from pyrococcus horikoshii ot3</p>
32	c1jxaA	Alignment	not modelled	99.5	15	<p>PDB header:transferase</p> <p>Chain: A: PDB Molecule:glucosamine 6-phosphate synthase;</p> <p>PDBTitle: glucosamine 6-phosphate synthase with glucose 6-phosphate</p>
33	c3odpA	Alignment	not modelled	99.5	17	<p>PDB header:isomerase</p> <p>Chain: A: PDB Molecule:putative tagatose-6-phosphate ketose/aldose isomerase;</p> <p>PDBTitle: crystal structure of a putative tagatose-6-phosphate ketose/aldose2 isomerase (nt01cx_0292) from clostridium novyi nt at 2.35 a3 resolution</p>
34	d1x9ia	Alignment	not modelled	99.5	16	<p>Fold:SIS domain</p> <p>Superfamily:SIS domain</p> <p>Family:double-SIS domain</p>
35	c2o3fc	Alignment		99.5	13	<p>PDB header:transcription</p> <p>Chain: C: PDB Molecule:putative hth-type transcriptional regulator ybbh;</p> <p>PDBTitle: structural genomics, the crystal structure of the n-2 terminal domain of the putative transcriptional regulator3 ybbh from bacillus subtilis subsp. subtilis str. 168.</p>
36	d2o3fa1	Alignment		99.5	13	<p>Fold:DNA/RNA-binding 3-helical bundle</p> <p>Superfamily:Homeodomain-like</p> <p>Family:RpiR-like</p>
37	c3i0zB	Alignment	not modelled	99.4	16	<p>PDB header:isomerase</p> <p>Chain: B: PDB Molecule:putative tagatose-6-phosphate ketose/aldose isomerase;</p> <p>PDBTitle: crystal structure of putative putative tagatose-6-phosphate2 ketose/aldose isomerase (np_344614.1) from streptococcus pneumoniae3 tigr4 at 1.70 a resolution</p>
38	c3c3jA	Alignment	not modelled	99.4	16	<p>PDB header:isomerase</p> <p>Chain: A: PDB Molecule:putative tagatose-6-phosphate ketose/aldose isomerase;</p> <p>PDBTitle: crystal structure of tagatose-6-phosphate ketose/aldose isomerase from2 escherichia coli</p>
39	c1zzgB	Alignment	not modelled	98.1	18	<p>PDB header:isomerase</p> <p>Chain: B: PDB Molecule:glucose-6-phosphate isomerase;</p> <p>PDBTitle: crystal structure of hypothetical protein tt0462 from thermus2 thermophilus hb8</p>
40	c2q8nB	Alignment	not modelled	97.9	17	<p>PDB header:isomerase</p> <p>Chain: B: PDB Molecule:glucose-6-phosphate isomerase;</p> <p>PDBTitle: crystal structure of glucose-6-phosphate isomerase (ec2 5.3.1.9) (tm1385) from thermotoga maritima at 1.82 a3 resolution</p>
41	c3ff1B	Alignment	not modelled	97.7	19	<p>PDB header:isomerase</p> <p>Chain: B: PDB Molecule:glucose-6-phosphate isomerase;</p> <p>PDBTitle: structure of glucose 6-phosphate isomerase from staphylococcus aureus</p>
42	d1c7qa	Alignment	not modelled	97.7	21	<p>Fold:SIS domain</p> <p>Superfamily:SIS domain</p> <p>Family:Phosphoglucose isomerase, PGI</p>
43	d1lnwa	Alignment	not modelled	97.2	13	<p>Fold:DNA/RNA-binding 3-helical bundle</p> <p>Superfamily:"Winged helix" DNA-binding domain</p> <p>Family:MarR-like transcriptional regulators</p>
44	d1gzda	Alignment	not modelled	97.1	14	<p>Fold:SIS domain</p> <p>Superfamily:SIS domain</p> <p>Family:Phosphoglucose isomerase, PGI</p>
45	c3nqoB	Alignment	not modelled	97.0	10	<p>PDB header:transcription</p> <p>Chain: B: PDB Molecule:marr-family transcriptional regulator;</p> <p>PDBTitle: crystal structure of a marr family transcriptional regulator (cd1569)2 from clostridium difficile 630 at 2.20 a resolution</p>
46	c2wu8A	Alignment	not modelled	96.9	12	<p>PDB header:isomerase</p> <p>Chain: A: PDB Molecule:glucose-6-phosphate isomerase;</p> <p>PDBTitle: structural studies of phosphoglucose isomerase from2 mycobacterium tuberculosis h37rv</p>
47	d1u0fa	Alignment	not modelled	96.9	14	<p>Fold:SIS domain</p> <p>Superfamily:SIS domain</p> <p>Family:Phosphoglucose isomerase, PGI</p>
48	c3ljkA	Alignment	not modelled	96.9	12	<p>PDB header:isomerase</p> <p>Chain: A: PDB Molecule:glucose-6-phosphate isomerase;</p> <p>PDBTitle: glucose-6-phosphate isomerase from francisella tularensis.</p>
49	d1hm5a	Alignment	not modelled	96.8	14	<p>Fold:SIS domain</p> <p>Superfamily:SIS domain</p>

					Family: Phosphoglucose isomerase, PGI
50	c3bj6B	Alignment	not modelled	96.8	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator, marr family; PDBTitle: crystal structure of marr family transcription regulator sp03579
51	d1iata	Alignment	not modelled	96.8	Fold: SIS domain Superfamily: SIS domain Family: Phosphoglucose isomerase, PGI
52	d1g50a	Alignment	not modelled	96.8	Fold: SIS domain Superfamily: SIS domain Family: Phosphoglucose isomerase, PGI
53	c3hjbA	Alignment	not modelled	96.8	PDB header: isomerase Chain: A: PDB Molecule: glucose-6-phosphate isomerase; PDBTitle: 1.5 angstrom crystal structure of glucose-6-phosphate isomerase from <i>vibrio cholerae</i> .
54	c3ujhB	Alignment	not modelled	96.7	PDB header: isomerase Chain: B: PDB Molecule: glucose-6-phosphate isomerase; PDBTitle: crystal structure of substrate-bound glucose-6-phosphate isomerase2 from <i>toxoplasma gondii</i>
55	c2fa5B	Alignment	not modelled	96.7	PDB header: transcription Chain: B: PDB Molecule: transcriptional regulator marr/emrr family; PDBTitle: the crystal structure of an unliganded multiple antibiotic-2 resistance repressor (marr) from <i>xanthomonas campestris</i>
56	c1t10A	Alignment	not modelled	96.7	PDB header: isomerase Chain: A: PDB Molecule: glucose-6-phosphate isomerase; PDBTitle: phosphoglucose isomerase from <i>leishmania mexicana</i> in complex with 2 substrate d-fructose-6-phosphate
57	c3jx9B	Alignment	not modelled	96.7	PDB header: isomerase Chain: B: PDB Molecule: putative phosphoheptose isomerase; PDBTitle: crystal structure of putative phosphoheptose isomerase2 (yp_001815198.1) from <i>exiguobacterium</i> sp. 255-15 at 1.95 a resolution
58	c3oopA	Alignment	not modelled	96.5	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: lin2960 protein; PDBTitle: the structure of a protein with unknown function from <i>listeria innocua</i> 2 clip11262
59	d3broa1	Alignment	not modelled	96.5	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
60	c2rdpA	Alignment	not modelled	96.5	PDB header: transcription Chain: A: PDB Molecule: putative transcriptional regulator marr; PDBTitle: the structure of a marr family protein from <i>bacillus</i> 2 stearotherophilus
61	c3kp3B	Alignment	not modelled	96.4	PDB header: transcription regulator/antibiotic Chain: B: PDB Molecule: transcriptional regulator tcar; PDBTitle: staphylococcus epidermidis in complex with ampicillin
62	d1yioa1	Alignment	not modelled	96.4	Fold: DNA/RNA-binding 3-helical bundle Superfamily: C-terminal effector domain of the bipartite response regulators Family: GerE-like (LuxR/UhpA family of transcriptional regulators)
63	c3deuB	Alignment	not modelled	96.3	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator slya; PDBTitle: crystal structure of transcription regulatory protein slya2 from <i>salmonella typhimurium</i> in complex with salicylate3 ligands
64	c3tgtA	Alignment	not modelled	96.3	PDB header: transcription Chain: A: PDB Molecule: adc operon repressor adcr; PDBTitle: crystal structure of the zinc-dependent marr family transcriptional2 regulator adcr in the zn(ii)-bound state
65	c2o2cB	Alignment	not modelled	96.3	PDB header: isomerase Chain: B: PDB Molecule: glucose-6-phosphate isomerase, glycosomal; PDBTitle: crystal structure of phosphoglucose isomerase from <i>t. brucei</i> 2 containing glucose-6-phosphate in the active site
66	c3boqB	Alignment	not modelled	96.3	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator, marr family; PDBTitle: crystal structure of marr family transcriptional regulator from <i>silicibacter pomeroyi</i>
67	d3deua1	Alignment	not modelled	96.2	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
68	d2fbha1	Alignment	not modelled	96.2	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
69	d2a61a1	Alignment	not modelled	96.2	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
70	c2nnnB	Alignment	not modelled	96.2	PDB header: transcription Chain: B: PDB Molecule: probable transcriptional regulator; PDBTitle: crystal structure of probable transcriptional regulator from <i>pseudomonas aeruginosa</i>
71	c3r0aB	Alignment	not modelled	96.1	PDB header: transcription regulator Chain: B: PDB Molecule: putative transcriptional regulator; PDBTitle: possible transcriptional regulator from <i>methanoscincina mazei</i> go1 (gi2 21227196)
72	c2gxgA	Alignment	not modelled	96.1	PDB header: transcription Chain: A: PDB Molecule: 146aa long hypothetical transcriptional regulator; PDBTitle: crystal structure of emrr homolog from hyperthermophilic archaea2 sulfolobus tokodaii strain7
73	d2etha1	Alignment	not modelled	96.0	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
74	c3nbuC	Alignment	not modelled	96.0	PDB header: isomerase Chain: C: PDB Molecule: glucose-6-phosphate isomerase; PDBTitle: crystal structure of pgi glucosephosphate isomerase

75	c1x3uA	Alignment	not modelled	96.0	16	PDB header: transcription Chain: A: PDB Molecule: transcriptional regulatory protein fixJ; PDBTitle: solution structure of the c-terminal transcriptional2 activator domain of fixJ from sinorhizobium meliloti
76	d1s3ja	Alignment	not modelled	95.9	18	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
77	c3k0IA	Alignment	not modelled	95.9	10	PDB header: transcription regulator Chain: A: PDB Molecule: repressor protein; PDBTitle: crystal structure of putative marr family transcriptional2 regulator from acinetobacter sp. adp
78	c2qwwB	Alignment	not modelled	95.9	13	PDB header: transcription Chain: B: PDB Molecule: transcriptional regulator, marr family; PDBTitle: crystal structure of multiple antibiotic-resistance repressor (marr)2 (yp_013417.1) from listeria monocytogenes 4b f2365 at 2.07 a3 resolution
79	c3cjnA	Alignment	not modelled	95.9	12	PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator, marr family; PDBTitle: crystal structure of transcriptional regulator, marr family, from2 silicibacter pomeroyi
80	c3s2wB	Alignment	not modelled	95.8	10	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator, marr family; PDBTitle: the crystal structure of a marr transcriptional regulator from2 methanoscincina mazei go1
81	c3frwF	Alignment	not modelled	95.8	24	PDB header: structural genomics, unknown function Chain: F: PDB Molecule: putative trp repressor protein; PDBTitle: crystal structure of putative trpr protein from ruminococcus obuum
82	c3ecoB	Alignment	not modelled	95.7	12	PDB header: transcription Chain: B: PDB Molecule: mepr; PDBTitle: crystal structure of mepr, a transcription regulator of the2 staphylococcus aureus multidrug efflux pump mepa
83	c3g3zA	Alignment	not modelled	95.7	13	PDB header: transcription Chain: A: PDB Molecule: transcriptional regulator, marr family; PDBTitle: the structure of nmb1585, a marr family regulator from neisseria2 meningitidis
84	d1jhfa1	Alignment	not modelled	95.7	18	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: LexA repressor, N-terminal DNA-binding domain
85	c2zkqb	Alignment	not modelled	95.6	8	PDB header: ribosomal protein/rna Chain: B: PDB Molecule: rna expansion segment es3; PDBTitle: structure of a mammalian ribosomal 40S subunit within an2 80S complex obtained by docking homology models of the rna3 and proteins into an 8.7 a cryo-em map
86	c3iz6A	Alignment	not modelled	95.6	7	PDB header: ribosome Chain: A: PDB Molecule: 40S ribosomal protein sa (s2p); PDBTitle: localization of the small subunit ribosomal proteins into a 5.5 a2 cryo-em map of triticum aestivum translating 80S ribosome
87	c3pr3B	Alignment	not modelled	95.6	15	PDB header: isomerase Chain: B: PDB Molecule: glucose-6-phosphate isomerase; PDBTitle: crystal structure of plasmodium falciparum glucose-6-phosphate2 isomerase (pf14_0341) in complex with fructose-6-phosphate
88	d1a04a1	Alignment	not modelled	95.6	14	Fold: DNA/RNA-binding 3-helical bundle Superfamily: C-terminal effector domain of the bipartite response regulators Family: GerE-like (LuxR/UhpA family of transcriptional regulators)
89	d1fsea	Alignment	not modelled	95.5	25	Fold: DNA/RNA-binding 3-helical bundle Superfamily: C-terminal effector domain of the bipartite response regulators Family: GerE-like (LuxR/UhpA family of transcriptional regulators)
90	d1lj9a	Alignment	not modelled	95.5	14	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
91	c3bchA	Alignment	not modelled	95.5	7	PDB header: cell adhesion, ribosomal protein Chain: A: PDB Molecule: 40S ribosomal protein sa; PDBTitle: crystal structure of the human laminin receptor precursor
92	c2e7zA	Alignment	not modelled	95.5	13	PDB header: lyase Chain: A: PDB Molecule: acetylene hydratase ahY; PDBTitle: acetylene hydratase from pelobacter acetylénicus
93	d2jioa2	Alignment	not modelled	95.5	16	Fold: Formate dehydrogenase/DMSO reductase, domains 1-3 Superfamily: Formate dehydrogenase/DMSO reductase, domains 1-3 Family: Formate dehydrogenase/DMSO reductase, domains 1-3
94	c2i4aA	Alignment	not modelled	95.4	12	PDB header: dna binding protein Chain: A: PDB Molecule: leucine responsive regulatory protein; PDBTitle: nmr structure of the dna-binding domain of e.coli lrp
95	d2cg4a1	Alignment	not modelled	95.3	22	Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: Lrp/AsnC-like transcriptional regulator N-terminal domain
96	c3ouuA	Alignment	not modelled	95.3	3	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: lin2118 protein; PDBTitle: the structure of a protein with unkown function from listeria innocua
97	c3cdhB	Alignment	not modelled	95.3	11	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator, marr family; PDBTitle: crystal structure of the marr family transcriptional regulator spo14532 from silicibacter pomeroyi dss-3
98	c3bpXB	Alignment	not modelled	95.3	13	PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator; PDBTitle: crystal structure of marr
99	c3kjxD	Alignment	not modelled	95.3	48	PDB header: transcription regulator Chain: D: PDB Molecule: transcriptional regulator, lacI family; PDBTitle: crystal structure of a transcriptional regulator, lacI2 family protein from silicibacter pomeroyi
100	d1h0ba2	Alignment	not modelled	95.3	16	Fold: Formate dehydrogenase/DMSO reductase, domains 1-3 Superfamily: Formate dehydrogenase/DMSO reductase, domains 1-

100	c1t1a_	Alignment	not modelled	95.3	10	3	Family: Formate dehydrogenase/DMSO reductase, domains 1-3
101	d2cfxa1	Alignment	not modelled	95.2	17		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: Lrp/AsnC-like transcriptional regulator N-terminal domain
102	c3mn2B_	Alignment	not modelled	95.2	8		PDB header: transcription regulator Chain: B: PDB Molecule: probable arac family transcriptional regulator; PDBTitle: the crystal structure of a probable arac family transcriptional2 regulator from rhodopseudomonas palustris cga009
103	c1zljE_	Alignment	not modelled	95.2	16		PDB header: transcription Chain: E: PDB Molecule: dormancy survival regulator; PDBTitle: crystal structure of the mycobacterium tuberculosis hypoxic2 response regulator dosr c-terminal domain
104	c3bjaaA_	Alignment	not modelled	95.2	10		PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator, marr family, putative; PDBTitle: crystal structure of putative marr-like transcription regulator2 (np_978771.1) from bacillus cereus atcc 10987 at 2.38 a resolution
105	c3lsgD_	Alignment	not modelled	95.2	12		PDB header: transcription regulator Chain: D: PDB Molecule: two-component response regulator yesn; PDBTitle: the crystal structure of the c-terminal domain of the two component response regulator yesn from fusobacterium3 nucleatum subsp. nucleatum atcc 25586
106	c3hrmA_	Alignment	not modelled	95.1	12		PDB header: transcription regulator Chain: A: PDB Molecule: hth-type transcriptional regulator sarz; PDBTitle: crystal structure of staphylococcus aureus protein sarz in sulfenic2 acid form
107	d2fbia1	Alignment	not modelled	95.1	14		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
108	c2k9sA_	Alignment	not modelled	95.1	11		PDB header: transcription Chain: A: PDB Molecule: arabinose operon regulatory protein; PDBTitle: solution structure of dna binding domain of e. coli arac
109	c2wteB_	Alignment	not modelled	95.1	18		PDB header: antiviral protein Chain: B: PDB Molecule: csa3; PDBTitle: the structure of the crispr-associated protein, csa3, from2 sulfolobus solfataricus at 1.8 angstrom resolution.
110	c3korD_	Alignment	not modelled	95.1	18		PDB header: transcription Chain: D: PDB Molecule: possible trp repressor; PDBTitle: crystal structure of a putative trp repressor from staphylococcus2 aureus
111	d2cyya1	Alignment	not modelled	95.0	17		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: Lrp/AsnC-like transcriptional regulator N-terminal domain
112	d2hr3a1	Alignment	not modelled	95.0	9		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
113	d1p4wa_	Alignment	not modelled	94.9	21		Fold: DNA/RNA-binding 3-helical bundle Superfamily: C-terminal effector domain of the bipartite response regulators Family: GerE-like (LuxR/UhpA family of transcriptional regulators)
114	c2rnjA_	Alignment	not modelled	94.9	16		PDB header: transcription Chain: A: PDB Molecule: response regulator protein vrar; PDBTitle: nmr structure of the s. aureus vrar dna binding domain
115	c3k2za_	Alignment	not modelled	94.8	13		PDB header: hydrolase Chain: A: PDB Molecule: lexa repressor; PDBTitle: crystal structure of a lexa protein from thermotoga maritima
116	d2iv2x2	Alignment	not modelled	94.8	24		Fold: Formate dehydrogenase/DMSO reductase, domains 1-3 Superfamily: Formate dehydrogenase/DMSO reductase, domains 1-3 Family: Formate dehydrogenase/DMSO reductase, domains 1-3
117	c1s1hb_	Alignment	not modelled	94.8	10		PDB header: ribosome Chain: B: PDB Molecule: 40s ribosomal protein s0-a; PDBTitle: structure of the ribosomal 80s-eef2-sordarin complex from2 yeast obtained by docking atomic models for rna and protein3 components into a 11.7 a cryo-em map. this file, 1s1h,4 contains 40s subunit. the 60s ribosomal subunit is in file 1s1i.
118	d1jgsa_	Alignment	not modelled	94.8	9		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: MarR-like transcriptional regulators
119	c1bdhA_	Alignment	not modelled	94.8	50		PDB header: transcription/dna Chain: A: PDB Molecule: protein (purine repressor); PDBTitle: purine repressor mutant-hypoxanthine-palindromic operator2 complex
120	d1stza1	Alignment	not modelled	94.8	16		Fold: DNA/RNA-binding 3-helical bundle Superfamily: "Winged helix" DNA-binding domain Family: Heat-inducible transcription repressor HrcA, N-terminal domain