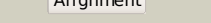
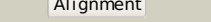
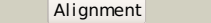
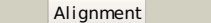
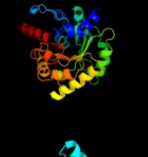
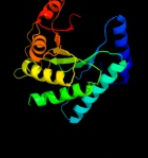


Phyre²

Email	I.a.kelley@imperial.ac.uk
Description	P0AB74
Date	Thu Jan 5 11:14:47 GMT 2012
Unique Job ID	edd5d0a97b9b6d6b

Detailed template information

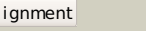
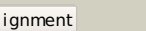
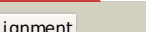
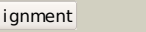
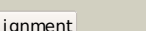
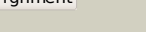
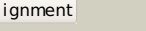
#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	d1gvfa_			100.0	100	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class II FBP aldolase
2	c3g94B_			100.0	40	PDB header: lyase Chain: B; PDB Molecule: fructose-bisphosphate aldolase, class ii; PDBTitle: the crystal structure of fructose 1,6-bisphosphate aldolase from2 bacillus anthracis str. 'ames ancestor'
3	d1rvga_			100.0	43	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class II FBP aldolase
4	c3c52B_			100.0	41	PDB header: lyase Chain: B; PDB Molecule: fructose-bisphosphate aldolase; PDBTitle: class ii fructose-1,6-bisphosphate aldolase from2 helicobacter pylori in complex with3 phosphoglycolohydroxamic acid, a competitive inhibitor
5	d1dosa_			100.0	25	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class II FBP aldolase
6	c3elfA_			100.0	26	PDB header: lyase Chain: A; PDB Molecule: fructose-bisphosphate aldolase; PDBTitle: structural characterization of tetrameric mycobacterium tuberculosis2 fructose 1,6-bisphosphate aldolase - substrate binding and catalysis3 mechanism of a class iia bacterial aldolase
7	c3qm3C_			100.0	26	PDB header: lyase Chain: C; PDB Molecule: fructose-bisphosphate aldolase; PDBTitle: 1.85 angstrom resolution crystal structure of fructose-bisphosphate2 aldolase (fba) from campylobacter jejuni
8	c2iswB_			100.0	44	PDB header: lyase Chain: B; PDB Molecule: putative fructose-1,6-bisphosphate aldolase; PDBTitle: structure of giardia fructose-1,6-bisphosphate aldolase in2 complex with phosphoglycolohydroxamate
9	c3pm6B_			100.0	32	PDB header: lyase Chain: B; PDB Molecule: putative fructose-bisphosphate aldolase; PDBTitle: crystal structure of a putative fructose-1,6-bisphosphate aldolase from2 coccidioides immitis solved by combined sad mr
10	d2figa1			99.4	16	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: GatZ-like
11	c2qjhH_			99.1	19	PDB header: lyase Chain: H; PDB Molecule: putative aldolase mj0400; PDBTitle: m. jannaschii adh synthase covalently bound to2 dihydroxyacetone phosphate

12	c2ze3A_	Alignment		98.7	15	PDB header: isomerase Chain: A: PDB Molecule: dfa0005; PDBTitle: crystal structure of dfa0005 complexed with alpha-ketoglutarate: a2 novel member of the icl/pepm superfamily from alkali-tolerant3 deinococcus fusicus
13	c3gr7A_	Alignment		98.6	17	PDB header: oxidoreductase Chain: A: PDB Molecule: nadph dehydrogenase; PDBTitle: structure of oye from geobacillus kaustophilus, hexagonal2 crystal form
14	d1thfd_	Alignment		98.6	14	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Histidine biosynthesis enzymes
15	c2c3zA_	Alignment		98.6	14	PDB header: lyase Chain: A: PDB Molecule: indole-3-glycerol phosphate synthase; PDBTitle: crystal structure of a truncated variant of indole-3-2-glycerol phosphate synthase from sulfolobus solfataricus
16	d1ps9a1	Alignment		98.5	20	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
17	c2h90A_	Alignment		98.4	17	PDB header: oxidoreductase Chain: A: PDB Molecule: xenobiotic reductase a; PDBTitle: xenobiotic reductase a in complex with coumarin
18	c3hf3A_	Alignment		98.4	15	PDB header: oxidoreductase Chain: A: PDB Molecule: chromate reductase; PDBTitle: old yellow enzyme from thermus scotoductus sa-01
19	d1j5ta_	Alignment		98.3	15	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
20	c3gndC_	Alignment		98.3	20	PDB header: lyase Chain: C: PDB Molecule: aldolase lsrf; PDBTitle: crystal structure of e. coli lsrf in complex with ribulose-5-phosphate
21	c1ps9A_	Alignment	not modelled	98.2	19	PDB header: oxidoreductase Chain: A: PDB Molecule: 2,4-dienoyl-coa reductase; PDBTitle: the crystal structure and reaction mechanism of e. coli 2,4-2 dienoyl coa reductase PDB header: lyase Chain: F: PDB Molecule: pa4872 oxaloacetate decarboxylase; PDBTitle: crystal structure of oxaloacetate decarboxylase from pseudomonas2 aeruginosa (pa4872) in complex with oxalate and mg2+.
22	c3b8iF_	Alignment	not modelled	98.2	18	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
23	d1a53a_	Alignment	not modelled	98.2	14	PDB header: oxidoreductase Chain: C: PDB Molecule: nadh:flavin oxidoreductase/nadh oxidase; PDBTitle: crystal structure of the thermostable old yellow enzyme from2 thermoanaerobacter pseudethanolicus e39
24	c3kruC_	Alignment	not modelled	98.2	17	PDB header: oxidoreductase Chain: C: PDB Molecule: nadh:flavin oxidoreductase/nadh oxidase; PDBTitle: crystal structure of the mycobacterium tuberculosis indole-3-glycerol 2 phosphate synthase (trpc) in apo form
25	c3qjaA_	Alignment	not modelled	98.2	17	PDB header: lyase Chain: A: PDB Molecule: indole-3-glycerol phosphate synthase; PDBTitle: crystal structure of the mycobacterium tuberculosis indole-3-glycerol 2 phosphate synthase (trpc) in apo form
26	c3igsB_	Alignment	not modelled	98.2	15	PDB header: isomerase Chain: B: PDB Molecule: n-acetylmannosamine-6-phosphate 2-epimerase 2; PDBTitle: structure of the salmonella enterica n-acetylmannosamine-6-phosphate 2-epimerase
27	d1xcfa_	Alignment	not modelled	98.1	15	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
28	d1h5ya_	Alignment	not modelled	98.1	15	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Histidine biosynthesis enzymes

29	d1y0ea	Alignment	not modelled	98.1	18	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: NanE-like
30	d1vyra	Alignment	not modelled	98.1	14	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
31	d1muma	Alignment	not modelled	98.0	18	Fold: TIM beta/alpha-barrel Superfamily: Phosphoenolpyruvate/pyruvate domain Family: Phosphoenolpyruvate mutase/Iscitrate lyase-like
32	c3eool	Alignment	not modelled	98.0	19	PDB header: lyase Chain: L: PDB Molecule: methylisocitrate lyase; PDBTitle: 2.9a crystal structure of methyl-isocitrate lyase from burkholderia pseudomallei
33	d1z41a1	Alignment	not modelled	98.0	15	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
34	d1i4na	Alignment	not modelled	97.9	15	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
35	c3q58A	Alignment	not modelled	97.9	13	PDB header: isomerase Chain: A: PDB Molecule: n-acetylmannosamine-6-phosphate 2-epimerase; PDBTitle: structure of n-acetylmannosamine-6-phosphate epimerase from salmonella2 enterica
36	c3k30B	Alignment	not modelled	97.9	15	PDB header: oxidoreductase Chain: B: PDB Molecule: histamine dehydrogenase; PDBTitle: histamine dehydrogenase from nocardiodes simplex
37	d1ujqa	Alignment	not modelled	97.9	18	Fold: TIM beta/alpha-barrel Superfamily: Phosphoenolpyruvate/pyruvate domain Family: Phosphoenolpyruvate mutase/Iscitrate lyase-like
38	c3ih1A	Alignment	not modelled	97.9	16	PDB header: lyase Chain: A: PDB Molecule: methylisocitrate lyase; PDBTitle: crystal structure of carboxyvinyl-carboxyphosphonate phosphorylmutase2 from bacillus anthracis
39	c2gq8A	Alignment	not modelled	97.8	14	PDB header: oxidoreductase Chain: A: PDB Molecule: oxidoreductase, fmn-binding; PDBTitle: structure of sye1, an oye homologue from s. ondeidensis, in complex2 with p-hydroxyacetophenone
40	d1ka9f	Alignment	not modelled	97.8	12	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Histidine biosynthesis enzymes
41	d1yxya1	Alignment	not modelled	97.8	11	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: NanE-like
42	c3bo9B	Alignment	not modelled	97.8	15	PDB header: oxidoreductase Chain: B: PDB Molecule: putative nitroalkan dioxygenase; PDBTitle: crystal structure of putative nitroalkan dioxygenase (tm0800) from thermotoga maritima at 2.71 a resolution
43	d1tqxa	Alignment	not modelled	97.8	11	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: D-ribulose-5-phosphate 3-epimerase
44	c2qiwa	Alignment	not modelled	97.7	13	PDB header: transferase Chain: A: PDB Molecule: pep phosphonomutase; PDBTitle: crystal structure of a putative phosphoenolpyruvate phosphonomutase2 (ncgl1015, cgl1060) from corynebacterium glutamicum atcc 13032 at 1.80 a resolution
45	d1wbha1	Alignment	not modelled	97.7	12	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
46	c2z6jB	Alignment	not modelled	97.7	18	PDB header: oxidoreductase Chain: B: PDB Molecule: trans-2-enoyl-acp reductase ii; PDBTitle: crystal structure of s. pneumoniae enoyl-acyl carrier2 protein reductase (fabk) in complex with an inhibitor
47	d1rd5a	Alignment	not modelled	97.7	14	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
48	c3l5aA	Alignment	not modelled	97.6	17	PDB header: oxidoreductase Chain: A: PDB Molecule: nahd/flavin oxidoreductase/nadh oxidase; PDBTitle: crystal structure of a probable nadh-dependent flavin oxidoreductase2 from staphylococcus aureus
49	d1qopa	Alignment	not modelled	97.6	13	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
50	d1gwja	Alignment	not modelled	97.6	13	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
51	c3fa4D	Alignment	not modelled	97.5	12	PDB header: lyase Chain: D: PDB Molecule: 2,3-dimethylmalate lyase; PDBTitle: crystal structure of 2,3-dimethylmalate lyase, a pep mutase/isocitrate2 lyase superfamily member, triclinic crystal form
52	d1piia2	Alignment	not modelled	97.5	16	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
53	c3tdmD	Alignment	not modelled	97.4	18	PDB header: de novo protein Chain: D: PDB Molecule: computationally designed two-fold symmetric tim-barrel PDBTitle: computationally designed tim-barrel protein, halfflr
54	d1vhca	Alignment	not modelled	97.4	12	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase

55	c3lyeA	Alignment	not modelled	97.3	11	PDB header: hydrolase Chain: A: PDB Molecule: oxaloacetate acetyl hydrolase; PDBTitle: crystal structure of oxaloacetate acetyl hydrolase
56	c2y85D	Alignment	not modelled	97.3	12	PDB header: isomerase Chain: D: PDB Molecule: phosphoribosyl isomerase a; PDBTitle: crystal structure of mycobacterium tuberculosis phosphoribosyl2 isomerase with bound rcdp
57	c2cw6B	Alignment	not modelled	97.3	17	PDB header: lyase Chain: B: PDB Molecule: hydroxymethylglutaryl-coa lyase, mitochondrial; PDBTitle: crystal structure of human hmg-coa lyase: insights into2 catalysis and the molecular basis for3 hydroxymethylglutaric aciduria
58	c2gjIA	Alignment	not modelled	97.3	16	PDB header: oxidoreductase Chain: A: PDB Molecule: hypothetical protein pa1024; PDBTitle: crystal structure of 2-nitropropane dioxygenase
59	c2w6rA	Alignment	not modelled	97.3	11	PDB header: lyase Chain: A: PDB Molecule: imidazole glycerol phosphate synthase subunit PDBTitle: crystal structure of an artificial (ba)8-barrel protein2 designed from identical half barrels
60	d1wa3a1	Alignment	not modelled	97.3	18	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
61	c1zlpA	Alignment	not modelled	97.2	14	PDB header: lyase Chain: A: PDB Molecule: petal death protein; PDBTitle: petal death protein psr132 with cysteine-linked glutaraldehyde forming2 a thiohemiacetal adduct
62	d1xm3a	Alignment	not modelled	97.2	18	Fold: TIM beta/alpha-barrel Superfamily: ThiG-like Family: ThiG-like
63	d1h1ya	Alignment	not modelled	97.2	12	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: D-ribulose-5-phosphate 3-epimerase
64	c2ftpA	Alignment	not modelled	97.2	16	PDB header: lyase Chain: A: PDB Molecule: hydroxymethylglutaryl-coa lyase; PDBTitle: crystal structure of hydroxymethylglutaryl-coa lyase from pseudomonas2 aeruginosa
65	c1djnB	Alignment	not modelled	97.2	19	PDB header: oxidoreductase Chain: B: PDB Molecule: trimethylamine dehydrogenase; PDBTitle: structural and biochemical characterization of recombinant wild type2 trimethylamine dehydrogenase from methylophilus methylotrophus (sp.3 w3a1)
66	c3thaB	Alignment	not modelled	97.1	15	PDB header: lyase Chain: B: PDB Molecule: tryptophan synthase alpha chain; PDBTitle: tryptophan synthase subunit alpha from campylobacter jejuni.
67	d1vzwa1	Alignment	not modelled	97.0	14	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Histidine biosynthesis enzymes
68	d1ojxa	Alignment	not modelled	97.0	19	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
69	c2rgfB	Alignment	not modelled	97.0	19	PDB header: lyase Chain: B: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from <i>hahella2 chejuensis</i> at 1.5a resolution
70	d1xi3a	Alignment	not modelled	97.0	21	Fold: TIM beta/alpha-barrel Superfamily: Thiamin phosphate synthase Family: Thiamin phosphate synthase
71	d1geqa	Alignment	not modelled	96.9	15	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
72	d1m3ua	Alignment	not modelled	96.9	16	Fold: TIM beta/alpha-barrel Superfamily: Phosphoenolpyruvate/pyruvate domain Family: Ketopantoate hydroxymethyltransferase PanB
73	c3daqB	Alignment	not modelled	96.9	17	PDB header: lyase Chain: B: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from methicillin-2 resistant staphylococcus aureus
74	d1oyaa	Alignment	not modelled	96.8	15	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
75	d1dqja1	Alignment	not modelled	96.8	18	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
76	d1wv2a	Alignment	not modelled	96.8	22	Fold: TIM beta/alpha-barrel Superfamily: ThiG-like Family: ThiG-like
77	c3jrkG	Alignment	not modelled	96.8	20	PDB header: lyase Chain: G: PDB Molecule: tagatose 1,6-diphosphate aldolase 2; PDBTitle: a putative tagatose 1,6-diphosphate aldolase from streptococcus2 pyogenes
78	c3lerA	Alignment	not modelled	96.7	19	PDB header: lyase Chain: A: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from campylobacter jejuni subsp. <i>jejuni</i> nctc 11168
79	d1s2wa	Alignment	not modelled	96.7	14	Fold: TIM beta/alpha-barrel Superfamily: Phosphoenolpyruvate/pyruvate domain Family: Phosphoenolpyruvate mutase/Isocitrate lyase-like
80	d1vvia	Alignment	not modelled	96.7	19	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases

81	d1o5ka	Alignment	not modelled	96.6	20	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
82	c3dx1B	Alignment	not modelled	96.6	18	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: putative aldolase; PDBTitle: crystal structure of the n-terminal domain of a putative aldolase (bvu_2661) from bacteroides vulgaris
83	c1ydoC	Alignment	not modelled	96.6	14	PDB header: lyase Chain: C: PDB Molecule: hmg-coa lyase; PDBTitle: crystal structure of the bacillus subtilis hmg-coa lyase, northeast2 structural genomics target sr181.
84	c3ajxA	Alignment	not modelled	96.6	15	PDB header: lyase Chain: A: PDB Molecule: 3-hexulose-6-phosphate synthase; PDBTitle: crystal structure of 3-hexulose-6-phosphate synthase
85	c2nx9B	Alignment	not modelled	96.5	22	PDB header: lyase Chain: B: PDB Molecule: oxaloacetate decarboxylase 2, subunit alpha; PDBTitle: crystal structure of the carboxyltransferase domain of the oxaloacetate decarboxylase na+ pump from vibrio cholerae
86	d1vc4a	Alignment	not modelled	96.5	16	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Tryptophan biosynthesis enzymes
87	d1icpa	Alignment	not modelled	96.4	14	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
88	c3inpA	Alignment	not modelled	96.4	15	PDB header: isomerase Chain: A: PDB Molecule: d-ribulose-phosphate 3-epimerase; PDBTitle: 2.05 angstrom resolution crystal structure of d-ribulose-phosphate 3-2 epimerase from francisella tularensis.
89	c1ydnA	Alignment	not modelled	96.4	17	PDB header: lyase Chain: A: PDB Molecule: hydroxymethylglutaryl-coa lyase; PDBTitle: crystal structure of the hmg-coa lyase from brucella melitensis, 2 northeast structural genomics target Ir35.
90	c3pueA	Alignment	not modelled	96.3	20	PDB header: lyase Chain: A: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of the complex of dihydrodipicolinate synthase from2 acinetobacter baumannii with lysine at 2.6a resolution
91	d2flia1	Alignment	not modelled	96.3	16	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: D-ribulose-5-phosphate 3-epimerase
92	c2zrvC	Alignment	not modelled	96.3	16	PDB header: isomerase Chain: C: PDB Molecule: isopentenyl-diphosphate delta-isomerase; PDBTitle: crystal structure of sulfolobus shibatae isopentenyl2 diphosphate isomerase in complex with reduced fmn.
93	d2mnra1	Alignment	not modelled	96.3	18	Fold: TIM beta/alpha-barrel Superfamily: Enolase C-terminal domain-like Family: D-glucarate dehydratase-like
94	c3ivuB	Alignment	not modelled	96.3	20	PDB header: transferase Chain: B: PDB Molecule: homocitrate synthase, mitochondrial; PDBTitle: homocitrate synthase lys4 bound to 2-og
95	c2vc6A	Alignment	not modelled	96.3	20	PDB header: lyase Chain: A: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: structure of mosa from s. meliloti with pyruvate bound
96	d1xkya1	Alignment	not modelled	96.3	17	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
97	d1mxsa	Alignment	not modelled	96.2	11	Fold: TIM beta/alpha-barrel Superfamily: Aldolase Family: Class I aldolase
98	d1w0ma	Alignment	not modelled	96.2	21	Fold: TIM beta/alpha-barrel Superfamily: Triosephosphate isomerase (TIM) Family: Triosephosphate isomerase (TIM)
99	d1rpxa	Alignment	not modelled	96.2	13	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: D-ribulose-5-phosphate 3-epimerase
100	d1qo2a	Alignment	not modelled	96.1	17	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: Histidine biosynthesis enzymes
101	c3noeA	Alignment	not modelled	96.1	21	PDB header: lyase Chain: A: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from pseudomonas2 aeruginosa
102	d1p0ka	Alignment	not modelled	96.1	20	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
103	c3f4wA	Alignment	not modelled	96.1	15	PDB header: synthase, lyase Chain: A: PDB Molecule: putative hexulose 6 phosphate synthase; PDBTitle: the 1.65a crystal structure of 3-hexulose-6-phosphate2 synthase from salmonella typhimurium
104	d1tza1	Alignment	not modelled	96.1	14	Fold: TIM beta/alpha-barrel Superfamily: Enolase C-terminal domain-like Family: D-glucarate dehydratase-like
105	c3gkaB	Alignment	not modelled	96.0	16	PDB header: oxidoreductase Chain: B: PDB Molecule: n-ethylmaleimide reductase; PDBTitle: crystal structure of n-ethylmaleimide reductase from burkholderia pseudomallei
106	d1nu5a1	Alignment	not modelled	96.0	19	Fold: TIM beta/alpha-barrel Superfamily: Enolase C-terminal domain-like Family: D-glucarate dehydratase-like
						PDB header: lyase

107	c3bi8A_		Alignment	not modelled	96.0	17	Chain: A: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: structure of dihydrodipicolinate synthase from clostridium2 botulinum
108	c2ps2A_		Alignment	not modelled	96.0	15	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: putative mandelate racemase/muconate lactonizing PDBTitle: crystal structure of putative mandelate racemase/muconate2 lactonizing enzyme from aspergillus oryzae
109	c1f9cA_		Alignment	not modelled	96.0	18	PDB header: isomerase Chain: A: PDB Molecule: protein (muconate cycloisomerase i); PDBTitle: crystal structure of mle d178n variant
110	d2a6na1		Alignment	not modelled	96.0	20	Fold: TIM beta/alpha-barrel Superfamily: Alkaline Family: Class I alkaline
111	d1jpma1		Alignment	not modelled	95.9	16	Fold: TIM beta/alpha-barrel Superfamily: Enolase C-terminal domain-like Family: D-glucarate dehydratase-like
112	c2ekcA_		Alignment	not modelled	95.9	16	PDB header: lyase Chain: A: PDB Molecule: tryptophan synthase alpha chain; PDBTitle: structural study of project id aq_1548 from aquifex aeolicus v5
113	d1tqja_		Alignment	not modelled	95.8	20	Fold: TIM beta/alpha-barrel Superfamily: Ribulose-phosphate binding barrel Family: D-ribulose-5-phosphate 3-epimerase
114	d1tb3a1		Alignment	not modelled	95.8	14	Fold: TIM beta/alpha-barrel Superfamily: FMN-linked oxidoreductases Family: FMN-linked oxidoreductases
115	c3d0cB_		Alignment	not modelled	95.8	16	PDB header: lyase Chain: B: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from oceanobacillus iheyensis at 1.9 a resolution PDB header: oxidoreductase
116	c2a7nA_		Alignment	not modelled	95.7	14	Chain: A: PDB Molecule: (+)-mandelate dehydrogenase; PDBTitle: crystal structure of the g81a mutant of the active chimera of (s)-2 mandelate dehydrogenase
117	c2ehhE_		Alignment	not modelled	95.7	17	PDB header: lyase Chain: E: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from2 aquifex aeolicus
118	c2gdqB_		Alignment	not modelled	95.7	16	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: ytf; PDBTitle: crystal structure of mandelate racemase/muconate lactonizing enzyme2 from bacillus subtilis at 1.8 a resolution
119	c3e96B_		Alignment	not modelled	95.6	20	PDB header: lyase Chain: B: PDB Molecule: dihydrodipicolinate synthase; PDBTitle: crystal structure of dihydrodipicolinate synthase from2 bacillus clausii
120	c3ewbX_		Alignment	not modelled	95.6	16	PDB header: transferase Chain: X: PDB Molecule: 2-isopropylmalate synthase; PDBTitle: crystal structure of n-terminal domain of putative 2-2 isopropylmalate synthase from listeria monocytogenes