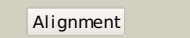
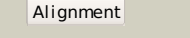
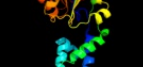
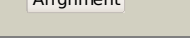
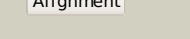


Phyre²

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Description	P77366
Date	Thu Jan 5 12:28:14 GMT 2012
Unique Job ID	10c0fbdb25e5042e5

Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c3dv9A_			100.0	19	PDB header: isomerase Chain: A; PDB Molecule: beta-phosphoglucomutase; PDBTitle: putative beta-phosphoglucomutase from bacteroides vulgatus.
2	c3iruA_			100.0	19	PDB header: structural genomics, unknown function Chain: A; PDB Molecule: phoshonoacetaldehyde hydrolase like protein; PDBTitle: crystal structure of phoshonoacetaldehyde hydrolase like protein from2 oleispira antarctica
3	d2hsza1			100.0	17	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
4	c3d6jA_			100.0	19	PDB header: hydrolase Chain: A; PDB Molecule: putative haloacid dehalogenase-like hydrolase; PDBTitle: crystal structure of putative haloacid dehalogenase-like hydrolase2 from bacteroides fragilis
5	c3nasA_			99.9	48	PDB header: isomerase Chain: A; PDB Molecule: beta-phosphoglucomutase; PDBTitle: the crystal structure of beta-phosphoglucomutase from bacillus2 subtilis
6	d1swva_			99.9	17	Fold: HAD-like Superfamily: HAD-like Family: Phosphonoacetaldehyde hydrolase-like
7	c2hi0B_			99.9	18	PDB header: hydrolase Chain: B; PDB Molecule: putative phosphoglycolate phosphatase; PDBTitle: crystal structure of putative phosphoglycolate phosphatase2 (yp_619066.1) from lactobacillus delbrueckii subsp. bulgaricus atcc3 baa-365 at 1.51 a resolution
8	d1o08a_			99.9	38	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
9	d2fdra1			99.9	20	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
10	c3s6jC_			99.9	23	PDB header: hydrolase Chain: C; PDB Molecule: hydrolase, haloacid dehalogenase-like family; PDBTitle: the crystal structure of a hydrolase from pseudomonas syringae
11	c3mc1A_			99.9	16	PDB header: hydrolase Chain: A; PDB Molecule: predicted phosphatase, had family; PDBTitle: crystal structure of a predicted phosphatase from2 clostridium acetobutylicum

12	d1te2a_			99.9	25	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
13	d2ah5a1			99.9	14	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
14	c2qltA_			99.9	20	PDB header: hydrolase Chain: A: PDB Molecule: (dl)-glycerol-3-phosphatase 1; PDBTitle: crystal structure of an isoform of dl-glycerol-3-phosphatase, rhr2p,2 from <i>saccharomyces cerevisiae</i>
15	c2pibA_			99.9	25	PDB header: hydrolase Chain: A: PDB Molecule: phosphorylated carbohydrates phosphatase tm_1254; PDBTitle: crystal structure of putative beta-phosphoglucomutase from2 <i>thermotoga maritima</i>
16	c3qnmA_			99.9	15	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: haloacid dehalogenase-like hydrolase; PDBTitle: haloalkane dehalogenase family member from <i>bacteroides2 thetaiotaomicron</i> of unknown function
17	c3e58A_			99.9	26	PDB header: isomerase Chain: A: PDB Molecule: putative beta-phosphoglucomutase; PDBTitle: crystal structure of putative beta-phosphoglucomutase from2 <i>streptococcus thermophilus</i>
18	d2hdoa1			99.9	15	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
19	d2go7a1			99.9	17	Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
20	c3sd7A_			99.9	14	PDB header: hydrolase Chain: A: PDB Molecule: putative phosphatase; PDBTitle: 1.7 angstrom resolution crystal structure of putative phosphatase from2 <i>clostridium difficile</i>
21	c3l5kA_		not modelled	99.9	25	PDB header: hydrolase Chain: A: PDB Molecule: haloacid dehalogenase-like hydrolase domain- PDBTitle: the crystal structure of human haloacid dehalogenase-like2 hydrolase domain containing 1a (<i>hdhd1a</i>)
22	c2yy6B_		not modelled	99.9	20	PDB header: hydrolase Chain: B: PDB Molecule: phosphoglycolate phosphatase; PDBTitle: crystal structure of the phosphoglycolate phosphatase from <i>aquifex2 aeolicus vF5</i>
23	d2fi1a1		not modelled	99.9	23	Fold: HAD-like Superfamily: HAD-like Family: Phosphonoacetaldehyde hydrolase-like
24	c3nuqA_		not modelled	99.9	16	PDB header: hydrolase Chain: A: PDB Molecule: putative nucleotide phosphatase; PDBTitle: structure of a putative nucleotide phosphatase from <i>saccharomyces2 cerevisiae</i>
25	c2om6A_		not modelled	99.9	16	PDB header: hydrolase Chain: A: PDB Molecule: probable phosphoserine phosphatase; PDBTitle: hypothetical protein (probable phosphoserine phosph (ph0253) from2 <i>pyrococcus horikoshii ot3</i>
26	c3ddhA_		not modelled	99.9	11	PDB header: hydrolase Chain: A: PDB Molecule: putative haloacid dehalogenase-like family hydrolase; PDBTitle: the structure of a putative haloacid dehalogenase-like family2 hydrolase from <i>bacteroides thetaiotaomicron vpi-5482</i>
27	d1qq5a_		not modelled	99.9	15	Fold: HAD-like Superfamily: HAD-like Family: HAD-related
28	c3m9IA_		not modelled	99.9	18	PDB header: hydrolase Chain: A: PDB Molecule: hydrolase, haloacid dehalogenase-like family;

					PDBTitle: crystal structure of probable had family hydrolase from2 pseudomonas fluorescens pf-5
29	d2hcfa1	Alignment	not modelled	99.9	15 Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
30	d1zrna_	Alignment	not modelled	99.9	14 Fold: HAD-like Superfamily: HAD-like Family: HAD-related
31	c2no5B_	Alignment	not modelled	99.9	20 PDB header: hydrolase Chain: B: PDB Molecule: (s)-2-haloacid dehalogenase iva; PDBTitle: crystal structure analysis of a dehalogenase with intermediate complex
32	c3k1zA_	Alignment	not modelled	99.9	19 PDB header: hydrolase Chain: A: PDB Molecule: haloacid dehalogenase-like hydrolase domain-containing PDBTitle: crystal structure of human haloacid dehalogenase-like hydrolase domain2 containing 3 (hdh3)
33	d1x42a1	Alignment	not modelled	99.9	14 Fold: HAD-like Superfamily: HAD-like Family: HAD-related
34	d1zs9a1	Alignment	not modelled	99.9	10 Fold: HAD-like Superfamily: HAD-like Family: Enolase-phosphatase E1
35	c3cnhA_	Alignment	not modelled	99.9	16 PDB header: hydrolase Chain: A: PDB Molecule: hydrolase family protein; PDBTitle: crystal structure of predicted hydrolase of haloacid dehalogenase-like2 superfamily (np_295428.1) from deinococcus radiodurans at 1.66 a3 resolution
36	c2hoqA_	Alignment	not modelled	99.9	19 PDB header: hydrolase Chain: A: PDB Molecule: putative had-hydrolase ph1655; PDBTitle: crystal structure of the probable haloacid dehalogenase (ph1655) from2 pyrococcus horikoshii ot3
37	d2gfh1	Alignment	not modelled	99.9	16 Fold: HAD-like Superfamily: HAD-like Family: beta-Phosphoglucomutase-like
38	c3ed5A_	Alignment	not modelled	99.9	15 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: yfnb; PDBTitle: the crystal structure of yfnb from bacillus subtilis subsp. subtilis2 str. 168
39	c2pkeA_	Alignment	not modelled	99.9	14 PDB header: hydrolase Chain: A: PDB Molecule: haloacid delahogenase-like family hydrolase; PDBTitle: crystal structure of haloacid delahogenase-like family hydrolase2 (np_639141.1) from xanthomonas campestris at 1.81 a resolution
40	c2g80C_	Alignment	not modelled	99.9	9 PDB header: hydrolase Chain: C: PDB Molecule: protein utr4; PDBTitle: crystal structure of utr4 protein (unknown transcript 4 protein)2 (yel038w) from saccharomyces cerevisiae at 2.28 a resolution
41	d2g80a1	Alignment	not modelled	99.9	8 Fold: HAD-like Superfamily: HAD-like Family: Enolase-phosphatase E1
42	d1zd3a1	Alignment	not modelled	99.9	16 Fold: HAD-like Superfamily: HAD-like Family: YihX-like
43	c2w11B_	Alignment	not modelled	99.9	12 PDB header: hydrolase Chain: B: PDB Molecule: 2-haloalkanoic acid dehalogenase; PDBTitle: structure of the l-2-haloacid dehalogenase from sulfolobus2 tokodaii
44	c2ho4A_	Alignment	not modelled	99.8	15 PDB header: hydrolase Chain: A: PDB Molecule: haloacid dehalogenase-like hydrolase domain PDBTitle: crystal structure of protein from mouse mm.236127
45	c2i6xA_	Alignment	not modelled	99.8	17 PDB header: hydrolase Chain: A: PDB Molecule: hydrolase, haloacid dehalogenase-like family; PDBTitle: the structure of a predicted had-like family hydrolase from2 porphyromonas gingivalis.
46	d2c4na1	Alignment	not modelled	99.8	13 Fold: HAD-like Superfamily: HAD-like Family: NagD-like
47	c3l8hC_	Alignment	not modelled	99.8	20 PDB header: hydrolase Chain: C: PDB Molecule: putative haloacid dehalogenase-like hydrolase; PDBTitle: crystal structure of d,d-heptose 1,7-bisphosphate phosphatase from b.2 bronchiseptica complexed with magnesium and phosphate
48	d2gmwa1	Alignment	not modelled	99.8	17 Fold: HAD-like Superfamily: HAD-like Family: Histidinol phosphatase-like
49	c3kzxA_	Alignment	not modelled	99.8	19 PDB header: hydrolase Chain: A: PDB Molecule: had-superfamily hydrolase, subfamily ia, variant 1; PDBTitle: crystal structure of a had-superfamily hydrolase from ehrlichia2 chaffeensis at 1.9a resolution
50	c3pdwA_	Alignment	not modelled	99.8	14 PDB header: hydrolase Chain: A: PDB Molecule: uncharacterized hydrolase yutf; PDBTitle: crystal structure of putative p-nitrophenyl phosphatase from bacillus2 subtilis
51	c3esqA_	Alignment	not modelled	99.8	17 PDB header: hydrolase Chain: A: PDB Molecule: d,d-heptose 1,7-bisphosphate phosphatase; PDBTitle: crystal structure of calcium-bound d,d-heptose 1,7-2 bisphosphate phosphatase from e. coli
					PDB header: hydrolase

52	c3qgmC	Alignment	not modelled	99.8	14	Chain: C: PDB Molecule: p-nitrophenyl phosphatase (pho2); PDBTitle: p-nitrophenyl phosphatase from archaeoglobus fulgidus
53	d2o2xa1	Alignment	not modelled	99.8	21	Fold: HAD-like Superfamily: HAD-like Family: Histidinol phosphatase-like
54	d1wvia	Alignment	not modelled	99.8	15	Fold: HAD-like Superfamily: HAD-like Family: NagD-like
55	d1ydfa1	Alignment	not modelled	99.8	15	Fold: HAD-like Superfamily: HAD-like Family: NagD-like
56	d2b0ca1	Alignment	not modelled	99.8	18	Fold: HAD-like Superfamily: HAD-like Family: YihX-like
57	c2p11A	Alignment	not modelled	99.8	19	PDB header: hydrolase Chain: A: PDB Molecule: hypothetical protein; PDBTitle: crystal structure of a putative haloacid dehalogenase-like hydrolase2 (bxe_b1342) from burkholderia xenovorans lb400 at 2.20 a resolution
58	c1cr6A	Alignment	not modelled	99.8	18	PDB header: hydrolase Chain: A: PDB Molecule: epoxide hydrolase; PDBTitle: crystal structure of murine soluble epoxide hydrolase2 complexed with cpu inhibitor
59	c2x4dB	Alignment	not modelled	99.8	12	PDB header: hydrolase Chain: B: PDB Molecule: phospholysine phosphohistidine inorganic pyrophosphate PDBTitle: crystal structure of human phospholysine phosphohistidine2 inorganic pyrophosphate phosphatase lhpp
60	c2cftA	Alignment	not modelled	99.8	17	PDB header: phosphatase Chain: A: PDB Molecule: pyridoxal phosphate phosphatase; PDBTitle: crystal structure of human pyridoxal 5'-phosphate2 phosphatase with its substrate
61	d1ys9a1	Alignment	not modelled	99.8	13	Fold: HAD-like Superfamily: HAD-like Family: NagD-like
62	d1vjra	Alignment	not modelled	99.8	15	Fold: HAD-like Superfamily: HAD-like Family: NagD-like
63	d1yv9a1	Alignment	not modelled	99.8	18	Fold: HAD-like Superfamily: HAD-like Family: NagD-like
64	d1u7pa	Alignment	not modelled	99.8	18	Fold: HAD-like Superfamily: HAD-like Family: Magnesium-dependent phosphatase-1, Mdp1
65	c3kd3A	Alignment	not modelled	99.8	11	PDB header: unknown function Chain: A: PDB Molecule: phosphoserine phosphohydrolase-like protein; PDBTitle: crystal structure of a phosphoserine phosphohydrolase-like protein2 from francisella tularensis subsp. tularensis schu s4
66	d1cr6a1	Alignment	not modelled	99.8	18	Fold: HAD-like Superfamily: HAD-like Family: YihX-like
67	c2zg6A	Alignment	not modelled	99.7	14	PDB header: hydrolase Chain: A: PDB Molecule: putative uncharacterized protein st2620; PDBTitle: crystal structure of hypothetical protein; probable 2-2 haloalkanoic acid dehalogenase from sulfobolbus tokodaii
68	c1zjJA	Alignment	not modelled	99.7	13	PDB header: hydrolase Chain: A: PDB Molecule: hypothetical protein ph1952; PDBTitle: crystal structure of hypothetical protein ph1952 from pyrococcus2 horikoshii ot3
69	c3m1yA	Alignment	not modelled	99.7	17	PDB header: hydrolase Chain: A: PDB Molecule: phosphoserine phosphatase (serb); PDBTitle: crystal structure of a phosphoserine phosphatase (serb) from2 helicobacter pylori
70	d2fpwa1	Alignment	not modelled	99.7	14	Fold: HAD-like Superfamily: HAD-like Family: Histidinol phosphatase-like
71	d1qyia	Alignment	not modelled	99.6	20	Fold: HAD-like Superfamily: HAD-like Family: Hypothetical protein MW1667 (SA1546)
72	d1j97a	Alignment	not modelled	99.6	15	Fold: HAD-like Superfamily: HAD-like Family: Phosphoserine phosphatase
73	d1nnla	Alignment	not modelled	99.6	16	Fold: HAD-like Superfamily: HAD-like Family: Phosphoserine phosphatase
74	c2odaB	Alignment	not modelled	99.6	19	PDB header: protein binding Chain: B: PDB Molecule: hypothetical protein pspto_2114; PDBTitle: crystal structure of pspto_2114
75	c3ib6B	Alignment	not modelled	99.6	15	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of an uncharacterized protein from listeria2 monocytogenes serotype 4b
76	d2feaa1	Alignment	not modelled	99.6	13	Fold: HAD-like Superfamily: HAD-like Family: MtnX-like
77	c2hx1D	Alignment	not modelled	99.6	14	PDB header: hydrolase Chain: D: PDB Molecule: predicted sugar phosphatases of the had PDBTitle: crystal structure of possible sugar phosphatase, had2 superfamily (zp_003110.1) from cytophaga hutchinsonii3 atcc 33406 at 2.10 a resolution
						Fold: HAD-like

78	d1rkua_	Alignment	not modelled	99.5	13	Superfamily: HAD-like Family: Homoserine kinase ThrH
79	c2pr7A_	Alignment	not modelled	99.5	17	PDB header: hydrolase Chain: A: PDB Molecule: haloacid dehalogenase/epoxide hydrolase family; PDBTitle: crystal structure of uncharacterized protein (np_599989.1) from2 corynebacterium glutamicum atcc 13032 kitasato at 1.44 a resolution
80	c3p96A_	Alignment	not modelled	99.2	16	PDB header: hydrolase Chain: A: PDB Molecule: phosphoserine phosphatase serb; PDBTitle: crystal structure of phosphoserine phosphatase serb from mycobacterium avium, native form
81	c2i7dB_	Alignment	not modelled	99.2	11	PDB header: hydrolase Chain: B: PDB Molecule: 5'(3')-deoxyribonucleotidase, cytosolic type; PDBTitle: structure of human cytosolic deoxyribonucleotidase in2 complex with deoxyuridine, alfa4 and mg2+
82	c3mn1B_	Alignment	not modelled	99.1	17	PDB header: hydrolase Chain: B: PDB Molecule: probable yrbi family phosphatase; PDBTitle: crystal structure of probable yrbi family phosphatase from pseudomonas2 syringae pv. phaseolica 1448a
83	d1q92a_	Alignment	not modelled	99.1	14	Fold: HAD-like Superfamily: HAD-like Family: 5'(3')-deoxyribonucleotidase (dNT-2)
84	c3fvvA_	Alignment	not modelled	99.1	11	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: the crystal structure of the protein with unknown function from2 bordetella pertussis tohama i
85	c3e8mD_	Alignment	not modelled	99.1	20	PDB header: transferase Chain: D: PDB Molecule: acylneuraminate cytidyltransferase; PDBTitle: structure-function analysis of 2-keto-3-deoxy-d-glycero-d-galacto-2 nononate-9-phosphate (kdn) phosphatase defines a new clad within the3 type c0 had subfamily
86	d1k1ea_	Alignment	not modelled	99.1	19	Fold: HAD-like Superfamily: HAD-like Family: Probable phosphatase Yrbl
87	c2r8zC_	Alignment	not modelled	99.1	20	PDB header: hydrolase Chain: C: PDB Molecule: 3-deoxy-d-manno-octulonate 8-phosphate phosphatase; PDBTitle: crystal structure of yrbi phosphatase from escherichia coli in complex2 with a phosphate and a calcium ion
88	c3n07B_	Alignment	not modelled	99.0	15	PDB header: hydrolase Chain: B: PDB Molecule: 3-deoxy-d-manno-octulonate 8-phosphate phosphatase; PDBTitle: structure of putative 3-deoxy-d-manno-octulonate 8-phosphate2 phosphatase from vibrio cholerae
89	d1wr8a_	Alignment	not modelled	99.0	13	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
90	c3n1uA_	Alignment	not modelled	99.0	22	PDB header: hydrolase Chain: A: PDB Molecule: hydrolase, had superfamily, subfamily iii a; PDBTitle: structure of putative had superfamily (subfamily iii a) hydrolase from2 legionella pneumophila
91	d1l6ra_	Alignment	not modelled	99.0	8	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
92	c3ewiB_	Alignment	not modelled	99.0	16	PDB header: transferase Chain: B: PDB Molecule: n-acylneuraminate cytidyltransferase; PDBTitle: structural analysis of the c-terminal domain of murine cmp-2 sialic acid synthetase
93	c2p9jH_	Alignment	not modelled	99.0	22	PDB header: structural genomics, unknown function Chain: H: PDB Molecule: hypothetical protein aq2171; PDBTitle: crystal structure of aq2171 from aquifex aeolicus
94	c3mmzA_	Alignment	not modelled	98.9	18	PDB header: hydrolase Chain: A: PDB Molecule: putative had family hydrolase; PDBTitle: crystal structure of putative had family hydrolase from streptomyces2 avermitilis ma-4680
95	c3n28A_	Alignment	not modelled	98.9	17	PDB header: hydrolase Chain: A: PDB Molecule: phosphoserine phosphatase; PDBTitle: crystal structure of probable phosphoserine phosphatase from vibrio2 cholerae, unliganded form
96	c3kc2A_	Alignment	not modelled	98.8	19	PDB header: hydrolase Chain: A: PDB Molecule: uncharacterized protein ykr070w; PDBTitle: crystal structure of mitochondrial had-like phosphatase from2 saccharomyces cerevisiae
97	c3zvmA_	Alignment	not modelled	98.7	19	PDB header: hydrolase/transferase/dna Chain: A: PDB Molecule: bifunctional polynucleotide phosphatase/kinase; PDBTitle: the structural basis for substrate recognition by mammalian2 polynucleotide kinase 3' phosphatase
98	c2qyhD_	Alignment	not modelled	98.6	14	PDB header: structural genomics, unknown function Chain: D: PDB Molecule: hypothetical conserved protein, gk1056; PDBTitle: crystal structure of the hypothetical protein (gk1056) from2 geobacillus kaustophilus hta426
99	d1s2oa1	Alignment	not modelled	98.6	12	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
100	c3fzqA_	Alignment	not modelled	98.6	11	PDB header: hydrolase Chain: A: PDB Molecule: putative hydrolase; PDBTitle: crystal structure of putative haloacid dehalogenase-like hydrolase2 (yp_001086940.1) from clostridium difficile 630 at 2.10 a resolution
101	d1rkqa_	Alignment	not modelled	98.5	11	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
						PDB header: hydrolase Chain: A: PDB Molecule: putative uncharacterized protein

102	c3l7yA	Alignment	not modelled	98.5	11	smu.1108c; PDBTitle: the crystal structure of smu.1108c from streptococcus mutans ua159
103	d1nf2a	Alignment	not modelled	98.5	9	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
104	c3niwA	Alignment	not modelled	98.5	15	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: haloacid dehalogenase-like hydrolase; PDBTitle: crystal structure of a haloacid dehalogenase-like hydrolase from2 bacteroides thetaiotaomicron
105	d2b30a1	Alignment	not modelled	98.5	15	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
106	c3da0B	Alignment	not modelled	98.4	9	PDB header: hydrolase Chain: B: PDB Molecule: putative phosphatase; PDBTitle: crystal structure of a putative phosphatase (eubrec_1417) from2 eubacterium rectale at 1.80 a resolution
107	c3r4cA	Alignment	not modelled	98.3	12	PDB header: hydrolase Chain: A: PDB Molecule: hydrolase, haloacid dehalogenase-like hydrolase; PDBTitle: divergence of structure and function among phosphatases of the2 haloalkanoate (had) enzyme superfamily: analysis of bt1666 from3 bacteroides thetaiotaomicron
108	d2rbka1	Alignment	not modelled	98.3	13	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
109	c3dnPA	Alignment	not modelled	98.3	15	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: stress response protein yhax; PDBTitle: crystal structure of stress response protein yhax from bacillus2 subtilis
110	d1z5ga1	Alignment	not modelled	98.3	14	Fold: HAD-like Superfamily: HAD-like Family: Class B acid phosphatase, AphA
111	d2vkqa1	Alignment	not modelled	98.2	10	Fold: HAD-like Superfamily: HAD-like Family: Pyrimidine 5'-nucleotidase (UMPH-1)
112	c3pgvB	Alignment	not modelled	98.2	25	PDB header: hydrolase Chain: B: PDB Molecule: haloacid dehalogenase-like hydrolase; PDBTitle: crystal structure of a haloacid dehalogenase-like hydrolase2 (kpn_04322) from klebsiella pneumoniae subsp. pneumoniae mgh 78578 at3 2.39 a resolution
113	d1nrwa	Alignment	not modelled	98.2	12	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
114	d1rlma	Alignment	not modelled	98.2	13	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
115	d1ltqa1	Alignment	not modelled	98.1	17	Fold: HAD-like Superfamily: HAD-like Family: phosphatase domain of polynucleotide kinase
116	d1wzca1	Alignment	not modelled	98.1	9	Fold: HAD-like Superfamily: HAD-like Family: Predicted hydrolases Cof
117	c3gygA	Alignment	not modelled	98.0	12	PDB header: hydrolase Chain: A: PDB Molecule: ntd biosynthesis operon putative hydrolase ntdb; PDBTitle: crystal structure of yhjk (haloacid dehalogenase-like hydrolase2 protein) from bacillus subtilis
118	d1yj5a1	Alignment	not modelled	98.0	17	Fold: HAD-like Superfamily: HAD-like Family: phosphatase domain of polynucleotide kinase
119	d2bdua1	Alignment	not modelled	97.9	11	Fold: HAD-like Superfamily: HAD-like Family: Pyrimidine 5'-nucleotidase (UMPH-1)
120	d2b82a1	Alignment	not modelled	97.9	16	Fold: HAD-like Superfamily: HAD-like Family: Class B acid phosphatase, AphA