



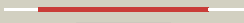





















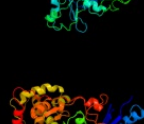



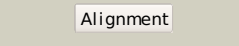
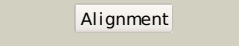
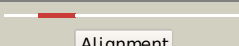
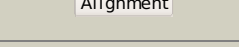
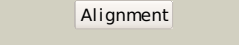


#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c3gbtA_	 Alignment		100.0	21	PDB header: transferase Chain: A: PDB Molecule: gluconate kinase; PDBTitle: crystal structure of gluconate kinase from lactobacillus acidophilus
2	c3jvpA_	 Alignment		100.0	22	PDB header: transferase Chain: A: PDB Molecule: ribulokinase; PDBTitle: crystal structure of ribulokinase from bacillus halodurans
3	c3ifrB_	 Alignment		100.0	24	PDB header: transferase Chain: B: PDB Molecule: carbohydrate kinase, fggy; PDBTitle: the crystal structure of xylulose kinase from rhodospirillum rubrum
4	c3gg4B_	 Alignment		100.0	18	PDB header: transferase Chain: B: PDB Molecule: glycerol kinase; PDBTitle: the crystal structure of glycerol kinase from yersinia2 pseudotuberculosis
5	c3flcX_	 Alignment		100.0	21	PDB header: transferase Chain: X: PDB Molecule: glycerol kinase; PDBTitle: crystal structure of the his-tagged h232r mutant of glycerol kinase2 from enterococcus casseliflavus with glycerol
6	c2d4wA_	 Alignment		100.0	21	PDB header: transferase Chain: A: PDB Molecule: glycerol kinase; PDBTitle: crystal structure of glycerol kinase from cellulomonas sp.2 nt3060
7	c2nlxA_	 Alignment		100.0	22	PDB header: transferase Chain: A: PDB Molecule: xylulose kinase; PDBTitle: crystal structure of the apo e. coli xylulose kinase
8	c3hz6A_	 Alignment		100.0	21	PDB header: transferase Chain: A: PDB Molecule: xylulokinase; PDBTitle: crystal structure of xylulokinase from chromobacterium violaceum
9	c2zf5O_	 Alignment		100.0	22	PDB header: transferase Chain: O: PDB Molecule: glycerol kinase; PDBTitle: crystal structure of highly thermostable glycerol kinase from a2 hyperthermophilic archaeon
10	c3g25B_	 Alignment		100.0	17	PDB header: transferase Chain: B: PDB Molecule: glycerol kinase; PDBTitle: 1.9 angstrom crystal structure of glycerol kinase (glpk) from2 staphylococcus aureus in complex with glycerol.
11	c2dnpB_	 Alignment		100.0	21	PDB header: transferase Chain: B: PDB Molecule: glycerol kinase; PDBTitle: crystal structure of the glycerol kinase from thermus2 thermophilus hb8

12	c2w40C_	Alignment		100.0	20	PDB header: transferase Chain: C: PDB Molecule: glycerol kinase, putative; PDBTitle: crystal structure of plasmodium falciparum glycerol kinase2 with bound glycerol
13	c1glbG_	Alignment		100.0	19	PDB header: phosphotransferase Chain: G: PDB Molecule: glycerol kinase; PDBTitle: structure of the regulatory complex of escherichia coli iiiglc with2 glycerol kinase
14	c3ezwD_	Alignment		100.0	20	PDB header: transferase Chain: D: PDB Molecule: glycerol kinase; PDBTitle: crystal structure of a hyperactive escherichia coli glycerol kinase2 mutant gly230 --> asp obtained using microfluidic crystallization3 devices
15	c1xupO_	Alignment		100.0	21	PDB header: transferase Chain: O: PDB Molecule: glycerol kinase; PDBTitle: enterococcus casseliflavus glycerol kinase complexed with glycerol
16	c2cgkB_	Alignment		100.0	16	PDB header: transferase Chain: B: PDB Molecule: l-rhamnulose kinase; PDBTitle: crystal structure of l-rhamnulose kinase from escherichia2 coli in an open uncomplexed conformation.
17	c3i8bA_	Alignment		100.0	18	PDB header: transferase Chain: A: PDB Molecule: xylulose kinase; PDBTitle: the crystal structure of xylulose kinase from2 bifidobacterium adolescentis
18	c3h6eB_	Alignment		100.0	17	PDB header: transferase Chain: B: PDB Molecule: carbohydrate kinase, fggy; PDBTitle: the crystal structure of a carbohydrate kinase from novosphingobium2 aromaticivorans
19	d2p3ra1	Alignment		100.0	22	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glycerol kinase
20	d1r59o1	Alignment		100.0	22	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glycerol kinase
21	d2p3ra2	Alignment	not modelled	100.0	18	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glycerol kinase
22	d1r59o2	Alignment	not modelled	100.0	21	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glycerol kinase
23	d1huxa_	Alignment	not modelled	99.4	16	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: BadF/BadG/BcrA/BcrD-like
24	c3h1qB_	Alignment	not modelled	99.4	17	PDB header: structural protein Chain: B: PDB Molecule: ethanolamine utilization protein eutj; PDBTitle: crystal structure of ethanolamine utilization protein eutj from2 carboxydotherrhus hydrogenoformans
25	c2e2pA_	Alignment	not modelled	99.2	18	PDB header: transferase Chain: A: PDB Molecule: hexokinase; PDBTitle: crystal structure of sulfolobus tokodaii hexokinase in2 complex with adp
26	c3eo3B_	Alignment	not modelled	99.0	20	PDB header: isomerase, transferase Chain: B: PDB Molecule: bifunctional udp-n-acetylglucosamine 2-epimerase/n- PDBTitle: crystal structure of the n-acetylmannosamine kinase domain of human2 gne protein
27	d2ews1	Alignment	not modelled	98.9	16	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Fumble-like
28	c2ch5D_	Alignment	not modelled	98.9	15	PDB header: transferase Chain: D: PDB Molecule: nagk protein; PDBTitle: crystal structure of human n-acetylglucosamine kinase in2 complex with n-acetylglucosamine

29	d1zc6a1	Alignment	not modelled	98.8	18	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: BadF/BadG/BcrA/BcrD-like
30	c3enoB	Alignment	not modelled	98.8	12	PDB header: hydrolase/unknown function Chain: B: PDB Molecule: putative o-sialoglycoprotein endopeptidase; PDBTitle: crystal structure of pyrococcus furiosus pcc1 in complex2 with thermoplasma acidophilum kae1
31	c2ivoC	Alignment	not modelled	98.8	14	PDB header: hydrolase Chain: C: PDB Molecule: up1; PDBTitle: structure of up1 protein
32	c1z05A	Alignment	not modelled	98.8	14	PDB header: transcription Chain: A: PDB Molecule: transcriptional regulator, rok family; PDBTitle: crystal structure of the rok family transcriptional regulator, homolog2 of e.coli mlc protein.
33	c2qm1D	Alignment	not modelled	98.8	24	PDB header: transferase Chain: D: PDB Molecule: glucokinase; PDBTitle: crystal structure of glucokinase from enterococcus faecalis
34	c3r8eA	Alignment	not modelled	98.7	21	PDB header: transferase Chain: A: PDB Molecule: hypothetical sugar kinase; PDBTitle: crystal structure of a hypothetical sugar kinase (chu_1875) from2 cytophaga hutchinsonii atcc 33406 at 1.65 a resolution
35	c1e4gT	Alignment	not modelled	98.7	15	PDB header: bacterial cell division Chain: T: PDB Molecule: cell division protein ftsa; PDBTitle: ftsa (atp-bound form) from thermotoga maritima
36	c1z6rC	Alignment	not modelled	98.7	15	PDB header: transcription Chain: C: PDB Molecule: mlc protein; PDBTitle: crystal structure of mlc from escherichia coli
37	c3cqyA	Alignment	not modelled	98.5	15	PDB header: transferase Chain: A: PDB Molecule: anhydro-n-acetylmuramic acid kinase; PDBTitle: crystal structure of a functionally unknown protein (so_1313) from2 shewanella oneidensis mr-1
38	d1z05a3	Alignment	not modelled	98.4	15	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
39	d2ch5a2	Alignment	not modelled	98.3	17	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: BadF/BadG/BcrA/BcrD-like
40	c1zc6A	Alignment	not modelled	98.3	17	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: probable n-acetylglucosamine kinase; PDBTitle: crystal structure of putative n-acetylglucosamine kinase from2 chromobacterium violaceum. northeast structural genomics target3 cvr23.
41	d2hoea3	Alignment	not modelled	98.3	8	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
42	c3en9B	Alignment	not modelled	98.2	18	PDB header: hydrolase Chain: B: PDB Molecule: o-sialoglycoprotein endopeptidase/protein kinase; PDBTitle: structure of the methanococcus jannaschii kae1-bud32 fusion2 protein
43	c1dkgD	Alignment	not modelled	98.0	17	PDB header: complex (hsp24/hsp70) Chain: D: PDB Molecule: molecular chaperone dnak; PDBTitle: crystal structure of the nucleotide exchange factor grpe2 bound to the atpase domain of the molecular chaperone dnak
44	d1z6ra2	Alignment	not modelled	98.0	16	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
45	c2ap1A	Alignment	not modelled	98.0	16	PDB header: transferase Chain: A: PDB Molecule: putative regulator protein; PDBTitle: crystal structure of the putative regulatory protein
46	c1ig8A	Alignment	not modelled	97.9	15	PDB header: transferase Chain: A: PDB Molecule: hexokinase pii; PDBTitle: crystal structure of yeast hexokinase pii with the correct2 amino acid sequence
47	c3d2fC	Alignment	not modelled	97.9	14	PDB header: chaperone Chain: C: PDB Molecule: heat shock protein homolog sse1; PDBTitle: crystal structure of a complex of sse1p and hsp70
48	c2v7zA	Alignment	not modelled	97.8	12	PDB header: chaperone Chain: A: PDB Molecule: heat shock cognate 71 kda protein; PDBTitle: crystal structure of the 70-kda heat shock cognate protein2 from rattus norvegicus in post-atp hydrolysis state
49	c1v4sA	Alignment	not modelled	97.8	15	PDB header: transferase Chain: A: PDB Molecule: glucokinase isoform 2; PDBTitle: crystal structure of human glucokinase
50	d1dkgd2	Alignment	not modelled	97.8	17	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
51	d1woqa1	Alignment	not modelled	97.8	14	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
52	c2v7yA	Alignment	not modelled	97.8	15	PDB header: chaperone Chain: A: PDB Molecule: chaperone protein dnak; PDBTitle: crystal structure of the molecular chaperone dnak from2 geobacillus kaustophilus hta426 in post-atp hydrolysis3 state
53	d2ap1a2	Alignment	not modelled	97.7	19	Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
54	c1hpmA	Alignment	not modelled	97.7	12	PDB header: hydrolase (acting on acid anhydrides) Chain: A: PDB Molecule: 44k atpase fragment (n-terminal) of 7o kd heat- PDBTitle: how potassium affects the activity of the molecular2

						chaperone hsc70. ii. potassium binds specifically in the3 atpase active site
55	c3htvA		Alignment	not modelled	97.7	21 PDB header: transferase Chain: A: PDB Molecule: d-allose kinase; PDBTitle: crystal structure of d-allose kinase (np_418508.1) from escherichia2 coli k12 at 1.95 a resolution
56	dljcea2		Alignment	not modelled	97.6	15 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
57	dlq18a1		Alignment	not modelled	97.6	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glucokinase
58	dlsz2a1		Alignment	not modelled	97.6	15 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Glucokinase
59	d1bupa2		Alignment	not modelled	97.5	12 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
60	c3vgkB		Alignment	not modelled	97.5	23 PDB header: transferase Chain: B: PDB Molecule: glucokinase; PDBTitle: crystal structure of a rok family glucokinase from streptomyces2 griseus
61	c2gupA		Alignment	not modelled	97.5	20 PDB header: transferase Chain: A: PDB Molecule: rok family protein; PDBTitle: structural genomics, the crystal structure of a rok family protein2 from streptococcus pneumoniae tigr4 in complex with sucrose
62	c2hoeA		Alignment	not modelled	97.5	8 PDB header: transferase Chain: A: PDB Molecule: n-acetylglucosamine kinase; PDBTitle: crystal structure of n-acetylglucosamine kinase (tm1224) from2 thermotoga maritima at 2.46 a resolution
63	d2e8aa2		Alignment	not modelled	97.5	13 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
64	d1e4ft1		Alignment	not modelled	97.5	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
65	c3iucC		Alignment	not modelled	97.5	15 PDB header: chaperone Chain: C: PDB Molecule: heat shock 70kda protein 5 (glucose-regulated PDBTitle: crystal structure of the human 70kda heat shock protein 52 (bip/grp78) atpase domain in complex with adp
66	c1jcgA		Alignment	not modelled	97.4	16 PDB header: structural protein Chain: A: PDB Molecule: rod shape-determining protein mreB; PDBTitle: mreB from thermotoga maritima, amppnp
67	c1sazA		Alignment	not modelled	97.4	11 PDB header: transferase Chain: A: PDB Molecule: probable butyrate kinase 2; PDBTitle: membership in the askha superfamily: enzymological2 properties and crystal structure of butyrate kinase 2 from3 thermotoga maritima
68	c2khoA		Alignment	not modelled	97.4	19 PDB header: chaperone Chain: A: PDB Molecule: heat shock protein 70; PDBTitle: nmr-rdc / xray structure of e. coli hsp70 (dnak) chaperone2 (1-605) complexed with adp and substrate
69	c1qhaA		Alignment	not modelled	97.4	13 PDB header: transferase Chain: A: PDB Molecule: protein (hexokinase); PDBTitle: human hexokinase type i complexed with atp analogue amp-pnp
70	d2aa4a1		Alignment	not modelled	97.3	14 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
71	c1bdgA		Alignment	not modelled	97.3	13 PDB header: hexokinase Chain: A: PDB Molecule: hexokinase; PDBTitle: hexokinase from schistosoma mansoni complexed with glucose
72	c2aa4B		Alignment	not modelled	97.3	14 PDB header: transferase Chain: B: PDB Molecule: putative n-acetylmannosamine kinase; PDBTitle: crystal structure of escherichia coli putative n-2 acetylmannosamine kinase, new york structural genomics3 consortium
73	c3mcpA		Alignment	not modelled	97.2	12 PDB header: transferase Chain: A: PDB Molecule: glucokinase; PDBTitle: crystal structure of glucokinase (bdi_1628) from parabacteroides2 distasonis atcc 8503 at 3.00 a resolution
74	c1xc3A		Alignment	not modelled	97.1	17 PDB header: transferase Chain: A: PDB Molecule: putative fructokinase; PDBTitle: structure of a putative fructokinase from bacillus subtilis
75	d2gupa1		Alignment	not modelled	97.0	19 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ROK
76	c1woqB		Alignment	not modelled	97.0	14 PDB header: transferase Chain: B: PDB Molecule: inorganic polyphosphate/atp-glucomannokinase; PDBTitle: crystal structure of inorganic polyphosphate/atp-glucomannokinase from2 arthrobacter sp. strain km at 1.8 a resolution
77	c3p4iA		Alignment	not modelled	96.9	16 PDB header: transferase Chain: A: PDB Molecule: acetate kinase; PDBTitle: crystal structure of acetate kinase from mycobacterium avium
78	c3tsuA		Alignment	not modelled	96.9	13 PDB header: transferase Chain: A: PDB Molecule: transcriptional regulatory protein; PDBTitle: crystal structure of e. coli hypf with amp-pnp and carbamoyl phosphate
79	d1xc3a1		Alianment	not modelled	96.8	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain

					Family:ROK
80	c3qbwA	Alignment	not modelled	96.8	16 PDB header: transferase Chain: A: PDB Molecule: anhydro-n-acetylmuramic acid kinase; PDBTitle: crystal structure of pseudomonas aeruginosa 1,6-anhydro-n-2 acetylmuramic acid kinase (anmk) bound to adenosine diphosphate
81	d1e4ft2	Alignment	not modelled	96.6	18 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
82	c3khyA	Alignment	not modelled	96.4	17 PDB header: transferase Chain: A: PDB Molecule: propionate kinase; PDBTitle: crystal structure of a propionate kinase from francisella2 tularensis subsp. tularensis schu s4
83	c1tuuA	Alignment	not modelled	96.4	18 PDB header: transferase Chain: A: PDB Molecule: acetate kinase; PDBTitle: acetate kinase crystallized with atpgs
84	c2q2rA	Alignment	not modelled	96.4	7 PDB header: transferase Chain: A: PDB Molecule: glucokinase 1, putative; PDBTitle: trypanosoma cruzi glucokinase in complex with beta-d-glucose and adp
85	c3lm2B	Alignment	not modelled	96.4	15 PDB header: transferase Chain: B: PDB Molecule: putative kinase; PDBTitle: crystal structure of putative kinase. (17743352) from agrobacterium2 tumefaciens str. c58 (dupont) at 1.70 a resolution
86	c2ychA	Alignment	not modelled	96.4	17 PDB header: cell cycle Chain: A: PDB Molecule: competence protein pilm; PDBTitle: pilm-piln type iv pilus biogenesis complex
87	d1bg3a3	Alignment	not modelled	96.3	11 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
88	d1bdga1	Alignment	not modelled	96.1	13 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
89	c2d0oA	Alignment	not modelled	96.0	17 PDB header: chaperone Chain: A: PDB Molecule: diol dehydratase-reactivating factor large PDBTitle: strcuture of diol dehydratase-reactivating factor complexed2 with adp and mg2+
90	d1ig8a1	Alignment	not modelled	95.9	19 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
91	d1v4sa1	Alignment	not modelled	95.8	13 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
92	c1zbsA	Alignment	not modelled	95.7	16 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: hypothetical protein pg1100; PDBTitle: crystal structure of the putative n-acetylglucosamine kinase (pg1100)2 from porphyromonas gingivalis, northeast structural genomics target3 pgr18
93	d3bzka5	Alignment	not modelled	95.7	22 Fold: Ribonuclease H-like motif Superfamily: Ribonuclease H-like Family: Tex RuvX-like domain-like
94	c3t69A	Alignment	not modelled	95.5	23 PDB header: transferase Chain: A: PDB Molecule: putative 2-dehydro-3-deoxygalactonokinase; PDBTitle: crystal structure of a putative 2-dehydro-3-deoxygalactonokinase2 protein from sinorhizobium meliloti
95	d1bg3a1	Alignment	not modelled	95.5	13 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
96	c3hm8D	Alignment	not modelled	95.2	11 PDB header: transferase Chain: D: PDB Molecule: hexokinase-3; PDBTitle: crystal structure of the c-terminal hexokinase domain of human hk3
97	d1czan3	Alignment	not modelled	95.2	10 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
98	d1czan1	Alignment	not modelled	94.9	15 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
99	d2i7na2	Alignment	not modelled	94.9	17 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Fumble-like
100	d1ig8a2	Alignment	not modelled	94.3	17 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
101	d2zgya2	Alignment	not modelled	94.0	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
102	d1g99a1	Alignment	not modelled	93.7	20 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Acetokinase-like
103	d1saza2	Alignment	not modelled	93.6	11 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Acetokinase-like
104	d1u6za2	Alignment	not modelled	93.6	11 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Ppx/GppA phosphatase
105	c1t6dB	Alignment	not modelled	93.3	12 PDB header: hydrolase Chain: B: PDB Molecule: exopolyphosphatase; PDBTitle: miras phasing of the aquifex aeolicus ppx/gppa

					phosphatase: crystal2 structure of the type ii variant
106	d2ch5a1	Alignment	not modelled	93.2	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: BadF/BadG/BcrA/BcrD-like
107	d2i7pa1	Alignment	not modelled	93.1	17 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Fumble-like
108	c1mwmA	Alignment	not modelled	92.7	17 PDB header: structural protein Chain: A: PDB Molecule: parm; PDBTitle: parm from plasmid r1 adp form
109	d2d0oa2	Alignment	not modelled	92.5	23 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: ATPase domain of dehydratase reactivase alpha subunit
110	d2e1za1	Alignment	not modelled	92.5	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Acetokinase-like
111	c2oceA	Alignment	not modelled	92.5	20 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: hypothetical protein pa5201; PDBTitle: crystal structure of tex family protein pa5201 from2 pseudomonas aeruginosa
112	c3dwlB	Alignment	not modelled	92.0	9 PDB header: structural protein Chain: B: PDB Molecule: actin-related protein 3; PDBTitle: crystal structure of fission yeast arp2/3 complex lacking the arp22 subunit
113	d1t6ca1	Alignment	not modelled	91.9	12 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Ppx/GppA phosphatase
114	d2e8aa1	Alignment	not modelled	91.9	32 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Actin/HSP70
115	c2h3gX	Alignment	not modelled	91.7	21 PDB header: biosynthetic protein Chain: X: PDB Molecule: biosynthetic protein; PDBTitle: structure of the type iii pantothenate kinase (coax) from bacillus2 anthracis
116	c1zxoB	Alignment	not modelled	91.5	13 PDB header: unknown function Chain: B: PDB Molecule: conserved hypothetical protein q8a1p1; PDBTitle: x-ray crystal structure of protein q8a1p1 from bacteroides2 thetaiotaomicron. northeast structural genomics consortium3 target btr25.
117	c3aapA	Alignment	not modelled	91.3	14 PDB header: hydrolase Chain: A: PDB Molecule: ectonucleoside triphosphate diphosphohydrolase i; PDBTitle: crystal structure of lp1ntpdase from legionella pneumophila
118	c3mdqA	Alignment	not modelled	90.7	10 PDB header: hydrolase Chain: A: PDB Molecule: exopolyphosphatase; PDBTitle: crystal structure of an exopolyphosphatase (chu_0316) from cytophaga2 hutchinsonii atcc 33406 at 1.50 a resolution
119	d1bdga2	Alignment	not modelled	90.7	16 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: Hexokinase
120	d3bexa1	Alignment	not modelled	90.5	20 Fold: Ribonuclease H-like motif Superfamily: Actin-like ATPase domain Family: CoaX-like