























| # | Template | Alignment Coverage | 3D Model | Confidence | % i.d. | Template Information |
|----|-------------------------|--|---|------------|--------|--|
| 1 | c3pvsA_ |  Alignment |  | 100.0 | 100 | PDB header: recombination Chain: A: PDB Molecule: replication-associated recombination protein a; PDBTitle: structure and biochemical activities of escherichia coli mgsa |
| 2 | d2r9ga1 |  Alignment |  | 100.0 | 38 | Fold: post-AAA+ oligomerization domain-like Superfamily: post-AAA+ oligomerization domain-like Family: MgsA/YrvN C-terminal domain-like |
| 3 | d3bgea1 |  Alignment |  | 100.0 | 79 | Fold: post-AAA+ oligomerization domain-like Superfamily: post-AAA+ oligomerization domain-like Family: MgsA/YrvN C-terminal domain-like |
| 4 | c3ctdB_ |  Alignment |  | 100.0 | 40 | PDB header: structural genomics, unknown function Chain: B: PDB Molecule: putative atpase, aaa family; PDBTitle: crystal structure of a putative aaa family atpase from2 prochlorococcus marinus subsp. pastoris |
| 5 | d3ctda1 |  Alignment |  | 100.0 | 43 | Fold: post-AAA+ oligomerization domain-like Superfamily: post-AAA+ oligomerization domain-like Family: MgsA/YrvN C-terminal domain-like |
| 6 | c1xxhB_ |  Alignment |  | 100.0 | 21 | PDB header: transferase Chain: B: PDB Molecule: dna polymerase iii subunit gamma; PDBTitle: atpgs bound e. coli clamp loader complex |
| 7 | c1sxjC_ |  Alignment |  | 100.0 | 22 | PDB header: replication Chain: C: PDB Molecule: activator 1 40 kda subunit; PDBTitle: crystal structure of the eukaryotic clamp loader2 (replication factor c, rfc) bound to the dna sliding clamp3 (proliferating cell nuclear antigen, pcna) |
| 8 | c1iqpF_ |  Alignment |  | 100.0 | 23 | PDB header: replication Chain: F: PDB Molecule: rfcs; PDBTitle: crystal structure of the clamp loader small subunit from2 pyrococcus furiosus |
| 9 | c2qw6A_ |  Alignment |  | 100.0 | 40 | PDB header: hydrolase Chain: A: PDB Molecule: aaa atpase, central region; PDBTitle: crystal structure of the c-terminal domain of an aaa atpase from2 enterococcus faecium do |
| 10 | d2qw6a1 |  Alignment |  | 100.0 | 40 | Fold: post-AAA+ oligomerization domain-like Superfamily: post-AAA+ oligomerization domain-like Family: MgsA/YrvN C-terminal domain-like |
| 11 | c3u5zM_ |  Alignment |  | 100.0 | 16 | PDB header: dna binding protein/dna Chain: M: PDB Molecule: dna polymerase accessory protein 44; PDBTitle: structure of t4 bacteriophage clamp loader bound to the t4 clamp,2 primer-template dna, and atp analog |

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|----|-------------------------|-----------|--------------|-------|----|--|
| 12 | clsxD_ | Alignment | | 100.0 | 19 | PDB header: replication Chain: D: PDB Molecule: activator 1 41 kda subunit; PDBTitle: crystal structure of the eukaryotic clamp loader2 (replication factor c, rfc) bound to the dna sliding clamp3 (proliferating cell nuclear antigen, pcna) |
| 13 | c2chgB_ | Alignment | | 100.0 | 22 | PDB header: dna-binding protein Chain: B: PDB Molecule: replication factor c small subunit; PDBTitle: replication factor c domains 1 and 2 |
| 14 | clsxE_ | Alignment | | 100.0 | 14 | PDB header: replication Chain: E: PDB Molecule: activator 1 40 kda subunit; PDBTitle: crystal structure of the eukaryotic clamp loader2 (replication factor c, rfc) bound to the dna sliding clamp3 (proliferating cell nuclear antigen, pcna) |
| 15 | clsxA_ | Alignment | | 100.0 | 18 | PDB header: replication Chain: A: PDB Molecule: activator 1 95 kda subunit; PDBTitle: crystal structure of the eukaryotic clamp loader2 (replication factor c, rfc) bound to the dna sliding clamp3 (proliferating cell nuclear antigen, pcna) |
| 16 | clsxB_ | Alignment | | 100.0 | 19 | PDB header: replication Chain: B: PDB Molecule: activator 1 37 kda subunit; PDBTitle: crystal structure of the eukaryotic clamp loader2 (replication factor c, rfc) bound to the dna sliding clamp3 (proliferating cell nuclear antigen, pcna) |
| 17 | dlsxb2 | Alignment | | 100.0 | 24 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 18 | dliqpa2 | Alignment | | 100.0 | 26 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 19 | c2chvE_ | Alignment | | 100.0 | 19 | PDB header: dna-binding protein Chain: E: PDB Molecule: replication factor c small subunit; PDBTitle: replication factor c adpnp complex |
| 20 | dlsxc2 | Alignment | | 100.0 | 26 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 21 | dlnjfa_ | Alignment | not modelled | 100.0 | 26 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 22 | clin8A_ | Alignment | not modelled | 100.0 | 23 | PDB header: dna binding protein Chain: A: PDB Molecule: holliday junction dna helicase ruvb; PDBTitle: thermotoga maritima ruvb t158v |
| 23 | dlsxd2 | Alignment | not modelled | 100.0 | 23 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 24 | clhqcB_ | Alignment | not modelled | 100.0 | 25 | PDB header: hydrolase Chain: B: PDB Molecule: ruvb; PDBTitle: structure of ruvb from thermus thermophilus hb8 |
| 25 | c3pfiB_ | Alignment | not modelled | 100.0 | 22 | PDB header: hydrolase Chain: B: PDB Molecule: holliday junction atp-dependent dna helicase ruvb; PDBTitle: 2.7 angstrom resolution crystal structure of a probable holliday2 junction dna helicase (ruvb) from campylobacter jejuni subsp. jejuni3 nctc 11168 in complex with adenosine-5'-diphosphate |
| 26 | dlsxa2 | Alignment | not modelled | 100.0 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 27 | dlsxe2 | Alignment | not modelled | 100.0 | 20 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 28 | dlin4a2 | Alignment | not modelled | 100.0 | 25 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases |

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|----|-------------------------|-----------|--------------|-------|----|---|
| | | | | | | Family: Extended AAA-ATPase domain Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 29 | d1ixsb2 | Alignment | not modelled | 100.0 | 26 | PDB header: hydrolase Chain: C: PDB Molecule: ruvb-like 1; PDBTitle: 3d structure of the human ruvb-like helicase ruvb1 |
| 30 | c2c9oC | Alignment | not modelled | 99.9 | 23 | PDB header: hydrolase Chain: C: PDB Molecule: ruvb-like 1; PDBTitle: 3d structure of the human ruvb-like helicase ruvb1 |
| 31 | c3b9pA | Alignment | not modelled | 99.9 | 21 | PDB header: hydrolase Chain: A: PDB Molecule: cg5977-pa, isoform a; PDBTitle: spastin |
| 32 | c1jr3E | Alignment | not modelled | 99.9 | 17 | PDB header: transferase Chain: E: PDB Molecule: dna polymerase iii, delta' subunit; PDBTitle: crystal structure of the processivity clamp loader gamma2 complex of e. coli dna polymerase iii |
| 33 | c3bosA | Alignment | not modelled | 99.9 | 20 | PDB header: hydrolase regulator,dna binding protein Chain: A: PDB Molecule: putative dna replication factor; PDBTitle: crystal structure of a putative dna replication regulator hda2 (sama_1916) from shewanella amazonensis sb2b at 1.75 a resolution |
| 34 | c3d8bB | Alignment | not modelled | 99.9 | 17 | PDB header: hydrolase Chain: B: PDB Molecule: fidgetin-like protein 1; PDBTitle: crystal structure of human fidgetin-like protein 1 in complex with adp |
| 35 | d1r6bx2 | Alignment | not modelled | 99.9 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 36 | c2dhrC | Alignment | not modelled | 99.9 | 23 | PDB header: hydrolase Chain: C: PDB Molecule: ftsh; PDBTitle: whole cytosolic region of atp-dependent metalloprotease2 ftsh (g399l) |
| 37 | d1um8a | Alignment | not modelled | 99.9 | 24 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 38 | c2ce7B | Alignment | not modelled | 99.9 | 19 | PDB header: cell division protein Chain: B: PDB Molecule: cell division protein ftsh; PDBTitle: edta treated |
| 39 | d2gnoa2 | Alignment | not modelled | 99.9 | 16 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 40 | d1g41a | Alignment | not modelled | 99.9 | 27 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 41 | d1a5ta2 | Alignment | not modelled | 99.9 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 42 | d2ce7a2 | Alignment | not modelled | 99.9 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 43 | c3h4mC | Alignment | not modelled | 99.9 | 16 | PDB header: hydrolase Chain: C: PDB Molecule: proteasome-activating nucleotidase; PDBTitle: aaa atpase domain of the proteasome- activating nucleotidase |
| 44 | c2z4rB | Alignment | not modelled | 99.9 | 19 | PDB header: dna binding protein Chain: B: PDB Molecule: chromosomal replication initiator protein dnaa; PDBTitle: crystal structure of domain iii from the thermotoga2 maritima replication initiation protein dnaa |
| 45 | c2zamA | Alignment | not modelled | 99.9 | 19 | PDB header: protein transport Chain: A: PDB Molecule: vacuolar protein sorting-associated protein 4b; PDBTitle: crystal structure of mouse skd1/vps4b apo-form |
| 46 | d1ofha | Alignment | not modelled | 99.9 | 29 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 47 | d1l8qa2 | Alignment | not modelled | 99.9 | 16 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 48 | c1xwiA | Alignment | not modelled | 99.9 | 18 | PDB header: protein transport Chain: A: PDB Molecule: skd1 protein; PDBTitle: crystal structure of vps4b |
| 49 | c3cf1C | Alignment | not modelled | 99.9 | 16 | PDB header: transport protein Chain: C: PDB Molecule: transitional endoplasmic reticulum atpase; PDBTitle: structure of p97/vcp in complex with adp/adp.alfx |
| 50 | c1r6bX | Alignment | not modelled | 99.8 | 19 | PDB header: hydrolase Chain: X: PDB Molecule: clpa protein; PDBTitle: high resolution crystal structure of clpa |
| 51 | d1w5sa2 | Alignment | not modelled | 99.8 | 21 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 52 | c3eihB | Alignment | not modelled | 99.8 | 20 | PDB header: protein transport Chain: B: PDB Molecule: vacuolar protein sorting-associated protein 4; PDBTitle: crystal structure of s.cerevisiae vps4 in the presence of atpgammas |
| 53 | d1qvra2 | Alignment | not modelled | 99.8 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |

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|----|-------------------------|-----------|--------------|------|----|--|
| 54 | d1ny5a2 | Alignment | not modelled | 99.8 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 55 | c1qvrB | Alignment | not modelled | 99.8 | 22 | PDB header: chaperone Chain: B: PDB Molecule: clpb protein; PDBTitle: crystal structure analysis of clpb |
| 56 | d1ixza | Alignment | not modelled | 99.8 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 57 | d1r6bx3 | Alignment | not modelled | 99.8 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 58 | d1fnna2 | Alignment | not modelled | 99.8 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 59 | c2qz4A | Alignment | not modelled | 99.8 | 17 | PDB header: hydrolase Chain: A: PDB Molecule: paraplegin; PDBTitle: human paraplegin, aaa domain in complex with adp |
| 60 | c2r65A | Alignment | not modelled | 99.8 | 22 | PDB header: hydrolase Chain: A: PDB Molecule: cell division protease ftsh homolog; PDBTitle: crystal structure of helicobacter pylori atp dependent protease, ftsh2 adp complex |
| 61 | c3pxiB | Alignment | not modelled | 99.8 | 23 | PDB header: protein binding Chain: B: PDB Molecule: negative regulator of genetic competence clpc/mecb; PDBTitle: structure of meca108:clpc |
| 62 | c3hteC | Alignment | not modelled | 99.8 | 26 | PDB header: motor protein Chain: C: PDB Molecule: atp-dependent clp protease atp-binding subunit clpx; PDBTitle: crystal structure of nucleotide-free hexameric clpx |
| 63 | d1jbka | Alignment | not modelled | 99.8 | 23 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 64 | d1qvra3 | Alignment | not modelled | 99.8 | 20 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 65 | c3hu2C | Alignment | not modelled | 99.8 | 19 | PDB header: transport protein Chain: C: PDB Molecule: transitional endoplasmic reticulum atpase; PDBTitle: structure of p97 n-d1 r86a mutant in complex with atpgs |
| 66 | d1g8pa | Alignment | not modelled | 99.8 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 67 | c3te6A | Alignment | not modelled | 99.8 | 14 | PDB header: gene regulation Chain: A: PDB Molecule: regulatory protein sir3; PDBTitle: crystal structure of the s. cerevisiae sir3 aaa+ domain |
| 68 | c1iy2A | Alignment | not modelled | 99.8 | 22 | PDB header: hydrolase Chain: A: PDB Molecule: atp-dependent metalloprotease ftsh; PDBTitle: crystal structure of the ftsh atpase domain from thermus2 thermophilus |
| 69 | d1lv7a | Alignment | not modelled | 99.8 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 70 | c1s3sA | Alignment | not modelled | 99.8 | 17 | PDB header: protein binding Chain: A: PDB Molecule: transitional endoplasmic reticulum atpase (ter PDBTitle: crystal structure of aaa atpase p97/vcp nd1 in complex with2 p47 c |
| 71 | c2p65A | Alignment | not modelled | 99.8 | 23 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: hypothetical protein pf08_0063; PDBTitle: crystal structure of the first nucleotide binding domain of2 chaperone clpb1, putative, (pv089580) from plasmodium vivax |
| 72 | d1e32a2 | Alignment | not modelled | 99.8 | 16 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 73 | c2hcbC | Alignment | not modelled | 99.8 | 16 | PDB header: replication Chain: C: PDB Molecule: chromosomal replication initiator protein dnaa; PDBTitle: structure of amppcp-bound dnaa from aquifex aeolicus |
| 74 | d1d2na | Alignment | not modelled | 99.8 | 22 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 75 | c1fnnB | Alignment | not modelled | 99.7 | 18 | PDB header: cell cycle Chain: B: PDB Molecule: cell division control protein 6; PDBTitle: crystal structure of cdc6p from pyrobaculum aerophilum |
| 76 | c2c99A | Alignment | not modelled | 99.7 | 15 | PDB header: transcription regulation Chain: A: PDB Molecule: psp operon transcriptional activator; PDBTitle: structural basis of the nucleotide driven conformational2 changes in the aaa domain of transcription activator pspf |
| 77 | c2gnoA | Alignment | not modelled | 99.7 | 15 | PDB header: replication Chain: A: PDB Molecule: dna polymerase iii, gamma subunit-related protein; PDBTitle: crystal structure of a dna polymerase iii, gamma subunit-related2 protein (tm0771) from thermotoga maritima msb8 at 2.00 a resolution |
| 78 | c1ojlF | Alignment | not modelled | 99.7 | 15 | PDB header: response regulator Chain: F: PDB Molecule: transcriptional regulatory protein zrar; PDBTitle: crystal structure of a sigma54-activator suggests the2 mechanism for the conformational switch necessary for3 sigma54 binding |

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|-----|-------------------------|-----------|--------------|------|----|--|
| 79 | c1nsfA | Alignment | not modelled | 99.7 | 22 | PDB header: protein transport Chain: A: PDB Molecule: n-ethylmaleimide sensitive factor; PDBTitle: d2 hexamerization domain of n-ethylmaleimide sensitive factor (nsf) |
| 80 | c2x8aA | Alignment | not modelled | 99.7 | 14 | PDB header: nuclear protein Chain: A: PDB Molecule: nuclear valosin-containing protein-like; PDBTitle: human nuclear valosin containing protein like (nvl), c-2 terminal aaa-atpase domain |
| 81 | c1ny5A | Alignment | not modelled | 99.7 | 16 | PDB header: transcription Chain: A: PDB Molecule: transcriptional regulator (ntrc family); PDBTitle: crystal structure of sigm54 activator (aaa+ atpase) in the inactive2 state |
| 82 | c1w5sB | Alignment | not modelled | 99.7 | 19 | PDB header: replication Chain: B: PDB Molecule: origin recognition complex subunit 2 orc2; PDBTitle: structure of the aeropyrum pernix orc2 protein (adp form) |
| 83 | c3kljA | Alignment | not modelled | 99.7 | 23 | PDB header: hydrolase Chain: A: PDB Molecule: atp-dependent protease lon; PDBTitle: crystal structure of lon protease from thermococcus onnurineus na1 |
| 84 | c3pxgA | Alignment | not modelled | 99.7 | 20 | PDB header: protein binding Chain: A: PDB Molecule: negative regulator of genetic competence clpc/mecb; PDBTitle: structure of meca121 and clpc1-485 complex |
| 85 | dlr7ra3 | Alignment | not modelled | 99.7 | 19 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 86 | c2qbyA | Alignment | not modelled | 99.7 | 21 | PDB header: replication/dna Chain: A: PDB Molecule: cell division control protein 6 homolog 1; PDBTitle: crystal structure of a heterodimer of cdc6/orc1 initiators2 bound to origin dna (from s. solfataricus) |
| 87 | c2v1uA | Alignment | not modelled | 99.7 | 19 | PDB header: replication Chain: A: PDB Molecule: cell division control protein 6 homolog; PDBTitle: structure of the aeropyrum pernix orc1 protein in complex2 with dna |
| 88 | c2r44A | Alignment | not modelled | 99.7 | 22 | PDB header: hydrolase Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a putative atpase (chu_0153) from cytophaga2 hutchinsonii atcc 33406 at 2.00 a resolution |
| 89 | c3dzdA | Alignment | not modelled | 99.7 | 21 | PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator (ntrc family); PDBTitle: crystal structure of sigma54 activator ntrc4 in the inactive2 state |
| 90 | c1xxiF | Alignment | not modelled | 99.6 | 11 | PDB header: transferase Chain: F: PDB Molecule: dna polymerase iii, delta subunit; PDBTitle: adp bound e. coli clamp loader complex |
| 91 | c3nbxX | Alignment | not modelled | 99.6 | 18 | PDB header: hydrolase Chain: X: PDB Molecule: atpase rava; PDBTitle: crystal structure of e. coli rava (regulatory atpase variant a) in2 complex with adp |
| 92 | c2qbyB | Alignment | not modelled | 99.6 | 19 | PDB header: replication/dna Chain: B: PDB Molecule: cell division control protein 6 homolog 3; PDBTitle: crystal structure of a heterodimer of cdc6/orc1 initiators2 bound to origin dna (from s. solfataricus) |
| 93 | c3m6aC | Alignment | not modelled | 99.6 | 19 | PDB header: hydrolase Chain: C: PDB Molecule: atp-dependent protease la 1; PDBTitle: crystal structure of bacillus subtilis lon c-terminal domain |
| 94 | dle94e | Alignment | not modelled | 99.5 | 26 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 95 | dl0z4a3 | Alignment | not modelled | 99.5 | 18 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 96 | c2kjqA | Alignment | not modelled | 99.5 | 21 | PDB header: replication Chain: A: PDB Molecule: dnaa-related protein; PDBTitle: solution structure of protein nmb1076 from neisseria meningitidis.2 northeast structural genomics consortium target mr101b. |
| 97 | c3ec2A | Alignment | not modelled | 99.4 | 18 | PDB header: replication Chain: A: PDB Molecule: dna replication protein dnac; PDBTitle: crystal structure of the dnac helicase loader |
| 98 | c3f8tA | Alignment | not modelled | 99.3 | 23 | PDB header: hydrolase Chain: A: PDB Molecule: predicted atpase involved in replication control, PDBTitle: crystal structure analysis of a full-length mcm homolog2 from methanopyrus kandleri |
| 99 | c2fnaA | Alignment | not modelled | 99.3 | 12 | PDB header: atp-binding protein Chain: A: PDB Molecule: conserved hypothetical protein; PDBTitle: crystal structure of an archaeal aaa+ atpase (sso1545) from sulfobolus2 solfataricus p2 at 2.00 a resolution |
| 100 | c2qgzA | Alignment | not modelled | 99.1 | 13 | PDB header: hydrolase Chain: A: PDB Molecule: putative primosome component; PDBTitle: crystal structure of a putative primosome component from2 streptococcus pyogenes serotype m3. northeast structural3 genomics target dr58 |
| 101 | c2w58B | Alignment | not modelled | 99.1 | 19 | PDB header: hydrolase Chain: B: PDB Molecule: primosome component (helicase loader); PDBTitle: crystal structure of the dnaI |
| 102 | d2fnaa2 | Alignment | not modelled | 99.1 | 16 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 103 | c2qenA | Alignment | not modelled | 99.1 | 16 | PDB header: unknown function Chain: A: PDB Molecule: walker-type atpase; PDBTitle: the walker-type atpase paby2304 of pyrococcus abyssi |
| | | | | | | PDB header: transcription regulator |

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|-----|-------------------------|-----------|--------------|------|----|--|
| 104 | c2p5tD_ | Alignment | not modelled | 98.9 | 13 | Chain: D: PDB Molecule: pez2; PDBTitle: molecular and structural characterization of the pezat chromosomal2 toxin-antitoxin system of the human pathogen streptococcus pneumoniae |
| 105 | d1gvnb_ | Alignment | not modelled | 98.9 | 13 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Plasmid maintenance system epsilon/zeta, toxin zeta subunit |
| 106 | d1jr3d2 | Alignment | not modelled | 98.9 | 12 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 107 | d1ye8a1 | Alignment | not modelled | 98.9 | 21 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: RecA protein-like (ATPase-domain) |
| 108 | d1svma_ | Alignment | not modelled | 98.8 | 11 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 109 | c3iytG_ | Alignment | not modelled | 98.8 | 13 | PDB header: apoptosis Chain: G: PDB Molecule: apoptotic protease-activating factor 1; PDBTitle: structure of an apoptosome-procaspase-9 card complex |
| 110 | c3co5B_ | Alignment | not modelled | 98.8 | 10 | PDB header: transcription regulator Chain: B: PDB Molecule: putative two-component system transcriptional response PDBTitle: crystal structure of sigma-54 interaction domain of putative2 transcriptional response regulator from neisseria gonorrhoeae |
| 111 | c3f9vA_ | Alignment | not modelled | 98.8 | 17 | PDB header: hydrolase Chain: A: PDB Molecule: minichromosome maintenance protein mcm; PDBTitle: crystal structure of a near full-length archaeal mcm: functional2 insights for an aaa+ hexameric helicase |
| 112 | c2a5yB_ | Alignment | not modelled | 98.7 | 19 | PDB header: apoptosis Chain: B: PDB Molecule: ced-4; PDBTitle: structure of a ced-4/ced-9 complex |
| 113 | d1tuea_ | Alignment | not modelled | 98.6 | 15 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Extended AAA-ATPase domain |
| 114 | c2gxaA_ | Alignment | not modelled | 98.5 | 18 | PDB header: replication/dna Chain: A: PDB Molecule: replication protein e1; PDBTitle: crystal structure of papillomavirus e1 hexameric helicase2 with ssdna and mgadp |
| 115 | c3qmzA_ | Alignment | not modelled | 98.5 | 17 | PDB header: motor protein Chain: A: PDB Molecule: cytoplasmic dynein heavy chain; PDBTitle: crystal structure of the cytoplasmic dynein heavy chain motor domain |
| 116 | d1kaga_ | Alignment | not modelled | 98.3 | 15 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Shikimate kinase (AroK) |
| 117 | c1w36G_ | Alignment | not modelled | 98.3 | 18 | PDB header: recombination Chain: G: PDB Molecule: exodeoxyribonuclease v alpha chain; PDBTitle: recbcd:dna complex |
| 118 | c2v9pH_ | Alignment | not modelled | 98.3 | 16 | PDB header: hydrolase Chain: H: PDB Molecule: replication protein e1; PDBTitle: crystal structure of papillomavirus e1 hexameric helicase2 dna-free form |
| 119 | c1z6tC_ | Alignment | not modelled | 98.2 | 14 | PDB header: apoptosis Chain: C: PDB Molecule: apoptotic protease activating factor 1; PDBTitle: structure of the apoptotic protease-activating factor 12 bound to adp |
| 120 | d1w36d1 | Alignment | not modelled | 98.2 | 18 | Fold: P-loop containing nucleoside triphosphate hydrolases Superfamily: P-loop containing nucleoside triphosphate hydrolases Family: Tandem AAA-ATPase domain |