

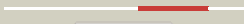






























| # | Template | Alignment Coverage | 3D Model | Confidence | % i.d. | Template Information |
|----|-------------------------|--------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|------------|--------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 1 | c3gbgA_ |  Alignment |  | 100.0 | 11 | PDB header: transcription regulator Chain: A: PDB Molecule: tcp pilus virulence regulatory protein; PDBTitle: crystal structure of tcp from vibrio cholerae o395 |
| 2 | c1d5yD_ |  Alignment |  | 99.9 | 25 | PDB header: transcription/dna Chain: D: PDB Molecule: rob transcription factor; PDBTitle: crystal structure of the e. coli rob transcription factor2 in complex with dna |
| 3 | c3oioA_ |  Alignment |  | 99.9 | 22 | PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator (arac-type dna-binding domain- PDBTitle: crystal structure of transcriptional regulator (arac-type dna-binding2 domain-containing proteins) from chromobacterium violaceum |
| 4 | c3mn2B_ |  Alignment |  | 99.9 | 25 | PDB header: transcription regulator Chain: B: PDB Molecule: probable arac family transcriptional regulator; PDBTitle: the crystal structure of a probable arac family transcriptional2 regulator from rhodopseudomonas palustris cga009 |
| 5 | c3oouA_ |  Alignment |  | 99.9 | 22 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: lin2118 protein; PDBTitle: the structure of a protein with unkown function from listeria innocua |
| 6 | c1bl0A_ |  Alignment |  | 99.9 | 23 | PDB header: transcription/dna Chain: A: PDB Molecule: protein (multiple antibiotic resistance protein); PDBTitle: multiple antibiotic resistance protein (mara)/dna complex |
| 7 | c2k9sA_ |  Alignment |  | 99.9 | 22 | PDB header: transcription Chain: A: PDB Molecule: arabinose operon regulatory protein; PDBTitle: solution structure of dna binding domain of e. coli arac |
| 8 | c3mklB_ |  Alignment |  | 99.9 | 28 | PDB header: transcription regulator Chain: B: PDB Molecule: hth-type transcriptional regulator gadx; PDBTitle: crystal structure of dna-binding transcriptional dual regulator from2 escherichia coli k-12 |
| 9 | c3lsgD_ |  Alignment |  | 99.8 | 24 | PDB header: transcription regulator Chain: D: PDB Molecule: two-component response regulator yesn; PDBTitle: the crystal structure of the c-terminal domain of the two-2 component response regulator yesn from fusobacterium3 nucleatum subsp. nucleatum atcc 25586 |
| 10 | d1bl0a2 |  Alignment |  | 99.4 | 21 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: AraC type transcriptional activator |
| 11 | d1d5ya2 |  Alignment |  | 99.4 | 25 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: AraC type transcriptional activator |

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|----|-------------------------|-----------|-------------------------------------------------------------------------------------|------|----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 12 | d1d5ya1 | Alignment |  | 99.1 | 26 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: AraC type transcriptional activator |
| 13 | d1bl0a1 | Alignment |  | 99.0 | 26 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: AraC type transcriptional activator |
| 14 | c1zgwA | Alignment |  | 99.0 | 24 | PDB header: transcription regulator/dna Chain: A: PDB Molecule: ada polypeptide; PDBTitle: nmr structure of e. coli ada protein in complex with dna |
| 15 | d2arca | Alignment |  | 98.6 | 7 | Fold: Double-stranded beta-helix Superfamily: Regulatory protein AraC Family: Regulatory protein AraC |
| 16 | c3ibmB | Alignment |  | 97.9 | 11 | PDB header: structural genomics, unknown function Chain: B: PDB Molecule: cupin 2, conserved barrel domain protein; PDBTitle: crystal structure of cupin 2 domain-containing protein hha1_0468 from2 halorhodospira halophila |
| 17 | c3jzvA | Alignment |  | 97.7 | 5 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein rru_a2000; PDBTitle: crystal structure of rru_a2000 from rhodospirillum rubrum: a cupin-22 domain. |
| 18 | c3kgzA | Alignment |  | 97.7 | 5 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: cupin 2 conserved barrel domain protein; PDBTitle: crystal structure of a cupin 2 conserved barrel domain protein from2 rhodopseudomonas palustris |
| 19 | c3h7yA | Alignment |  | 97.7 | 8 | PDB header: biosynthetic protein Chain: A: PDB Molecule: bacilysin biosynthesis protein bacb; PDBTitle: crystal structure of bacb, an enzyme involved in bacilysin synthesis,2 in tetragonal form |
| 20 | c3cewA | Alignment |  | 97.6 | 17 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized cupin protein; PDBTitle: crystal structure of a cupin protein (bf4112) from bacteroides2 fragilis. northeast structural genomics consortium target bfr205 |
| 21 | c3l2hD | Alignment | not modelled | 97.5 | 10 | PDB header: isomerase Chain: D: PDB Molecule: putative sugar phosphate isomerase; PDBTitle: crystal structure of putative sugar phosphate isomerase (afe_0303)2 from acidithiobacillus ferrooxidans atcc 23270 at 1.85 a resolution |
| 22 | d1o4ta | Alignment | not modelled | 97.5 | 10 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1287-like |
| 23 | c2vpvA | Alignment | not modelled | 97.4 | 6 | PDB header: cell cycle Chain: A: PDB Molecule: protein mif2; PDBTitle: dimerization domain of mif2p |
| 24 | d1vj2a | Alignment | not modelled | 97.4 | 12 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1459-like |
| 25 | d1y9qa2 | Alignment | not modelled | 97.4 | 11 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Probable transcriptional regulator VC1968, C-terminal domain |
| 26 | c2d40C | Alignment | not modelled | 97.4 | 11 | PDB header: oxidoreductase Chain: C: PDB Molecule: putative gentisate 1,2-dioxygenase; PDBTitle: crystal structure of z3393 from escherichia coli o157:h7 |
| 27 | c3i7dB | Alignment | not modelled | 97.4 | 11 | PDB header: isomerase Chain: B: PDB Molecule: sugar phosphate isomerase; PDBTitle: crystal structure of sugar phosphate isomerase from a cupin2 superfamily spo2919 from silicibacter pomeroyi (yp_168127.1) from3 silicibacter pomeroyi dss-3 at 2.30 a resolution |
| 28 | d1yfua1 | Alignment | not modelled | 97.3 | 16 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins |

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|----|-------------------------|-----------|--------------|------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | | | | | Family: 3-hydroxyanthranilic acid dioxygenase-like |
| 29 | c2gu9B | Alignment | not modelled | 97.3 | 10 PDB header: immune system Chain: B: PDB Molecule: tetracenomycin polyketide synthesis protein; PDBTitle: crystal structure of xc5357 from xanthomonas campestris: a2 putative tetracenomycin polyketide synthesis protein3 adopting a novel cupin subfamily structure |
| 30 | d1sefa | Alignment | not modelled | 97.3 | 15 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: YlbA-like |
| 31 | c1sefA | Alignment | not modelled | 97.3 | 15 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: conserved hypothetical protein; PDBTitle: crystal structure of cupin domain protein ef2996 from enterococcus2 faecalis |
| 32 | d1sfna | Alignment | not modelled | 97.3 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: YlbA-like |
| 33 | c3fjsC | Alignment | not modelled | 97.2 | 14 PDB header: biosynthetic protein Chain: C: PDB Molecule: uncharacterized protein with rmlc-like cupin fold; PDBTitle: crystal structure of a putative biosynthetic protein with rmlc-like2 cupin fold (reut_b4087) from ralstonia eutropha jmp134 at 1.90 a3 resolution |
| 34 | d1v70a | Alignment | not modelled | 97.2 | 12 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1287-like |
| 35 | d1rc6a | Alignment | not modelled | 97.2 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: YlbA-like |
| 36 | d2phda1 | Alignment | not modelled | 97.2 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Gentisate 1,2-dioxygenase-like |
| 37 | d1lr5a | Alignment | not modelled | 97.2 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 38 | c2pfbB | Alignment | not modelled | 97.2 | 3 PDB header: unknown function Chain: B: PDB Molecule: cupin 2, conserved barrel domain protein; PDBTitle: crystal structure of a rmlc-like cupin (sfri_3105) from shewanella2 frigidimarina ncimb 400 at 1.90 a resolution |
| 39 | d3bu7a1 | Alignment | not modelled | 97.1 | 7 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Gentisate 1,2-dioxygenase-like |
| 40 | c3bu7A | Alignment | not modelled | 97.1 | 7 PDB header: oxidoreductase Chain: A: PDB Molecule: gentisate 1,2-dioxygenase; PDBTitle: crystal structure and biochemical characterization of gdosp,2 a gentisate 1,2-dioxygenase from silicibacter pomeroyi |
| 41 | d2f4pa1 | Alignment | not modelled | 97.1 | 8 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1287-like |
| 42 | c2ozjB | Alignment | not modelled | 97.1 | 5 PDB header: unknown function Chain: B: PDB Molecule: cupin 2, conserved barrel; PDBTitle: crystal structure of a cupin superfamily protein (dsy2733) from2 desulfitobacterium hafniense dcb-2 at 1.60 a resolution |
| 43 | d1dgwa | Alignment | not modelled | 97.1 | 16 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 44 | c3lwcA | Alignment | not modelled | 97.1 | 6 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of structural genomics, unknown function2 (yp_766765.1) from rhizobium leguminosarum bv. viciae 3841 at 1.40 a3 resolution |
| 45 | d2b8ma1 | Alignment | not modelled | 97.0 | 6 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: MJ0764-like |
| 46 | d1y3ta1 | Alignment | not modelled | 97.0 | 5 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Quercetin 2,3-dioxygenase-like |
| 47 | d1sq4a | Alignment | not modelled | 97.0 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: YlbA-like |
| 48 | d1zvfa1 | Alignment | not modelled | 97.0 | 19 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: 3-hydroxyanthranilic acid dioxygenase-like |
| 49 | c2o8qA | Alignment | not modelled | 97.0 | 3 PDB header: metal binding protein Chain: A: PDB Molecule: hypothetical protein; PDBTitle: crystal structure of a protein with a cupin-like fold and unknown2 function (bx_c0505) from burkholderia xenovorans lb400 at 1.55 a3 resolution |
| 50 | d2d40a1 | Alignment | not modelled | 97.0 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Gentisate 1,2-dioxygenase-like |
| 51 | d1uika1 | Alignment | not modelled | 97.0 | 10 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 52 | d1yhfa1 | Alignment | not modelled | 97.0 | 12 Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1287-like |
| 53 | c2oa2A | Alignment | not modelled | 97.0 | 11 PDB header: structural genomics, unknown function Chain: A: PDB Molecule: bh2720 protein; PDBTitle: crystal structure of bh2720 (10175341) from bacillus |

| | | | | | | |
|----|-------------------------|-----------|--------------|------|----|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | | | | | | halodurans at2 1.41 a resolution |
| 54 | c3rnsA | Alignment | not modelled | 97.0 | 15 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: cupin 2 conserved barrel domain protein; PDBTitle: cupin 2 conserved barrel domain protein from leptotrichia buccalis |
| 55 | c3h8uA | Alignment | not modelled | 97.0 | 13 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized conserved protein with double-stranded PDBTitle: crystal structure of uncharacterized conserved protein with double-2 stranded beta-helix domain (yp_001338853.1) from klebsiella3 pneumoniae subsp. pneumoniae mgh 78578 at 1.80 a resolution |
| 56 | d2bnma2 | Alignment | not modelled | 96.9 | 7 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: TM1459-like |
| 57 | c3d82A | Alignment | not modelled | 96.9 | 12 | PDB header: metal binding protein Chain: A: PDB Molecule: cupin 2, conserved barrel domain protein; PDBTitle: crystal structure of a cupin-2 domain containing protein (sfri_3543)2 from shewanella frigidimarina ncimb 400 at 2.05 a resolution |
| 58 | c2opkC | Alignment | not modelled | 96.9 | 11 | PDB header: isomerase Chain: C: PDB Molecule: hypothetical protein; PDBTitle: crystal structure of a putative mannose-6-phosphate isomerase2 (reut_a1446) from ralstonia eutropha jmp134 at 2.10 a resolution |
| 59 | c3ht2A | Alignment | not modelled | 96.9 | 10 | PDB header: lyase Chain: A: PDB Molecule: remf protein; PDBTitle: zink containing polyketide cyclase remf from streptomyces2 resistomycificus |
| 60 | c2i45C | Alignment | not modelled | 96.7 | 7 | PDB header: structural genomics, unknown function Chain: C: PDB Molecule: hypothetical protein; PDBTitle: crystal structure of protein nmb1881 from neisseria meningitidis |
| 61 | d1uija1 | Alignment | not modelled | 96.6 | 8 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 62 | c1y9qA | Alignment | not modelled | 96.5 | 11 | PDB header: transcription regulator Chain: A: PDB Molecule: transcriptional regulator, hth_3 family; PDBTitle: crystal structure of hth_3 family transcriptional regulator2 from vibrio cholerae |
| 63 | c3bcwB | Alignment | not modelled | 96.5 | 10 | PDB header: unknown function Chain: B: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a duf861 family protein with a rmlc-like cupin2 fold (bb1179) from bordetella bronchiseptica rb50 at 1.60 a3 resolution |
| 64 | d1j3pa | Alignment | not modelled | 96.5 | 13 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Glucose-6-phosphate isomerase, GPI |
| 65 | c2fqpD | Alignment | not modelled | 96.4 | 7 | PDB header: metal binding protein Chain: D: PDB Molecule: hypothetical protein bp2299; PDBTitle: crystal structure of a cupin domain (bp2299) from bordetella pertussis2 tohama i at 1.80 a resolution |
| 66 | d1juha | Alignment | not modelled | 96.3 | 7 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Quercetin 2,3-dioxygenase-like |
| 67 | d1x82a | Alignment | not modelled | 96.3 | 6 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Glucose-6-phosphate isomerase, GPI |
| 68 | d1o5ua | Alignment | not modelled | 96.3 | 4 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Hypothetical protein TM1112 |
| 69 | c2oziA | Alignment | not modelled | 96.2 | 9 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: hypothetical protein rpa4178; PDBTitle: structural genomics, the crystal structure of a putative2 protein rpa4178 from rhodopseudomonas palustris cga009 |
| 70 | d1fxza1 | Alignment | not modelled | 96.2 | 11 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 71 | c2bnoA | Alignment | not modelled | 96.2 | 7 | PDB header: oxidoreductase Chain: A: PDB Molecule: epoxidase; PDBTitle: the structure of hydroxypropylphosphonic acid epoxidase2 from s. wedmorenis. |
| 72 | d1zrra1 | Alignment | not modelled | 96.2 | 11 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Acireductone dioxygenase |
| 73 | c2vqaC | Alignment | not modelled | 96.1 | 16 | PDB header: metal-binding protein Chain: C: PDB Molecule: sll1358 protein; PDBTitle: protein-folding location can regulate mn versus cu- or zn-2 binding. crystal structure of mnca. |
| 74 | d2et1a1 | Alignment | not modelled | 96.0 | 14 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 75 | d1j58a | Alignment | not modelled | 95.9 | 15 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 76 | c2q30C | Alignment | not modelled | 95.9 | 14 | PDB header: unknown function Chain: C: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a rmlc-like cupin protein (dde_2303) from2 desulfovibrio desulfuricans subsp. at 1.94 a resolution |
| 77 | d2pyta1 | Alignment | not modelled | 95.8 | 13 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: EutQ-like |
| | | | | | | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein pspto 0244; |

| | | | | | | |
|-----|-------------------------|-----------|--------------|------|----|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 78 | c3myxA | Alignment | not modelled | 95.6 | 5 | PDBTitle: crystal structure of a pspto_0244 (protein with unknown function which2 belongs to pfam duf861 family) from pseudomonas syringae pv. tomato3 str. dc3000 at 1.30 a resolution |
| 79 | c1uijA | Alignment | not modelled | 95.5 | 11 | PDB header: sugar binding protein Chain: A: PDB Molecule: beta subunit of beta conglycinin; PDBTitle: crystal structure of soybean beta-conglycinin beta2 homotrimer (i122m/k124w) |
| 80 | c3es1A | Alignment | not modelled | 95.3 | 6 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: cupin 2, conserved barrel domain protein; PDBTitle: crystal structure of protein with a cupin-like fold and unknown2 function (yp_001165807.1) from novosphingobium aromaticivorans dsm3 12444 at 1.91 a resolution |
| 81 | c2cauA | Alignment | not modelled | 95.3 | 12 | PDB header: plant protein Chain: A: PDB Molecule: protein (canavalin); PDBTitle: canavalin from jack bean |
| 82 | c2eaaB | Alignment | not modelled | 95.0 | 8 | PDB header: plant protein Chain: B: PDB Molecule: 7s globulin-3; PDBTitle: crystal structure of adzuki bean 7s globulin-3 |
| 83 | d2hsga1 | Alignment | not modelled | 95.0 | 21 | Fold: lambda repressor-like DNA-binding domains Superfamily: lambda repressor-like DNA-binding domains Family: GalR/LacI-like bacterial regulator |
| 84 | d2phla2 | Alignment | not modelled | 94.7 | 22 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 85 | d2phla1 | Alignment | not modelled | 94.7 | 13 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 86 | c2qtqB | Alignment | not modelled | 94.5 | 11 | PDB header: transcription Chain: B: PDB Molecule: transcriptional regulator, tetr family; PDBTitle: crystal structure of a predicted dna-binding transcriptional regulator2 (saro_1072) from novosphingobium aromaticivorans dsm at 1.85 a3 resolution |
| 87 | c2gfnA | Alignment | not modelled | 94.5 | 16 | PDB header: transcription Chain: A: PDB Molecule: hth-type transcriptional regulator pksa related protein; PDBTitle: crystal structure of hth-type transcriptional regulator pksa related2 protein from rhodococcus sp. rha1 |
| 88 | d2bjca1 | Alignment | not modelled | 94.5 | 16 | Fold: lambda repressor-like DNA-binding domains Superfamily: lambda repressor-like DNA-binding domains Family: GalR/LacI-like bacterial regulator |
| 89 | d1efaa1 | Alignment | not modelled | 94.5 | 14 | Fold: lambda repressor-like DNA-binding domains Superfamily: lambda repressor-like DNA-binding domains Family: GalR/LacI-like bacterial regulator |
| 90 | c3kglB | Alignment | not modelled | 94.4 | 7 | PDB header: plant protein Chain: B: PDB Molecule: cruciferin; PDBTitle: crystal structure of procruciferin, 11s globulin from2 brassica napus |
| 91 | c1cauB | Alignment | not modelled | 94.4 | 16 | PDB header: seed storage protein Chain: B: PDB Molecule: canavalin; PDBTitle: determination of three crystal structures of canavalin by molecular2 replacement |
| 92 | c3iwfA | Alignment | not modelled | 94.3 | 16 | PDB header: transcription regulator Chain: A: PDB Molecule: transcription regulator rpir family; PDBTitle: the crystal structure of the n-terminal domain of a rpir2 transcriptional regulator from staphylococcus epidermidis to 1.4a |
| 93 | d2gfnal | Alignment | not modelled | 94.3 | 18 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: Tetracyclin repressor-like, N-terminal domain |
| 94 | c3bruA | Alignment | not modelled | 94.2 | 18 | PDB header: transcription Chain: A: PDB Molecule: regulatory protein, tetr family; PDBTitle: crystal structure of regulatory protein tetr from rhodobacter2 sphaeroides |
| 95 | c3on4D | Alignment | not modelled | 94.1 | 11 | PDB header: transcription Chain: D: PDB Molecule: transcriptional regulator, tetr family; PDBTitle: crystal structure of tetr transcriptional regulator from legionella2 pneumophila |
| 96 | c1vi0B | Alignment | not modelled | 94.1 | 13 | PDB header: transcription Chain: B: PDB Molecule: transcriptional regulator; PDBTitle: crystal structure of a transcriptional regulator |
| 97 | d1qpza1 | Alignment | not modelled | 94.1 | 24 | Fold: lambda repressor-like DNA-binding domains Superfamily: lambda repressor-like DNA-binding domains Family: GalR/LacI-like bacterial regulator |
| 98 | d1uija2 | Alignment | not modelled | 94.0 | 18 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 99 | c3s7eB | Alignment | not modelled | 93.9 | 11 | PDB header: allergen Chain: B: PDB Molecule: allergen ara h 1, clone p41b; PDBTitle: crystal structure of ara h 1 |
| 100 | d1sgma1 | Alignment | not modelled | 93.8 | 10 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: Tetracyclin repressor-like, N-terminal domain |
| 101 | c2ibdB | Alignment | not modelled | 93.7 | 14 | PDB header: structural genomics, unknown function Chain: B: PDB Molecule: possible transcriptional regulator; PDBTitle: crystal structure of probable transcriptional regulatory protein2 rha5900 |
| 102 | c3geuC | Alignment | not modelled | 93.7 | 10 | PDB header: cell adhesion Chain: C: PDB Molecule: intercellular adhesion protein r; PDBTitle: crystal structure of icar from staphylococcus aureus, a member of the2 tetracycline repressor protein family |
| 103 | c1jumB | Alignment | not modelled | 93.7 | 6 | PDB header: transcription Chain: B: PDB Molecule: hypothetical transcriptional regulator in qaca PDBTitle: crystal structure of the multidrug binding |

| | | | | | | |
|-----|-------------------------|-----------|--------------|------|----|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| | | | | | | transcriptional2 repressor qacr bound to the natural drug berberine |
| 104 | d1t56a1 | Alignment | not modelled | 93.7 | 18 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: Tetracyclin repressor-like, N-terminal domain |
| 105 | c3djpB | Alignment | not modelled | 93.7 | 9 | PDB header: dna binding protein Chain: B: PDB Molecule: transcription regulator, tetr family; PDBTitle: the crystal structure of a tetr transcription regulator2 from silicibacter pomeroyi dss |
| 106 | c1sgmA | Alignment | not modelled | 93.7 | 10 | PDB header: transcription Chain: A: PDB Molecule: putative hth-type transcriptional regulator yxaf; PDBTitle: crystal structure of hypothetical protein yxaf |
| 107 | c3g7rB | Alignment | not modelled | 93.5 | 9 | PDB header: transcription regulator Chain: B: PDB Molecule: putative transcriptional regulator; PDBTitle: crystal structure of sco4454, a tetr-family transcriptional2 regulator from streptomyces coelicolor |
| 108 | c3dcfB | Alignment | not modelled | 93.5 | 12 | PDB header: transcription regulator Chain: B: PDB Molecule: transcriptional regulator of the tetr/acrr PDBTitle: crystal structure of transcriptional regulator of the2 tetr/acrr family (yp_290855.1) from thermobifida fusca yx-3 er1 at 2.50 a resolution |
| 109 | d2fbqa1 | Alignment | not modelled | 93.5 | 9 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: Tetracyclin repressor-like, N-terminal domain |
| 110 | c3kscD | Alignment | not modelled | 93.5 | 14 | PDB header: plant protein Chain: D: PDB Molecule: lega class; PDBTitle: crystal structure of pea prolegumin, an 11s seed globulin2 from pisum sativum.l. |
| 111 | c2hytA | Alignment | not modelled | 93.5 | 14 | PDB header: transcription Chain: A: PDB Molecule: tetr-family transcriptional regulator; PDBTitle: crystal structure of a tetr-family transcriptional regulator (eca1819)2 from pectobacterium atrosepticum at 1.64 a resolution |
| 112 | c2o3fC | Alignment | not modelled | 93.4 | 20 | PDB header: transcription Chain: C: PDB Molecule: putative hth-type transcriptional regulator ybbh; PDBTitle: structural genomics, the crystal structure of the n-2 terminal domain of the putative transcriptional regulator3 ybbh from bacillus subtilis subsp. subtilis str. 168. |
| 113 | d2o3fa1 | Alignment | not modelled | 93.4 | 20 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: RpiR-like |
| 114 | c3anpD | Alignment | not modelled | 93.3 | 10 | PDB header: transcription Chain: D: PDB Molecule: transcriptional repressor, tetr family; PDBTitle: crystal structure of thermus thermophilus fadr, a tetr family2 transcriptional repressor, in complex with lauroyl-coa. |
| 115 | c3lwiA | Alignment | not modelled | 93.2 | 12 | PDB header: transcription regulator Chain: A: PDB Molecule: putative tetr-family transcriptional regulator; PDBTitle: crystal structure of putative tetr-family transcriptional regulator2 (yp_752756.1) from syntrophomonas wolfei str. goettingen at 2.07 a3 resolution |
| 116 | c1fxzC | Alignment | not modelled | 93.2 | 11 | PDB header: plant protein Chain: C: PDB Molecule: glycinin g1; PDBTitle: crystal structure of soybean proglycinin a1ab1b homotrimer |
| 117 | d2o7ta1 | Alignment | not modelled | 93.2 | 20 | Fold: DNA/RNA-binding 3-helical bundle Superfamily: Homeodomain-like Family: Tetracyclin repressor-like, N-terminal domain |
| 118 | d1od5a1 | Alignment | not modelled | 93.2 | 12 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 119 | d1od5a2 | Alignment | not modelled | 93.1 | 10 | Fold: Double-stranded beta-helix Superfamily: RmlC-like cupins Family: Germin/Seed storage 7S protein |
| 120 | c3es4B | Alignment | not modelled | 93.1 | 6 | PDB header: structural genomics, unknown function Chain: B: PDB Molecule: uncharacterized protein duf861 with a rmlc-like cupin fold; PDBTitle: crystal structure of protein of unknown function (duf861) with a rmlc-2 like cupin fold (17741406) from agrobacterium tumefaciens str. c583 (dupont) at 1.64 a resolution |