
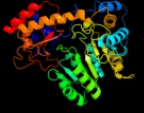






















Phyre2

| | |
|---------------|-----------------------------|
| Email | I.a.kelley@imperial.ac.uk |
| Description | P0A959 |
| Date | Thu Jan 5 11:09:27 GMT 2012 |
| Unique Job ID | ce1925147becd214 |

Detailed template information

| # | Template | Alignment Coverage | 3D Model | Confidence | % i.d. | Template Information |
|----|-------------------------|-----------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|------------|--------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 1 | c3g0tA_ |  Alignment |  | 100.0 | 18 | PDB header: transferase Chain: A: PDB Molecule: putative aminotransferase; PDBTitle: crystal structure of putative aspartate aminotransferase (np_905498.1)2 from porphyromonas gingivalis w83 at 1.75 a resolution |
| 2 | d1xi9a_ |  Alignment |  | 100.0 | 40 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 3 | c3h14A_ |  Alignment |  | 100.0 | 24 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase, classes i and ii; PDBTitle: crystal structure of a putative aminotransferase from silicibacter2 pomeroyi |
| 4 | d1j32a_ |  Alignment |  | 100.0 | 29 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 5 | c3ihjA_ |  Alignment |  | 100.0 | 26 | PDB header: transferase Chain: A: PDB Molecule: alanine aminotransferase 2; PDBTitle: human alanine aminotransferase 2 in complex with plp |
| 6 | d1o4sa_ |  Alignment |  | 100.0 | 27 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 7 | d1w7la_ |  Alignment |  | 100.0 | 20 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 8 | c3e2yB_ |  Alignment |  | 100.0 | 21 | PDB header: transferase, lyase Chain: B: PDB Molecule: kynurenine-oxoglutarate transaminase 3; PDBTitle: crystal structure of mouse kynurenine aminotransferase iii in complex2 with glutamine |
| 9 | c1ynuA_ |  Alignment |  | 100.0 | 16 | PDB header: lyase Chain: A: PDB Molecule: 1-aminocyclopropane-1-carboxylate synthase; PDBTitle: crystal structure of apple acc synthase in complex with l-vinylglycine |
| 10 | c3b46B_ |  Alignment |  | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: aminotransferase bna3; PDBTitle: crystal structure of bna3p, a putative kynurenine2 aminotransferase from saccharomyces cerevisiae |
| 11 | c2zc0C_ |  Alignment |  | 100.0 | 18 | PDB header: transferase Chain: C: PDB Molecule: alanine glyoxylate transaminase; PDBTitle: crystal structure of an archaeal alanine:glyoxylate aminotransferase |

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|----|-------------------------|-----------|--------------|-------|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 12 | d1bw0a_ | Alignment | | 100.0 | 23 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 13 | d2gb3a1 | Alignment | | 100.0 | 23 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 14 | d1b5pa_ | Alignment | | 100.0 | 27 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 15 | c2o1bA_ | Alignment | | 100.0 | 23 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase, class i; PDBTitle: structure of aminotransferase from staphylococcus aureus |
| 16 | d1u08a_ | Alignment | | 100.0 | 20 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 17 | d1m7ya_ | Alignment | | 100.0 | 16 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 18 | d2r5ea1 | Alignment | | 100.0 | 18 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 19 | c3eibB_ | Alignment | | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: II-diaminopimelate aminotransferase; PDBTitle: crystal structure of k270n variant of II-diaminopimelate2 aminotransferase from arabidopsis thaliana |
| 20 | c3jtbB_ | Alignment | | 100.0 | 20 | PDB header: transferase Chain: B: PDB Molecule: aminotransferase; PDBTitle: crystal structure of aminotransferase (np_283882.1) from neisseria2 meningitidis z2491 at 1.91 a resolution |
| 21 | d1wsta1 | Alignment | not modelled | 100.0 | 20 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 22 | d1iaya_ | Alignment | not modelled | 100.0 | 19 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 23 | c2o0rA_ | Alignment | not modelled | 100.0 | 21 | PDB header: transferase Chain: A: PDB Molecule: rv0858c (n-succinyl diaminopimelate aminotransferase); PDBTitle: the three-dimensional structure of n-succinyl diaminopimelate2 aminotransferase from mycobacterium tuberculosis |
| 24 | c3dzzB_ | Alignment | not modelled | 100.0 | 16 | PDB header: transferase Chain: B: PDB Molecule: putative pyridoxal 5'-phosphate-dependent c-s lyase; PDBTitle: crystal structure of a putative plp-dependent aminotransferase2 (lbul_1103) from lactobacillus delbrueckii subsp. at 1.61 a3 resolution |
| 25 | c3qguB_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: II-diaminopimelate aminotransferase; PDBTitle: I,I-diaminopimelate aminotransferase from chlamydomonas reinhardtii |
| 26 | d1gdea_ | Alignment | not modelled | 100.0 | 29 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 27 | c3l8aB_ | Alignment | not modelled | 100.0 | 19 | PDB header: lyase Chain: B: PDB Molecule: putative aminotransferase, probable beta-cystathionase; PDBTitle: crystal structure of metc from streptococcus mutans |
| | | | | | | PDB header: transferase |

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|----|-------------------------|-----------|--------------|-------|----|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 28 | c3if2B_ | Alignment | not modelled | 100.0 | 17 | Chain: B: PDB Molecule: aminotransferase; PDBTitle: crystal structure of putative amino-acid aminotransferase2 (yp_265399.1) from psychrobacter arcticum 273-4 at 2.50 a resolution |
| 29 | d1vp4a_ | Alignment | not modelled | 100.0 | 18 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 30 | c3pplB_ | Alignment | not modelled | 100.0 | 16 | PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of an aspartate transaminase (ncgl0237, cgl0240)2 from corynebacterium glutamicum atcc 13032 kitasato at 1.25 a3 resolution |
| 31 | d1x0ma1 | Alignment | not modelled | 100.0 | 21 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 32 | c2x5dD_ | Alignment | not modelled | 100.0 | 24 | PDB header: transferase Chain: D: PDB Molecule: probable aminotransferase; PDBTitle: crystal structure of a probable aminotransferase from2 pseudomonas aeruginosa |
| 33 | d1c7na_ | Alignment | not modelled | 100.0 | 17 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 34 | c3fdbA_ | Alignment | not modelled | 100.0 | 16 | PDB header: transferase Chain: A: PDB Molecule: putative plp-dependent beta-cystathionase; PDBTitle: crystal structure of a putative plp-dependent beta-cystathionase2 (aecd, dip1736) from corynebacterium diphtheriae at 1.99 a resolution |
| 35 | c3ezsB_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: aminotransferase aspb; PDBTitle: crystal structure of aminotransferase aspb (np_207418.1) from2 helicobacter pylori 26695 at 2.19 a resolution |
| 36 | c3eleB_ | Alignment | not modelled | 100.0 | 23 | PDB header: transferase Chain: B: PDB Molecule: amino transferase; PDBTitle: crystal structure of amino transferase (rer070207001803) from2 eubacterium rectale at 2.10 a resolution |
| 37 | c3dydB_ | Alignment | not modelled | 100.0 | 23 | PDB header: transferase Chain: B: PDB Molecule: tyrosine aminotransferase; PDBTitle: human tyrosine aminotransferase |
| 38 | c3nraA_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of an aspartate aminotransferase (yp_354942.1) from2 rhodobacter sphaeroides 2.4.1 at 2.15 a resolution |
| 39 | c3kaxB_ | Alignment | not modelled | 100.0 | 18 | PDB header: lyase Chain: B: PDB Molecule: aminotransferase, classes i and ii; PDBTitle: crystal structure of a putative c-s lyase from bacillus anthracis |
| 40 | c2z61A_ | Alignment | not modelled | 100.0 | 28 | PDB header: transferase Chain: A: PDB Molecule: probable aspartate aminotransferase 2; PDBTitle: crystal structure of mj0684 from methanococcus jannaschii2 reveals its similarity in the active site to kynurenine3 aminotransferases |
| 41 | c3pdxA_ | Alignment | not modelled | 100.0 | 23 | PDB header: transferase Chain: A: PDB Molecule: tyrosine aminotransferase; PDBTitle: crystal structural of mouse tyrosine aminotransferase |
| 42 | c2douA_ | Alignment | not modelled | 100.0 | 20 | PDB header: transferase Chain: A: PDB Molecule: probable n-succinyldiaminopimelate aminotransferase; PDBTitle: probable n-succinyldiaminopimelate aminotransferase (ttha0342) from2 thermus thermophilus hb8 |
| 43 | c3asaA_ | Alignment | not modelled | 100.0 | 21 | PDB header: transferase Chain: A: PDB Molecule: II-di-aminopimelate aminotransferase; PDBTitle: crystal structure of apo-II-di-aminopimelate aminotransferase from2 chlamydia trachomatis |
| 44 | c2x5fB_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: aspartate_tyrosine_phenylalanine pyridoxal-5' PDBTitle: crystal structure of the methicillin-resistant2 staphylococcus aureus sar2028, an3 aspartate_tyrosine_phenylalanine pyridoxal-5'-phosphate4 dependent aminotransferase |
| 45 | c3ez1A_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase mocr family; PDBTitle: crystal structure of putative aminotransferase (mocr family)2 (yp_604413.1) from deinococcus geothermalis dsm 11300 at 2.60 a3 resolution |
| 46 | d1d2fa_ | Alignment | not modelled | 100.0 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 47 | c1d2fB_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: B: PDB Molecule: maly protein; PDBTitle: x-ray structure of maly from escherichia coli: a pyridoxal-5'-2 phosphate-dependent enzyme acting as a modulator in mal gene3 expression |
| 48 | c3dc1A_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: A: PDB Molecule: kynurenine/alpha-aminoadipate aminotransferase PDBTitle: crystal structure of kynurenine aminotransferase ii complex with2 alpha-ketoglutarate |
| 49 | d1v2da_ | Alignment | not modelled | 100.0 | 23 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 50 | c3t18D_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: D: PDB Molecule: aminotransferase class i and ii; PDBTitle: crystal structure of aminotransferase from anaerococcus prevotii dsm2 20548. |
| | | | | | | Fold: PLP-dependent transferase-like |

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|----|-------------------------|-----------|--------------|-------|----|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 51 | d2csta_ | Alignment | not modelled | 100.0 | 13 | Superfamily: PLP-dependent transferases Family: AAT-like |
| 52 | c3k7yA_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: aspartate aminotransferase of plasmodium falciparum |
| 53 | c3d6kB_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: B: PDB Molecule: putative aminotransferase; PDBTitle: the crystal structure of a putative aminotransferase from2 corynebacterium diphtheriae |
| 54 | c3rq1A_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase class i and ii; PDBTitle: crystal structure of aminotransferase class i and ii from veillonella2 parvula |
| 55 | c3g7qA_ | Alignment | not modelled | 100.0 | 21 | PDB header: transferase Chain: A: PDB Molecule: valine-pyruvate aminotransferase; PDBTitle: crystal structure of valine-pyruvate aminotransferase avta2 (np_462565.1) from salmonella typhimurium lt2 at 1.80 a resolution |
| 56 | d7aata_ | Alignment | not modelled | 100.0 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 57 | d1ajsa_ | Alignment | not modelled | 100.0 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 58 | c3cbfA_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: A: PDB Molecule: alpha-aminodipate aminotransferase; PDBTitle: crystal structure of lysn, alpha-aminodipate2 aminotransferase, from thermus thermophilus hb27 |
| 59 | d2ay1a_ | Alignment | not modelled | 100.0 | 14 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 60 | d1yaaa_ | Alignment | not modelled | 100.0 | 14 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 61 | c2zy3A_ | Alignment | not modelled | 100.0 | 15 | PDB header: lyase Chain: A: PDB Molecule: l-aspartate beta-decarboxylase; PDBTitle: dodecameric l-aspartate beta-decarboxylase |
| 62 | d2q7wa1 | Alignment | not modelled | 100.0 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 63 | c3op7A_ | Alignment | not modelled | 100.0 | 25 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase class i and ii; PDBTitle: crystal structure of a plp-dependent aminotransferase (zp_03625122.1)2 from streptococcus suis 89-1591 at 1.70 a resolution |
| 64 | c3f6tA_ | Alignment | not modelled | 100.0 | 21 | PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of aspartate aminotransferase (e.c. 2.6.1.1)2 (yp_194538.1) from lactobacillus acidophilus ncfm at 2.15 a3 resolution |
| 65 | d3tata_ | Alignment | not modelled | 100.0 | 17 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 66 | c3mebB_ | Alignment | not modelled | 100.0 | 11 | PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: structure of cytoplasmic aspartate aminotransferase from giardia2 lamblia |
| 67 | c3ly1C_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: C: PDB Molecule: putative histidinol-phosphate aminotransferase; PDBTitle: crystal structure of putative histidinol-phosphate aminotransferase2 (yp_050345.1) from erwinia carotovora atroseptica scri1043 at 1.80 a3 resolution |
| 68 | d1lc5a_ | Alignment | not modelled | 100.0 | 20 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 69 | c3getA_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: crystal structure of putative histidinol-phosphate aminotransferase2 (np_281508.1) from campylobacter jejuni at 2.01 a resolution |
| 70 | c3cq6E_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: E: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: histidinol-phosphate aminotransferase from corynebacterium2 glutamicum holo-form (plp covalently bound) |
| 71 | c3eucB_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: B: PDB Molecule: histidinol-phosphate aminotransferase 2; PDBTitle: crystal structure of histidinol-phosphate aminotransferase2 (yp_297314.1) from ralstonia eutropha jmp134 at 2.05 a resolution |
| 72 | c3ftbA_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: the crystal structure of the histidinol-phosphate2 aminotransferase from clostridium acetobutylicum |
| 73 | c3bwnF_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: F: PDB Molecule: l-tryptophan aminotransferase; PDBTitle: l-tryptophan aminotransferase |
| 74 | c3hdoB_ | Alignment | not modelled | 100.0 | 17 | PDB header: transferase Chain: B: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: crystal structure of a histidinol-phosphate aminotransferase from2 geobacter metallireducens |
| 75 | c3fkdC_ | Alignment | not modelled | 100.0 | 18 | PDB header: lyase Chain: C: PDB Molecule: l-threonine-o-3-phosphate decarboxylase; PDBTitle: the crystal structure of l-threonine-o-3-phosphate2 decarboxylase from porphyromonas gingivalis |

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|-----|-------------------------|-----------|--------------|-------|----|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 76 | d1fg7a_ | Alignment | not modelled | 100.0 | 18 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 77 | d2f8ja1 | Alignment | not modelled | 100.0 | 20 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 78 | c3ffhA_ | Alignment | not modelled | 100.0 | 16 | PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: the crystal structure of histidinol-phosphate aminotransferase from2 listeria innocua clip11262. |
| 79 | d2hoxa1 | Alignment | not modelled | 100.0 | 16 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 80 | c3p1tB_ | Alignment | not modelled | 100.0 | 18 | PDB header: transferase Chain: B: PDB Molecule: putative histidinol-phosphate aminotransferase; PDBTitle: crystal structure of a putative aminotransferase (bpsl1724) from2 burkholderia pseudomallei k96243 at 2.60 a resolution |
| 81 | d2aeua1 | Alignment | not modelled | 100.0 | 16 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: SelA-like |
| 82 | c3f0hA_ | Alignment | not modelled | 100.0 | 9 | PDB header: transferase Chain: A: PDB Molecule: aminotransferase; PDBTitle: crystal structure of aminotransferase (rer070207000802) from2 eubacterium rectale at 1.70 a resolution |
| 83 | d1tpla_ | Alignment | not modelled | 100.0 | 10 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Beta-eliminating lyases |
| 84 | c2z9wA_ | Alignment | not modelled | 100.0 | 12 | PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of pyridoxamine-pyruvate aminotransferase complexed2 with pyridoxal |
| 85 | c3hqtB_ | Alignment | not modelled | 100.0 | 12 | PDB header: transferase Chain: B: PDB Molecule: cai-1 autoinducer synthase; PDBTitle: plp-dependent acyl-coa transferase cqsa |
| 86 | c3a2bA_ | Alignment | not modelled | 100.0 | 13 | PDB header: transferase Chain: A: PDB Molecule: serine palmitoyltransferase; PDBTitle: crystal structure of serine palmitoyltransferase from sphingobacterium2 multivorum with substrate l-serine |
| 87 | d1m6sa_ | Alignment | not modelled | 100.0 | 15 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 88 | d1vjoa_ | Alignment | not modelled | 100.0 | 10 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 89 | c2w8wA_ | Alignment | not modelled | 100.0 | 11 | PDB header: transferase Chain: A: PDB Molecule: serine palmitoyltransferase; PDBTitle: n100y spt with plp-ser |
| 90 | d1v72a1 | Alignment | not modelled | 100.0 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 91 | c3islA_ | Alignment | not modelled | 100.0 | 11 | PDB header: transferase Chain: A: PDB Molecule: purine catabolism protein pucg; PDBTitle: crystal structure of ureidoglycine-glyoxylate aminotransferase (pucg)2 from bacillus subtilis |
| 92 | d2e7ja1 | Alignment | not modelled | 100.0 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: SepSecS-like |
| 93 | c2dr1A_ | Alignment | not modelled | 100.0 | 15 | PDB header: transferase Chain: A: PDB Molecule: 386aa long hypothetical serine aminotransferase; PDBTitle: crystal structure of the ph1308 protein from pyrococcus horikoshii ot3 |
| 94 | c3nnkC_ | Alignment | not modelled | 100.0 | 12 | PDB header: transferase Chain: C: PDB Molecule: ureidoglycine-glyoxylate aminotransferase; PDBTitle: biochemical and structural characterization of a ureidoglycine2 aminotransferase in the klebsiella pneumoniae uric acid catabolic3 pathway |
| 95 | d2bwna1 | Alignment | not modelled | 100.0 | 11 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 96 | c3tqxA_ | Alignment | not modelled | 100.0 | 11 | PDB header: transferase Chain: A: PDB Molecule: 2-amino-3-ketobutyrate coenzyme a ligase; PDBTitle: structure of the 2-amino-3-ketobutyrate coenzyme a ligase (kbl) from2 coxiella burnetii |
| 97 | d1svva_ | Alignment | not modelled | 100.0 | 16 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like |
| 98 | c3mc6C_ | Alignment | not modelled | 100.0 | 12 | PDB header: lyase Chain: C: PDB Molecule: sphingosine-1-phosphate lyase; PDBTitle: crystal structure of scdpl1 |
| 99 | c2po3B_ | Alignment | not modelled | 100.0 | 16 | PDB header: transferase Chain: B: PDB Molecule: 4-dehydrase; PDBTitle: crystal structure analysis of desi in the presence of its2 tdp-sugar product |
| 100 | c2huuA_ | Alignment | not modelled | 100.0 | 13 | PDB header: transferase Chain: A: PDB Molecule: alanine glyoxylate aminotransferase; PDBTitle: crystal structure of aedes aegypti alanine glyoxylate2 aminotransferase in complex with alanine |
| | | | | | | Fold: PLP-dependent transferase-like |

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|-----|-------------------------|-----------|--------------|-------|----|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| 101 | d1t3ia_ | Alignment | not modelled | 100.0 | 16 | Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 102 | d1jf9a_ | Alignment | not modelled | 100.0 | 15 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 103 | c3caiA_ | Alignment | not modelled | 100.0 | 11 | PDB header: transferase Chain: A: PDB Molecule: possible aminotransferase; PDBTitle: crystal structure of mycobacterium tuberculosis rv3778c2 protein |
| 104 | c3ffrA_ | Alignment | not modelled | 100.0 | 9 | PDB header: transferase Chain: A: PDB Molecule: phosphoserine aminotransferase serc; PDBTitle: crystal structure of a phosphoserine aminotransferase serc (chu_0995)2 from cytophaga hutchinsonii atcc 33406 at 1.75 a resolution |
| 105 | d1bs0a_ | Alignment | not modelled | 100.0 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 106 | c3i4jC_ | Alignment | not modelled | 100.0 | 12 | PDB header: transferase Chain: C: PDB Molecule: aminotransferase, class iii; PDBTitle: crystal structure of aminotransferase, class iii from2 deinococcus radiodurans |
| 107 | c2ordA_ | Alignment | not modelled | 100.0 | 14 | PDB header: transferase Chain: A: PDB Molecule: acetylornithine aminotransferase; PDBTitle: crystal structure of acetylornithine aminotransferase (ec 2.6.1.11)2 (acoat) (tm1785) from thermotoga maritima at 1.40 a resolution |
| 108 | c2hdyA_ | Alignment | not modelled | 100.0 | 14 | PDB header: structural genomics, unknown function Chain: A: PDB Molecule: selenocysteine lyase; PDBTitle: structure of human selenocysteine lyase |
| 109 | d1c7ga_ | Alignment | not modelled | 100.0 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Beta-eliminating lyases |
| 110 | d2ch1a1 | Alignment | not modelled | 99.9 | 11 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 111 | d1sffa_ | Alignment | not modelled | 99.9 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 112 | d1b9ha_ | Alignment | not modelled | 99.9 | 14 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 113 | d1vefa1 | Alignment | not modelled | 99.9 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 114 | d2c0ra1 | Alignment | not modelled | 99.9 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 115 | c3mafB_ | Alignment | not modelled | 99.9 | 13 | PDB header: lyase Chain: B: PDB Molecule: sphingosine-1-phosphate lyase; PDBTitle: crystal structure of stspl (asymmetric form) |
| 116 | d1fc4a_ | Alignment | not modelled | 99.9 | 13 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like |
| 117 | c3nx3A_ | Alignment | not modelled | 99.9 | 10 | PDB header: transferase Chain: A: PDB Molecule: acetylornithine aminotransferase; PDBTitle: crystal structure of acetylornithine aminotransferase (argd) from2 campylobacter jejuni |
| 118 | c2eh6A_ | Alignment | not modelled | 99.9 | 16 | PDB header: transferase Chain: A: PDB Molecule: acetylornithine aminotransferase; PDBTitle: crystal structure of acetylornithine aminotransferase from aquifex2 aeolicus vf5 |
| 119 | d1eg5a_ | Alignment | not modelled | 99.9 | 11 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |
| 120 | d1h0ca_ | Alignment | not modelled | 99.9 | 12 | Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like |