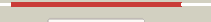
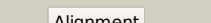
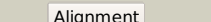
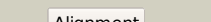
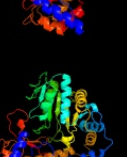


Phyre²

Email	mdejesus@rockefeller.edu
Description	RVBD1178 (-) _1309369_1310457
Date	Wed Jul 31 22:05:26 BST 2019
Unique Job ID	d0a846a75beacc28

Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c3tcmb			100.0	20	PDB header: transferase Chain: B; PDB Molecule: alanine aminotransferase 2; PDBTitle: crystal structure of alanine aminotransferase from hordeum vulgare
2	c3g0tA			100.0	17	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
3	c3h14A			100.0	25	PDB header: transferase Chain: A; PDB Molecule: aminotransferase, classes i and ii; PDBTitle: crystal structure of a putative aminotransferase from silicibacter2 pomeroyi
4	c3ezsB			100.0	24	PDB header: transferase Chain: B; PDB Molecule: aminotransferase aspb; PDBTitle: crystal structure of aminotransferase aspb (np_207418.1) from helicobacter pylori 26695 at 2.19 a resolution
5	c3e2yB			100.0	21	PDB header: transferase, lyase Chain: B; PDB Molecule: kynurene-oxoglutarate transaminase 3; PDBTitle: crystal structure of mouse kynurenine aminotransferase iii in complex2 with glutamine
6	c5yhvA			100.0	24	PDB header: transferase Chain: A; PDB Molecule: aminotransferase; PDBTitle: crystal structure of an aminotransferase from mycobacterium2 tuberculosis
7	c3b1dD			100.0	15	PDB header: lyase Chain: D; PDB Molecule: betac-s lyase; PDBTitle: crystal structure of betac-s lyase from streptococcus anginosus in2 complex with l-serine: external aldimine form
8	c4my5C			100.0	23	PDB header: transferase Chain: C; PDB Molecule: putative amino acid aminotransferase; PDBTitle: crystal structure of the aromatic amino acid aminotransferase from2 streptococcus mutants
9	d1o4sa			100.0	22	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
10	d1u08a			100.0	23	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
11	c3gguB			100.0	24	PDB header: transferase Chain: B; PDB Molecule: l-l-diaminopimelate aminotransferase; PDBTitle: l,l-diaminopimelate aminotransferase from chalmydomonas reinhardtii

12	d2r5ea1	Alignment		100.0	21	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferses Family: AAT-like
13	d1w7la	Alignment		100.0	21	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferses Family: AAT-like
14	c3eibB	Alignment		100.0	21	PDB header: transferase Chain: B: PDB Molecule: ll-diaminopimelate aminotransferase; PDBTitle: crystal structure of k270n variant of ll-diaminopimelate2 aminotransferase from arabidopsis thaliana
15	c4fl0A	Alignment		100.0	21	PDB header: transferase Chain: A: PDB Molecule: aminotransferase ald1; PDBTitle: crystal structure of ald1 from arabidopsis thaliana
16	c3jtxB	Alignment		100.0	29	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
17	c2douA	Alignment		100.0	29	PDB header: transferase Chain: A: PDB Molecule: probable n-succyldiaminopimelate aminotransferase; PDBTitle: probable n-succyldiaminopimelate aminotransferase (ttha0342) from2 thermus thermophilus hb8
18	c3b46B	Alignment		100.0	20	PDB header: transferase Chain: B: PDB Molecule: aminotransferase bna3; PDBTitle: crystal structure of bna3p, a putative kynurenone2 aminotransferase from saccharomyces cerevisiae
19	c1ynuA	Alignment		100.0	20	PDB header: lyase Chain: A: PDB Molecule: 1-aminocyclopropane-1-carboxylate synthase; PDBTitle: crystal structure of apple acc synthase in complex with L-vinylglycine
20	c2x5dD	Alignment		100.0	32	PDB header: transferase Chain: D: PDB Molecule: probable aminotransferase; PDBTitle: crystal structure of a probable aminotransferase from pseudomonas2 aeruginosa
21	c3pplB	Alignment	not modelled	100.0	17	PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of an aspartate transaminase (ncgl0237, cgj02402) from corynebacterium glutamicum atcc 13032 kitasato at 1.25 a3 resolution
22	c2o0rA	Alignment	not modelled	100.0	28	PDB header: transferase Chain: A: PDB Molecule: rv0858c (n-succyldiaminopimelate aminotransferase); PDBTitle: the three-dimensional structure of n-succyldiaminopimelate2 aminotransferase from mycobacterium tuberculosis
23	c3dzzB	Alignment	not modelled	100.0	17	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
24	d2gb3a1	Alignment	not modelled	100.0	23	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferses Family: AAT-like
25	c6f35B	Alignment	not modelled	100.0	19	PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase b; PDBTitle: crystal structure of the aspartate aminotranferase from rhizobium2 meliloti
26	c2o1bA	Alignment	not modelled	100.0	27	PDB header: transferase Chain: A: PDB Molecule: aminotransferase, class i; PDBTitle: structure of aminotransferase from staphylococcus aureus
27	c3if2B	Alignment	not modelled	100.0	18	PDB header: transferase 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
28	c3l8aB	Alignment	not modelled	100.0	PDB header: lyase Chain: B: PDB Molecule: putative aminotransferase, probable beta-cystathionase; PDBTitle: crystal structure of metc from streptococcus mutans
29	d1j32a	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
30	d1b5pa	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
31	c3eleB	Alignment	not modelled	100.0	PDB header: transferase Chain: B: PDB Molecule: amino transferase; PDBTitle: crystal structure of amino transferase (rer070207001803) from2 eubacterium rectale at 2.10 a resolution
32	c3asaA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: ll-diaminopimelate aminotransferase; PDBTitle: crystal structure of apo-ll-diaminopimelate aminotransferase from2 chlamydia trachomatis
33	c2zc0C	Alignment	not modelled	100.0	PDB header: transferase Chain: C: PDB Molecule: alanine glyoxylate transaminase; PDBTitle: crystal structure of an archaeal alanine:glyoxylate aminotransferase
34	c4je5C	Alignment	not modelled	100.0	PDB header: transferase Chain: C: PDB Molecule: aromatic/amino adipate aminotransferase 1; PDBTitle: crystal structure of the aromatic aminotransferase aro8, a putative2 alpha-amino adipate aminotransferase in saccharomyces cerevisiae
35	d1iaya	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like
36	c6f77D	Alignment	not modelled	100.0	PDB header: transferase Chain: D: PDB Molecule: aspartate aminotransferase a; PDBTitle: crystal structure of the prephenate aminotransferase from rhizobium2 melliloti
37	d1wsta1	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
38	c5c6uA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: aminotransferase; PDBTitle: rv3722c aminotransferase from mycobacterium tuberculosis
39	d1m7ya	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: GABA-aminotransferase-like
40	c4dgta	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: putative pyridoxal phosphate-dependent transferase; PDBTitle: crystal structure of plp-bound putative aminotransferase from2 clostridium difficile 630 crystallized with magnesium formate
41	c6hnuA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: aromatic amino acid aminotransferase i; PDBTitle: crystal structure of the aminotransferase aro8 from c. albicans with2 ligands
42	d1vp4a	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
43	c6hndA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: aromatic-amino-acid:2-oxoglutarate transaminase; PDBTitle: crystal structure of the aromatic aminotransferase aro9 from c.2 albicans
44	d1c7na	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathionine synthase-like
45	d1xi9a	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
46	c5wmIA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: bifunctional aspartate aminotransferase and PDBTitle: arabidopsis thaliana prephenate aminotransferase mutant- t84v
47	c3ihjA	Alignment	not modelled	100.0	PDB header: transferase Chain: A: PDB Molecule: alanine aminotransferase 2; PDBTitle: human alanine aminotransferase 2 in complex with plp
48	c3kaxB	Alignment	not modelled	100.0	PDB header: lyase Chain: B: PDB Molecule: aminotransferase, classes i and ii; PDBTitle: crystal structure of a putative c-s lyase from bacillus anthracis
49	c3ez1A	Alignment	not modelled	100.0	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
50	d1gdea	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
51	c3t18D	Alignment	not modelled	100.0	PDB header: transferase Chain: D: PDB Molecule: aminotransferase class i and ii; PDBTitle: crystal structure of aminotransferase from anaerococcus prevotii dsm2 20548.
52	d1x0ma1	Alignment	not modelled	100.0	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like

53	c4cvqB	Alignment	not modelled	100.0	19	PDB header: transferase Chain: B: PDB Molecule: glutamate-pyruvate aminotransferase alaa; PDBTitle: crystal structure of an aminotransferase from escherichia coli at 2.2 1 angstroem resolution
54	c2z61A	Alignment	not modelled	100.0	18	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
55	c5z0qG	Alignment	not modelled	100.0	21	PDB header: transferase Chain: G: PDB Molecule: aminotransferase, class i and ii; PDBTitle: crystal structure of ovob
56	c3k7yA	Alignment	not modelled	100.0	17	PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: aspartate aminotransferase of plasmodium falciparum
57	c3nraA	Alignment	not modelled	100.0	20	PDB header: transferase Chain: A: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of an aspartate aminotransferase (yp_354942.1) from2 rhodobacter sphaerooides 2.4.1 at 2.15 a resolution
58	c3fdbA	Alignment	not modelled	100.0	17	PDB header: transferase Chain: A: PDB Molecule: putative plp-dependent beta-cystathionease; PDBTitle: crystal structure of a putative plp-dependent beta-cystathionease2 (aecd, dip1736) from corynebacterium diphtheriae at 1.99 a resolution
59	d1bw0a	Alignment	not modelled	100.0	20	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
60	d2csta	Alignment	not modelled	100.0	18	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
61	c4ix8B	Alignment	not modelled	100.0	18	PDB header: transferase Chain: B: PDB Molecule: tyrosine aminotransferase; PDBTitle: crystal structure of tyrosine aminotransferase from leishmania2 infantum
62	c3rq1A	Alignment	not modelled	100.0	16	PDB header: transferase Chain: A: PDB Molecule: aminotransferase class i and ii; PDBTitle: crystal structure of aminotransferase class i and ii from veillonella2 parvula
63	c6c3aB	Alignment	not modelled	100.0	15	PDB header: biosynthetic protein Chain: B: PDB Molecule: uncharacterized protein; PDBTitle: o2-, plp-dependent l-arginine hydroxylase rohp 4-hydroxy-2-2 ketoarginine complex
64	d1ajsa	Alignment	not modelled	100.0	18	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
65	c4h51B	Alignment	not modelled	100.0	15	PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of a putative aspartate aminotransferase from2 leishmania major friedlin
66	c3d6kB	Alignment	not modelled	100.0	17	PDB header: transferase Chain: B: PDB Molecule: putative aminotransferase; PDBTitle: the crystal structure of a putative aminotransferase from2 corynebacterium diphtheriae
67	c4rkda	Alignment	not modelled	100.0	16	PDB header: transferase Chain: A: PDB Molecule: aromatic amino acid aminotransferase; PDBTitle: psychrophilic aromatic amino acids aminotransferase from psychrobacter2 sp. b6 cocrystallized with aspartic acid
68	d7aata	Alignment	not modelled	100.0	17	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
69	c4n0bA	Alignment	not modelled	100.0	15	PDB header: transcription activator Chain: A: PDB Molecule: hth-type transcriptional regulatory protein gabr; PDBTitle: crystal structure of bacillus subtilis gabr, an autorepressor and2 transcriptional activator of gabt
70	c2x5fB	Alignment	not modelled	100.0	17	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
71	c4wd2A	Alignment	not modelled	100.0	20	PDB header: transferase Chain: A: PDB Molecule: aromatic-amino-acid transaminase tyrb; PDBTitle: crystal structure of an aromatic amino acid aminotransferase from2 burkholderia cenocepacia j2315
72	c1d2fB	Alignment	not modelled	100.0	17	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
73	c3dc1A	Alignment	not modelled	100.0	21	PDB header: transferase Chain: A: PDB Molecule: kynurenine/alpha-amino adipate aminotransferase PDBTitle: crystal structure of kynurenine aminotransferase ii complex with2 alpha-ketoglutarate
74	c3g7qA	Alignment	not modelled	100.0	19	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
75	d1v2da	Alignment	not modelled	100.0	29	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
						Fold: PLP-dependent transferase-like

76	d2ay1a_	Alignment	not modelled	100.0	18	Superfamily: PLP-dependent transferases Family: AAT-like
77	d1yaaa_	Alignment	not modelled	100.0	16	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
78	d1d2fa_	Alignment	not modelled	100.0	17	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: Cystathione synthase-like
79	c4effA_	Alignment	not modelled	100.0	22	PDB header: transferase Chain: A: PDB Molecule: aromatic-amino-acid aminotransferase; PDBTitle: crystal structure of aromatic-amino-acid aminotransferase from2 burkholderia pseudomallei
80	c4eu1A_	Alignment	not modelled	100.0	19	PDB header: transferase Chain: A: PDB Molecule: mitochondrial aspartate aminotransferase; PDBTitle: structure of a mitochondrial aspartate aminotransferase from2 trypanosoma brucei
81	c6ezlB_	Alignment	not modelled	100.0	20	PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: crystal structure of aspartate aminotransferase from trypanosoma cruzi2 at 2.07 angstrom resolution
82	c3ly1C_	Alignment	not modelled	100.0	20	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
83	d1lc5a_	Alignment	not modelled	100.0	18	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
84	d2q7wa1	Alignment	not modelled	100.0	18	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
85	c4r8dB_	Alignment	not modelled	100.0	19	PDB header: transferase Chain: B: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: crystal structure of rv1600 encoded aminotransferase in complex with2 plp-mes from mycobacterium tuberculosis
86	c2zy4F_	Alignment	not modelled	100.0	21	PDB header: lyase Chain: F: PDB Molecule: l-aspartate beta-decarboxylase; PDBTitle: dodecameric l-aspartate beta-decarboxylase
87	c3mebB_	Alignment	not modelled	100.0	18	PDB header: transferase Chain: B: PDB Molecule: aspartate aminotransferase; PDBTitle: structure of cytoplasmic aspartate aminotransferase from giardia lamblia
88	d3tata_	Alignment	not modelled	100.0	20	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
89	c3bwnF_	Alignment	not modelled	100.0	18	PDB header: transferase Chain: F: PDB Molecule: l-tryptophan aminotransferase; PDBTitle: l-tryptophan aminotransferase
90	c3cbfA_	Alignment	not modelled	100.0	23	PDB header: transferase Chain: A: PDB Molecule: alpha-aminodipate aminotransferase; PDBTitle: crystal structure of lysn, alpha-aminodipate2 aminotransferase, from thermus thermophilus hb27
91	c3getA_	Alignment	not modelled	100.0	16	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
92	d2f8ja1	Alignment	not modelled	100.0	18	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
93	c3pdxA_	Alignment	not modelled	100.0	20	PDB header: transferase Chain: A: PDB Molecule: tyrosine aminotransferase; PDBTitle: crystal structural of mouse tyrosine aminotransferase
94	c2zy3A_	Alignment	not modelled	100.0	19	PDB header: lyase Chain: A: PDB Molecule: l-aspartate beta-decarboxylase; PDBTitle: dodecameric l-aspartate beta-decarboxylase
95	c3dydB_	Alignment	not modelled	100.0	20	PDB header: transferase Chain: B: PDB Molecule: tyrosine aminotransferase; PDBTitle: human tyrosine aminotransferase
96	c4r2nA_	Alignment	not modelled	100.0	23	PDB header: transferase Chain: A: PDB Molecule: putative phenylalanine aminotransferase; PDBTitle: crystal structure of rv3772 in complex with its substrate
97	c3cq6E_	Alignment	not modelled	100.0	20	PDB header: transferase Chain: E: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: histidinol-phosphate aminotransferase from corynebacterium2 glutamicum holo-form (plp covalently bound)
98	c3f6tA_	Alignment	not modelled	100.0	19	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
99	c5dj3B_	Alignment	not modelled	100.0	16	PDB header: transferase Chain: B: PDB Molecule: plp-dependent l-arginine hydroxylase mppp; PDBTitle: structure of the plp-dependent l-arginine hydroxylase mppp with d-2 arginine bound
100	d1fg7a_	Alignment	not modelled	100.0	21	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transferases Family: AAT-like
						PDB header: transferase Chain: B: PDB Molecule: histidinol-phosphate aminotransferase 2;

101	c3eucB	Alignment	not modelled	100.0	22	PDBTitle: crystal structure of histidinol-phosphate aminotransferase2 (yp_297314.1) from ralstonia eutropha jmp134 at 2.05 a resolution PDB header: transferase Chain: B: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: crystal structure of a histidinol-phosphate aminotransferase from2 geobacter metallireducens
102	c3hdoB	Alignment	not modelled	100.0	24	PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase class i and ii; PDBTitle: crystal structure of a plp-dependent aminotransferase (zp_03625122.12) from streptococcus suis 89-1591 at 1.70 a resolution
103	c3op7A	Alignment	not modelled	100.0	20	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transfers Family: AAT-like
104	d2hoxa1	Alignment	not modelled	100.0	16	PDB header: transferase Chain: A: PDB Molecule: probable histidinol-phosphate aminotransferase; PDBTitle: crystal structure of histidinol-phosphate aminotransferase from2 sinorhizobium meliloti in complex with pyridoxal-5'-phosphate
105	c4wbtA	Alignment	not modelled	100.0	18	PDB header: transferase Chain: A: PDB Molecule: aminotransferase; PDBTitle: crystal structure of plp-dependent cyclase orfr in complex with au
106	c4m2jA	Alignment	not modelled	100.0	21	PDB header: transferase Chain: A: PDB Molecule: aminotransferase; PDBTitle: crystal structure of plp-dependent cyclase orfr in complex with au
107	c3ffhA	Alignment	not modelled	100.0	19	PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: the crystal structure of histidinol-phosphate aminotransferase from2 listeria innocua clip11262.
108	c3ftbA	Alignment	not modelled	100.0	13	PDB header: transferase Chain: A: PDB Molecule: histidinol-phosphate aminotransferase; PDBTitle: the crystal structure of the histidinol-phosphate aminotransferase2 from clostridium acetobutylicum
109	c3fkdc	Alignment	not modelled	100.0	15	PDB header: lyase Chain: C: PDB Molecule: l-threonine-o-3-phosphate decarboxylase; PDBTitle: the crystal structure of l-threonine-o-3-phosphate decarboxylase from2 porphyromonas gingivalis
110	c3p1tB	Alignment	not modelled	100.0	23	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
111	c6ouxB	Alignment	not modelled	100.0	17	PDB header: lyase Chain: B: PDB Molecule: threonine phosphate decarboxylase-like enzyme; PDBTitle: structure of smul_1544, a decarboxylase from sulfurospirillum2 multivorans
112	d2aeua1	Alignment	not modelled	100.0	14	Fold: PLP-dependent transferase-like Superfamily: PLP-dependent transfers Family: SelA-like
113	c5jayB	Alignment	not modelled	100.0	17	PDB header: transferase Chain: B: PDB Molecule: 8-amino-7-oxononanoate synthase; PDBTitle: crystal structure of an 8-amino-7-oxononanoate synthase from2 burkholderia xenovorans
114	c2eh6A	Alignment	not modelled	100.0	17	PDB header: transferase Chain: A: PDB Molecule: acetylornithine aminotransferase; PDBTitle: crystal structure of acetylornithine aminotransferase from aquifex2 aeolicus vf5
115	c3wy7D	Alignment	not modelled	100.0	21	PDB header: transferase Chain: D: PDB Molecule: 8-amino-7-oxononanoate synthase; PDBTitle: crystal structure of mycobacterium smegmatis 7-keto-8-aminopalargonic2 acid (kapa) synthase biof
116	c3hqtB	Alignment	not modelled	100.0	14	PDB header: transferase Chain: B: PDB Molecule: cai-1 autoinducer synthase; PDBTitle: plp-dependent acyl-coa transferase cqsA
117	c3a2bA	Alignment	not modelled	100.0	18	PDB header: transferase Chain: A: PDB Molecule: serine palmitoyltransferase; PDBTitle: crystal structure of serine palmitoyltransferase from sphingobacterium2 multivorans with substrate l-serine
118	c2w8wA	Alignment	not modelled	100.0	20	PDB header: transferase Chain: A: PDB Molecule: serine palmitoyltransferase; PDBTitle: n100y spt with plp-ser
119	c4iw7A	Alignment	not modelled	100.0	14	PDB header: transferase Chain: A: PDB Molecule: 8-amino-7-oxononanoate synthase; PDBTitle: crystal structure of 8-amino-7-oxononanoate synthase (biof) from2 francisella tularensis.
120	c3nx3A	Alignment	not modelled	100.0	13	PDB header: transferase Chain: A: PDB Molecule: acetylornithine aminotransferase; PDBTitle: crystal structure of acetylornithine aminotransferase (argd) from2 campylobacter jejuni