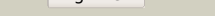
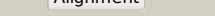
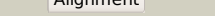
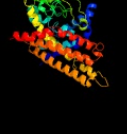


Phyre²

| | |
|---------------|-----------------------------------|
| Email | mdejesus@rockefeller.edu |
| Description | RVBD3140_(fadE23)_3506787_3507992 |
| Date | Thu Aug 8 16:20:32 BST 2019 |
| Unique Job ID | 3a152c3c0f74677d |

Detailed template information

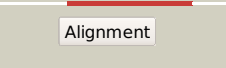
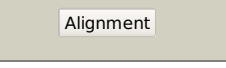
| # | Template | Alignment Coverage | 3D Model | Confidence | % i.d. | Template Information |
|----|-------------------------|---|---|------------|--------|--|
| 1 | c2a1tC_ |  |  | 100.0 | 30 | PDB header: oxidoreductase/electron transport Chain: C: PDB Molecule: acyl-coa dehydrogenase, medium-chain specific, PDBTitle: structure of the human mcad:etf e165betaa complex |
| 2 | c2ix5A_ |  |  | 100.0 | 22 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coenzyme a oxidase 4, peroxisomal; PDBTitle: short chain specific acyl-coa oxidase from arabidopsis thaliana, acx42 in complex with acetoacetyl-coa |
| 3 | c6es9A_ |  |  | 100.0 | 28 | PDB header: flavoprotein Chain: A: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: methylsuccinyl-coa dehydrogenase of paracoccus denitrificans with2 bound flavin adenine dinucleotide |
| 4 | c5zw2A_ |  |  | 100.0 | 22 | PDB header: biosynthetic protein Chain: A: PDB Molecule: l-prolyl-[peptidyl-carrier protein] dehydrogenase; PDBTitle: fad complex of piga |
| 5 | c2z1qA_ |  |  | 100.0 | 27 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of acyl coa dehydrogenase |
| 6 | clegcB_ |  |  | 100.0 | 29 | PDB header: electron transfer Chain: B: PDB Molecule: medium chain acyl-coa dehydrogenase; PDBTitle: structure of t255e, e376g mutant of human medium chain acyl-2 coa dehydrogenase complexed with octanoyl-coa |
| 7 | c1rx0B_ |  |  | 100.0 | 21 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase family member 8, mitochondrial; PDBTitle: crystal structure of isobutyryl-coa dehydrogenase complexed with2 substrate/ligand. |
| 8 | c3sf6A_ |  |  | 100.0 | 23 | PDB header: oxidoreductase Chain: A: PDB Molecule: glutaryl-coa dehydrogenase; PDBTitle: crystal structure of glutaryl-coa dehydrogenase from mycobacterium2 smegmatis |
| 9 | c3swoA_ |  |  | 100.0 | 23 | PDB header: oxidoreductase Chain: A: PDB Molecule: glutaryl-coa dehydrogenase; PDBTitle: crystal structure of a glutaryl-coa dehydrogenase from mycobacterium2 smegmatis in complex with fadh2 |
| 10 | c3owaC_ |  |  | 100.0 | 25 | PDB header: oxidoreductase Chain: C: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of acyl-coa dehydrogenase complexed with fad from2 bacillus anthracis |
| 11 | c4n5fA_ |  |  | 100.0 | 32 | PDB header: oxidoreductase Chain: A: PDB Molecule: putative acyl-coa dehydrogenase; PDBTitle: crystal structure of a putative acyl-coa dehydrogenase with bound2 fadh2 from burkholderia cenocepacia j235 |

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|----|-------------------------|-----------|---|-------|----|--|
| 12 | c4hr3A_ | Alignment |  | 100.0 | 23 | PDB header: oxidoreductase Chain: A: PDB Molecule: putative acyl-coa dehydrogenase; PDBTitle: structure of a putative acyl-coa dehydrogenase from mycobacterium2 abscessus |
| 13 | c1siqA_ | Alignment |  | 100.0 | 20 | PDB header: oxidoreductase Chain: A: PDB Molecule: glutaryl-coa dehydrogenase; PDBTitle: the crystal structure and mechanism of human glutaryl-coa2 dehydrogenase |
| 14 | c4irnF_ | Alignment |  | 100.0 | 26 | PDB header: oxidoreductase Chain: F: PDB Molecule: prolyl-acp dehydrogenase; PDBTitle: crystal structure of the prolyl acyl carrier protein oxidase anab |
| 15 | c2cx9C_ | Alignment |  | 100.0 | 29 | PDB header: oxidoreductase Chain: C: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of acyl-coa dehydrogenase |
| 16 | c1vhD_ | Alignment |  | 100.0 | 23 | PDB header: oxidoreductase Chain: D: PDB Molecule: isovaleryl-coa dehydrogenase; PDBTitle: structure of human isovaleryl-coa dehydrogenase at 2.62 angstroms resolution: structural basis for substrate3 specificity |
| 17 | c2pg0B_ | Alignment |  | 100.0 | 25 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of acyl-coa dehydrogenase from geobacillus2 kaustophilus |
| 18 | c5ahsB_ | Alignment |  | 100.0 | 22 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: 3-sulfinopropionyl-coenzyme a (3sp-coa) desulfinase from advenella2 mimgardefordensis dpn7t: holo crystal structure with the substrate3 analog succinyl-coa |
| 19 | c1ukwA_ | Alignment |  | 100.0 | 30 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of medium-chain acyl-coa dehydrogenase2 from thermus thermophilus hb8 |
| 20 | c6cy8B_ | Alignment |  | 100.0 | 28 | PDB header: biosynthetic protein Chain: B: PDB Molecule: butyryl-coa dehydrogenase; PDBTitle: crystal structure of fad-dependent dehydrogenase |
| 21 | c5ol2F_ | Alignment | not modelled | 100.0 | 33 | PDB header: flavoprotein Chain: F: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: the electron transferring flavoprotein/butyryl-coa dehydrogenase2 complex from clostridium difficile |
| 22 | c3oibB_ | Alignment | not modelled | 100.0 | 26 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of a putative acyl-coa dehydrogenase from2 mycobacterium smegmatis, iodide soak |
| 23 | c3r7kB_ | Alignment | not modelled | 100.0 | 25 | PDB header: oxidoreductase Chain: B: PDB Molecule: probable acyl coa dehydrogenase; PDBTitle: crystal structure of a probable acyl coa dehydrogenase from2 mycobacterium abscessus atcc 19977 / dsm 44196 |
| 24 | c4l1fB_ | Alignment | not modelled | 100.0 | 31 | PDB header: electron transport Chain: B: PDB Molecule: acyl-coa dehydrogenase domain protein; PDBTitle: electron transferring flavoprotein of acidaminococcus fermentans2 towards a mechanism of flavin-based electron bifurcation |
| 25 | c2jifA_ | Alignment | not modelled | 100.0 | 26 | PDB header: oxidoreductase Chain: A: PDB Molecule: short/branched chain specific acyl-coa dehydrogenase; PDBTitle: structure of human short-branched chain acyl-coa dehydrogenase2 (acadsb) |
| 26 | c2ebal_ | Alignment | not modelled | 100.0 | 21 | PDB header: oxidoreductase Chain: I: PDB Molecule: putative glutaryl-coa dehydrogenase; PDBTitle: crystal structure of the putative glutaryl-coa dehydrogenase from2 thermus thermophilus |
| 27 | c3eomD_ | Alignment | not modelled | 100.0 | 19 | PDB header: oxidoreductase Chain: D: PDB Molecule: glutaryl-coa dehydrogenase; PDBTitle: 2.4 a crystal structure of native glutaryl-coa dehydrogenase from2 burkholderia pseudomallei |
| | | | | | | PDB header: oxidoreductase Chain: A: PDB Molecule: very-long-chain specific acyl-coa |

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|----|------------------------|-----------|--------------|-------|----|--|
| 28 | c2uxwA | Alignment | not modelled | 100.0 | 27 | dehydrogenase; PDBTitle: crystal structure of human very long chain acyl-coa dehydrogenase2 (acadv1) |
| 29 | c3mkhC | Alignment | not modelled | 100.0 | 22 | PDB header: oxidoreductase Chain: C: PDB Molecule: nitroalkane oxidase; PDBTitle: podospora anserina nitroalkane oxidase |
| 30 | c1bucB | Alignment | not modelled | 100.0 | 31 | PDB header: oxidoreductase Chain: B: PDB Molecule: butyryl-coa dehydrogenase; PDBTitle: three-dimensional structure of butyryl-coa dehydrogenase from2 megasphaera elsdenii |
| 31 | c6fahD | Alignment | not modelled | 100.0 | 30 | PDB header: flavoprotein Chain: D: PDB Molecule: caffeyl-coa reductase-etf complex subunit carc; PDBTitle: molecular basis of the flavin-based electron-bifurcating caffeyl-coa2 reductase reaction |
| 32 | c2vigC | Alignment | not modelled | 100.0 | 32 | PDB header: oxidoreductase Chain: C: PDB Molecule: short-chain specific acyl-coa dehydrogenase,; PDBTitle: crystal structure of human short-chain acyl coa dehydrogenase |
| 33 | c4iv6A | Alignment | not modelled | 100.0 | 28 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase fade3; PDBTitle: x-ray crystal structure of an isovaleryl-coa dehydrogenase from2 mycobacterium smegmatis |
| 34 | c3mpjG | Alignment | not modelled | 100.0 | 26 | PDB header: oxidoreductase Chain: G: PDB Molecule: glutaryl-coa dehydrogenase; PDBTitle: structure of the glutaryl-coenzyme a dehydrogenase |
| 35 | c6ijcA | Alignment | not modelled | 100.0 | 24 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase family protein; PDBTitle: structure of mmpa-coa dehydrogenase from roseovarius rubinhibens lsm |
| 36 | c2rehD | Alignment | not modelled | 100.0 | 18 | PDB header: oxidoreductase Chain: D: PDB Molecule: nitroalkane oxidase; PDBTitle: mechanistic and structural analyses of the roles of arg4092 and asp402 in the reaction of the flavoprotein nitroalkane3 oxidase |
| 37 | c4u83A | Alignment | not modelled | 100.0 | 31 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: structure of brucella abortus butyryl-coa dehydrogenase |
| 38 | c3nf4B | Alignment | not modelled | 100.0 | 31 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of acyl-coa dehydrogenase from mycobacterium2 thermoresistibile bound to flavin adenine dinucleotide |
| 39 | c4ktob | Alignment | not modelled | 100.0 | 23 | PDB header: oxidoreductase Chain: B: PDB Molecule: isovaleryl-coa dehydrogenase; PDBTitle: crystal structure of a putative isovaleryl-coa dehydrogenase (psi-2 nysgrc-012251) from sinorhizobium meliloti 1021 |
| 40 | c2dvlB | Alignment | not modelled | 100.0 | 30 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of project tt0160 from thermus thermophilus hb8 |
| 41 | c5iduB | Alignment | not modelled | 100.0 | 29 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase domain protein; PDBTitle: crystal structure of an acyl-coa dehydrogenase domain protein from2 burkholderia phymatum bound to fad |
| 42 | c4w9uD | Alignment | not modelled | 100.0 | 20 | PDB header: oxidoreductase Chain: D: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of an acyl-coa dehydrogenase from brucella2 melitensis |
| 43 | c3pfdB | Alignment | not modelled | 100.0 | 31 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of an acyl-coa dehydrogenase from mycobacterium2 thermoresistibile bound to reduced flavin adenine dinucleotide solved3 by combined iodide ion sad mr |
| 44 | c2wbIB | Alignment | not modelled | 100.0 | 20 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase family member 11; PDBTitle: crystal structure of human acyl-coa dehydrogenase 11 |
| 45 | c1r2jA | Alignment | not modelled | 100.0 | 22 | PDB header: oxidoreductase Chain: A: PDB Molecule: protein fkbi; PDBTitle: fkbi for biosynthesis of methoxymalonyl extender unit of fk5202 polyketide immunosupresant |
| 46 | c4x28B | Alignment | not modelled | 100.0 | 22 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of the chse4-chse5 complex from mycobacterium2 tuberculosis |
| 47 | c4rm7A | Alignment | not modelled | 100.0 | 19 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: the crystal structure of acyl-coa dehydrogenase from slackia2 heliotrinireducens dsm 20476 |
| 48 | c5lnxC | Alignment | not modelled | 100.0 | 29 | PDB header: oxidoreductase Chain: C: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of mmgc, an acyl-coa dehydrogenase from bacillus2 subtilis. |
| 49 | c5gj7A | Alignment | not modelled | 100.0 | 19 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa dehydrogenase type 2 domain protein; PDBTitle: putative acyl-coa dehydrogenase |
| 50 | c4doyE | Alignment | not modelled | 100.0 | 18 | PDB header: oxidoreductase Chain: E: PDB Molecule: dibenzothiophene desulfurization enzyme c; PDBTitle: crystal structure of dibenzothiophene desulfurization enzyme c |
| 51 | c4m6zB | Alignment | not modelled | 100.0 | 18 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-[acyl-carrier-protein] dehydrogenase mbtn; PDBTitle: crystal structure of an acyl-acp dehydrogenase |
| 52 | c5o73B | Alignment | not modelled | 100.0 | 21 | PDB header: oxidoreductase Chain: B: PDB Molecule: acyl-coa dehydrogenase; |

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|----|-------------------------|-----------|--------------|-------|----|---|
| 52 | c5e2D | Alignment | not modelled | 100.0 | 21 | PDBTitle: crystal structure acyl-coa dehydrogenase from brucella melitensis in2 complex with fad PDB header: oxidoreductase Chain: B: PDB Molecule: thermophilic dibenzothiophene desulfurization enzyme c; PDBTitle: crystal structure of indole-bound tdsc from paenibacillus sp. a11-2 |
| 53 | c5xdcB | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: A: PDB Molecule: p-hydroxyphenylacetate hydroxylase c2:oxygenase PDBTitle: structure of the monooxygenase component of p-2 hydroxyphenylacetate hydroxylase from acinetobacter3 baumannii |
| 54 | c2jbtA | Alignment | not modelled | 100.0 | 15 | PDB header: oxidoreductase Chain: A: PDB Molecule: fad-dependent oxidoreductase; PDBTitle: x-ray structure of a kjjd3 in complex with dtdp |
| 55 | c3m9vA | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: D: PDB Molecule: acyl-coa dehydrogenase; PDBTitle: crystal structure of the chse4-chse5 complex from mycobacterium2 tuberculosis |
| 56 | c4x28D | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: R: PDB Molecule: xiaf protein; PDBTitle: xiaf from streptomyces sp. in complex with fadh2 and glycerol |
| 57 | c5mr6R | Alignment | not modelled | 100.0 | 13 | PDB header: oxidoreductase Chain: B: PDB Molecule: protein acdh-11, isoform b; PDBTitle: crystal structure of caenorhabditis elegans acdh-11 |
| 58 | c4y9IB | Alignment | not modelled | 100.0 | 22 | PDB header: oxidoreductase Chain: A: PDB Molecule: protein aidb; PDBTitle: crystal structure of alkylation response protein e. coli aidb |
| 59 | c3djIA | Alignment | not modelled | 100.0 | 20 | PDB header: oxidoreductase Chain: A: PDB Molecule: 3-hsa hydroxylase, oxygenase; PDBTitle: crystal structure of 3-hsa hydroxylase from rhodococcus sp. rha1 |
| 60 | c2rfqA | Alignment | not modelled | 100.0 | 13 | PDB header: oxidoreductase Chain: B: PDB Molecule: hydroxylase; PDBTitle: structural genomics, the crystal structure of a putative hydroxylase2 from rhodococcus sp. rha1 |
| 61 | c2or0B | Alignment | not modelled | 100.0 | 15 | PDB header: oxidoreductase Chain: B: PDB Molecule: dnmz; PDBTitle: streptomyces peucetius nitrososynthase dnmz in ligand-free state |
| 62 | c4zxvB | Alignment | not modelled | 100.0 | 16 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa oxidase; PDBTitle: crystal structure of acyl-coa oxidase complexed with 3-oh-dodecanoate |
| 63 | c2ddhA | Alignment | not modelled | 100.0 | 19 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coa oxidase; PDBTitle: arabidopsis thaliana acyl-coa oxidase 1 |
| 64 | c1w07A | Alignment | not modelled | 100.0 | 23 | PDB header: oxidoreductase Chain: H: PDB Molecule: acyl-coenzyme a oxidase; PDBTitle: crystal structure of acyl-coa oxidase-1 in caenorhabditis elegans2 bound with fad, ascaroside-coa, and atp |
| 65 | c5k3jA | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coenzyme a oxidase; PDBTitle: crystals structure of acyl-coa oxidase-2 in caenorhabditis elegans2 bound with fad, ascaroside-coa, and atp |
| 66 | c2fonA | Alignment | not modelled | 100.0 | 23 | PDB header: oxidoreductase Chain: A: PDB Molecule: peroxisomal acyl-coa oxidase 1a; PDBTitle: x-ray crystal structure of leacx1, an acyl-coa oxidase from2 lycopersicon esculentum (tomato) |
| 67 | c5k3iH | Alignment | not modelled | 100.0 | 19 | PDB header: oxidoreductase Chain: H: PDB Molecule: acyl-coenzyme a oxidase; PDBTitle: crystal structure of acyl-coa oxidase-1 in caenorhabditis elegans2 complexed with fad and atp |
| 68 | c3mxLB | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: B: PDB Molecule: nitrososynthase; PDBTitle: crystal structure of nitrososynthase from micromonospora carbonacea2 var. africana |
| 69 | c5y9dA | Alignment | not modelled | 100.0 | 21 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coenzyme a oxidase 1; PDBTitle: crystal structure of acyl-coa oxidase1 from yarrowia lipolytica |
| 70 | c5ys9A | Alignment | not modelled | 100.0 | 21 | PDB header: oxidoreductase Chain: A: PDB Molecule: acyl-coenzyme a oxidase 3; PDBTitle: crystal structure of acyl-coa oxidase3 from yarrowia lipolytica |
| 71 | d1egda2 | Alignment | not modelled | 100.0 | 24 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 72 | d2d29a2 | Alignment | not modelled | 100.0 | 25 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 73 | d1rx0a2 | Alignment | not modelled | 100.0 | 21 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 74 | d3mdea2 | Alignment | not modelled | 100.0 | 26 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 75 | d1ukwa2 | Alignment | not modelled | 100.0 | 29 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 76 | d1jqia2 | Alignment | not modelled | 100.0 | 30 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 77 | d2c12a2 | Alignment | not modelled | 100.0 | 16 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |

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|-----|--------------------------|--|-----------|--------------|-------|----|---|
| 78 | d1buca2 | | Alignment | not modelled | 100.0 | 29 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 79 | d1ivha2 | | Alignment | not modelled | 100.0 | 23 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 80 | d1r2ja2 | | Alignment | not modelled | 100.0 | 24 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 81 | d1siqa2 | | Alignment | not modelled | 100.0 | 20 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and middle) domains |
| 82 | d2ddha3 | | Alignment | not modelled | 100.0 | 23 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: acyl-CoA oxidase N-terminal domains |
| 83 | c3hwcd | | Alignment | not modelled | 100.0 | 16 | PDB header: oxidoreductase Chain: D: PDB Molecule: chlorophenol-4-monoxygenase component 2; PDBTitle: crystal structure of chlorophenol 4-monoxygenase (tftd) of 2' burkholderia cepacia ac1100 |
| 84 | d1w07a3 | | Alignment | not modelled | 100.0 | 26 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: acyl-CoA oxidase N-terminal domains |
| 85 | c4g5eD | | Alignment | not modelled | 100.0 | 17 | PDB header: oxidoreductase Chain: D: PDB Molecule: 2,4,6-trichlorophenol 4-monoxygenase; PDBTitle: 2,4,6-trichlorophenol 4-monoxygenase |
| 86 | c2yyjA | | Alignment | not modelled | 100.0 | 14 | PDB header: oxidoreductase Chain: A: PDB Molecule: 4-hydroxyphenylacetate-3-hydroxylase; PDBTitle: crystal structure of the oxygenase component (hpab) of 4-2 hydroxyphenylacetate 3-monoxygenase complexed with fad and 4-3 hydroxyphenylacetate |
| 87 | c1u8vA | | Alignment | not modelled | 100.0 | 17 | PDB header: lyase, isomerase Chain: A: PDB Molecule: gamma-aminobutyrate metabolism dehydratase/isomerase; PDBTitle: crystal structure of 4-hydroxybutyryl-coa dehydratase from clostridium2 aminobutyricum: radical catalysis involving a [4fe-4s] cluster and3 flavin |
| 88 | d1siqa1 | | Alignment | not modelled | 100.0 | 17 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 89 | d3mdea1 | | Alignment | not modelled | 99.9 | 35 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 90 | d1legda1 | | Alignment | not modelled | 99.9 | 34 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 91 | d1ivha1 | | Alignment | not modelled | 99.9 | 19 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 92 | c6eb0A | | Alignment | not modelled | 99.9 | 13 | PDB header: oxidoreductase Chain: A: PDB Molecule: 4-hydroxyphenylacetate 3-monoxygenase, oxygenase subunit; PDBTitle: structure of 4-hydroxyphenylacetate 3-monoxygenase (hpab), oxygenase2 component from escherichia coli |
| 93 | d1buca1 | | Alignment | not modelled | 99.9 | 33 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 94 | d1qjial | | Alignment | not modelled | 99.9 | 32 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 95 | d1rx0a1 | | Alignment | not modelled | 99.9 | 20 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 96 | d1ukwa1 | | Alignment | not modelled | 99.9 | 32 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 97 | d2d29a1 | | Alignment | not modelled | 99.9 | 34 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 98 | d2c12a1 | | Alignment | not modelled | 99.9 | 20 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 99 | d2ddha1 | | Alignment | not modelled | 99.9 | 15 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: acyl-CoA oxidase C-terminal domains |
| 100 | d1r2ja1 | | Alignment | not modelled | 99.9 | 19 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: Medium chain acyl-CoA dehydrogenase-like, C-terminal domain |
| 101 | d1w07a1 | | Alignment | not modelled | 99.9 | 17 | Fold: Bromodomain-like Superfamily: Acyl-CoA dehydrogenase C-terminal domain-like Family: acyl-CoA oxidase C-terminal domains |
| 102 | d1u8va2 | | Alignment | not modelled | 99.8 | 15 | Fold: Acyl-CoA dehydrogenase NM domain-like Superfamily: Acyl-CoA dehydrogenase NM domain-like Family: Medium chain acyl-CoA dehydrogenase, NM (N-terminal and |

| | | | | | |
|-----|-------------------------|---|-----------|--------------|-----------------|
| | | | | | middle) domains |
| 103 | c4oo2D_ |  | Alignment | not modelled | 99.5 |
| 104 | d1u8va1 |  | Alignment | not modelled | 95.3 |