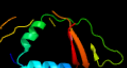
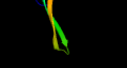


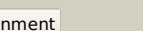
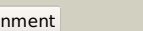
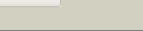
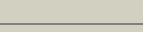
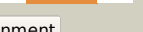
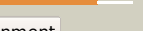
Phyre²

Email	mdejesus@rockefeller.edu
Description	RVBD1973 (-)_2217172_2217654
Date	Mon Aug 5 13:25:07 BST 2019
Unique Job ID	2162bc4cbe4b4bfcc

Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	d2ux0a1	Alignment		97.2	12	Fold: Cystatin-like Superfamily: NTF2-like Family: Association domain of calcium/calmodulin-dependent protein kinase type II alpha subunit, CAMK2A
2	c4i4kb_	Alignment		97.1	7	PDB header: structural genomics, unknown function Chain: B: PDB Molecule: uncharacterized protein sgcj; PDBTitle: streptomyces globisporus c-1027 9-membered enediyne conserved protein2 sgc6
3	c5d9rA_	Alignment		97.1	15	PDB header: biosynthetic protein Chain: A: PDB Molecule: protein accumulation and replication of chloroplasts 6, PDBTitle: crystal structure of a conserved domain in the intermembrane space2 region of the plastid division protein arc6
4	d3b8la1	Alignment		96.9	11	Fold: Cystatin-like Superfamily: NTF2-like Family: BaiE/LinA-like
5	d1hkxa_	Alignment		96.8	12	Fold: Cystatin-like Superfamily: NTF2-like Family: Association domain of calcium/calmodulin-dependent protein kinase type II alpha subunit, CAMK2A
6	c3h51A_	Alignment		96.5	15	PDB header: protein binding Chain: A: PDB Molecule: putative calcium/calmodulin dependent protein kinase ii PDBTitle: crystal structure of putative calcium/calmodulin dependent protein2 kinase ii association domain (np_636218.1) from xanthomonas3 campestris at 1.70 a resolution
7	c5cnlA_	Alignment		96.5	8	PDB header: protein transport Chain: A: PDB Molecule: icml-like; PDBTitle: crystal structure of an icml-like type iv secretion system protein2 (lpg0120) from legionella pneumophila subsp. pneumophila str.3 philadelphia 1 at 2.65 a resolution
8	c4ovmE_	Alignment		96.3	9	PDB header: unknown function Chain: E: PDB Molecule: uncharacterized protein sgcj; PDBTitle: crystal structure of sgcj protein from streptomyces carzinostaticus
9	d2rfra1	Alignment		96.3	17	Fold: Cystatin-like Superfamily: NTF2-like Family: BaiE/LinA-like
10	d1idpa_	Alignment		96.2	18	Fold: Cystatin-like Superfamily: NTF2-like Family: Scytalone dehydratase
11	c5u9oD_	Alignment		96.2	15	PDB header: cell cycle Chain: D: PDB Molecule: plastid division protein cdp1, chloroplastic,plastid PDBTitle: cocrystal structure of the intermembrane space region of the plastid2 division proteins parc6 and pdv1

12	d3cu3a1			96.1	8	Fold: Cystatin-like Superfamily: NTF2-like Family: BaiE/LinA-like
13	d2f86b1			96.1	11	Fold: Cystatin-like Superfamily: NTF2-like Family: Association domain of calcium/calmodulin-dependent protein kinase type II alpha subunit, CAMK2A
14	c3wz4F			95.9	13	PDB header: unknown function Chain: F; PDB Molecule: doti; PDBTitle: structure of the periplasmic domain of doti (crystal form i)
15	c5ig4A			95.9	13	PDB header: transferase Chain: A; PDB Molecule: predicted protein; PDBTitle: crystal structure of n. vectensis camkii-a hub
16	c3wz3A			95.8	11	PDB header: unknown function Chain: A; PDB Molecule: tram protein; PDBTitle: structure of a periplasmic fragment of tram
17	d3stda			95.6	16	Fold: Cystatin-like Superfamily: NTF2-like Family: Scytalone dehydratase
18	c5ig5E			95.4	13	PDB header: transferase Chain: E; PDB Molecule: camkii-b hub; PDBTitle: crystal structure of n. vectensis camkii-b hub at ph 4.2
19	d2rgqa1			95.3	20	Fold: Cystatin-like Superfamily: NTF2-like Family: Rv3472-like
20	d2chca1			95.2	13	Fold: Cystatin-like Superfamily: NTF2-like Family: Rv3472-like
21	c3a76B		not modelled	94.6	12	PDB header: lyase Chain: B; PDB Molecule: gamma-hexachlorocyclohexane dehydrochlorinase; PDBTitle: the crystal structure of lina
22	c3robC		not modelled	94.1	13	PDB header: structural genomics, unknown function Chain: C; PDB Molecule: uncharacterized conserved protein; PDBTitle: the crystal structure of a conserved protein from planctomycetes2 limnophilus dsm 3776
23	c6of9G		not modelled	93.9	20	PDB header: unknown function Chain: G; PDB Molecule: camkii hub; PDBTitle: structure of the chlamydamonas reinhardtii camkii hub homology domain
24	d3ef8a1		not modelled	93.7	17	Fold: Cystatin-like Superfamily: NTF2-like Family: BaiE/LinA-like
25	c4nhfF		not modelled	93.2	12	PDB header: protein transport Chain: F; PDB Molecule: trwg protein; PDBTitle: crystal structure of the soluble domain of trwg type iv secretion2 machinery from bartonella grahamii
26	c4gb5A		not modelled	93.1	13	PDB header: structural genomics, unknown function Chain: A; PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of kfia4162 protein from kribbella flava
27	d3ejva1		not modelled	93.1	27	Fold: Cystatin-like Superfamily: NTF2-like Family: BaiE/LinA-like
28	c3bb9D		not modelled	93.0	15	PDB header: unknown function Chain: D; PDB Molecule: putative orphan protein; PDBTitle: crystal structure of a putative ketosteroid isomerase (sfrI_1973) from2 shewanella frigidimarina ncimb 400 at 1.80 a resolution

29	d3d9ra1		Alignment	not modelled	92.9	21	Fold: Cystatin-like Superfamily: NTF2-like Family: ECA1476-like
30	c3soyA_		Alignment	not modelled	92.7	9	PDB header: membrane protein Chain: A: PDB Molecule: ntf2-like superfamily protein; PDBTitle: nuclear transport factor 2 (ntf2-like) superfamily protein from2 salmonella enterica subsp. enterica serovar typhimurium str. lt2
31	c3hx8A_		Alignment	not modelled	92.5	17	PDB header: isomerase Chain: A: PDB Molecule: putative ketosteroid isomerase; PDBTitle: crystal structure of putative ketosteroid isomerase (np_103587.1)from2 mesorhizobium loti at 1.45 a resolution
32	d2cc3a1		Alignment	not modelled	92.4	14	Fold: Cystatin-like Superfamily: NTF2-like Family: VirB8-like
33	c3f7sA_		Alignment	not modelled	92.3	16	PDB header: unknown function Chain: A: PDB Molecule: uncharacterized ntf2-like protein; PDBTitle: crystal structure of a ntf2-like protein of unknown function (pp_4556)2 from pseudomonas putida kt2440 at 2.11 a resolution
34	c2gbxF_		Alignment	not modelled	92.2	9	PDB header: oxidoreductase Chain: F: PDB Molecule: biphenyl 2,3-dioxygenase beta subunit; PDBTitle: crystal structure of biphenyl 2,3-dioxygenase from sphingomonas2 yanoikuya b1 bound to biphenyl
35	c3k7cC_		Alignment	not modelled	91.9	21	PDB header: protein binding Chain: C: PDB Molecule: putative ntf2-like transpeptidase; PDBTitle: crystal structure of putative ntf2-like transpeptidase (np_281412.1)2 from campylobacter jejuni at 2.00 a resolution
36	c4lehA_		Alignment	not modelled	91.8	10	PDB header: lyase Chain: A: PDB Molecule: bile acid 7-alpha dehydratase, baie; PDBTitle: crystal structure of a bile-acid 7-alpha dehydratase (closci_03134)2 from clostridium scindens atcc 35704 at 2.90 a resolution
37	c3ub1C_		Alignment	not modelled	91.8	13	PDB header: transport protein Chain: C: PDB Molecule: orf13-like protein; PDBTitle: ntf2 like protein involved in plasmid conjugation
38	c3gzrA_		Alignment	not modelled	91.7	8	PDB header: unknown function Chain: A: PDB Molecule: uncharacterized protein with a ntf2-like fold; PDBTitle: crystal structure of an uncharacterized protein with a cystatin-like2 fold (cc_2572) from caulobacter vibrioides at 1.40 a resolution
39	c3gwrA_		Alignment	not modelled	91.2	14	PDB header: protein binding Chain: A: PDB Molecule: putative calcium/calmodulin-dependent protein kinase type PDBTitle: crystal structure of putative calcium/calmodulin-dependent protein2 kinase type ii association domain (yp_315894.1) from thiobacillus3 denitrificans atcc 25259 at 2.00 a resolution
40	c5i97C_		Alignment	not modelled	90.3	11	PDB header: protein transport Chain: C: PDB Molecule: conjugal transfer protein; PDBTitle: structural analysis and inhibition of trae from the pkm101 type iv2 secretion system
41	d3ebya1		Alignment	not modelled	90.1	9	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
42	c2qguA_		Alignment	not modelled	89.1	12	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: probable signal peptide protein; PDBTitle: three-dimensional structure of the phospholipid-binding protein from2 ralstonia solanacearum q8xv73_ralsq in complex with a phospholipid at3 the resolution 1.53 a. northeast structural genomics consortium4 target rsr89
43	d3e99a1		Alignment	not modelled	88.9	15	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
44	c4fczB_		Alignment	not modelled	88.1	10	PDB header: transport protein Chain: B: PDB Molecule: toluene-tolerance protein; PDBTitle: crystal structure of toluene-tolerance protein from pseudomonas putida2 (strain kt2440), northeast structural genomics consortium (nsg)3 target ppr99
45	c4hzB_E		Alignment	not modelled	86.1	25	PDB header: hydrolase Chain: E: PDB Molecule: putative periplasmic protein; PDBTitle: crystal structure of the type vi semet effector-immunity complex tae3-2 tai3 from ralstonia picketti
46	c2bhmE		Alignment	not modelled	86.1	14	PDB header: bacterial protein Chain: E: PDB Molecule: type iv secretion system protein virb8; PDBTitle: crystal structure of virb8 from brucella suis
47	d2bhma1		Alignment	not modelled	84.5	15	Fold: Cystatin-like Superfamily: NTF2-like Family: VirB8-like
48	c5uwab_		Alignment	not modelled	84.1	15	PDB header: transport protein Chain: B: PDB Molecule: probable phospholipid-binding protein mlac; PDBTitle: structure of e. coli phospholipid binding protein mlac
49	d2b1xb1		Alignment	not modelled	83.9	19	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
50	d3cnxa1		Alignment	not modelled	83.3	14	Fold: Cystatin-like Superfamily: NTF2-like Family: SAV4671-like
51	c4meiA_		Alignment	not modelled	82.4	8	PDB header: protein transport Chain: A: PDB Molecule: virb8 protein; PDBTitle: crystal structure of a virb8 type iv secretion system machinery2 soluble domain from bartonella tribocorum
52	d2r4ia1		Alignment	not modelled	82.3	13	Fold: Cystatin-like Superfamily: NTF2-like Family: CHU142-like

53	d3blz1		not modelled	81.1	13	Fold: Cystatin-like Superfamily: NTF2-like Family: Sbal0622-like
54	c5ig0A_		not modelled	79.7	16	PDB header: transferase Chain: A: PDB Molecule: camk/camk2 protein kinase; PDBTitle: crystal structure of s. rosetta camkii hub
55	d3b7ca1		not modelled	74.0	15	Fold: Cystatin-like Superfamily: NTF2-like Family: SO0125-like
56	d2owpa1		not modelled	71.2	9	Fold: Cystatin-like Superfamily: NTF2-like Family: BxeB1374-like
57	d1vqqa1		not modelled	69.4	7	Fold: Cystatin-like Superfamily: NTF2-like Family: Penicillin binding protein 2a (PBP2A), N-terminal domain
58	d3bb9a1		not modelled	68.4	18	Fold: Cystatin-like Superfamily: NTF2-like Family: SO0125-like
59	c3kspA_		not modelled	66.9	12	PDB header: unknown function Chain: A: PDB Molecule: calcium/calmodulin-dependent kinase ii association domain; PDBTitle: crystal structure of a putative ca/calmodulin-dependent kinase ii2 association domain (exig_1688) from exiguobacterium sibiricum 255-153 at 2.59 a resolution
60	c2rsxA_		not modelled	65.6	29	PDB header: hydrolase inhibitor Chain: A: PDB Molecule: uncharacterized protein yoeb; PDBTitle: solution structure of iseaa, an inhibitor protein of dl-endopeptidases2 from bacillus subtilis
61	c3hzpA_		not modelled	63.0	11	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: ntf2-like protein of unknown function; PDBTitle: crystal structure of ntf2-like protein of unknown function mn2a_05052 from prochlorococcus marinus (yp_291699.1) from prochlorococcus sp.3 natl2a at 1.40 a resolution
62	c3fkad_		not modelled	61.0	33	PDB header: unknown function Chain: D: PDB Molecule: uncharacterized ntf-2 like protein; PDBTitle: crystal structure of a ntf-2 like protein of unknown function2 (spo1084) from silicibacter pomeroyi dss-3 at 1.69 a resolution
63	c3fsdA_		not modelled	59.5	13	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: ntf2-like protein of unknown function in nutrient uptake; PDBTitle: crystal structure of ntf2-like protein of unknown function in nutrient2 uptake (yp_427473.1) from rhodospirillum rubrum atcc 11170 at 1.70 a3 resolution
64	c3ke7A_		not modelled	54.5	11	PDB header: isomerase Chain: A: PDB Molecule: putative ketosteroid isomerase; PDBTitle: crystal structure of putative ketosteroid isomerase (yp_001303366.1)2 from parabacteroides distasonis atcc 8503 at 1.45 a resolution
65	c1mwuA_		not modelled	50.1	8	PDB header: biosynthetic protein Chain: A: PDB Molecule: penicillin-binding protein 2a; PDBTitle: structure of methicillin acyl-penicillin binding protein 2a from2 methicillin resistant staphylococcus aureus strain 27r at 2.60 a3 resolution.
66	c3dukD_		not modelled	45.6	18	PDB header: unknown function Chain: D: PDB Molecule: ntf2-like protein of unknown function; PDBTitle: crystal structure of a ntf2-like protein of unknown function2 (mfla_0564) from methyllobacillus flagellatus kt at 2.200 a resolution
67	d1ulib_		not modelled	42.2	8	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
68	c3gzxB_		not modelled	40.2	11	PDB header: oxidoreductase Chain: B: PDB Molecule: biphenyl dioxygenase subunit beta; PDBTitle: crystal structure of the biphenyl dioxygenase in complex with biphenyl2 from commamonas testosteroni sp. strain b-356
69	c6bjuD_		not modelled	38.0	9	PDB header: unknown function Chain: D: PDB Molecule: atzh; PDBTitle: the structure of atzh: a little known member of the atrazine breakdown2 pathway
70	c4r4gA_		not modelled	29.1	17	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: putative lipoprotein ycda; PDBTitle: crystal structure of a putative lipoprotein (ycda) from bacillus2 subtilis subsp. subtilis str. 168 at 2.62 a resolution
71	d1wqlb1		not modelled	26.9	15	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
72	c5ve9C_		not modelled	26.2	25	PDB header: protein binding Chain: C: PDB Molecule: microtubule-actin cross-linking factor 1, isoforms 1/2/3/5; PDBTitle: structure of hacf7 ef1-ef2-gar domains
73	c5dqjP_		not modelled	26.1	35	PDB header: oxidoreductase Chain: P: PDB Molecule: putative uncharacterized protein; PDBTitle: the structure of cbb3 cytochrome oxidase.
74	c5dqjN_		not modelled	26.1	35	PDB header: oxidoreductase Chain: N: PDB Molecule: putative uncharacterized protein; PDBTitle: the structure of cbb3 cytochrome oxidase.
75	c5dqjQ_		not modelled	26.1	35	PDB header: oxidoreductase Chain: Q: PDB Molecule: putative uncharacterized protein; PDBTitle: the structure of cbb3 cytochrome oxidase.
76	c5dqjO_		not modelled	26.1	35	PDB header: oxidoreductase Chain: O: PDB Molecule: putative uncharacterized protein; PDBTitle: the structure of cbb3 cytochrome oxidase.
77	c17aoY		not modelled	25.4	10	PDB header: metal binding protein Chain: X: PDB Molecule: cation efflux system protein cusf;

77	c1zeyA	Alignment	not modelled	23.4	10	PDBTitle: 1.5 a structure of apo-cusf residues 6-88 from escherichia coli
78	d1v5ra1	Alignment	not modelled	24.1	25	Fold: N domain of copper amine oxidase-like Superfamily: GAS2 domain-like Family: GAS2 domain
79	c4o3vA	Alignment	not modelled	23.7	8	PDB header: protein transport Chain: A: PDB Molecule: virb8-like protein of type iv secretion system; PDBTitle: crystal structure of a virb8-like protein of type iv secretion system2 from rickettsia typhi
80	c4ce4j	Alignment	not modelled	22.2	9	PDB header: ribosome Chain: I: PDB Molecule: mrpl9; PDBTitle: 39s large subunit of the porcine mitochondrial ribosome
81	d2bmob1	Alignment	not modelled	19.2	8	Fold: Cystatin-like Superfamily: NTF2-like Family: Ring hydroxylating beta subunit
82	d2rcda1	Alignment	not modelled	16.8	20	Fold: Cystatin-like Superfamily: NTF2-like Family: BxeB1374-like
83	c2ml6A	Alignment	not modelled	15.5	29	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: nmr structure of protein zp_02069618.1 from bacteroides uniformis atcc2 8492
84	c5up5A	Alignment	not modelled	13.2	50	PDB header: de novo protein Chain: A: PDB Molecule: ehee_rd1_0284; PDBTitle: solution structure of the de novo mini protein ehee_rd1_0284
85	c4ouqA	Alignment	not modelled	12.8	14	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a duf4783 family protein (bf1468) from2 bacteroides fragilis ych46 at 1.55 a resolution
86	c3iuuA	Alignment	not modelled	12.6	18	PDB header: lyase Chain: A: PDB Molecule: phosphothreonine lyase ospf; PDBTitle: structure of the type iii effector/phosphothreonine lyase ospf from2 shigella flexneri
87	c4orlA	Alignment	not modelled	11.6	24	PDB header: structural genomics, unknown function Chain: A: PDB Molecule: uncharacterized protein; PDBTitle: crystal structure of a duf4783 family protein (bacova_04304) from2 bacteroides ovatus atcc 8483 at 1.40 a resolution
88	d2gxfa1	Alignment	not modelled	10.9	16	Fold: Cystatin-like Superfamily: NTF2-like Family: YybH-like
89	d1m56d	Alignment	not modelled	10.7	44	Fold: Single transmembrane helix Superfamily: Bacterial aa3 type cytochrome c oxidase subunit IV Family: Bacterial aa3 type cytochrome c oxidase subunit IV
90	c2micB	Alignment	not modelled	9.7	12	PDB header: membrane protein Chain: B: PDB Molecule: tumor necrosis factor receptor superfamily member 16; PDBTitle: nmr structure of p75 transmembrane domain in dpc micelles
91	c2micA	Alignment	not modelled	9.7	12	PDB header: membrane protein Chain: A: PDB Molecule: tumor necrosis factor receptor superfamily member 16; PDBTitle: nmr structure of p75 transmembrane domain in dpc micelles
92	c5ts4A	Alignment	not modelled	9.3	14	PDB header: de novo protein Chain: A: PDB Molecule: denovo ntf2; PDBTitle: crystal structure of a de novo designed protein with curved beta-sheet
93	c2z8nB	Alignment	not modelled	8.9	22	PDB header: lyase Chain: B: PDB Molecule: 27.5 kda virulence protein; PDBTitle: structural basis for the catalytic mechanism of phosphothreonine lyase
94	c2l55A	Alignment	not modelled	8.6	15	PDB header: metal binding protein Chain: A: PDB Molecule: silb,silver efflux protein, mfp component of the three PDBTitle: solution structure of the c-terminal domain of silb from cupriavidus2 metallidurans
95	c3efyB	Alignment	not modelled	8.4	24	PDB header: cell cycle Chain: B: PDB Molecule: cif (cell cycle inhibiting factor); PDBTitle: structure of the cyclomodulin cif from pathogenic2 escherichia coli
96	d2cw9a1	Alignment	not modelled	8.2	12	Fold: Cystatin-like Superfamily: NTF2-like Family: TIM44-like
97	c2luvA	Alignment	not modelled	8.0	15	PDB header: cell adhesion Chain: A: PDB Molecule: integrin alpha-x; PDBTitle: structure and binding interface of the cytosolic tails of axb22 integrin
98	c3qk9B	Alignment	not modelled	7.9	14	PDB header: protein transport Chain: B: PDB Molecule: mitochondrial import inner membrane translocase subunit PDBTitle: yeast tim44 c-terminal domain complexed with cymal-3
99	c2na9A	Alignment	not modelled	7.6	13	PDB header: signaling protein Chain: A: PDB Molecule: cytokine receptor common subunit beta; PDBTitle: transmembrane structure of the p441a mutant of the cytokine receptor2 common subunit beta