
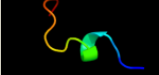
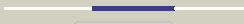




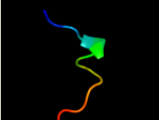


Phyre2

Email	l.a.kelley@imperial.ac.uk
Description	Q47272
Date	Thu Jan 5 12:36:43 GMT 2012
Unique Job ID	41ce00c30883272e

Detailed template information

#	Template	Alignment Coverage	3D Model	Confidence	% i.d.	Template Information
1	c3b8fB_	 Alignment		6.9	21	PDB header: hydrolase Chain: B: PDB Molecule: putative blastidicin s deaminase; PDBTitle: crystal structure of the cytidine deaminase from bacillus anthracis
2	d1rq0a_	 Alignment		6.1	47	Fold: Release factor Superfamily: Release factor Family: Release factor
3	d1tafa_	 Alignment		5.8	32	Fold: Histone-fold Superfamily: Histone-fold Family: TBP-associated factors, TAFs
4	c3eudE_	 Alignment		5.6	58	PDB header: nuclear protein Chain: E: PDB Molecule: protein shq1; PDBTitle: structure of the cs domain of the essential h/aca rnp2 assembly protein shq1p