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Description	P0A7W1
Date	Thu Jan 5 11:06:20 GMT 2012
Unique Job ID	e8a63f2fa9bd3c95

Sequence MAHI EKQAGELQEKLIAVNRVSKTVKGGRI F SFTALT VVGDNGRVGF GYGKAREVPAAI QKAMEKARRNMINVALNNGTLQHPVKGVHTGSRVFMQPA SEGTGI I AGGAMRAVLEEVAGV HNVLA KAYGSTNPINVV RATI DGL ENMNSPEMVA AKRGKSVEEILGK


Secondary structure: Alpha helices (green) and beta strands (blue) are shown. Helices are located at residues 13-20, 24-31, 35-42, 46-53, 57-64, 71-78, 82-89, 93-100, 104-111, 115-122, 129-136, 140-147, 151-158, 162-169, 173-180, 184-191, 195-202, 206-213, 217-224, 228-235, 239-246, 250-257, 261-268, 272-279, 283-290, 294-301, 305-312, 316-323, 327-334, 338-345, 349-356, 360-367, 371-378, 382-389, 393-400, 404-411, 415-422, 426-433, 437-444, 448-455, 459-466, 470-477, 481-488, 492-499, 503-510, 514-521, 525-532, 536-543, 547-554, 558-565, 569-576, 580-587, 591-598, 602-609, 613-620, 624-631, 635-642, 646-653, 657-664, 668-675, 679-686, 690-697, 704-711, 715-722, 726-733, 737-744, 748-755, 759-766, 770-777, 781-788, 792-799, 803-810, 814-821, 825-832, 836-843, 847-854, 858-865, 869-876, 880-887, 891-898, 899-906, 909-916, 919-926, 929-936, 939-946, 949-956, 959-966, 969-976, 979-986, 989-996, 999-1006.

SS: Solvent accessibility is shown as a bar chart. High accessibility (red) is observed in the first 100 residues, while lower accessibility (blue/green) is seen in the second 100 residues.


confidence: Confidence scores are shown as a bar chart. High confidence (red) is observed in the first 100 residues, while lower confidence (blue/green) is seen in the second 100 residues.


Disorder: Disorder predictions are shown as a bar chart. High disorder (red) is observed in the first 100 residues, while lower disorder (blue/green) is seen in the second 100 residues.

Confidence Key

High(9)  Low (0)

? Disordered

 Alpha helix

 Beta strand