


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Date	Thu Jan 5 12:21:47 GMT 2012
Unique Job ID	c3753b9c3cbe846c


Figure 1: Sequence and structural analysis of the protein. The figure displays three segments of the protein sequence (1-60, 70-120, and 130-170) with corresponding secondary structure predictions (SS), confidence scores, disorder predictions, and disorder confidence scores. The sequence is color-coded by amino acid type. Secondary structure is shown as green alpha-helices and blue beta-strands. Confidence scores are represented by colored bars, and disorder predictions are shown as question marks.

Confidence Key

High(9)  Low (0)

? Disordered

 Alpha helix

 Beta strand