



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| Email | l.a.kelley@imperial.ac.uk |
| Description | P77564 |
| Date | Thu Jan 5 12:30:37 GMT 2012 |
| Unique Job ID | dcd22edb48601eda |


Figure 1: Protein structure and disorder analysis of the human protein. The figure displays the protein sequence (MGKMNHQDELPLAKVSEVDEAKRQWLQGM RHPVDTVTEPEPAEILAEFIRQHSAAGQLVA) and its predicted secondary structure (SS) and disorder. The protein is shown in three segments: 1-60, 70-120, and 130-180. The secondary structure is represented by green cylinders (alpha-helices) and blue arrows (beta-strands). The disorder is indicated by a red bar (disordered) and a blue bar (ordered). The disorder analysis shows that the protein is highly disordered, with only a few regions of order.

Confidence Key

High(9)  Low (0)

? Disordered

 Alpha helix

 Beta strand