## **Secondary Structure Prediction**

The Secondary Structure Prediction plugin provides a set of algorithms for the protein secondary structure (alpha-helix, beta-sheet) prediction from a raw sequence.

Currently, available algorithms are:

- GORIV Jean Garnier, Jean-Francois Gibrat, and Barry Robson,"GOR Method for Predicting Protein Secondary Structure from Amino Acid Sequence", in Methods in Enzymology, vol.266, pp. 540 553, (1996).
- The improved version of the GOR method in J. Garnier, D. Osguthorpe, and B. Robson, J. Mol. Biol., vol. 120, p. 97 (1978).
- PsiPred Bryson K, McGuffin LJ, Marsden RL, Ward JJ, Sodhi JS. & Jones DT. (2005) Protein structure prediction servers at University College London. Nucl. Acids Res. 33(Web Server issue): W36-38.

Jones DT. (1999) Protein secondary structure prediction based on position-specific scoring matrices. J. Mol. Biol. 292: 195-202.

You can access these analysis capabilities for a protein sequence using the Analyze Predict secondary structure... context menu item. The dialog will appear:

Secondary Structure Prediction			
Algorithm	GORIV  Range End: 82 -		
Range Start: 1 🚊 Results:	Range End.  82		
Region	Structure Type		
, Total predicted: 0			
Help	Save Cancel Predict		

It supports the following options:

Algorithm -- you can choose the preferred algorithm. Currently, "GORIV" and "PsiPred" algorithms are available.

Range start / Range end — select the sequence range for prediction.

Results - the visual representation of the prediction results, for example:

Secondary Structure Prediction			
Algorithm		GORIV	
Range Start: 1 🚊		Range End: 82 💼	
Results:			
Region		Structure Type	
1 [57]	beta_strand		
2 [915]	beta_strand		
3 [2432]	alpha_helix		
4 [3975]	alpha_helix		
5 [7581]	beta_strand		
Total predicted: 5			
Help	S	ave Cancel Predict	

Save as annotation — select this button to save the results as annotations of the current protein sequence.