Cancer Research Center, Department of Epidemiology and Biostatistics, University at Albany, One Discovery Drive, Rensselaer, NY 12144, USA; 2 Department of Pharmacology and Systems Therapeutics, Mount Sinai School of Medicine, One Gustave L Levy Place, New York, NY 10029, USA; Igor B. Kuznetsov – Email: ikuznetsov@uamail.albany.edu; *Corresponding author

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Abstract:
Many proteins contain conformationally flexible segments that undergo significant changes in the backbone conformation or completely lack a well-defined conformation. Previously, we have developed the generalized local propensity (GLP), a quantitative sequence-based measure of the protein backbone flexibility. In this paper, we present the CFP (Conformational Flexibility Profile) web-server that constructs the GLP flexibility profile for a user-submitted sequence and uses this profile to identify segments with high backbone flexibility. The statistical significance of a flexible sequence segment is assessed using the discrete scan statistics based on the density of flexible residues observed in this segment.

Availability: CFP is publicly available at http://cfp.rit.albany.edu

Keywords: conformational variability, protein backbone, flexibility, local propensity, sequence

Background:
Many proteins contain conformationally flexible segments. These segments undergo significant changes in backbone conformation, or are completely disordered (lack a well-defined structure) [1-3]. A quantitative representation of the conformational flexibility of the protein backbone is important for many applications. Previously, we developed generalized local propensity (GLP), a quantitative sequence-based measure of backbone flexibility [4]. The GLP can be used to construct sequence-based protein flexibility profiles, and provides an objective numeric threshold for defining conformationally flexible segments [5]. For a given sequence position \( k \), the GLP measures the width of the context-dependent distribution of backbone conformations accessible to this position, \( \text{glp}(k) \) (see references [4-5] for details). If \( \text{glp}(k) \geq 1 \), it indicates that sequence position \( k \) is conformationally flexible.

Here, we present the CFP (Conformational Flexibility Profile) web server that constructs the GLP flexibility profile for a user-submitted sequence and uses this profile to identify segments with high conformational flexibility. Below is a brief outline of the steps implemented in CFP: The GLP flexibility profile is constructed for the query sequence and then smoothed using a sliding window of size \( W_1 \). Consecutive positions which have GLP above a threshold \( T_1 \) are merged into seed flexible segments. Each seed flexible segment is extended by adding extension windows of size \( W_2 \) until its average GLP drops below this threshold \( T_2 \). An extension window is added only if its average GLP is above a certain threshold \( T_3 \). This extension procedure is similar to that used in the SEG program [6].

The extended flexible segments are reported in the final table. If the number of flexible residues observed in a given final flexible segment is unusually high (p-value < 0.05), then this segment is marked as statistically significant. The significance of the number of flexible residues is estimated using the discrete scan statistic. This statistical procedure is the same as the one we previously implemented in the BIAS software to identify statistically significant clusters of user-specified amino acid types [7-8]. The web-server is publicly available at http://cfp.rit.albany.edu.
Figure 1: (a) First part of the output - the GLP flexibility plot; (b) Second part of the output - the summary of all flexible segments.
Flexible residues: A set of flexible residues used in the scan statistics to estimate the statistical significance of flexible segments (G, H, D, N by default).

SWIS-PROT or PDB background frequencies: The amino acid frequencies of the SwissProt or Protein Databank are used to estimate the statistical significance.

X axis size, Y axis size: The size of X and Y axis of the plot in pixels.

Create a plot: Display the smoothed GLP profile in web-browser.

Create a text file: Save the raw and smoothed GLP profiles in a text file.

Output:
The CFP output consists of two parts. The first part shows the smoothed GLP plot of the input sequence (Figure 1A). The second part shows the detailed information for every flexible segment found in the input sequence and the p-values that provide the estimates of the statistical significance (Figure 1B). If the p-value for a given segment is less than 0.05, this segment has an unusually high density of residues with high degree of backbone flexibility.

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