Computation of Conformational Entropy from Protein Sequences Using the Machine-Learning Method—Application to the Study of the Relationship between Structural Conservation and Local Structural Stability

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ABSTRACT A complete protein sequence can usually determine a unique conformation; however, the situation is different for shorter subsequencessome of them are able to adopt unique conformations, independent of context; while others assume diverse conformations in different contexts. The conformations of subsequences are determined by the interplay between local and nonlocal interactions. A quantitative measure of such structural conservation or variability will be useful in the understanding of the sequence-structure relationship. In this report, we developed an approach using the support vector machine method to compute the conformational variability directly from sequences, which is referred to as the sequence structural entropy. As a practical application, we studied the relationship between sequence structural entropy and the hydrogen exchange for a set of well-studied proteins. We found that the slowest exchange cores usually comprise amino acids of the lowest sequence structural entropy. Our results indicate that structural conservation is closely related to the local structural stability. This relationship may have interesting implications in the protein folding processes, and may be useful in the study of the sequence-structure relationship. Proteins 2005;59:802-809. © 2005 Wiley-Liss, Inc.

Key words: structural conservation; sequence structural entropy; structural profile; support vector machines; hydrogen exchange

INTRODUCTION

Studies¹⁻⁴ showed that both designed and naturally occurring subsequences may assume different secondary structures in different contexts. For example, Minor and Kim² have designed an 11-amino acid sequence that forms an α -helix in one context while a β -sheet in another; systematic search^{1,3,4} for protein fragments in the Protein Data Bank⁵ revealed that identical subsequences could have very different conformations—for example, the pentapeptide AVLAE forms an α -helix in a potassium channel but forms a β -sheet in cytidine deaminase. On the other

hand, a similar pentapeptide AALAE (the second amino acid is changed from V from A) remains the same α -helical conformation in different proteins. Such structural conservation or variability is dictated by the interplay between local and nonlocal interactions. Studies⁶⁻⁹ showed that protein folding is facilitated by the existence of some structured regions, dominated by stronger local interactions that are compatible with the native conformation. Hydrogen isotope exchange rates provide a useful probe of the equilibrium folding pathway of proteins.^{10–12} The slow hydrogen exchange regions usually consist of the amino acids that are either buried or involved in the hydrogenbonding network.^{13,14} Hence, it is reasonable to assume that the amino acids involved in the slow amide proton exchange will have more conserved local structures. In this work, we developed an approach based on the machine learning method to compute conformational entropy from sequences. We applied this approach to a set of proteins with known hydrogen exchange data, and we found a close correlation between structural conservation and the slow hydrogen exchange.

METHODS

Theory

A protein sequence α of length *L* is denoted by $\alpha = \alpha_1 \alpha_2 \dots \alpha_L$, where α_i is the amino acid at the *i*th position. The structural profile of α can be expressed in terms of an $L \times N$ matrix **M**,¹⁵ where *N* is the number of structural descriptors (to be given later)

$$\mathbf{M} = (\mathbf{P}_1, \mathbf{P}_2, \ldots, \mathbf{P}_L), \qquad (1)$$

where the column vector \mathbf{P}_i is the probability distribution of the structural descriptors at α_i , that is,

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$$\mathbf{P}_{i} = (p_{i1}, p_{i2}, \dots, p_{iN}), \qquad (2)$$

where p_{ij} is the probability of the *j*th structural descriptor at α_i .¹⁵ Using Shannon's information theory, ¹⁶ we compute the conformational entropy of α_i by

$$S_i = -\sum_{j}^{N} p_{ij} \log p_{ij}$$
(3)

We will refer to S_i as sequence structural entropy (SSE), because it is derived directly from sequences. SSE provides a straightforward measure of the structural conservation or variability of the residues. Lower SSE indicates a more conserved structure (or a context-independent conformation), stabilized mainly by local interactions, while higher SSE indicates a higher degree of structural variability. Note that the definition of the structural descriptor is not unique, and in principle, it may be any structure-related properties like the secondary structural elements, the backbone torsion angles, or the solvent-accessible areas. Previous study¹⁵ showed that the secondary structural element is a more convenient and useful structure descriptor in the computation of conformational entropy. In this work we use the secondary structures defined by the widely used DSSP method,¹⁷ whose assignment of secondary structures is based on the hydrogen bonding patterns of protein structures. The DSSP method defines eight secondary structures: α -helix (H), 3_{10} -helix (G), π -helix (I), extended β -strand (E), isolated β -strand (B), turn (T), bend (S), and coil (U). Indeed, there are a number of secondary structure assignment methods (see the review by Andersen and Rost¹⁸) based on various algorithms such as the aforementioned hydrogen bonding patterns,17 backbone dihedral angles,¹⁹ or combination of both,^{20,21} quantification of backbone curvature,²² or the distance matrices of structural fragments.²³ In general, these methods make similar secondary structure assignments for the more common conformations such as α -helix or β -strand; however, due to difference in assigning secondary structures, these methods may define different secondary structure types for particular conformations. For example, the DSSP method does not define the polyproline II helix (PII) but assigns it to T, S, or U according to the corresponding hydrogen-bonding patterns; on the other hand, the tool PROSS,¹⁹ which assigns secondary structures based solely on backbone torsional angles, defines PII but does not define G, I, S, or B. However, as long as these methods provide a complete, consistent description of the protein conformations, Equation (3) will provide a useful quantitative measure of the structural conservation or variation of a protein.

The Support Vector Machines

The support vector machine method $(SVM)^{24}$ has been successfully applied to secondary structure prediction,^{25–27} protein fold assignment,^{28,29} subcellular localization prediction^{30,31} and other biological pattern classification problems.^{32–35} The original idea of the SVM is to find the separating hyperplane with the largest distance between two classes. However, because the data to be classified may not always be linearly separable, the SVM overcomes this difficulty by nonlinearly transforming the original input space into a higher dimensional feature space by the so-called "kernel functions," so that the SVM may effectively separate these data in the higher dimensional space. SVMs perform well compared with other machine-learning methods because of convenient classifier's capacity control and effective avoidance of overfitting. In this work, the software package LIBSVM³⁶ was used in this work.

Generation of Structural Profiles and Sequence Structural Entropy

We compute the probability distribution of the secondary structures using the method illustrated in Figure 1. The inputs to the SVM are in the form of W imes 20 PSI-BLAST profiles or the position-specific scoring matrix (PSSM),³⁷ where *W* is the size of the sliding window of the sequence. The window size W is chosen to be an odd number so that the target residue is always centered in the sliding window. In this work, a 15×20 scoring matrix is used as an input to the SVM. The PSI-BLAST profile was obtained after three iterations with the E-value threshold set to 1×10^{-3} against the nonredundant protein sequence database.³⁸ Each element of the matrix represents the log-odds score of a particular residue substitution at that position, and its value is usually in the range ± 7 . These matrix elements are normalized to the range [0,1] by the following scaling function.²⁶

$$\pi(x) = \begin{cases} 0.0 & \text{if } x \le -5\\ 0.5 + 0.1x & \text{if } -5 \le x \le 5\\ 1.0 & \text{if } x \ge 5 \end{cases}$$
(4)

where x is the original value of the matrix element. The SVM output for the target residue (i.e., the central amino acid of the sliding window) is an eight-element vector $\mathbf{O} =$ (o_1, o_2, \ldots, o_8) , where o_i is the decision value of secondary structure type *i*. Because the SVM does not provide estimates of the posterior probability of class membership, we transform this decision value o_i by the function $[\arctan(o_i) + \pi]$ to the range [0,1]. We then normalize the resultant values to obtain p_i , that is, the probability of secondary structure type i of the target residue. With $\mathbf{P} =$ (p_1, p_2, \ldots, p_8) , we compute SSE of the target residue using Equation (3). We train the SVM model using the standard data set RS126,³⁹ a nonhomologous data set that the pair wise sequence identity is less than 25% over a length of more than 80 residues. An important issue of optimizing SVMs is the selection of parameters such as the penalty parameters and the kernel parameters of the kernel function, which must be determined in advance. We use the cross validation on different parameters for the model selection.40

RESULTS Hen Egg-White Lysozyme

Hen egg-white lysozyme (HEWL) has two subdomains: the α -domain composed of four α -helices (A, B, C, and D), two 3₁₀ helices (E and F), and the β -domain formed by

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The PSSM of the protein sequence





Fig. 1. The method to compute SSE from the protein sequence. The protein sequence is shown on the left and the target residue (in this example, a leucine residue) is in the shaded area. In step 1, the position-specific scoring matrix of the protein sequence is generated by the iterative PSI-BLAST. In step 2, each element of the window of 15 rows (in the dotted frame) centered on the leucine residue is rescaled to the range [0,1] [using Eq. (4)] as input to the SVM. In step 3, the outputs of the SVM are mapped to posterior probabilities P_L of secondary structures. Finally, in step 4, the SSE of the target residue *L* is evaluated from P_L using Equation (3).

three contiguous β -strands. In the native form of HEWL, the study⁴¹ showed that slowest exchange amide protons are located in the α -domain: helix A (M12), helix B (W28–A31), and helix C (A95, K96, I98); and the next slowest in strand β 3 (I58). Figure 2 shows the SSE profile of HEWL. The lowest entropy regions are A11–K13 (helix A), W28–K33 (helix B), and N93–V99 (helix C), respec-

tively. These amino acids overlap well with the slow exchange amide proton in helices A, B, and C. Note the residues in helix D (V109–R114) have relatively higher entropy. This is consistent with the experiment that helix D amide protons exchange much more rapidly.⁴¹ Figure 3 compares the spatial arrangements of the low SSE and the slow exchange regions in the ribbon diagrams of HEWL.



Fig. 2. The SSE profile of HEWL. The secondary structural elements of HEWL together with their notations are shown on the top of the figure. On the bottom, we label the positions of the low SSE and the slow exchange regions by the red and blue circles, respectively. The entropy is rescaled to the range [0, 1]. The boundary of the SSE region is set by allowing deviations in SSE of 0.2 around the local minimum.



Fig. 3. Comparison of the slow exchange and the low SSE regions of HEWL. The slow exchange regions are colored in blue (left), and the low SSE regions in red (right). The secondary structure elements are also labeled for the corresponding slow exchange regions. Ribbon diagrams are drawn with Molscript.⁵⁶

Chymotrypsin Inhibitor 2

Chymotrypsin inhibitor 2 (CI2) has a unique α -helix that packs against the β -sheets to form the hydrophobic core of the protein. The hydrogen exchange studies^{42,43} showed that the exchange behavior of CI2 is determined by its native structure. Figure 4 shows that two residues (I20–L21) in the helix, which are the most buried in the helix, and the residues (V47, L49–V51) on the central strand of the β -sheet (β 4) have the slowest hydrogen exchange rates; the other slowest exchange amide protons are K11 (β 2), I30 and L32 (β 3).^{14,42,43} These residues are among those of the lowest SSE (with the exception of K11), most of which are in sheet β 3 and β 4, and helix α . Figure 5 compares the relative spatial distributions of the slow exchanging regions and the low entropy regions on the ribbon diagrams of CI2.

Cytochrome c

In the native horse heart cytochrome c (Cyt c), the slowest exchange regions are located in the N- and



Fig. 4. The SSE profile of Cl2. The notations of the secondary structural elements follows those of Otzen et al. $^{\rm 57}$



Fig. 5. Comparison of the slow exchange and the low SSE regions of CI2 in ribbon diagrams.



C-helices.⁴⁴ Specifically, F10 (N-helix) and L94–K99 (C-helix) carry the slowest exchanging amide protons.⁴⁴ Figure 6 shows that I9–V11 and D93–L98 have the lowest SSE values, which coincide well with those with the slowest exchange amide protons. Note that the amino acids in the 60s helix also have relatively low SSE, which are consistent with the experiment that the next slowest exchange amide protons are in the 60s helix. Figure 7 compares the spatial orientations of the observed and the calculated exchange regions on the ribbon diagrams of Cyt c. Note that the two helices that carry the residues of the lowest SSE are close to each other in space.



Fig. 7. Comparison of the slow exchange and the low SSE regions of Cyt *c* in ribbon diagrams.



Fig. 8. The SSE profiles of Cyt *c* with the unfolding structural units (R, Y, G, and B) colored in their respective colors (red for R, yellow for Y, green for G, and blue for B). The average SSE of each structural unit is indicated by the solid line. (The dotted lines are visual aids for comparing the relative magnitude of the average SSE of each structural unit.)

Equilibrium Protein Folding

Cyt c provides a good model system for folding studies.^{45–47} Under native conditions, protein molecules, continually folding and unfolding, explore all possible unfolded conformation states in accordance with thermodynamic principles. The native state hydrogen exchanges studies⁴⁵⁻⁴⁷ have identified four cooperative structural unfolding units in the Cyt c molecule, which are designated as the Blue bihelix (B), the Green Ω loop and helix (G), the Yellow (Y) and the Red Ω loops (R), respectively, in the order of decreasing free energy of unfolding. These structural units, through various combinations, may produce the intermediates that define the folding and unfolding pathways of Cyt c. Figure 8 shows the SSE profiles of these structural units together with their respective average structural entropies. The B structural unit has the lowest average SSE, and G has the second lowest average SSE, and then Y, and R, which contains the residues of the largest SSE (A83–I85) in Cyt c and is also the least stable structural unit. The order of increasing average SSE follows that of decreasing free energy of unfolding.^{45,46} The SSE appears to be closely related to the local structural stability.

Other Examples

We compute the SSE for the following proteins: Ribonuclease T1 (RNase T1),⁴⁸ cardiotoxin analogue (CTX



Fig. 9. A summary of the comparison of the low SSE and the slow exchange regions for RNase T1, CTX III, Cyt *c*, HEWL, T4 Lysozyme, Cl2, BPTI, barnase, and RNase HI. The solid lines indicate the relative lengths of these sequences. The slow exchange and the low SSE regions are shown in red and blue circles, respectively.

III), 49 T4 Lysozyme, 50 bovine pancreatic trypsin inhibitor (BPTI), 51 barnase, 52,53 and Ribonuclease HI (RNase HI). 54 All exchange data are for the native proteins.¹⁴ Figure 9 summarizes the results for these proteins as well as the examples presented in the previous sections. Figure 10 compares the spatial arrangements of the computed and observed slow exchange regions of these proteins. Specifically, Figure 10(A) shows the native form of RNase T1 whose slowest exchange amide protons are located in strands $\beta 2-\beta 4$ of the central β -pleated sheet.⁴⁸ As shown in the figure, the agreement between the observed and the computed is quite good. Figure 10(B) shows the snake venom CTX III, a small protein with a two-stranded and a three-stranded β -sheet (β 3– β 5). Under native conditions, β 5 and β 3 are the most protected strands, followed by β 4.⁴⁹ Our calculations show that $\beta 5$ and $\beta 3$ are indeed of the lowest SSE among the five strands, and that the residues near the C-terminal β4 strand also have low SSE. Figure 10(C) shows that T4 lysozyme has two domains: the N-domain is mainly a β -sheet structure, and the Cterminal domain is predominantly α -helical. In the native T4 lysozyme, slow exchanges are observed in helices $\alpha 1$, α 3- α 5, and α 10. The slow exchange amide protons are mostly in $\alpha 5$ and $\alpha 10$. Our results show that most residues in $\alpha 5$ and $\alpha 10$ have low SSE. Figure 10(D) shows that most slow exchange regions of the native BPTI are in the antiparallel strands, that is, strand $\beta 1$ and $\beta 2$,⁵¹ which also have the lowest SSE. Figure 10(E) and 10(F) shows barnase and RNase HI, respectively. The observed slow

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Fig. 10. Comparison of the slow exchange (left, in blue) and the low SSE regions (right, in red) in ribbon diagrams of (A) RNase T1, (B) CTX III, (C) T4 lysozyme, (D) BPTI, (E) barnase, and (F) RNase HI.

exchange regions and the computed low SSE regions are in good agreement with each other.

DISCUSSION

Previously, Hilser and Freire¹¹ developed a combinatorial algorithm to generate a large ensemble of conformational states by partitioning the protein into blocks along the sequence. By assigning each block an arbitrary conformational state, they generated all possible combinational states and computed the Gibbs energy for each state. Using this approach, they were able to compute the hydrogen exchange protection factors for five proteins. Hespenheide et al.⁵⁵ simulated the incremental thermal denaturation of protein structures by breaking the tertiary network of hydrogen bonds and salt bridges one by one. Using these procedures, they were able to identify the structural stable and flexible regions, and found that they were closely related to the slow hydrogen exchanging regions for proteins with different architectures. Both approaches are structure-based and require well-parameterized energy functions (or scoring functions). In this report, we develop a sequence-based approach to compute the SSE of protein sequences, which provides a straightforward measure of structure diversity of local sequences. Using this approach, we found a close relationship between the low SSE and the slow hydrogen exchange regions. These results suggest that SSE is closely related to the local structural stability, as revealed by the hydrogen exchange data. These findings are consistent with our recent results that conformational variability is related to thermal stability.¹⁵ In summary, we have developed a sequence-based machine-learning method to compute conformational entropy directly from sequences, which may provide a useful tool in predicting local structural stability on the sequence level. Such information should be useful in the studies of sequence–structural relationship such as folding processes and protein de novo design.

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