

Prediction of Protein Secondary Structures from Conformational Biases

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ABSTRACT We use LINUS (the “Local Independently Nucleated Units of Structure”), a procedure developed by Srinivasan and Rose, to provide a physical interpretation of and predict the secondary structures of proteins. The secondary structure type at a given site is identified by the largest conformational bias during short simulations. We examine the rate of successful prediction as a function of temperature and the interaction window. At high temperatures, there is a large propensity for the establishment of β -strands whereas α -helices appear only when the temperature is lower than a certain threshold value. It is found that there exists an optimal temperature at which the correct secondary structures are predicted most accurately. We find that this temperature is close to the peak temperature of the specific heat. Changing the interaction window or carrying out longer simulations approaching equilibrium lead to little change in the optimal success rate. Our findings are in accord with the observation by Srinivasan and Rose that the secondary structures are mainly determined by local interactions and appear in the early stage of folding. *Proteins* 2002;48:558–565.

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Key words: protein folding; secondary structures; α -helix; β -strand; protein structure prediction

INTRODUCTION

A knowledge of the 3-D structure of a protein is crucial for understanding its biological functionality. Unfortunately, the rate at which protein structures can be experimentally solved is far behind the speed at which the sequences are determined. With progress in the Human Genome Project, a good computer-based method for the prediction of protein structures from their sequences would be an invaluable tool for modern microbiology as well as for drug design. The existing methods for structure prediction can be divided into two classes: (1) template-based methods that compare a sequence with unknown structure against the library of solved structures and (2) ab initio methods that seek to identify the native fold, usually defined as the lowest energy point in conformational space. The latter are especially useful when a target sequence has a low similarity with the existing protein sequences of

known structures. It should be noted, though, that many so-called ab initio methods use information derived from the protein database as input.

Significant progress has been achieved in the ab initio approach to protein structure prediction as witnessed in the CASP competitions,^{1–3} wherein the structures of large protein fragments, comprising as many as 100 residues, were predicted with an accuracy of 4–7 Å in root mean square deviation (RMSD). A notable success reported was that of the Baker group and entailed the assembly of protein conformations from fragments of known structures in the protein database, which have local sequences similar to that of the target sequence, using statistically derived scoring functions.^{4–6} In Levitt’s approach,^{7,8} secondary structures, which were predicted by using several existing secondary structure prediction methods,^{9–11} are fitted to best scoring compact conformations obtained on a simplified tetrahedral lattice. Scheraga and coworkers^{12,13} used an off-lattice C_α -based model with interactions imposed on virtual side-chains and virtual peptide groups. The lowest-energy C_α trace obtained by extensive conformational space annealing is then converted to an all-atom backbone for further refinements. Ortiz et al.¹⁴ built discretized protein conformations using predicted secondary structures and a number of tertiary restraints derived from multiple sequence alignments. The success of these ab initio methods relies to a large extent on knowledge-based information, that is, data derived from known protein structures, such as that used in the scoring functions, secondary structure prediction, or in the choice of fragments to incorporate in the model.

This work deals with secondary structure prediction and builds on a truly ab initio protein structure prediction procedure called LINUS (the “Local Independently Nucleated Units of Structure”) by Srinivasan and Rose.¹⁵ LINUS does not use any knowledge-based information and thus provides a clear picture of the role played by the different factors in folding. Further, the algorithm for

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determining the structure is not based on energy minimization—LINUS captures the interplay between energy and entropy in determining the local secondary structure.

The most powerful aspect of LINUS is its simplicity—it is based on just four essential aspects of protein behavior: (1) excluded volume, (2) preferred occupancies of the dihedral angles in certain regions in the Ramachandran plot,¹⁶ (3) hydrophobic interactions and hydrogen bonding, and (4) the hierarchical organization of protein structures.^{17,18} Despite this simplicity, LINUS has already proved effective in predicting the secondary and supersecondary structures of protein fragments.¹⁵ Note that the hierarchical algorithm steers folding along some specific pathways and the resulting structure does not necessarily correspond to the global energy minimum.

In a subsequent study,¹⁹ Srinivasan and Rose used LINUS to propose a physical basis for secondary structures, which showed that protein secondary structures are mainly determined by steric effects and local interactions. This conclusion recently obtained strong support from experimental evidence that unfolded protein conformations, under highly denaturing conditions and thus in the absence of long-range contacts, are still characterized by local native-like topology.^{20,21} In LINUS, the conformational bias toward a type of secondary structure is determined through the probability of being in this conformation during simulation.

We have found this idea to be intriguing and worthy of a careful reexamination. Here, we make an assessment of how well the secondary structures can be predicted based on the analysis of conformational biases. In particular we concentrate on the role played by the temperature, T , in determining the success rate and find that there is an optimal T at which the secondary structure prediction is the best. For most of the proteins studied, this temperature coincides with the one that Rose and Srinivasan used in their studies and is found to be near the peak in the specific heat where the conformational conversion in the system is largest. The optimal conditions for the structure prediction do not depend much on whether the window in the interactions allowed for purely local or also for nonlocal interactions. They are also insensitive to the duration of the simulations. We obtained similar results when long, nearly equilibrium, simulations were considered.

The aim of this study is to elucidate how LINUS works and what its strengths and weaknesses are. The ultimate goal would be to determine what kinds of improvements could be made in this physically appealing framework to move toward first principles tertiary structure prediction.

METHODS

A detailed description of LINUS can be found in the original articles of Srinivasan and Rose.^{15,19} We developed our own version of LINUS that strictly follows the improved development as described in the PNAS article.¹⁹ Briefly, in LINUS the coordinates of all backbone atoms are considered whereas a side-chain is represented in a simplified manner. Specifically, glycine has no side-chain, alanine's side-chain is made of a C_β , and the remaining

amino acids are represented by C_β and one or two pseudo- C_γ atoms, depending on whether the side-chain is branched out or not. The atoms are modeled as hard spheres that are not allowed to overlap. The sizes of the spheres depend on the type of the atom and the sizes of the pseudoatoms depend on the size of the side-chains that they represent.

Apart from steric interactions, the Hamiltonian consists of just a few terms that provide attraction between atoms: hydrogen bonding (H-bond), hydrophobic interaction, and salt bridges. All backbone nitrogens, except for those that belong to a proline, are considered to be H-bond donors and participate in no more than one H-bond but the nitrogen at the N-terminus may participate in up to three H-bonds. The backbone oxygens and the side-chains of some amino acids (Ser, Thr, Asn, Asp, Gln, Glu) are acceptors. A backbone-to-backbone hydrogen bond is assumed to be formed between residues i and j when they are at least three residues apart in the sequence and when the distance between a donor and an acceptor is smaller than 5 Å. An energy of -0.5ϵ is assigned, where ϵ is an energy unit, and the energy is scaled quasilinearly from 0 to its minimal value as the distance decreases to 3.5 Å. It is also required that the out-of-plane dihedral angle $O(j)-N(i)-C_\alpha(i)-C(i-1)$ should be larger than 40° . A side-chain-to-backbone hydrogen bond is formed when the donor-to-acceptor distance is smaller than 4 Å and the acceptor must be not further than four residues away from the donor in the sequence. In this case, an energy of -1.0ϵ is assigned and no scaling of the energy is involved.

Hydrophobic attraction is postulated to occur for contacts between the side-chain atoms of hydrophobic (Cys, Ile, Leu, Met, Phe, Trp, Val) and amphipathic (Ala, His, Thr, Tyr) residues. The minimal value of the contact energy is -0.5ϵ when both residues are hydrophobic and -0.25ϵ when one of them is hydrophobic and the other is amphipathic. A contact between two atoms i and j is said to form when the distance between them is smaller than $R(i) + R(j) + 1.4$ Å, where $R(i)$ and $R(j)$ are the contact radii of the two atoms. The contact radii of the atoms¹⁹ depend on the kind of atoms and are larger than their hard sphere radii. The energy of a contact scales from 0 to its minimal value as the distance between two atoms decreases from its cutoff value to $R(i) + R(j)$. A salt bridge is assigned to contacts between oppositely charged groups (namely, the side-chains of Arg or Lys with Glu or Asp). The minimal energy of a salt bridge is -0.5ϵ . In LINUS, there is also an energy function to chase residues away from the right side of the Ramachandran plot. When a residue has a positive torsional angle ϕ it is punished with an energy of 1.0ϵ if the residue is not a glycine; otherwise, it is rewarded with an energy of -1.0ϵ .

The main degrees of freedom used in LINUS are the Ramachandran torsional angles ϕ and ψ and the torsional χ , which corresponds to rotation of the side-chains. In addition, the torsional angle ω about the peptide bond and the $N-C_\alpha-C$ bond angle are allowed to be perturbed slightly during the simulation. All other bond angles and bond lengths are kept fixed. Three consecutive residues (i , $i+1$, $i+2$) are perturbed at a time and the movements

advance from the N- to the C-terminus. The moves at an i th residue are repeatedly chosen until a move is obtained in which there are no steric clashes within the three residue fragments considered. At the next stage, the whole protein chain is checked for the presence of steric clashes. Up to 50 such attempts are performed to find a conformation without any steric clashes. If the new conformation found still has steric clashes it is rejected; otherwise, it is accepted with a probability $\mathbf{P} = \min\{1, e^{-\Delta E/k_B T}\}$, where k_B is the Boltzmann constant, T is the temperature measured in the units of ϵ/k_B , and ΔE is the energy difference. A complete progression from N to C is called a cycle.

LINUS uses a smart move set that consists of the following, equally probable, move types:

1. α -Helix: three consecutive residues ($i - 1, i, i + 1$) are set to having $\phi = -64 \pm 7^\circ, \psi = -43 \pm 7^\circ$.
2. β -Strand: three residues ($i - 1, i, i + 1$) are set to having $\phi = -130 \pm 15^\circ, \psi = 135 \pm 15^\circ$. If a residue is a proline then ϕ is reset to $70 \pm 15^\circ$.
3. Turn: there are four types of turns, namely I, I', II, and II'. For each turn type, there are two possibilities: (a) setting residues ($i - 1, i$) to have turn ϕ and ψ values while residue $i + 1$ is set to random coil and (b) setting $i - 1$ to random coil and ($i + 1, i$) to have turn ϕ and ψ values. Overall, there are eight such possibilities. The turn ϕ and ψ values for two consecutive residues are given below for each type of a turn move (the notations used for the residues are $i - 1$ and i but they can also be i and $i + 1$).
 Type I—residue ($i - 1$): $\phi = -60 \pm 15^\circ, \psi = -30 \pm 15^\circ$;
 residue (i): $\phi = -90 \pm 15^\circ, \psi = 0 \pm 15^\circ$.
 Type I'—residue ($i - 1$): $\phi = 55 \pm 15^\circ, \psi = 40 \pm 15^\circ$;
 residue (i): $\phi = 80 \pm 15^\circ, \psi = 5 \pm 15^\circ$.
 Type II—residue ($i - 1$): $\phi = -60 \pm 15^\circ, \psi = 110 \pm 15^\circ$;
 residue (i): $\phi = 90 \pm 15^\circ, \psi = -5 \pm 15^\circ$.
 Type II'—residue ($i - 1$): $\phi = 60 \pm 15^\circ, \psi = -120 \pm 15^\circ$;
 residue (i): $\phi = -80 \pm 15^\circ, \psi = 0 \pm 15^\circ$.
 For all turn moves, if a residue is a proline then its ϕ is reset to $70 \pm 15^\circ$.
4. Random coil: ϕ and ψ are chosen randomly in one of the favorite regions of the Ramachandran plot. For nonglycine and nonproline residues, $(\phi, \psi) \in \{(-135 \pm 45^\circ, 135 \pm 45^\circ), (-75 \pm 30^\circ, -30 \pm 30^\circ), (75 \pm 15^\circ, 30 \pm 15^\circ)\}$. For glycine, $\phi \in \{90 \pm 30^\circ, 180 \pm 30^\circ\}$ and $\psi \in \{0 \pm 30^\circ, 180 \pm 30^\circ\}$. For proline, $\phi = -70 \pm 15^\circ$ and $\psi \in \{135 \pm 30^\circ, -45 \pm 30^\circ\}$.

For the first three move types, $\omega = 180 \pm 5^\circ$ whereas for the coil move $\omega = 180 \pm 10^\circ$. For all move types, the side-chain torsional angles χ_s are chosen at random in 10° windows around $-60, 60,$ and 180° .

The conformational bias, P , of a given type of secondary structure is defined as the probability of being in this structure during the simulation. P is usually computed as a function of residue in the sequence. The computation of P requires a procedure of secondary structure assignment, which allows one to determine to which type of secondary motifs a residue belongs at a given instant. We use an

assignment procedure in the most recent unpublished development of LINUS (R. Srinivasan and G.D. Rose, personal communication), which proceeds through the following steps:

1. Set all residues to the coil conformation (c).
2. For i running from 1 through $N - 3$, where N is the number of residues, compute the torsion Θ between four consecutive $C_{\alpha s}$ ($i, i + 1, i + 2, i + 3$).
 a. If $|\Theta| \geq 135^\circ$, then residues ($i + 1$) and ($i + 2$) are set to the strand conformation (s).
 b. If $45^\circ \leq \Theta \leq 65^\circ$, then residues ($i + 1$) and ($i + 2$) are set to the helix conformation (h).
 c. If $-50^\circ \leq \Theta \leq 45^\circ$, then residues ($i + 1$) and ($i + 2$) are set to the turn conformation (t).
3. Check again all residues from 1 through N :
 a. If a segment of less than five residues with an h assignment is found, then all residues in this segment are set to t .
 b. If a segment of less than three residues with an s assignment is found, then all residues in this segment are set to c .

To compute P , one starts from an open conformation and makes a simulation of 1000 cycles. After each cycle a conformation assignment is determined to gather statistics on P . The average is taken over 10 simulations for each T .

To make comparisons with the DSSP-based native assignments²² used in the Protein Databank (PDB),²³ we adopt a simplified correspondence in which the $3_{10}, \pi,$ and α -helix correspond to h , the isolated β -bridges and extended β -strands to s , the hydrogen bonded turn to t , and bends and undefined segments to c . It should be noted that the native-state secondary structure assignment used by Srinivasan and Rose¹⁶ for the proteins studied does not fully agree with the one used in the PDB. In the following, our results are benchmarked against the PDB-based assignment.

To explore the role of local and nonlocal interactions, we consider two choices for the interaction window, Δ , of 6 and N . The interaction window restricts interactions along the sequence. $\Delta = 6$ means that all interactions between two residues i and j with $|i - j| > 6$ are switched off, whereas in the case of $\Delta = N$ all interactions are present.

RESULTS AND DISCUSSION

We begin our discussions with protein G (PDB code 1GB1), the protein showing the best conformational biases toward native secondary structures in the set of proteins studied by Srinivasan and Rose. Figure 1 shows P as a function of residue at three different temperatures $T = 0.8\epsilon/k_B, 0.5\epsilon/k_B,$ and $0.2\epsilon/k_B$. $T = 0.5\epsilon/k_B$ is the temperature Srinivasan and Rose used in their simulations. The interaction window is set to 6. The conformational biases toward α -helices (h), β -strands (s), turns (t), and coils (c) are shown. Note that at $T = 0.8\epsilon/k_B$ the strands dominate over all other structures. Thus, the whole protein chain prefers to be in the strand conformation at high tempera-

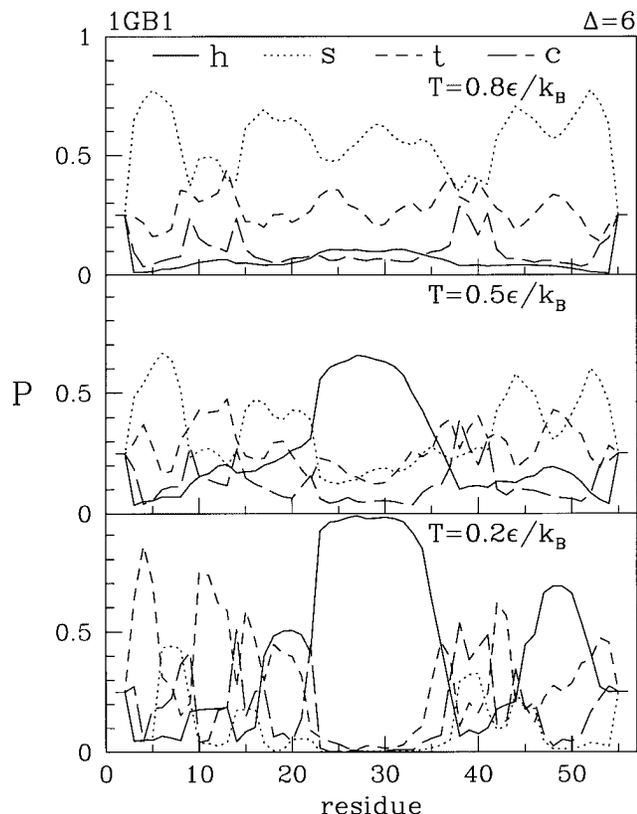


Fig. 1. Conformational biases towards secondary structures, P , as functions of residues determined for protein G at three different temperatures $T = 0.8, 0.5$, and $0.2\epsilon/k_B$. The types of secondary structures are denoted as h for α -helices (continuous line), s for β -strands (dotted line), t for turns (broken line), and c for coils (long broken line). The biases are computed by averaging over 10 trajectories each of 1000 cycles starting from an open conformation. The interaction window is set to 6.

tures. This follows from the simple observation that the entropy is largest in the strand conformation and is the dominant factor in the free energy at high temperatures. At low temperatures, such as $T = 0.2\epsilon/k_B$, the bias toward the strands vanishes while the highest biases belong to helices and turns. This is because helices and turns involve favorable interactions, which are predominantly local and are thus stabilized at low temperatures. At the intermediate temperature, $T = 0.5\epsilon/k_B$, the dominating structure varies as one proceeds along the sequence. Some parts of the protein prefer to be in a strand conformation while others form helices and turns.

Because the biases strongly depend on T , one may ask what the temperature is at which the native secondary structure can be most reliably predicted. To answer this question, we carried out an extensive analysis of the biases over a wide range of temperatures. Figure 2(a) shows two sequences of secondary structure assignments. The first corresponds to the known native conformation of protein G and the second is obtained from the biases given in the middle panel of Figure 1, that is, at $T = 0.5\epsilon/k_B$. In the latter case, an assignment at a given site is set to the type of secondary structure showing the highest bias. We introduce a parameter η that estimates overlaps between

a) 1GB1, $T=0.5\epsilon/k_B$, $\Delta=6$

```
PDB      -sssssssc ccsssssssc cchhhhhhhh
Predicted --ssssstt ttttssssss sshhhhhhhh

        hhhhhhtttcc csssstttt sssss-
        hhhhhhtttt tssssstts ssss--
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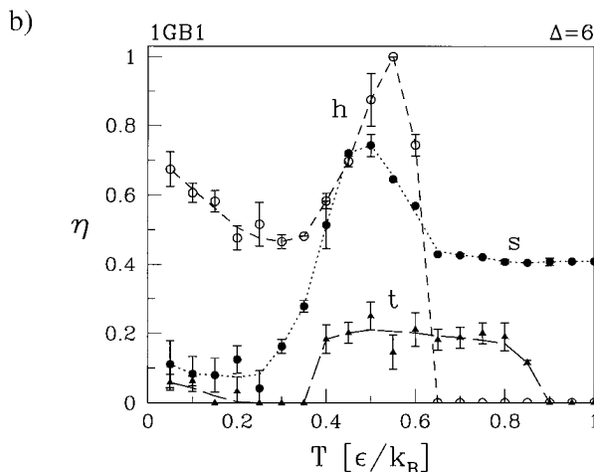


Fig. 2. **a**: Assignment of the secondary structures for protein G extracted from the PDB structure using the DSSP method of Kabsch and Sander²² (box) and predicted from the analysis of the conformational biases at $T = 0.5\epsilon/k_B$ and for $\Delta = 6$. **b**: Rate of success in prediction, η , as a function of temperature for three kinds of secondary structures: helix (h), strand (s), and turn (t). The simulations are performed for protein G with $\Delta = 6$. At each temperature studied, the conformational biases are computed by averaging over 10 trajectories each of 1000 cycles starting from an open conformation. The error bars are determined from three simulations at each temperature.

the two sets of assignments for each kind of secondary structure. For a given type of conformation, x ($x \in \{s, h, t, c\}$), η is defined as the number of sites at which both assignments (from PDB and from the biases) are x divided by the number of sites at which at least one of the assignments is x . Specifically, if A is a set of sites of type x in the PDB assignment and B is a set of sites of the same type of conformation predicted by the biases then

$$\eta \equiv \frac{f(A \cap B)}{f(A \cup B)},$$

where $f(X)$ is a function that returns the number of elements in X . Thus, if $A \equiv B$ then $\eta = 1$. We call η the rate of successful prediction.

Figure 2(b) shows η as a function of T for the strands, helices, and turns for protein G. Note that the values of η are the largest around $T = 0.5\epsilon/k_B$. At this temperature the rate of prediction is the highest for helices and exceeds 90% (it is 100% at $T = 0.5\epsilon/k_B$), while strands and turns are predicted at 74% and 23% levels, respectively. The values of η were obtained with the reference to PDB assignment. If the Srinivasan and Rose assignment is used instead, the corresponding success rates are 90, 63, and 35%, respectively. As T increases the rate for the strands first decreases and then remains roughly at a constant

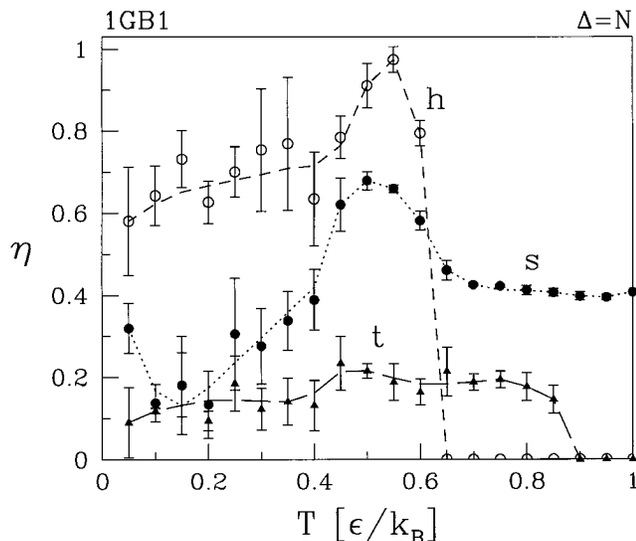


Fig. 3. The rate of success in prediction, η , as a function of temperature for three kinds of secondary structures—helix (h), strand (s), and turn (t)—for protein G with $\Delta = N$ or with no restriction on the range of interactions. The details are the same as in the lower part of Fig. 2.

value while the rate for helices drops rapidly and vanishes at $T = 0.7\epsilon/k_B$. A nearly opposite scenario is observed as T becomes smaller than 0.5—the rate for strands drops rapidly while it remains high for the helices.

Figure 3 shows η as a function of T for protein G but as calculated with $\Delta = N$. We still observe the same picture as for $\Delta = 6$, except that at low temperatures the prediction rates for helices and strands become somewhat higher. The optimal temperature, however, remains close to $T = 0.5\epsilon/k_B$.

Figures 4 and 5 show η as functions of T for six other proteins in the set studied by Rose for $\Delta = 6$ and N , respectively. For $\Delta = 6$, as in protein G, the best prediction rates are obtained at about $T = 0.5\epsilon/k_B$ for all of the proteins except for plastocyanin (6PCY) and myo hemerythrin (2HMQ). The latter proteins are special because they consist of only one kind of secondary structure in addition to the turn. The native conformation of plastocyanin is built only of β -strands and that of hemerythrin is a four-helix bundle. The best prediction rates are obtained for a range of temperatures that corresponds to $T \geq 0.4\epsilon/k_B$ for plastocyanin and to $T \leq 0.4\epsilon/k_B$ for hemerythrin. For $\Delta = N$, the optimal temperature varies a little but for most of the proteins it remains in the range from 0.4–0.6 ϵ/k_B . It should also be noted that, when $\Delta = N$, the rates for the strands at low temperatures become significantly larger than in the $\Delta = 6$ case for all proteins. The reason for this behavior is that, at low T , the strands can be stabilized only by nonlocal interactions, which are absent when $\Delta = 6$. The results in Figures 2–5 are in general similar even when the Srinivasan–Rose secondary structure assignment is used underscoring the robustness of our results. The only difference is that the predictions pertaining to the turns are improved compared to the PDB-based secondary structure assignment.

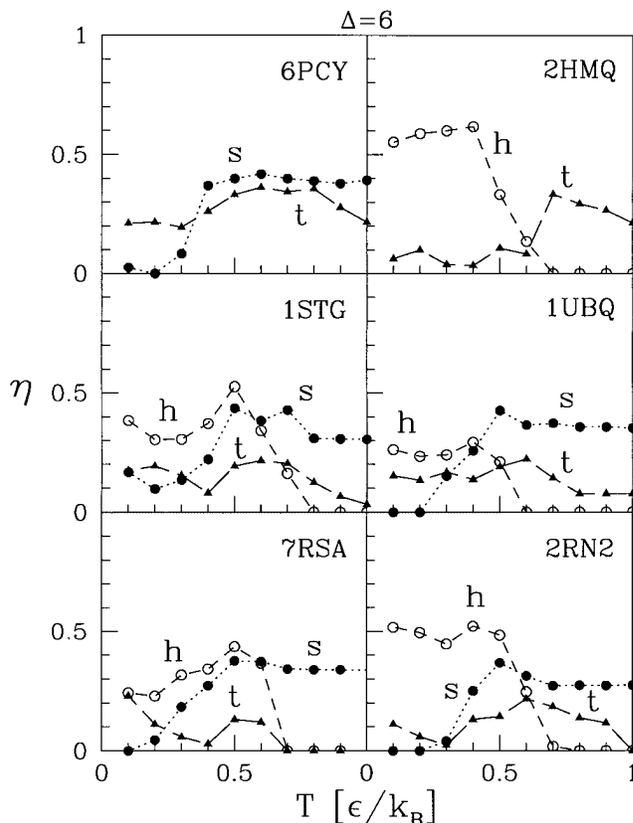
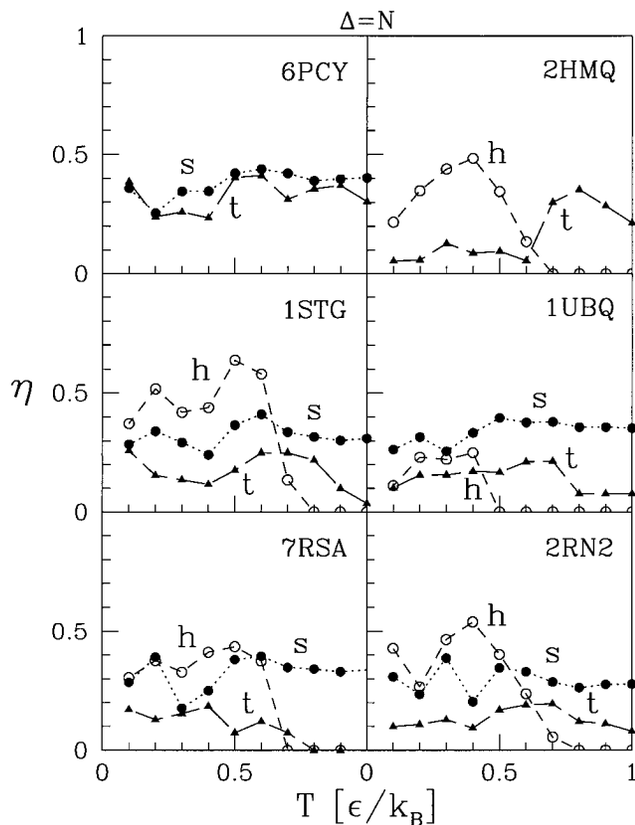
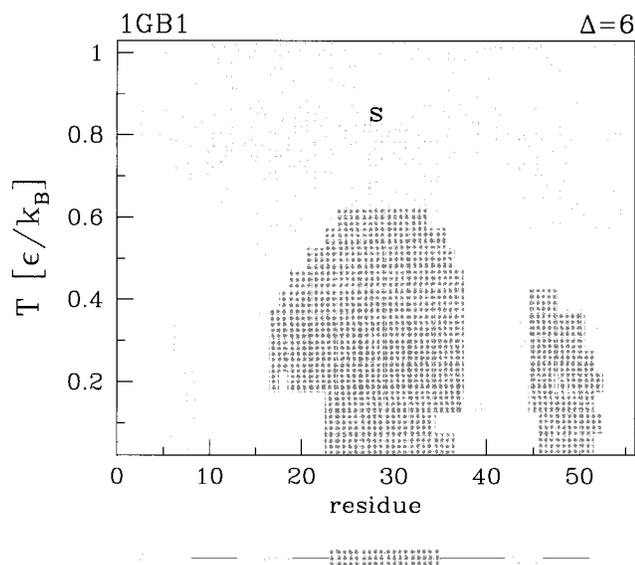


Fig. 4. Rate of successful prediction of secondary structures as a function of temperature for plastocyanin (6PCY), myo hemerythrin (2HMQ), staphylococcal nuclease (1STG), ubiquitin (1UBQ), ribonuclease A (7RSA), and ribonuclease H (2RN2). The simulations are performed in the same way as for protein G as described in the legend of Fig. 2. The interaction window is set equal to 6.

What is the principle that governs the choice of the optimal temperature? Figure 6 shows how conformational changes occur with respect to temperature for each residue of protein G for the case of $\Delta = 6$. It is seen clearly that most of the strands are destabilized at low temperatures whereas helices are absent at high temperatures. Thus, to have both kinds of structures predicted the optimal temperature for the prediction should be in a range of intermediate temperatures where helices have started to form but strands have not vanished. It can be seen in Figure 6 that as the temperature is lowered the strands undergo a transition to helices or other kinds of structures such as turns or coils. A helix can also be formed from a coil as the temperature continues to decrease. Because helices and turns are associated with the establishment of H-bonds while strands and coils usually have no contacts, such transitions entail a change in energy that is reflected in the specific heat. This suggests that the optimal temperature for secondary structure prediction ought to be in the vicinity of the peak of the specific heat, and likely a bit higher than the temperature of its maximum to not discriminate against the strands.

The connection between the thermodynamics of the system and the optimal temperature for the prediction are


 Fig. 5. Same as Fig. 4 but with $\Delta = N$.

 Fig. 6. Top: Conformational diagram plotted as a function of temperature and residue in protein G. The dark and light grey areas correspond to the helix (h) and strand (s) conformations, respectively. In these calculations, $\Delta = 6$. Bottom: strand (thin box) and helix (thick box) fragments found in the native conformation of protein G.

shown in Figures 7 and 8 for protein G and plastocyanin and hemerythrin, respectively. The specific heat C as a function of temperature is calculated using the histogram

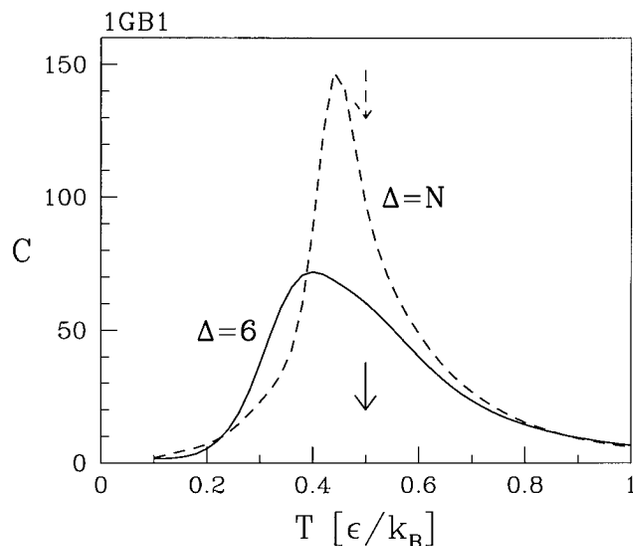
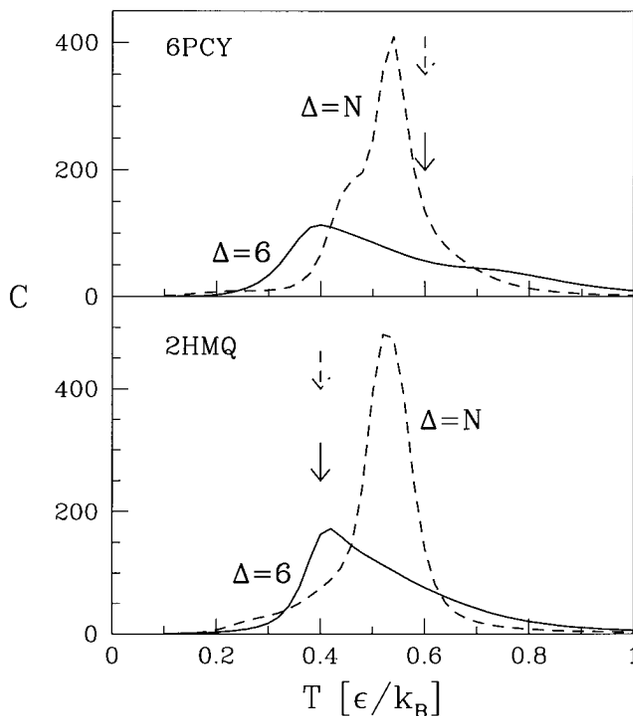

 Fig. 7. Specific heat as a function of temperature for protein G. The thermodynamic averages were carried out by performing a long simulation of 200,000 cycles at $T = 0.5\epsilon/k_B$ and then using the histogram method to extract quantities at other temperatures. The lower peak (continuous line) and the higher one (broken line) correspond to the interaction window equal to 6 and N , respectively. The arrows show the temperatures at which the secondary structures are best predicted for the two values of Δ .


Fig. 8. Same as Fig. 7 but for plastocyanin (top) and myo hemerythrin (bottom).

technique.²⁴ For each protein we performed a long simulation of 200,000 cycles at $T = 0.5\epsilon/k_B$ to deduce the thermodynamic behavior at that temperature and other temperatures in its vicinity. The results are shown for the

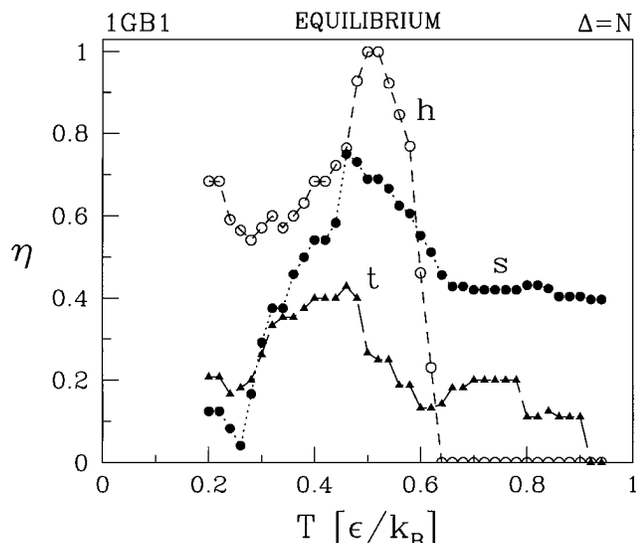


Fig. 9. Same as Fig. 3 but the rates of successful prediction are computed from an equilibrium simulation. The secondary structure biases as a function of temperature are computed using the histogram method. A long simulation of 200,000 cycles at $T = 0.5\epsilon/k_B$ is performed to extract quantities at other nearby temperatures.

two values of Δ . For $\Delta = 6$, the maximum in C occurs roughly at $T = 0.4\epsilon/k_B$ for the three proteins. For $\Delta = N$, the magnitude of the peak in C is higher and it also occurs at a slightly higher temperature due to the presence of the long-range interactions. (It is interesting to note that the experimentally determined specific heat of plastocyanin²⁵ shows a sharp maximum around 68°C.) Note that for protein G the optimal temperature for the secondary structure prediction is found in the vicinity of the peak in the specific heat—just to the right of the maximum. A similar behavior is observed in the case of plastocyanin except that the optimal temperature at $\Delta = 6$ is farther from the maximum in C . This is due to the fact that plastocyanin consists only of β -sheets and the strands are more favored at high temperatures. However, in the case of myo hemerythrin, whose native state contains mainly of α -helices, the behavior is just the opposite. The optimal temperature for the prediction is now on the low-temperature side of the peak in the specific heat. A comparison of Figures 4 and 5 shows that at the temperature corresponding to the maximum in C the rates of successful prediction are already close to their best values.

The results described so far are based on short simulations that last for 1000 cycles at each T . Figure 9 shows η as a function of T for protein G when the conformational biases are determined from nearly equilibrium simulations. These simulations are performed similar to the calculation of the specific heat: We make a long simulation of 200,000 cycles at $T = 0.5\epsilon/k_B$ and then use the histogram method to obtain the biases at other temperatures. The profiles of η over T are surprisingly similar to those shown in Figure 3 and the peaks seem to be even more pronounced. The predictions given by the conformational biases are found to be insensitive to the length of simula-

tions, at least for a range of temperatures that are close to the optimal value.

CONCLUSIONS

Our results confirm that the analysis of conformational biases is a fast and useful tool to get information about native protein secondary structures. We find that the most common secondary motifs, α -helices and β -strands, can be predicted with an accuracy ranging from roughly 40% up to 100%. Our analysis shows that while the rate of successful prediction is insensitive to the interaction window, as well as to the length of the simulations, the choice of temperature appears to be critical. The optimal run temperature is found to be related to the peak temperature in the specific heat. Unlike commonly used algorithms in which one attempts to minimize an energy function to determine the native-state structure, LINUS is an algorithm that relies on a delicate interplay between the entropy favoring the strands and energetic considerations favoring turns and helices. Because there is no procedure in LINUS that allows for an assembly of strands through the appropriate nonlocal interactions into a sheet, secondary structure prediction in essence depends on the persistence of a strand conformation down to intermediate temperatures in regions corresponding to strands in the native structure, while other regions adopt the helix and turn conformations due to the energy gain through the local contacts. An improvement in the prediction might be expected on extending LINUS to some judiciously chosen nonlocal interactions for the assembly of β -sheets.

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