Are Protein Folds Atypical?

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Abstract

Protein structures are a very special class among all possible structures. It was suggested that a “designability principle” plays a crucial role in nature’s selection of protein sequences and structures. Here we provide a theoretical base for such a selection principle, using a novel formulation of the protein folding problem based on hydrophobic interactions. A structure is reduced to a string of 0’s and 1’s which represent the surface and core sites, respectively, as the backbone is traced. Each structure is therefore associated with one point in a high dimensional space. Sequences are represented by strings of their hydrophobicities and thus can be mapped into the same space. A sequence which lies closer to a particular structure in this space than to any other structures will have that structure as its ground state. Atypical structures, namely those far away from other structures in the high dimensional space, have more sequences which fold into them, and are thermodynamically more stable. We argue that the most common folds of proteins are the most atypical in the space of possible structures.
Protein structures seem to be a very special class among all possible folded configurations of a polypeptide chain. There are preferred secondary structures and motifs as well as striking regularities in the geometries of protein structures. Two proteins are said to have a common fold if they have the same major secondary structures in the same arrangement with the same topological connections. Common folds occur even for proteins with different biological functions. Indeed there are “superfolds” and “fold space attractors” which account for the structures of many proteins. It has been estimated that the total number of natural protein folds is only about 1000, an extremely small number compared with both the number of proteins and the number of all possible structures. One may ask: is there anything special about natural protein structures – are they merely an arbitrary outcome of evolution or is there some fundamental reason behind their selection?

Many features of folded proteins can be understood from an energetic point of view. Close packing of secondary structures favors some geometrical patterns. It was argued that certain empirical rules for connections between secondary structures can be explained by lower bending energies for the connecting loops. It was further conjectured that a structure with lower energy would stabilize more sequences and would be more likely a protein fold. However, the number of low energy structures is huge, and overall low energy does not necessarily imply thermodynamic stability since other low energy structures still compete with the ground state. Clearly, purely energetic considerations are not the whole story.

A recent study on a simple model of protein folding suggested that a rather different mechanism – the so-called “designability principle” – should play a crucial role in nature’s selection of protein sequences and structures. The designability of a structure is defined as the number of sequences that possess the structure as their nondegenerate ground state. It was demonstrated in the model that structures with the same low energy (when averaged over sequences) differ drastically in terms of their designability; highly designable structures emerge with a number of associated sequences much larger than the average. These highly designable structures are relatively stable against mutation and are more thermodynamically
stable than other structures. In addition, they possess “proteinlike” secondary structures and motifs. A number of questions arise: Among the large number of low energy structures, why are some structures highly designable? Why does designability also guarantee thermodynamic stability? Why do highly designable structures have geometrical regularities? Here we address these questions using a novel formulation of protein folding problem based on hydrophobic interactions.

Among the various forces involved in the folding of a polypeptide chain – van der Waals force, electrostatic force, hydrogen bonding, hydrophobic force – there is strong and increasing evidence that the hydrophobic force is the dominant one in determining the overall folded structure \[13,14\]. The hydrophobic force originates from the contact of nonpolar groups with water, which disrupts the hydrogen bonding exchange pattern between water molecules. Thus nonpolar groups in water tend to coalesce to minimize their contact with water. For a nonpolar amino acid, the free energy reduction from the hydrophobic interaction is proportional to the total area of the side chain protected from water \[13,18\]. For an amino acid with a polar side chain, there is a smaller reduction because of the possibility of hydrogen bonding between the polar side chain and water molecules \[13\]. To model the hydrophobic force in protein folding, one can assign parameters \( h_\sigma \) to characterize the hydrophobicities of each of the 20 amino acids \[18\]. Each sequence of amino acids then has an associated vector \( h = (h_\sigma_1, h_\sigma_2, \ldots, h_\sigma_i, \ldots, h_\sigma_N) \), where \( \sigma_i \) specifies the amino acid at position \( i \) of the sequence. We take the energy of a sequence folded into a particular structure to be the sum of the contributions from each amino acid upon burial away from water:

\[
H = -\sum_i s_i h_\sigma_i, \tag{1}
\]

where \( s_i \) is a structure-dependent number characterizing the degree of burial of the \( i \)-th amino acid in the peptide chain \[13\]. Here we ignore all other forces which, undoubtedly, help determine the details of a protein’s structure. The advantage of considering only the hydrophobic force is that it drastically simplifies the analysis and thereby elucidates some
essential features of the folding problem.

To simplify the application of Eq. (1), let us consider only globular compact structures and let $s_i$ take only two values: 0 and 1, depending on whether the amino acid is on the surface or in the core of the structure, respectively. Therefore, each compact structure can be represented by a string $\{s_i\}$ of 0’s and 1’s: $s_i = 0$ if the $i$-th amino acid is on the surface and $s_i = 1$ if it is in the core (see Fig. 1 for an example on a lattice). Assuming every compact structure of a given size has the same numbers of surface and core sites and noting that the term $\sum_i h_{\sigma_i}^2$ is a constant for a fixed sequence of amino acids and does not play any role in determining the relative energies of structures folded by the sequence, Eq. (1) is equivalent to:

$$H = \sum_{i=1}^{N} (h_{\sigma_i} - s_i)^2. \quad (2)$$

Having formulated the protein folding problem in terms of Eq. (2), we now proceed to make a few observations. The problem involves two spaces: the sequence space and the structure space. We represent a sequence by the vector of its hydrophobicities $h = (h_{\sigma_1}, h_{\sigma_2}, \ldots, h_{\sigma_N})$, and the sequence space $\{h\}$ consists of $20^N$ sequences since there can be any of 20 amino acids at each site. A structure is also represented by a vector $s = (s_1, s_2, \ldots, s_N)$, where $s_i = 0$ or 1, and the structure space $\{s\}$ consists of the $2^N$ possible strings of 0’s and 1’s. However, only a small subset of the strings of 0’s and 1’s represent realizable structures. If two or more structures map into the same string, we say that these structures are degenerate (see Fig. 1(a)). It is evident that a degenerate structure cannot be the unique ground state for any sequence within this formulation. The fraction of all structures which are nondegenerate depends on the ratio of surface sites to core sites. This fraction approaches zero in the limits of very large and very small surface-to-core ratios. It is worthwhile noting that for natural proteins the surface-to-core ratio is of the order one.

Now imagine embedding both the sequence space $\{h\}$ and the structure space $\{s\}$ in an $N$-dimensional euclidean space (Fig. 2). This is simplest to picture if one normalizes the $h_{\sigma}$ so that $0 \leq h_{\sigma} \leq 1$. Since the energy for a sequence $h$ folded into a structure $s$ is the square
of the distance between $h$ and $s$ (Eq. (2)), it is evident that $h$ will have $s$ as its unique ground state if and only if $h$ is closer to $s$ than to any other structure. Therefore, the set of all sequences $\{h(s)\}$ which uniquely design a structure $s$ can be found by the following geometrical construction: Draw bisector planes between $s$ and all of its neighboring structures in the $N$-dimensional space (see Fig. 2). The volume enclosed by these planes is called the Voronoi polytope around $s$. $\{h(s)\}$ then consists of all sequences within the Voronoi polytope. Hence, the designabilities of structures are directly related to the distribution of the structures in the structure space $\{s\}$. A structure closely surrounded by many neighbors will have a small Voronoi polytope and hence a low designability; while a structure far away from others will have a large Voronoi polytope and hence a high designability. Furthermore, the thermodynamic stability of a folded structure is directly related to the size of its Voronoi polytope. For a sequence $h$, the energy gap between the ground state and an excited state is the difference of the squared distances between $h$ and the two states (Eq. (4)). A larger Voronoi polytope implies, on average, a larger gap as excited states can only lie outside of the Voronoi polytope of the ground state. Thus, this geometrical representation of the problem naturally explains the positive correlation between the thermodynamic stability and the designability, an observation made in Ref. [12].

To further illustrate and elaborate on the above ideas, let us proceed with a simple example: a two-dimensional lattice HP model [20]. Instead of 20, we use only two amino acids: H (hydrophobic) and P (polar). The vector representing a sequence is now $h = (h_1, h_2, \ldots, h_i, \ldots, h_N)$, where $h_i = 1$ if the $i$-th amino acid is an H and $h_i = 0$ if it is a P. The sequence space now coincides with the structure space, both consisting of all the possible strings of 0’s and 1’s of length $N$. To obtain a set of allowed structure strings, we focus on the compact $6 \times 6$ two-dimensional lattice structures (Fig. 1), which can be easily enumerated. We divide the 36 sites into 20 surface sites and 16 core sites; the surface/core ratio is 1.25. There are 57,337 compact structures not related by symmetries. These structures map into 30,408 distinct strings, among which 18,213 ($\sim 30\%$ of all structures) represent nondegenerate structures. To obtain a histogram of the designability for all structures, we
randomly sampled the sequence space. We randomly chose \( \sim 20 \) million sequences (enough to suppress statistical fluctuations), and for each of these calculated its energy on all the structure strings using Eq. (1). We found that \( \sim 8.8\% \) of all the sequences chosen have unique ground states, i.e., they fall inside of the Voronoi polytope of some nondegenerate structure. The designability histogram obtained in the sampling is plotted in Fig. 3. Recall that the designability of a structure \( D(s) \) is the number of sequences found to have that structure as their unique ground state. The designabilities obtained from this model are very well correlated with those found in the previous work [12] which were obtained from a different HP model, and with those calculated using the Miyazawa-Jernigan interaction matrix [21] for all 20 amino acids. The sets of highly designable structures are essentially the same for all three models.

Certain features of Fig. 3 can be understood from the following simple consideration. If the structure strings were randomly distributed in structure space, then one would expect the distribution of the volumes \( V \) of the Voronoi polytopes constructed around every structure to be \( \propto \exp(-cV) \), for \( cV \gg 1 \), where \( c \) is proportional to the overall density of structure strings in structure space [22]. Moreover, the probability of finding very small Voronoi volumes \( V \) would also vanish because small \( V \)'s require a close clustering of randomly distributed points [23]. The distribution shown in Fig. 3 indeed has a rising part for small designability \( D \) and a decaying part which is roughly exponential. However, one should not conclude from this that the structures are randomly distributed in structure space. In fact, the structures are highly correlated and clustered. For example, in Fig. 3 there are structures (with \( D(s) > 500 \)) which are highly designable and which do not follow the exponential decay.

In order to more fully understand the designabilities of structures, one has to know the distribution of structures in structure space. Each realizable string \( s \) represents one (sometimes more than one) globular compact structure; its components \( s_i \) are determined by the geometry of the represented structure. One trivial correlation between the realizable \( s \)'s comes from the constraint that \( \sum_i s_i = n_c \), where \( n_c \) is the number of core sites and is a constant depending only on \( N \). Therefore the allowed structures reside in a subspace
consisting of those strings whose Hamming distance \(d\) to the origin equals \(n_c\). Other more subtle correlations are related to the properties of compact self-avoiding walks of length \(N\). To characterize the ensemble properties of the allowed structures, we measure the correlation function \(c(i,j) = \langle s_i s_j \rangle - \langle s_i \rangle \langle s_j \rangle\), where the average is taken over all 57,337 compact structures. In Fig. 4, \(c(i,j)\) is plotted vs. \(j - i\) for three different starting positions: \(i = 1, i = 9,\) and \(i = 18\). A common feature of these correlation functions is that \(s_i\) has positive correlation with nearby sites and that the correlation turns negative as \(j - i\) increases. This feature simply reflects a property of self-avoiding random walks in a confined geometry: if a site is in the core then most likely the next few sites are also in the core, and similarly if a site is on the surface the next few sites are most likely on the surface. From Fig. 4 we see that the transition between core and surface takes, on average, about 4 to 6 steps. The residual negative correlation at large distances is a result of the constraint that there is a fixed number of core (and surface) sites.

Another way to measure the correlation of the allowed structures in structure space is to measure the number of distinct structure strings, \(n_s(d)\), at a Hamming distance \(d\) from a given string \(s\). In Fig. 5, \(n_s(d)\) is plotted for three different structures with low, intermediate, and high designabilities, respectively. A highly designable structure typically has fewer neighbors than a less designable structure, not only at the smallest \(d\)'s but out to \(d\)'s of order 10 - 12. This indicates that there exists a local “density” around each allowed structure in the structure space.

From the above discussion, we see that structure strings are not randomly distributed in the structure space. Their distribution is highly correlated. There are high density regions as well as regions with very low density. High density regions consist of “typical” structures or typical random walks (see Fig. 1(a) for an example) whose correlation functions are similar to the correlation function of the ensemble (Fig. 4). These structures usually have small Voronoi polytopes because they are closely surrounded by many neighbors. “Atypical” or “rare” structures reside in regions with very low local density, and their correlation functions are generally very different from Fig. 4. Atypical structures usually have high designabilities.
because they are relatively far away from other structures in the structure space. One way for a structure to have an atypical correlation function is to have many surface to core transitions along its backbone. This leads to geometrically regular pleated patterns such as the one shown in Fig. 1(b), which is in fact the most designable structure. Thus, the emergence of the “proteinlike” sub-structures and motifs in the highly designable structures [12] occurs not because they are common, but precisely because they are rare. A related property of rare structures, which follows from the large Voronoi polytope, is that it is hard to make any local change to them. For the “typical” structure in Fig. 1(a), one can make local changes as indicated by the dotted lines to transform the structure to other compact structures. It is impossible to perform any such local changes for the most designable – and therefore most “atypical” – structure in Fig. 1(b).

In conclusion, we have formulated the protein folding problem in a novel way using only hydrophobic interactions. A very simple and transparent picture emerges from the formulation in which the designability of a structure, i.e., the number of sequences for which the structure is a unique ground state, is directly related to the volume of the Voronoi polytope around the structure. Structures with atypical patterns of surface and core sites have high designability and are generally more stable thermodynamically. These atypical patterns produce geometrically regular structures with “proteinlike” motifs. According to this picture, the most common folds in natural proteins are the most atypical in the space of possible structures. Note that highly designable structures do not have lower energies than other structures. It is easy to see from Eq. (1) that the average energy and the energy spectrum (over all the sequences) are the same for all the compact globular structures. Thus a structure is highly designable in our model, not because it has a lower energy or an unusual energy spectrum as conjectured by Finkelstein and colleagues [11], but because it is far away from neighboring competing structures in the structure space. In our discussion we considered only the hydrophobic interaction and compact globular structures. The formulation can be generalized, at least conceptually, without much difficulty to include non-compact structures and main-chain hydrogen bonding, both of which are purely structural proper-
ties. The overall picture is unchanged. Our results can be tested on real protein structures.

For a globular protein there is a natural division into core and surface sites and thus a string representation. The strings of natural proteins can be compared with the ensemble of strings of random structures. The knowledge of the distribution of allowed structures in structure space can be used to aid protein structure classification, protein design, and structure prediction.

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FIG. 1. Structures are represented by strings $s$ of 0’s and 1’s, according to whether a site is on the surface or in the core, respectively. Shown are two examples of compact $6 \times 6$ lattice structures. (a) A typical structure. Dotted lines indicate local changes that can be performed to transform it to other compact structures. Note that the change at the lower right corner does not change the string pattern, so this structure is a degenerate one. (b) The most designable structure.
FIG. 2. Schematic plot of the sequence and the structure spaces and the Voronoi construction.

The Voronoi polytope is the shaded region.

FIG. 3. Histogram of the designability obtained by random sampling using 19,492,200 sequences.
FIG. 4. Correlation functions for the structures.

FIG. 5. Number of structures vs. the Hamming distance for three typical structures with: low (circles), intermediate (triangles), and high (squares) designability.
REFERENCES

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[1] Levitt, M. & Chothia, C. Structural patterns in globular proteins. *Nature* **261**, 552-558 (1976).

[2] Richardson, J.S. Handedness of crossover connections in \(\beta\)-sheets. *Proc. Nat. Acad. Sci., U.S.A.* **73**, 2619-2623 (1976).

[3] Richardson, J.S. The anatomy and taxonomy of protein structures. *Advan. Protein Chem.* **34**, 167-339 (1981).

[4] Murzin, A.G., Brenner, S.E., Hubbard, T. & Chothia, C. SCOP: a structural classification of protein database for the investigation of sequences and structures. *J. Mol. Biol.* **247**, 536-540 (1995).

[5] Orengo, C.A., Jones, D.T., & Thornton, J.M. Protein superfamilies and domain superfolds. *Nature* **372**, 631-634 (1994).

[6] Holm, L. & Sander, C. Mapping the protein universe. *Science* **273**, 595-603 (1996).

[7] Chothia, C. One thousand families for the molecular biologist. *Nature* **357**, 543-544 (1992).

[8] Chothia, C. & Finkelstein, A.V. The classification and origins of protein folding patterns. *Annu. Rev. Biochem.* **59**, 1007-1039 (1990).

[9] Chothia, C., Levitt, M. & Richardson, D. Structure of proteins: packing of \(\alpha\)-helices and \(\beta\)-sheets. *Proc. Natl. Acad. Sci. USA* **74**, 4130-4134 (1977).

[10] Finkelstein, A.V. & Ptitsyn, O.B. Why do globular proteins fit the limited set of folding patterns? *Prog. Biophys. Mol. Biol.* **50**, 171-190 (1987).

[11] Finkelstein, A.V., Gutun, A.M. & Badretdinov, A.Ya. Why are the same protein folds used to perform different functions? *FEBS* **325**, 23-28 (1993).
[12] Li, H., Helling, R., Tang, C. & Wingreen, N. Emergence of preferred structures in a simple model of protein folding. *Science* **273**, 666-669 (1996).

[13] Kauzmann, W. Some factors in the interaction of protein denaturation. *Adv. Protein Chem.* **14**, 1-63 (1959).

[14] Dill, K.A. Dominant forces in protein folding. *Biochemistry* **29**, 7133-7155 (1990).

[15] Chothia, C. Hydrophobic bonding and accessible surface area in proteins. *Nature* **248**, 338-339 (1974).

[16] Hermann, R.B. Theory of hydrophobic bonding. II. Correlation of hydrocarbon solubility in water with solvent cavity surface area. *J. Phys. Chem.* **76**, 2754-2759 (1972).

[17] Richards, F.M. Areas, volume, packing, and protein structure. *Ann. Rev. Biophys. Bioeng.* **6**, 151-176 (1977).

[18] Eisenberg, D. & McLachlan, A.D. Solvation energy in protein folding and binding. *Nature* **319**, 199-203 (1986).

[19] It has recently been shown that the Miyazawa-Jernigan matrix [21] contains a dominant hydrophobic interaction of the form $E_{\alpha \beta} = h_\alpha + h_\beta$ (Li H., Tang, C. & Wingreen, N.S. Nature of driving force for protein folding: a result from analyzing the statistical potential. *Phys. Rev. Lett.* **79**, 765 (1997)). Eq. (1) can be derived from this form of the contact energy with $s_i$ being the number of non-covalent nearest neighbor contacts for the $i$-th amino acid.

[20] Lau, K.F. & Dill, K.A. A lattice statistical mechanics model of the conformational and sequence spaces of proteins. *Macromolecules* **22**, 3986-3997 (1989).

[21] Miyazawa, S. & Jernigan, R.L. Residue-residue potentials with a favorable contact pair term and an unfavorable high packing density term, for simulation and threading. *J. Mol. Biol.* **256**, 623-644 (1996).
[22] Gilbert, E.N. Random Subdivisions of space into crystals. *Ann. Math. Stat.* **33**, 958-972 (1962).

[23] In one dimension, the distribution function for Voronoi segments $l$ from randomly distributed points is $c^2 l \exp(-cl)$. For two and three dimensions, numerical studies show that the distribution can be best fit by a gamma distribution, $V^{a-1} \exp(-bV)/b^a \Gamma(a)$ (Hinde, A.L. & Miles, R.E. Monte Carlo estimates of the distributions of the random polygons of the Voronoi tessellation with respect to a Poisson process. *J. Stat. Comput. Simul.* **10**, 205-223 (1980); Kumar, S., Kurtz, S.K., Banavar, J.R. & Sharma, M.G. Properties of a three-dimensional Poisson-Voronoi tessellation: a Monte Carlo study. *J. Stat. Phys.* **67**, 523-551 (1992).) We have found that the designabilities for a randomly distributed set of structure strings of length 36 can also be fit by the gamma distribution.

[24] The Hamming distance between $s$ and $s'$, two strings of 0’s and 1’s, is defined as $d = \sum_i |s_i - s'_i|$. 

