Fast tree search for enumeration of a lattice model of protein folding

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Using a fast tree-searching algorithm and a Pentium cluster, we enumerated all the sequences and compact conformations (structures) for a protein folding model on a cubic lattice of size $4 \times 3 \times 3$. We used two types of amino acids—hydrophobic (H) and polar (P)—to make up the sequences, so there were $2^{36} \approx 6.87 \times 10^{10}$ different sequences. The total number of distinct structures was 84,731,192. We made use of a simple solvation model in which the energy of a sequence folded into a structure is minus the number of hydrophobic amino acids in the “core” of the structure. For every sequence, we found its ground state or ground states, i.e., the structure or structures for which its energy is lowest. About 0.3% of the sequences have a unique ground state. The number of structures that are unique ground states of at least one sequence is 2,662,050, about 3% of the total number of structures. However, these “designable” structures differ drastically in their designability, defined as the number of sequences whose unique ground state is that structure. To understand this variation in designability, we studied the distribution of structures in a high-dimensional space in which each structure is represented by a string of 1’s and 0’s, denoting core and surface sites, respectively.

I. INTRODUCTION

The protein folding problem has long attracted the attention of scientists from various disciplines. The relationship between the amino-acid sequence and the three-dimensional structure of a protein is not only an extremely important and practical problem in biology, but also a fundamental problem in science. Despite a tremendous amount of effort and progress over many decades, the problem remains essentially unsolved. At least part of the difficulty arises from the intrinsic complexity of the protein-folding problem. Since the seminal work of Anfinsen about 40 years ago, it has been demonstrated that the native state of a small, single domain protein is the global minimum of the free energy. However, the minimum-free-energy conformation of a polypeptide chain is “hiding” in a large space of $z^N$ conformations, where $N$ is the length of the chain and $z$ is the effective coordination number. Even if we count only the compact conformations for which $z \approx 2$ (see below) and take $N = 100$ for typical small proteins, the number of conformations is huge, $z^N \approx 2^{100} \sim 10^{30}$. On top of this huge conformational space lies the heterogeneity of amino acids. The 20 natural amino acids differ in size, hydrophobicity, and other physical and chemical properties. This heterogeneity is coupled with two intrinsic features of polymers: chain connectivity and the excluded volume effect. The free energy is a sensitive and complicated function in this huge conformational space with complex constraints.

In the last decade or so, there has been increasing interest in studying simple lattice models of protein folding. In these models, polypeptide chains are represented by self-avoiding walks on a regular lattice (e.g., Fig. 1), greatly simplifying the conformational space. Very often the sequence space and hence the heterogeneity is also simplified by using only two types of amino acids: hydrophobic (H) and polar (P). These so-called “HP lattice models” nonetheless capture some essential features of the protein folding problem. Simple lattice models have been applied to a wide range of problems including collapse and folding transitions, the influence of packing on secondary-structure formation, and differences in the designability of structures. The advantage of HP lattice models is that they are simple enough to be amenable to thorough theoretical study. These studies can provide fruitful insights to feed back to or test against realistic models and experiments.

One approach for calculating thermodynamic and other properties of lattice models is to enumerate all possible sequences and conformations. Since native globular protein structures and presumably most of the low-energy states are compact, enumeration studies are usually done on compact conformations only. The number of compact conformations $C_N$ scales with the chain length $N$ as $C_N \approx z^N$, where $z$ is an effective coordination number. Numerical estimation gives $z \approx 1.47$ for 2D square lattices and $z \approx 1.86$ for 3D cubic lattices, in good agreement with mean-field calculations of $Z/e$ and $(Z-1)/e$, respectively.

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where \( Z \) is the coordination number of the lattice and \( e = 2.718 \cdots \) is the base of the natural logarithm. (For real peptide chains the number of “distinct” states an amino acid can take, as estimated very roughly from Ramachandran plots of dihedral-angle frequencies, is about 5 or 6, which gives \( Z = 2 \).) For HP models in which there are only two types of amino acids, the number of sequences is \( 2^N \). If in the enumeration study the energies of every sequence folded into every compact conformation are evaluated, the total number of energy calculations is \( 2^N \times C_N \). The largest system previously evaluated in this way is an \( N = 27 \) (3 \( \times \) 3 \( \times \) 3) cubic lattice model,\(^{13} \) where \( 2^N \times C_N = 2^{27} \times 103 346 = 10^{13} \). Several interesting results were found in the enumeration of the 27-mer, in particular the idea of designability and its relation to thermodynamic stability.\(^{13} \) However, \( N = 27 \) is still small compared with typical protein sizes. It would be very desirable to enumerate larger systems if at all possible. In this paper, we report results of a complete enumeration of an \( N = 36 \) (4 \( \times \) 3 \( \times \) 3) cubic-lattice model. The number of compact conformations is\(^{16} \) 84 731 192, so, naively, the total number of energy calculations is \( 2^{36} \times 84 731 192 \approx 6 \times 10^{18} \). The task was made possible by using a binary model, a fast tree-search algorithm yielding a speed-up factor of 1600, and a 53-processor 200 MHz Pentium Pro cluster.

II. THE MODEL

The protein folding model we use in the enumeration study is the solvation model discussed in Li et al.\(^{20} \) There is considerable evidence that hydrophobic solvation forces are primarily responsible for the folding of a protein into a particular configuration.\(^{21-23} \) The hydrophobicity of each amino acid can be determined experimentally\(^{24-26} \) or by statistical analysis.\(^{23,27} \) It is energetically favorable for the more hydrophobic amino acids to occupy core sites,\(^{28} \) where there is low exposure to water. In the model, we denote a sequence of amino acids by \{\( \sigma_i \)\}. We take only two types of amino acids: hydrophobic H (\( \sigma = 1 \)) and polar P (\( \sigma = 0 \)). A “structure” is the set of all reflections and rotations of a given compact conformation. The energy of a sequence folded into a structure is taken to be the sum of the contributions from each amino acid upon burial away from water,

\[
E = -\sum_i \sigma_i s_i,
\]

where \( s_i \) is a structure-dependent number characterizing the degree of burial of the \( i \)th amino acid in the chain. Larger \( s_i \) corresponds to a smaller surface area accessible to the solvent. For a 4 \( \times \) 3 \( \times \) 3 structure on a cubic lattice there are four different kinds of sites: center, face, edge, and corner (see Fig. 1). So, in principle, there could be four different values of \( s_i \). To simplify the calculation, we take only two values for \( s_i \); we define a string \{\( s_i \)\} for each structure with \( s_i = 1 \) if the \( i \)th site is a “core” (center or face) and \( s_i = 0 \) if it is a “surface” (edge or corner). Thus each compact structure of 4 \( \times \) 3 \( \times \) 3 is mapped into a string of 1’s and 0’s, \{\( s_i \)\}, with 12 1’s (“cores”) and 24 0’s (“surfaces”). The surface-to-core ratio is 2, close to the values for small natural proteins. With this simplification, the energy in Eq. (1) is just minus the dot product of two binary strings. For a given sequence \{\( \sigma_i \)\}, a ground-state structure is one that minimizes Eq. (1). A sequence may have more than one ground state structure, but we will be primarily interested in sequences with unique ground states. Out of the 84 731 192 compact structures, the number of distinct structure strings is 14 062 236, among which 2 662 050 (corresponding to 1 331 025 lattice conformations) each represent exactly one structure. Each of the remaining 11 400 186 36-bit strings represents multiple structures. We also analyzed a 3 \( \times \) 3 \( \times \) 3 model with 7 “cores” (1 center and 6 faces) and 20 “surfaces” (12 edges and 8 corners). In this case, there are 103 346 structures and 6 291 distinct structure strings, among which only 120 (corresponding to 60 lattice conformations) represent exactly one structure apiece.

III. TREE SEARCHING ALGORITHM

Because the protein chains in our model are considered to be directed, there are generally two structures for each geometrical conformation, related by reversal of the direction of the chain (see Fig. 1). A small subset of structures are their own reversals. A structure string can be its own reversal even if its associated structure is not reversal symmetric. We found that among the 14 062 236 structure strings in the 4 \( \times \) 3 \( \times \) 3 model, there are 2 850 which are their own reversals. The remaining 14 059 386 strings form 7 029 693 pairs, with the two members of each pair being the reversals of each other. To reduce memory use, we keep only one member of each such pair, with an extra bit tagged on the string to indicate that it actually represents two strings: itself and its reversal. There are thus 7 029 693 + 2 850 = 7 032 543 distinct strings which we keep in the calculation.

Each of the 7 032 543 distinct 36-bit structure strings \{\( s_i \)\} has exactly 12 1’s and 24 0’s. Our goal is to find a way, given a 36-bit sequence string \{\( \sigma_i \)\}, to find if there is a unique entry in the table which maximizes the dot product of the structure string with \{\( \sigma_i \)\}, or, equivalently, which minimizes the energy of the sequence \{\( \sigma_i \)\} according to Eq. (1).
all strings in the node the total dot product is at least as big as the dot product of \( \{ \sigma_i \} \) with known-ones. On the other hand, the total dot product is at most \( \{ \sigma_i \} \) dotted with known-ones plus the dot product of \( \{ \sigma_i \} \) with undecided. Another upper bound for the total dot product is \( \{ \sigma_i \} \) dotted with known-ones plus the integer missing-ones.

Given such a tree, where the root corresponds to all of the 7032 543 entries, and a 36-bit sequence string \( \{ \sigma_i \} \), here is how we search the table.

Compute an upper bound for the total dot product using the smaller of the two upper bounds described above. Call this the “goal” \( G \). If there are any entries which achieve this goal, we are done. If not, repeat with the reversed version of the sequence. (This is necessary because our table contains only one member of each reversed pairs of structure strings. Whenever a ground state structure string is found for a sequence, the reversed sequence necessarily has the reversed structure string as a ground state.) Again, if we achieve the goal \( G \), we are done. If not, decrease the goal \( G \) by 1 and try again. Repeat until the goal is satisfied.

Given a goal \( G \), we search the tree as follows, starting at the root node:

If the bound on the node indicates that goal \( G \) is unachievable, return failure.

If the node is a leaf node, check each entry. If one is found that satisfies the goal, return success. If not, then return failure.

If the node is not a leaf node, try each of the children. We first try the child that matches \( \{ \sigma_i \} \). That is, if the children split on the value at the \( j \)th position, then we refer to bit \( j \) of \( \{ \sigma_i \} \). If it is a 1, then we first do the child having all 1’s at position \( j \), and second do the child having all 0’s at position \( j \). Similarly, if bit \( j \) of \( \{ \sigma_i \} \) is a 0, then we first do the child having all 0’s at position \( j \).

The essential advantage of the tree structure is that, typically, we do not have to check many structure strings for each sequence because nodes high up in the tree get eliminated by the upper bound. An additional advantage accrues because we are only interested in sequences with unique ground states. Therefore, as soon as two strings are found that satisfy the goal \( G \), the search can be stopped for that sequence. Our protocol of following the branches that match the sequence \( \{ \sigma_i \} \) is intended to quickly identify strings which satisfy \( G \).

We now discuss how the tree is actually built. In order to take advantage of the natural clustering of structure strings, we choose to split each node at the position that makes each of the two child nodes as tightly clustered as possible. We measure this clustering for each child as the sum of the “entropies” for each bit, with the tightest clustering corresponding to the minimum entropy. Specifically, for each child we evaluate the total entropy \( S \) of its set of structure strings as

\[
S = -N_{\text{child}} \sum_i (p_i \ln p_i + q_i \ln q_i),
\]

where \( p_i \) is the probability of the \( i \)th position being a 1, and \( q_i \) is the probability of the \( i \)th position being 0, averaged
over all \( N_{\text{child}} \) structure strings in the child node. We choose to split each node at the position that minimizes the combined entropy of the two children.

The only remaining decision is when to stop splitting. When a node contains sufficiently few entries, it is fastest to just examine each entry in the node. Different stopping sizes were tried, and 16 seemed to be optimal to minimize search time per sequence. Another consideration was memory. The memory used by the entries themselves is 7,032,543 bytes = 33.5 megabytes, but each node takes additional space. With the given stopping criterion, there were 1,384,679 nodes in the tree, and the total space required was 70.5 megabytes.

Constructing a tree in this manner took a few hours on a 200 MHz Pentium Pro. Once the tree was constructed, carrying out each search took, on average, about 800 μs per sequence, but this time varied widely from sequence to sequence.

For comparison, we also implemented a naive search algorithm in which the energy of a sequence is computed for each structure string. The tree-search algorithm ran approximately 1600 times faster than our best variant of the naive approach.

IV. COMPUTING ALL THE GROUND STATES

The tree-search algorithm allows us to quickly compute ground states for each protein sequence in turn, and to record those sequences with unique ground states, together with the corresponding structure string and energy value. When all this is done, we would also like additional statistics, such as the designability of each structure, i.e., for how many sequences it is the unique ground state. This and other statistics can be computed, after the fact, if the unique ground-state solutions are stored.

Because our protein chains are directed, i.e., the two ends are not considered identical, both a structure and its oppositely directed partner are allowed. (Sometimes these are the same structure.) As a result, if a particular sequence \( \{ \sigma_i \} \) has a particular structure as a unique ground state, then the reversed sequence must have the reversed structure as its unique ground state. For the \( 4 \times 3 \times 3 \) problem, there are therefore \((2^{35}) + (2^{17}) = 34,359,869,440 \) possible sequences that need to be considered, counting all 36-bit binary strings but rejecting reversed strings. There are 7,032,543 distinct structure strings. Since we are interested in unique ground states, each structure string is tagged with an additional bit to indicate whether it represents exactly one, or more than one, geometrical structure.

The overall computation is trivially parallelizable because calculating the ground state for each sequence can be done independently. In order to manage the calculation of ground states for all sequences in parallel, it proved useful to divide the space of sequences into “bundles,” and use these as the unit of parallelism. Instead of organizing these bundles by fixing some high-order bits of the 36-bit sequence \( \{ \sigma_i \} \) and varying the rest, we fixed an equal number of low- and high-order bits, varying the bits in the middle. This way entire bundles could be eliminated as reversals of the sequences in another bundle. We performed our computations with 14 bits fixed and 22 bits varying, which produced a total of \((2^{13}) + (2^6) = 8,256 \) bundles.

These 8,256 bundles were executed on the NECI Large Array Multiple Processors (LAMP) system, which is a collection of 28 computers, all containing 200 MHz Intel Pentium Pro microprocessors. Three machines were uniprocessors, the rest had two processors each. Every machine ran the Linux operating system and had at least 128 MB of memory. Each computation typically required about 10 MB of memory for itself, plus about 70 MB for the (read-only) tree of protein structure information, which was shared in memory on the multiprocessors. The dual processors handled two independent bundles concurrently, and sharing the tree was necessary to keep the total memory “footprint” of the jobs small enough to fit together without conflict.

The distribution of bundles to “workers” was handled by a single “master” machine running shell and AWK scripts to poll the others, start new bundles, collect results, and detect any crashes that might occur. The scripts were written to be restartable with minimal lost effort in the event of a failure affecting their own operation. As each bundle was completed, a compactly coded binary output file was produced. At the end, all 8,256 output files were merged into a single 450 MB result file. Auxiliary programs were written to extract human-readable data from these binary files.

The complete computation ran for about 198 h, with an average of 39 processors running at any one time, giving a total of 7,805 CPU hours. Bundle execution times varied from 23 s to almost 19 h, with a mean of 56.7 min.

V. RESULTS

Using the tree-searching algorithm and the LAMP system, we were able to completely enumerate the \( 4 \times 3 \times 3 \) HP lattice protein model. We found that 114,572,949 sequences have unique ground states, which is about 0.3% of all sequences. In comparison, 0.09% of the sequences in the \( 3 \times 3 \times 3 \) model have unique ground states.

We associate to each structure a quantity called designability.\(^{13,20}\) The designability of a structure is the number of sequences having that structure as the unique ground state. By this definition, if two or more structures share the same bit-string representation then they have zero designability since those structures can never be the unique ground state of any sequence. Thus, only the 1331,025 structures (and their reverse paths) each of which has its own, unshared string representation can have nonzero designability. For these structures, the average designability is 114,572,949/1331,025 = 86. However, the designability of these structures has a very broad range: from 1 to 4,466. (The minimum designability is 1 because the sequence with the same bit-string as the structure is guaranteed to have that structure as a unique ground state.) In Fig. 3, we plot the number of structures with a given designability versus the designability. One sees a long tail in the high designability region, consistent with previous results on the \( 3 \times 3 \times 3 \) model\(^{12}\) and on various two-dimensional models.\(^{13,20}\)

Also consistent with these previous works, there are noticeable geometrical differences between the highly design-
able $4 \times 3 \times 3$ structures and the less designable ones. In Fig. 4, we have plotted the average structure string $\langle s_i \rangle$ for highly designable structures, and for all structures. Since $s_i = 1$ for a core site, and $s_i = 0$ for a surface site, the ensemble average $\langle s_i \rangle$ gives the probability that the $i$th monomer on the chain occupies a core site. It is seen in Fig. 4 that for highly designable structures the first few monomers on the chain tend to occupy core sites, while there is no such tendency for the average compact structure. The average of $s_i$ for each chain is exactly $1/3$ because every $4 \times 3 \times 3$ structure has exactly 12 core sites ($s_i = 1$) and 24 surface sites ($s_i = 0$). Therefore the tendency of the ends of highly designable structures to occupy core sites must be balanced by a tendency for the rest of the structure to occupy surface sites. This is also seen in Fig. 4—the central third of the chain for highly designable structures has an increased probability to occupy surface sites, on average.

In Fig. 5 we have plotted the two-point correlation function of structure strings, $C(i,j) = \langle s_i s_j \rangle - \langle s_i \rangle \langle s_j \rangle$, averaged over highly designable structures and over all structures. There is a clear correlation of site types, with a range of roughly one monomer in either direction along the chain. That is, if the $i$th monomer of a chain occupies a core site, there is an enhanced probability for monomers $i \pm 1$ to occupy core sites. This simply represents a general geometrical property of compact, self-avoiding structures. The correlation length is slightly shorter for highly designable structures [Fig. 5(b)], implying more frequent transitions between surface and core sites. In Fig. 6, we plot the number of transitions $t$ between surface and core sites versus designability $N_s$. For clarity, only structures with selected $N_s$’s are included in the plot. We see a weak positive correlation between $t$ and $N_s$, with a large variance for a given $N_s$. A much stronger positive correlation between surface-core transitions and designability was found in a two-dimensional $6 \times 6$ lattice model, possibly reflecting the larger more compact core in the two-dimensional model.

As an example, the topmost designable $4 \times 3 \times 3$ structure is plotted in Fig. 7. The geometry of this structure is consistent with the results shown in Figs. 4 and 5. The 12 core sites are equally divided between the two ends of the chain, with the center part of the chain, $i = 9 - 26$, consisting entirely of surface sites. Moreover, while the core sites tend
that a hydrophobic monomer occupies position $i$.

The complete enumeration of sequences and structures allows us to identify all sequences with a given structure as their unique ground state. We can therefore analyze the statistical properties of sequences that design a particular structure. For example, in Fig. 8, we have plotted the probability that a hydrophobic monomer occupies position $i$, averaged over all 4466 sequences that design the structure in Fig. 7. Figure 8 therefore represents the complete mutation pattern of the topmost designable structure. The 12 core sites are easily identified since the probability of a hydrophobic monomer at these sites is nearly one. That is, nearly all of the 4466 sequences that design this structure have 1’s at these 12 positions. Similarly, the first three surface sites at the beginning of the chain and the last four surface sites at the end of the chain are always occupied by polar monomers. Interestingly, the monomers in the central parts of the chains, $i = 10–26$, have a roughly 1/3 chance of being hydrophobic even though all of these are surface sites. The mutation pattern therefore has the nontrivial feature that the monomers at some sites are critical for the stability of the ground state while the monomers at other sites are freely mutable.

VI. DISCUSSION

The existence of highly designable structures emerged from a study of $3 \times 3 \times 3$ and $6 \times 6$ HP lattice protein models with interaction energies that included both solvation and segregation components. A later study verified the existence of highly designable structures for HP lattice models in two dimensions including only solvation energies. Moreover, in two dimensions, there was little change in the qualitative behavior of designability with increasing structure size, suggesting that highly designable structures persist up to realistic protein chain lengths. Within the $3 \times 3 \times 3$ solvation model, however, only 120 (0.1%) out of the 103 346 total structures have nonzero designability. These 120 (60 lattice conformations) stand out as highly designable structures with the largest average gaps in the solvation plus segregation model of Li et al. It has remained unclear whether the solvation model in three dimensions can produce a significant fraction of highly designable structures at realistic protein chain lengths. The current study, extending the solvation model up to $4 \times 3 \times 3$ structures, offers strong evidence that the existence of highly designable structures is a general feature of solvation models in three dimensions.

To understand the ubiquitous appearance of highly designable structures, it is helpful to review the geometrical interpretation of designability for the solvation model. To this end, the energy in Eq. (1) is rewritten as

$$E = \frac{1}{2} \sum_{i=1}^{N} \left[ |\sigma_i - s_i| - |\sigma_i| - |s_i| \right].$$

The last term is constant for a given sequence, and the second-to-last term is constant for all compact structures. Therefore, the ground state for a given sequence is determined by the first term alone, which is one-half the Hamming distance between the sequence string and the structure string. Simply put, the structure nearest to a sequence is its ground state. The designability of a structure is thus equal to the number of sequences that lie closer to it than to any other structure.

This geometrical interpretation suggests that highly designable structures are those with few nearby competing structures. To test this, we have plotted in Fig. 9 the number of neighboring structures, as a function of Hamming distance between structure strings, for the topmost designable structure and for structures of intermediate ($N_S = 100$) and low ($N_S = 1$) designability. It is seen that high designability implies a reduced number of neighbors, and this correlation persists out to distance 10 (structures whose strings differ by interchange of five surface and core sites). For comparison, we have also plotted in Fig. 9 the expected $n(d)$ if the 14 062 236 structure strings were uniformly distributed on the hyperplane given by the constraints $\Sigma s_i = 12$ and $\Sigma (-1)^{d} \times s_i = 0$. The second constraint comes from the fact that the $4 \times 3 \times 3$ cubic lattice is a bipartite lattice, so that for any structure there are six core sites with $i$ even and six core sites with $i$ odd. The total number of points in the hyperplane is $(C_{18}^6)^2 = 344 622 096$, where $C_{18}^6 = n! / m! (n-m)!$. The number of points in the plane at a Hamming distance $d = 2 \times 1^l$ from a given point is simply: $N(d) = \sum_{k=0}^{l} C_6^k C_{12}^{l-k} C_{12}^k$, with $C_{12}^0 = 0$ if $m > n$. The expected $n(d)$ is then $\rho \times N(d)$, where $\rho = 14 062 236 / 344 622 096 \approx 0.04$ is the average density of real structure strings in the
The existence of highly designable structures can therefore be viewed as a geometrical property of the space of structure strings. For the two-dimensional solvation model, it has been stressed\(^\text{20}\) that the most highly designable structures fall in regions of low density in the space of structure strings. In this sense, highly designable structures have "atypical" patterns of surface and core sites. The current work extends this conclusion to a three-dimensional solvation model with chain lengths \((N = 36)\) approaching those of real proteins.

Many aspects of real proteins are not addressed by our simple lattice model. Interactions such as hydrogen bonds, salt bridges, disulfide bonds, and detailed sidechain packing are beyond the model’s minimal representation of hydrophobicity and excluded volume. Moreover, real proteins contain secondary structures (\(\alpha\) helices and \(\beta\) strands), and come in a variety of sizes and shapes. What aspects of our model study pertain to the more complex world of real proteins? First, the overall notion of designability—that some structures have intrinsically better folding properties than others—has potentially broad application. Studies of RNA folding\(^\text{22}\) and off-lattice protein models\(^\text{33,34}\) both yield highly designable structures. Second, insofar as hydrophobic patterning is a dominant factor in the folding of real proteins, the model result that high designability correlates with atypical hydrophobicity patterns may have real significance.

In summary, we have employed a fast tree-search algorithm to find the ground states of all sequences for a \(4 \times 3 \times 3\) lattice HP model of proteins. The results confirm the existence of highly designable structures in a three-dimensional solvation model with a surface-to-core ratio of \(2\)-to-\(1\), close to the values for small natural proteins. Highly designable structures are found to differ geometrically from other structures. Interestingly, structures are found to have nontrivial mutation patterns with some sites strictly conserved and others mutable. The fast tree-search algorithm is particularly well suited to lattice HP solvation models, and we hope our detailed description of the method will prove useful.

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31 M. Kardar and M. Yahyanejad (private communication).