Predicting the Secondary Structure of Proteins by the Use of Hamming Distances and Alignment Scores

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Abstract
Researchers are confident about the validity of the basic hypothesis that the secondary and tertiary structures of a protein are uniquely determined by its sequence of amino acids, that is its primary structure. In this article we use a database of 200 proteins. To find the secondary structure of a new protein, the first thirteen residues of this protein are taken as a substring. Then the conformations of the central amino acids of thirteen residue substrings of the proteins in the database, whose hamming distances are less than a given threshold or alignment scores exceed a given limit are collected in a basin. The commonest conformation in this basin is attached as the conformation of the central amino acid of the substring of the unknown protein. Using this technique, for MHsim threshold 3.0, a correct estimation rate of 53.4% is obtained with 4.74% indecisives and for MHsim threshold 5.0, the success was 56.93% with 76.59% indecisives. When the half of the proteins, whose secondary structure estimations are higher, subjected to same calculation the following results are obtained; for MHsim threshold 3.0, correct estimation rate is 79.52% with 58.87% indecisives and for MHsim threshold 5.0, correct estimation rate is 65.52% with 5.02% indecisives. Average correct estimation rate for the alignment scores was %54.

1. INTRODUCTION
For more than four decades, the protein folding problem has been among the most challenging problems in the biological sciences. In 1994, a protein structure prediction contest was organized with the aim of assessing the real virtues and defects of several well known methodologies. Analysis of the structures predicted by the contestants (Moult et al., 1995) has generally shown that even the most promising techniques need considerable improvement, and that the protein folding problem should still be considered unresolved. Briefly, preliminary calculations, although promising, are feasible only for small-size proteins; there have been no major breakthroughs in the molecular modeling techniques and threading techniques need further development. During this contest, protein secondary structure prediction was reevaluated and recognized as a useful tool for establishing starting points for tertiary structure calculation determination of protein structures. Early approaches to
protein secondary structure prediction from the primary sequence had prediction accuracy, which is the percentage of correctly predicted residues in the three states: α-helix, β-strand, and coil, of about 57% (Chou & Fasman, 1978; Garnier et al., 1978). Various later attempts to improve the accuracy (Gibrat et al., 1987; Biou et al., 1988; Levin & Garnier, 1988; Holley & Karplus, 1989; Qian & Sejnowski, 1989; King & Sternberg, 1990; Salzberg & Cost, 1992; Stolorz et al., 1992; Zhang et al., 1992; Munson et al., 1994) with innovative artificial intelligence techniques, such as neural networks, machine learning, nearest neighbors, and combined approaches, have not achieved prediction accuracies greater than 66%. The inclusion of evolutionarily related sequences into the prediction scheme has given a significant boost in prediction accuracy, up to values of about 68-72% (Zvelebil et al., 1987; Levin et al., 1993; Rost & Sander, 1993, 1994; Rost et al., 1994a; Di Francesco et al., 1995; Salamov & Solovyev, 1995). In general, the suggested explanation for these improvements in prediction accuracy is that sequence alignments of homologous proteins should emulate as closely as possible the structural alignment. Thus, aligned residues, in particular those in the core proteins, should belong to the same secondary structure elements. Sequence alignments may be utilized to obtain a consensus from the predictions based on each homologous sequence, or they may be used to build sequence profiles at each aligned position. In addition to the identity of the aligned residues, which is a feature exploited by all the predictive schemes, other information is available from sequence alignments, such as the location of gaps or the patterns of residue mutation in the aligned protein families. Some authors have used such information to refine their prediction models (Zvelebil et al., 1987; Rost & Sander, 1993; Rost et al., 1994a; Salamov & Solovyev, 1995). However, the reasons why the inclusion of this additional information improves the quality of the prediction have not been understood.

In his extensive review Rost (Rost, 2001) asks the following question: 88% is a limit, but shall we ever reach close to there?

A database of 200 random proteins with known secondary structure formations is prepared. To find the secondary structure of a protein, test substrings of consecutive residues of length 13 of this protein are formed. Then in proteins in the database, substrings of length 13 with high enough similarities to the test string are collected in a pool. The most common secondary structure formation corresponding to the central amino acid of substrings in the pool is attached to the central amino acid of the test substring as secondary structure formation.

2. FORMULATION OF THE PROBLEM

To estimate the conformation of the protein at a given residue, we consider 6 right and 6 left neighbors of this residue. Our hypothesis is that the conformation at the central residue is determined by these neighbors and by itself.

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**Table 1** Names and symbols of 20 amino acids

| # | Amino acid | Chemical | alphabet |
|---|------------|----------|----------|
| 1 | Alanine    | Ala      | A        |
| 2 | Arginine   | Arg      | R        |
| 3 | Asparagine | Asn      | N        |
| 4 | Aspartic acid | Asp   | D        |
| 5 | Cysteine   | Cys      | C        |
| 6 | Glutamine  | Gln      | Q        |
| 7 | Glutamic acid | Glu  | E        |
| 8 | Glycine    | Gly      | G        |
| 9 | Histidine  | His      | H        |
|10 | Isoleucine | Ile      | I        |
|11 | Leucine    | Leu      | L        |
|12 | Lysine     | Lys      | K        |
|13 | Methionine | Met      | M        |
|14 | Phenylalanine | Phe | F        |
|15 | Proline    | Pro      | P        |
|16 | Serine     | Ser      | S        |
|17 | Threonine  | Thr      | T        |
|18 | Tryptophan | Trp      | W        |
|19 | Tyrosine   | Tyr      | Y        |
|20 | Valine     | Val      | V        |
Based on the protein chain it is easy to create its relevant sequence of amino acids replacing an amino acid in chain by its code in Latin alphabet. As a result a word on the amino acids’ alphabet is received. This word can be called a protein primary structure on the condition that letters in this word are in the same order as amino acids in the protein chain are.

A secondary structure of a protein is a subsequence of amino acids coming from the relevant protein. These subchains form in the three dimensional space regular structures which are the same in shape for different proteins. In the analysis, a similar representation for the secondary structures as for the primary ones has been used. A secondary structure is represented by a word on the relevant alphabet of secondary structures – each kind of a secondary structure has its own unique letter α-helix, H; β-sheet S, and coil C. An alphabet of secondary structures consisting of three different secondary structures has been considered in the analysis.

(c) Coding the Data
In this paper, data corresponding to an amino acid consists of 6 right, and 6 left neighboring amino acids of this amino acid in the primary chain of the protein as in Table 2. In the second row, secondary structure conformations of these neighboring amino acids are given.

| A | E | E | K | E | A | V | L | G | I | W | G | K |
|---|---|---|---|---|---|---|---|---|---|---|---|---|
| H | H | H | H | E | E | E | E | C | C | C | C | E |

Table 2 Six right, and six left neighboring amino acids of the central amino acid V.

Secondary structure letters H, E, and C are coded as in the table below;

| H | E | C |
|---|---|---|
| 1 | 0 | 0 |
| 0 | 1 | 0 |
| 0 | 0 | 1 |

Table 3 Codes for secondary structure letters H, E, and C.

(d) Similarity Measures
To find the secondary structure of a protein, test substrings of consecutive residues of length 13 of this protein are cut. Then in proteins in the database, substrings of consecutive residues of length 13 are cut as well. To infer the conformation of the central amino acid of test substring, we search for similar substrings of the same length of 13 from the proteins in the database. For this purpose, two similarity measures are used.

(1) Modified Hamming Similarity
Hamming distance of two substrings of the same length is the number of the mismatches as seen in Table 4.

\[
H_{sim} = 13 - H_{dis}. \quad (1)
\]

There is a consensus about the affect of amino acid composition of the primary sequence on the secondary structure of a protein. But clearly this affect is local. That is amino acids far away of the central amino acid have less affect on the conformation at the central amino acid, compared to the nearer ones. It means that the match “VV” at the 8th position is more important than the match “AA” at 13th position.

To weight matches we propose a Gaussian curve

\[
F(x) = e^{-\frac{(x-\gamma)^2}{s^2}}, \quad (2)
\]

where \(s\) is a measure for the spread of the curve.

For the substrings in Table 4, hamming similarity is \(H_{sim} = 4\), modified similarity is \(MH_{sim} = 2.75\).

(2) Alignment Score
The alignment of two strings

GRLPACVVDCGTG
MLSPADKTNVKAA \( (3) \)

is obtained as

\[
\begin{array}{cccccccccccc}
G & R & L & P & A & C & V & V & D & C & G & T & A \\
M & L & S & P & A & D & K & V & N & K & A & A & \\
0 & 0 & 0 & 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 1 \\
\end{array}
\]

Table 4 Hamming distance of two substrings of the same length is the number of the mismatches

Now a similarity measure of the given two strings can be defined as

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0 & 0 & 0 & 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 1 \\
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0 & 0 & 0 & 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 1 \\
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proteins can be performed? We believe that answering this question is the structure of the similar regions in highly similar proteins. In what bases correct structural classification of proteins is going to cover some fundamental questions such as; what is the similarity of structure, physiology based upon a genetic factor. The protein homology most recognized by similarity; the more closely related are the scaffold structures. Based on this approach, proteins primary sequence similarity was investigated with searching for similar substrings of the same length of 13 from the database is found and collected into the basket with some certain proteins in database has been detected. This protein’s structure, physiological characters and their physicochemical properties are going to be analyzed in order to reveal information about the influence of this parameter. The essence of particular parameter aimed to be found which make this protein structure unique. We believe that it will provide us a new attribute in order to increase the prediction capacity of our algorithms.

For each test substring of length 13, around 16000 comparisons are made with 13tuples of amino acid residues of target proteins. In an average desktop computer this operation is performed in around five seconds. Therefore it is not feasible to increase the number proteins in the database. For this reason, for high thresholds for the similarity, some of the test 13tuples may not have similar enough 13tuples in target proteins. In such a case, the conformation of the central amino acid of the test 13tuple remains undefined. On the other hand, high similarity brings high accuracy in the secondary structure estimation. In Table 4, for certain values of the modified hamming similarity threshold, the percentage of the indecisive residues, and accuracy in the three conformations α-helices, β-sheets, and c-coils are given.

Table 5 Similarity thresholds vs. accuracy in MHsim

| MHsim | Indecisives % | Accuracy % |
|-------|---------------|------------|
| ≥ 5.0 | 76.59         | 56.93      |
| ≥ 4.5 | 54.15         | 54.15      |
| ≥ 3.0 | 4.74          | 53.41      |

Table 6 Similarity thresholds vs. accuracy in MHsim for better half.

| MHsim | Indecisives % | Accuracy % |
|-------|---------------|------------|
| ≥ 5.0 | 58.87         | 79.52      |
| ≥ 4.5 | 40.74         | 67.06      |
| ≥ 3.0 | 5.02          | 65.56      |

For the half of the proteins in database whose correct secondary structure estimations are better, the correct estimation rates are as in Table 6.

For the similarity measure computed by the use of the alignment score, the built in function Needleman Wunsch Similarity in MATHEMATICA is used. As scoring matrix, PAM70 is chosen. Test and target substrings are of length 17 to give more stable and reliable alignment scores.

In Table 7, for the values 1, and −5 of the alignment score thresholds, the percentages of the indecisive residues, and

### Database of Proteins

200 proteins of known structures, with a total 169 026 amino acid residues collected from PDB almost randomly. To test the accuracy of the method each time one of the proteins is chosen as the testing protein, and other 199 proteins are taken as target proteins.

### 4. RESULTS AND DISCUSSION

In a biological context, the term homology, defines similarity of structure, physiology based upon a genetic factor. The protein homology most recognized by similarities in their amino acid sequence. There is a widely accepted hypothesis that: “the greater the sequence similarity; the more closely related are the scaffold structure”. Based on this approach, proteins primary sequence similarity was investigated with searching for similar substrings of the same length of 13 from the proteins in the database. Each of the similar 13 tuples in the database is found and collected into the basket with modified hamming similarity threshold 5 and 3 separately. The proteins that have high similarity and high accuracy with some certain proteins in database has been detected and separated for further analysis. Further analyses is going to cover some fundamental questions such as; what is the structure of the similar regions in highly similar proteins? In what bases correct structural classification of proteins can be performed? We believe that answering these questions will enable us to classify proteins, existed protein classification approaches are going to be analyzed and methodology is going to be strengthening. The following question: “what is the advantage of the structural classification of proteins over randomly chosen proteins” will be addressed. In the other hand proteins that have no similar or low similar sequences in the database also detected. This protein’s structure, physiological characters and their physicochemical properties are going to be analyzed in order to reveal information about the influence of this parameter. The essence of particular parameter aimed to be found which make this protein structure unique. We believe that it will provide us a new attribute in order to increase the prediction capacity of our algorithms.
average accuracies in the three conformations $\alpha$-helices, $\beta$-sheets, and $\epsilon$-coils are given.

| A. Score | Indecisives % | Accuracy % |
|----------|---------------|------------|
| $\geq 1$ | 69.69         | 63.65      |
| $\geq -5$ | 6.31          | 54.01      |

Table 7 Alignment score thresholds vs. accuracy in ASsim

For the half of the proteins in database whose correct secondary structure estimations are better, the correct estimation rates are as in Table 8.

| A. Score | Indecisives % | Accuracy % |
|----------|---------------|------------|
| $\geq 1$ | 51.78         | 93.24      |
| $\geq -5$ | 7.31          | 62.16      |

Table 8 Alignment score thresholds vs. accuracy in ASsim for better half.

These results show that the analysis which relies on a database of 200 proteins has an estimation power that is comparable with the famous online estimation tools. Table 6, and Table 8 display the correct estimation rates of the half of the proteins in database whose correct secondary structure estimations are better.

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