Research Article

A Study of Residue Correlation within Protein Sequences and Its Application to Sequence Classification

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We investigate methods of estimating residue correlation within protein sequences. We begin by using mutual information (MI) of adjacent residues, and improve our methodology by defining the mutual information vector (MIV) to estimate long range correlations between nonadjacent residues. We also consider correlation based on residue hydropathy rather than protein-specific interactions. Finally, in experiments of family classification tests, the modeling power of MIV was shown to be significantly better than the classic MI method, reaching the level where proteins can be classified without alignment information.

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1. INTRODUCTION

A protein can be viewed as a string composed from the 20-symbol amino acid alphabet or, alternatively, as the sum of their structural properties, for example, residue-specific interactions or hydropathy (hydrophilic/hydrophobic) interactions. Protein sequences contain sufficient information to construct secondary and tertiary protein structures. Most methods for predicting protein structure rely on primary sequence information by matching sequences representing unknown structures to those with known structures. Thus, researchers have investigated the correlation of amino acids within and across protein sequences [1–3]. Despite all this, in terms of character strings, proteins can be regarded as slightly edited random strings [1].

Previous research has shown that residue correlation can provide biological insight, but that MI calculations for protein sequences require careful adjustment for sampling errors. An information-theoretic analysis of amino acid contact potential pairings with a treatment of sampling biases has shown that the amount of amino acid pairing information is small, but statistically significant [2]. Another recent study by Martin et al. [3] showed that normalized mutual information can be used to search for coevolving residues.

From the literature surveyed, it was not clear what significance the correlation of amino acid pairings holds for protein structure. To investigate this question, we used the family and sequence alignment information from Pfam-A [4]. To model sequences, we defined and used the mutual information vector (MIV) where each entry represents the MI estimation for amino acid pairs separated by a particular distance in the primary structure. We studied two different properties of sequences: amino acid identity and hydropathy.

In this paper, we report three important findings.

(1) MI scores for the majority of 1000 real protein sequences sampled from Pfam are statistically significant (as defined by a P value cutoff of .05) as compared to random sequences of the same character composition, see Section 4.1.

(2) MIV has significantly better modeling power of proteins than MI, as demonstrated in the protein sequence classification experiment, see Section 5.2.

(3) The best classification results are provided by MIVs containing scores generated from both the amino acid alphabet and the hydropathy alphabet, see Section 5.2.

In Section 2, we briefly summarize the concept of MI and a method for normalizing MI content. In Section 3, we formally define the MIV and its use in characterizing protein sequences. In Section 4, we test whether MI scores for protein sequences sampled from the Pfam database are statistically significant compared to random sequences of the
same residue composition. We test the ability of MIV to classify sequences from the Pfam database in Section 5, and in Section 6, we examine correlation with MIVs and further investigate the effects of alphabet size in terms of information theory. We conclude with a discussion of the results and their implications.

2. MUTUAL INFORMATION (MI) CONTENT

We use MI content to estimate correlation in protein sequences to gain insight into the prediction of secondary and tertiary structures. Measuring correlation between residues is problematic because sequence elements are symbolic variables that lack a natural ordering or underlying metric [5]. Residues can be ordered in certain properties such as hydrophathy, charge, and molecular weight. Weiss and Herzel [6] analyzed several such correlation functions.

MI is a measure of correlation from information theory [7] based on entropy, which is a function of the probability distribution of residues. We can estimate entropy by counting residue frequencies. Entropy is maximal when all residues appear with the same frequency. MI is calculated by systematically extracting pairs of residues from a sequence and calculating the distribution of pair frequencies weighted by the frequencies of the residues composing the pairs.

By defining a pair as adjacent residues in the protein sequence, MI estimates the correlation between the identities of adjacent residues. We later define pairs using nonadjacent residues, and physical properties rather than residue identities.

MI has been proven useful in multiple studies of biological sequences. It has been used to predict coding regions in DNA [8], and has been used to detect coevolving residue pairs in protein multiple sequence alignments [3].

2.1. Mutual information

The entropy of a random variable $X$, $H(X)$, represents the uncertainty of the value of $X$. $H(X)$ is 0 when the identity of $X$ is known, and $H(X)$ is maximal when all possible values of $X$ are equally likely. The mutual information of two variables $MI(X, Y)$ represents the reduction in uncertainty of $X$ given $Y$, and conversely, $MI(Y, X)$ represents the reduction in uncertainty of $Y$ given $X$:

$$MI(X, Y) = H(X) - H(X \mid Y) = H(Y) - H(Y \mid X).$$

(1)

When $X$ and $Y$ are independent, $H(X \mid Y)$ simplifies to $H(X)$, so $MI(X, Y)$ is 0. The upper bound of $MI(X, Y)$ is the lesser of $H(X)$ and $H(Y)$, representing complete correlation between $X$ and $Y$:

$$H(X \mid Y) = H(Y \mid X) = 0.$$  

(2)

We can measure the entropy of a protein sequence $S$ as

$$H(S) = - \sum_{i \in \Sigma_A} P(x_i) \log_2 P(x_i),$$

(3)

where $\Sigma_A$ is the alphabet of amino acid residues and $P(x_i)$ is the marginal probability of residue $i$. In Section 3.3, we discuss several methods for estimating this probability.

From the entropy equations above, we derive the MI equation for a protein sequence $X = (x^1, \ldots, x^N)$:

$$MI = \sum_{i \in \Sigma_A} \sum_{j \in \Sigma_A} P(x_i, x_j) \log_2 \left( \frac{P(x_i, x_j)}{P(x_i)P(x_j)} \right),$$

(4)

where the pair probability $P(x_i, x_j)$ is the frequency of two residues being adjacent in the sequence.

2.2. Normalization by joint entropy

Since $MI(X, Y)$ represents a reduction in $H(X)$ or $H(Y)$, the value of $MI(X, Y)$ can be altered significantly by the entropy in $X$ and $Y$. The MI score we calculate for a sequence is also affected by the entropy in that sequence. Martin et al. [3] propose a method of normalizing the MI score of a sequence using the joint entropy of a sequence. The joint entropy, or $H(X, Y)$, can be defined as

$$H(X, Y) = - \sum_{i \in \Sigma_A} \sum_{j \in \Sigma_A} P(x_i, x_j) \log_2 P(x_i, x_j)$$

(5)

and is related to $MI(X, Y)$ by the equation

$$MI(X, Y) = H(X) + H(Y) - H(X, Y).$$

(6)

The complete equation for our normalized MI measurement is

$$\frac{MI(X, Y)}{H(X, Y)} = \frac{\sum_{i \in \Sigma_A} \sum_{j \in \Sigma_A} P(x_i, x_j) \log_2 (P(x_i, x_j)/P(x_i)P(x_j))}{\sum_{i \in \Sigma_A} \sum_{j \in \Sigma_A} P(x_i, x_j) \log_2 P(x_i, x_j)}.$$  

(7)

3. MUTUAL INFORMATION VECTOR (MIV)

We calculate the MI of a sequence to characterize the structure of the resulting protein. The structure is affected by different types of interactions, and we can modify our methods to consider different biological properties of a protein sequence. To improve our characterization, we combine these different methods to create a vector of MI scores.

Using the flexibility of MI and existing knowledge of protein structures, we investigate several methods for generating MI scores from a protein sequence. We can calculate the pair probability $P(x_i, x_j)$ using any relationship that is defined for all amino acid identities $i, j \in \Sigma_A$. In particular, we examine distance between residue pairings, different types of residue-residue interactions, classical and normalized MI scores, and three methods of interpreting gap symbols in Pfam alignments.

3.1. Distance MI vectors

Protein exists as a folded structure, allowing nonadjacent residues to interact. Furthermore, these interactions help to determine that structure. For this reason, we use MIV to characterize nonadjacent interactions. Our calculation of MI for adjacent pairs of residues is a specific case of a more general relationship, separation by exactly $d$ residues in the sequence.
amino acid residues, \[ \Sigma \] in Section 6.2.

Unfortunately, the quality of the estimation suffers from the short length of protein sequences. Our second method is to use a common prior probability distribution for all sequences. Since all of our sequences are part of the Pfam database, we use residue frequencies calculated from Pfam as our prior. In our results, we refer to this method as the Pfam prior. The large sample size allows the frequency to more accurately estimate the probability. However, since Pfam contains sequences from many organisms, the probability distribution is less accurate.

### 3.4. Interpreting gap symbols

The Pfam sequence alignments contain gap information, which presents a challenge for our MIV calculations. The gap character does not represent a physical element of the sequence, but it does provide information on how to view the sequence and compare it to others. Because of this contradiction, we compared three strategies for processing gap characters in the alignments.

#### The strict method

This method removes all gap symbols from a sequence before performing any calculations, operating on the protein sequence rather than an alignment.

#### The literal method

Gaps are a proven tool in creating alignments between related sequences and searching for relationships between sequences. This method expands the sequence alphabet to include the gap symbol. For \( \Sigma_A \) we define and use a new alphabet:

\[
\Sigma_A^* = \Sigma_A \cup \{-\}. \tag{10}
\]

MI is then calculated for \( \Sigma_A^* \). \( \Sigma_H \) is transformed to \( \Sigma_G^* \) using the same method.

#### The hybrid method

This method is a compromise of the previous two methods. Gap symbols are excluded from the sequence alphabet when calculating MI. Occurrences of the gap symbol are still considered when calculating the total number of symbols. For a sequence containing one or more gap symbols,

\[
\sum_{i \in \Sigma_A} P_i < 1. \tag{11}
\]

Pairs containing any gap symbols are also excluded, so for a gapped sequence,

\[
\sum_{i,j \in \Sigma_A} P_{ij} < 1. \tag{12}
\]

These adjustments result in a negative MI score for some sequences, unlike classical MI where a minimum score of 0 represents independent variables.
Table 3: MIVs’ examples calculated for four sequences from Pfam. All methods used literal gap interpretation.

|     | Globin MI(d) | Ferrochelatase MI(d) | DUF629 MI(d) | Big2 MI(d) |
|-----|--------------|----------------------|--------------|------------|
| d   | Σ_A          | Σ_H                 | Σ_A          | Σ_H        | Σ_A          | Σ_H        | Σ_A          | Σ_H        |
| 0   | 1.34081      | 0.42600              | 0.95240      | 0.13820    | 0.70611      | 0.04752    | 1.26794      | 0.21026    |
| 1   | 1.20553      | 0.23740              | 0.93240      | 0.03837    | 0.63171      | 0.00856    | 0.92824      | 0.05522    |
| 2   | 1.07361      | 0.12164              | 0.90004      | 0.02497    | 0.63330      | 0.00367    | 0.95326      | 0.07424    |
| 3   | 0.92912      | 0.02704              | 0.87380      | 0.03133    | 0.66955      | 0.00575    | 0.96360      | 0.04962    |
| 4   | 0.97230      | 0.00380              | 0.90400      | 0.02153    | 0.62328      | 0.00587    | 1.00100      | 0.08373    |
| 5   | 0.91082      | 0.00392              | 0.78479      | 0.02944    | 0.68383      | 0.00674    | 0.98737      | 0.03664    |
| 6   | 0.90658      | 0.01581              | 0.81559      | 0.00588    | 0.63120      | 0.00782    | 0.95326      | 0.07424    |
| 7   | 0.87965      | 0.02435              | 0.91757      | 0.00822    | 0.67433      | 0.00172    | 1.04627      | 0.12002    |
| 8   | 0.83376      | 0.01860              | 0.87615      | 0.01247    | 0.6319      | 0.00495    | 0.90784      | 0.05221    |
| 9   | 0.88404      | 0.01000              | 0.90823      | 0.00721    | 0.61597      | 0.00411    | 0.97119      | 0.04002    |
| 10  | 0.88685      | 0.01353              | 0.89673      | 0.00611    | 0.60790      | 0.00718    | 1.02660      | 0.02240    |
| 11  | 0.90792      | 0.01719              | 0.94314      | 0.02195    | 0.66750      | 0.00867    | 0.92858      | 0.02261    |
| 12  | 0.95955      | 0.00231              | 0.87247      | 0.01027    | 0.64879      | 0.00805    | 0.98879      | 0.03156    |
| 13  | 0.88584      | 0.01387              | 0.85914      | 0.00733    | 0.66959      | 0.00607    | 1.09997      | 0.04766    |
| 14  | 0.93670      | 0.01490              | 0.88250      | 0.00335    | 0.66033      | 0.00106    | 1.06989      | 0.01286    |
| 15  | 0.86407      | 0.02052              | 0.94592      | 0.00548    | 0.62171      | 0.01363    | 1.27002      | 0.06204    |
| 16  | 0.89004      | 0.04024              | 0.92664      | 0.01398    | 0.63445      | 0.00314    | 1.05699      | 0.03154    |
| 17  | 0.91409      | 0.01706              | 0.80241      | 0.00108    | 0.67801      | 0.00536    | 1.06677      | 0.02136    |
| 18  | 0.89522      | 0.01691              | 0.85366      | 0.00719    | 0.65903      | 0.00898    | 1.05439      | 0.03310    |
| 19  | 0.92742      | 0.03319              | 0.90928      | 0.01334    | 0.70176      | 0.00151    | 1.17621      | 0.01902    |

3.5. MIV examples

Table 3 shows eight examples of MIVs calculated from the Pfam database. A sequence was taken from four random families, and the MIV was calculated using the literal gap method for both Σ_H and Σ_A. All scores are in bits. The scores generated from Σ_A are significantly larger than those from Σ_H. We investigate this observation further in Sections 4.1 and 6.2.

3.6. MIV concatenation

The previous sections have introduced several methods for scoring sequences that can be used to generate MIVs. Just as we combined MI scores to create MIV, we can further concatenate MIVs. Any number of vectors calculated by any methods can be concatenated in any order. However, for two vectors to be comparable, they must be the same length, and must agree on the feature stored at every index.

Definition 3. Any two MIVs, MIV_j(A) and MIV_k(B), can be concatenated to form MIV_j+k(C).

4. ANALYSIS OF CORRELATION IN PROTEIN SEQUENCES

In theory, a random string contains no correlation between characters. So, we expect a “slightly edited random string” to exhibit little correlation. In practice, noninfinite random strings usually have a nonzero MI score. This overestimation of MI in finite sequences is a factor of the length of the string, alphabet size, and frequency of the characters that make up the string. We investigated the significance of this error for our calculations and methods for reducing or correcting for the error.

To confirm the significance of our MI scores, we used a permutation-based technique. We compared known coding sequences to random sequences in order to generate a P value signifying the chance that our observed MI score or higher would be obtained from a random sequence of residues. Since MI scores are dependent on sequence length and residue frequency, we used the shuffle command from the HMMER package to conserve these parameters in our random sequences.

We sampled 1000 sequences from our subset of Pfam-A. A simple random sample was performed without replacement from all sequences between 100 and 1000 residues in length. We calculated MI(0) for each sequence sampled. We then generated 10 000 shuffled versions of each sequence and calculated MI(0) for each.

We used three scoring methods to calculate MI(0):

1. Σ_A with literal gap interpretation,
2. Σ_A normalized by joint entropy with literal gap interpretation,
3. Σ_H with literal gap interpretation.
In all three cases, the MI(0) score for a shuffled sequence of infinite length would be 0; therefore, the calculated scores represent the error introduced by sample-size effects. Figure 1, mean MI(0) of shuffled sequences, shows the average shuffled sequence scores (i.e., sampling error) in bits for each method. This figure shows that, as expected, the sampling error tends to decrease as the sequence length increases.

4.1. Significance of MI(0) for protein sequences

To compare the amount of error, in each method we normalized the mean MI(0) scores from Figure 1 by dividing the mean MI(0) score by the MI(0) score of the sequence used to generate the shuffles. This ratio estimates the amount of the sequence MI(0) score attributed to sample-size effects.

Figure 2, normalized MI(0) of shuffled sequences, compares the effectiveness of our two corrective methods in minimizing the sample-size effects. This figure shows that normalization by joint entropy is not as effective as Figure 1 suggests. Despite a large reduction in bits, in most cases, the portion of the score attributed to sampling effects shows only a minor improvement. $\Sigma_H$ still shows a significant reduction in sample-size effects for most sequences.

Figures 1 and 2 provide insight into trends for the three methods, but do not answer our question of whether or not the MI scores are significant. For a given sequence $S$, we estimated the $P$ value as

$$P = \frac{x}{N},$$

where $N$ is the number of random shuffles and $x$ is the number of shuffles whose MI(0) was greater than or equal to MI(0) for $S$. For this experiment, we choose a significance cutoff of .05. For a sequence to be labeled significant, no more than 50 of the 10 000 shuffled versions may have an MI(0) score equal or larger than the original sequence. We repeated this experiment for MI(1), MI(5), MI(10), and MI(15) and summarized the results in Table 4.

These results suggest that despite the low MI content of protein sequences, we are able to detect significant MI in a majority of our sampled sequences at MI(0). The number of significant sequences decreases for MI(d) as $d$ increases. The results for the classic MI method are significantly affected by sampling error. Normalization by joint entropy reduces this error slightly for most sequences, and using $\Sigma_H$ is a much more effective correction.

5. MEASURING MIV PERFORMANCE THROUGH PROTEIN CLASSIFICATION

We used sequence classification to evaluate the ability of MI to characterize protein sequences and to test our hypothesis that MIV characterizes a protein sequence better MI. As such, our objective is to measure the difference in accuracy between the methods, rather than to reach a specific classification accuracy.

We used the Pfam-A dataset to carry out this comparison. The families contained in the Pfam database vary in sequence count and sequence length. We removed all families containing any sequence of less than 100 residues due to complications with calculating MI for small strings. We also limited our study to families with more than 10 sequences and less than or equal to 200 sequences. After filtering Pfam-A based on our requirements, we were left with 2392 families to consider in the experiment.

Sequence similarity is the most widely used method of family classification. BLAST [10] is a popular tool incorporating this method. Our method differs significantly, in that classification is based on a vector of numerical features, rather than the protein’s residue sequence.
Table 4: Sequence significance calculated for significance cutoff .05.

| Scoring method | Number of significant sequences (of 1000) |
|----------------|------------------------------------------|
|                | MI(0) | MI(1) | MI(5) | MI(10) | MI(15) |
| Literal-ΣA     | 762   | 630   | 277   | 103    | 54     |
| Normalized literal-ΣA | 777   | 657   | 309   | 106    | 60     |
| Literal-ΣH     | 894   | 783   | 368   | 162    | 117    |

Classification of feature vectors is a well-studied problem with many available strategies. A good introduction to many methods is available in [11], and the method chosen can significantly affect performance. Since the focus of this experiment is to compare methods of calculating MIV, we only used the well-established and versatile nearest neighbor classifier in conjunction with Euclidean distance [12].

5.1. Classification implementation

For classification, we used the WEKA package [11]. WEKA uses the instance based 1 (IB1) algorithm [13] to implement nearest neighbor classification. This is an instance-based learning algorithm derived from the nearest neighbor pattern classifier and is more efficient than the naive implementation.

The results of this method can differ from the classic nearest neighbor classifier in that the range of each attribute is normalized. This normalization ensures that each attribute contributes equally to the calculation of the Euclidean distance. As shown in Table 3, MI scores calculated from ΣA have a larger magnitude than those calculated from ΣH. This normalization allows the two alphabets to be used together.

5.2. Sequence classification with MIV

In this experiment, we explore the effectiveness of classifications made using the correlation measurements outlined in Section 3.

Each experiment was performed on a random sample of 50 families from our subset of the Pfam database. We then used leave-one-out cross-validation [14] to test each of our classification methods on the chosen families.

In leave-one-out validation, the sequences from all 50 families are placed in a training pool. In turn, each sequence is extracted from this pool and the remaining sequences are used to build a classification model. The extracted sequence is then classified using this model. If the sequence is placed in the correct family, the classification is counted as a success. Accuracy for each method is measured as

\[
\text{Accuracy} = \frac{\text{no. of correct classifications}}{\text{no. of classification attempts}}
\]

We repeated this process 100 times, using a new sampling of 50 families from Pfam each time. Results are reported for each method as the mean accuracy of these repetitions. For each of the 24 combinations of scoring options outlined in Section 3, we evaluated classification based on MI(0), as well as MIV20. The results for these experiments are summarized in Table 5, classification Results for MI(0) and MIV20.

All MIV20 methods were more accurate than their MI(0) counterparts. The best method was ΣH with hybrid gap scoring with a mean accuracy of 85.14%. The eight best performing methods used ΣH, with the best method based on ΣA having a mean accuracy of only 66.69%. Another important observation is that strict gap interpretation performs poorly in sequence classification. The best strict method had a mean accuracy of 29.96%—much lower than the other gap methods.

Our final classification attempts were made using concatenations of previously generated MIV20 scores. We evaluated all combinations of methods. The five combinations most accurate at classification are shown in Table 6. The best method combinations are over 90% accurate, with the best being 90.99%. The classification power of ΣH with hybrid gap interpretation is demonstrated, as this method appears in all five results. Surprisingly, two strict scoring methods appear in the top 5, despite their poor performance when used alone.

Based on our results, we made the following observations.

1. The correlation of non-adjacent pairs as measured by MIV is significant. Classification based on every method improved significantly for MIV compared to MI(0). The highest accuracy achieved for MI(0) was 26.73% and for MIV it was 85.14% (see Table 5).
2. Normalized MI had an insignificant effect on scores generated from ΣH. Both methods reduce the sample-size error in estimating entropy and MI for sequences. A possible explanation for the lack of further improvement through normalization is that ΣH is a more effective corrective measure than normalization. We explore this possibility further in Section 6.2, were we consider entropy for both alphabets.
3. For the most accurate methods, using the Pfam prior decreased accuracy. Despite our concerns about using the frequency of a short sequence to estimate the marginal residue probabilities, the results show that these estimations better characterize the sequences than the Pfam prior probability distribution. However, four of the five best combinations contain a method utilizing the Pfam prior, showing that the two methods for estimating marginal probabilities are complimentary.
4. As with sequence-based classification, introducing gaps improves accuracy. For all methods, removing gap characters with the strict method drastically reduced accuracy. Despite this, two of the five best combinations included a strict scoring method.
5. The best scoring concatenated MIVs included both alphabets. The inclusion of ΣA is significant—all eight nonstrict ΣH methods scored better than any ΣA method (see Table 5). The inclusion shows that ΣA provides information not included in the ΣH and strengthens our assertion that the different alphabets characterize different forces affecting protein structure.
Table 5: Classification results for MI(0) and MIV20 methods. SD represents the standard deviation of the experiment accuracies.

| Rank | Method                        | MI(0) accuracy | MIV20 accuracy |
|------|-------------------------------|----------------|----------------|
|      |                               | Mean  | SD  | Mean  | SD  |
| 1    | Hybrid-Σ_H                   | 26.73%| 2.59| 85.14%| 2.06|
| 2    | Normalized hybrid-Σ_H        | 26.20%| 4.16| 85.01%| 2.19|
| 3    | Literal-Σ_H                  | 22.92%| 3.41| 79.51%| 2.79|
| 4    | Normalized literal-Σ_H       | 22.45%| 3.88| 78.86%| 2.79|
| 5    | Normalized Hybrid-Σ_H w/Pfam prior | 26.31% | 3.95 | 77.21% | 2.94 |
| 6    | Literal-Σ_H w/Pfam prior     | 22.81%| 2.97| 71.57%| 3.15|
| 7    | Normalized literal-Σ_A       | 17.76%| 3.21| 66.69%| 4.14|
| 8    | Hybrid-Σ_A                   | 17.16%| 3.06| 64.09%| 4.32|
| 9    | Normalized literal-Σ_A w/Pfam prior | 22.73% | 4.90 | 76.89% | 2.91 |
| 10   | Literal-Σ_A w/Pfam prior     | 19.60%| 3.67| 63.39%| 4.05|
| 11   | Normalized Hybrid-Σ_A w/Pfam prior | 17.16% | 3.06 | 64.09% | 4.32 |
| 12   | Literal-Σ_A w/Pfam prior     | 16.36%| 2.84| 61.97%| 4.32|
| 13   | Normalized strict-Σ_A w/Pfam prior | 19.25% | 3.38 | 28.44% | 3.91 |
| 14   | Strict-Σ_A                   | 12.97%| 2.63| 24.09%| 3.87|
| 15   | Normalized strict-Σ_A w/Pfam prior | 19.25% | 3.38 | 28.44% | 3.91 |
| 16   | Normalized strict-Σ_A w/Pfam prior | 19.95% | 3.65 | 40.46% | 4.04 |
| 17   | Normalized strict-Σ_A w/Pfam prior | 19.60% | 3.65 | 40.46% | 4.04 |
| 18   | Normalized literal-Σ_A w/Pfam prior | 22.45% | 4.89 | 76.29% | 2.96 |
| 19   | Normalized literal-Σ_A       | 22.81%| 2.97| 71.57%| 3.15|
| 20   | Normalized literal-Σ_A w/Pfam prior | 22.81% | 2.97 | 71.57% | 3.15 |
| 21   | Normalized literal-Σ_A w/Pfam prior | 17.76% | 3.21 | 66.69% | 4.14 |
| 22   | Normalized literal-Σ_A       | 17.16%| 3.06| 64.09%| 4.32|
| 23   | Normalized literal-Σ_A w/Pfam prior | 17.16% | 3.06 | 64.09% | 4.32 |
| 24   | Normalized literal-Σ_A       | 16.36%| 2.84| 61.97%| 4.32|

6. FURTHER MIV ANALYSIS

In this section, we examine the results of our different methods of calculating MIVs for Pfam sequences. We first use correlation within the MIV as a metric to compare several of our scoring methods. We then take a closer look at the effect of reducing our alphabet size when translating from Σ_A to Σ_H.

6.1. Correlation within MIVs

We calculated MIVs for 120 276 Pfam sequences using each of our methods and measured the correlation within each method using Pearson’s correlation. The results of this analysis are presented in Figure 3. Each method is represented by a 20 × 20 grid containing each pairing of entries within that MIV.

The results strengthen our observations from the classification experiment. Methods that performed well in classification exhibit less redundancy between MIV indexes. In particular, the advantage of methods using Σ_H is clear. In each case, correlation decreases as the distance between indexes increases. For short distances, Σ_A methods exhibit this to a lesser degree; however, after index 10, the scores are highly correlated.

6.2. Effect of alphabets

Not all intraprotein interactions are residue specific. Cline [2] explored information attributed to hydrophathy, charge, disulfide bonding, and burial. Hydrophathy, an alphabet composed of two symbols, was found to contain half as much information as the 20-element amino acid alphabet. However,
with only two symbols, the alphabet should be more resistant to the underestimation of entropy and overestimation of MI caused by finite sequence effects [15].

For this method, a protein sequence is translated using the process given in Section 3.2. It is important to remember that the scores generated for entropy and MI are actually estimates based on finite samples. Because of the reduced alphabet size of $\Sigma_H$, we expected to see increased accuracy in entropy and MI estimations. To confirm this, we examined the effects of converting random sequences of 100 residues (a length representative of those found in the Pfam database) into $\Sigma_H$.

We generated each sequence from a Bernoulli scheme. Each position in the sequences is selected independently of any residues selected before it, and all selections are made randomly from a uniform distribution. Therefore, for every position in the sequence, all residues are equally likely to occur.

By sampling residues from a uniform distribution, the Bernoulli scheme maximizes entropy for the alphabet size ($N$):

$$H = -\log_2 \frac{1}{N}. \quad (15)$$

Since all positions are independent of others, MI is 0. Knowing the theoretical values of both entropy and MI, we can compare the calculated estimates for a finite sequence to the theoretical values to determine the magnitude of finite sequence effects.

We estimated entropy and MI for each of these sequences and then translated the sequences to $\Sigma_H$. The translated sequences are no longer Bernoulli sequences because the residue partitioning is not equal—eight residues fall into one category and twelve into the other. Therefore, we estimated the entropy for the new alphabet using this probability distribution. The positions remain independent, so the expected MI remains 0.

Table 7 shows the measured and expected entropies for both alphabets. The entropy for $\Sigma_A$ is underestimated by .144, and the entropy for $\Sigma_H$ is underestimated by only .007. The effect of $\Sigma_H$ on MI estimation is much more pronounced. Figure 4 shows the dramatic overestimation of MI in $\Sigma_A$ and high standard deviation around the mean. The overestimation of MI for $\Sigma_H$ is negligible in comparison.

### Table 7: Comparison of measured entropy to expected entropy values for 1000 amino acid sequences. Each sequence is 100 residues long and was generated by a Bernoulli scheme.

| Alphabet | Alphabet size | Theoretical entropy | Mean measured entropy |
|----------|---------------|---------------------|-----------------------|
| $\Sigma_A$ | 20            | 4.322               | 4.178                 |
| $\Sigma_H$ | 2             | 0.971               | 0.964                 |

7. CONCLUSIONS

We have shown that residue correlation information can be used to characterize protein sequences. To model sequences, we defined and used the mutual information vector (MIV) where each entry represents the mutual information content between two amino acids for the corresponding distance. We have shown that MIV of proteins is significantly different from random sequences of the same character composition when the distance between residues is considered. Furthermore, we have shown that the MIV values of proteins are significant enough to determine the family membership of a protein sequence with an accuracy of over 90%. What we have shown is simply that the MIV score of a protein is significant enough
for family classification—MIV is not a practical alternative to similarity-based family classification methods.

There are a number of interesting questions to be answered. In particular, it is not clear how to interpret a vector of mutual information values. It would also be interesting to study the effect of distance in computing mutual information in relation to protein structures, especially in terms of secondary structures. In our experiment (see Table 4), we have observed that normalized MIV scores exhibit more information content than nonnormalized MIV scores. However, in the classification task, normalized MIV scores did not always achieve better classification accuracy than non-normalized MIV scores. We hope to investigate this issue in the future.

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