Domain-Based Predictive Models for Protein-Protein Interaction Prediction

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Protein interactions are of biological interest because they orchestrate a number of cellular processes such as metabolic pathways and immunological recognition. Recently, methods for predicting protein interactions using domain information are proposed and preliminary results have demonstrated their feasibility. In this paper, we develop two domain-based statistical models (neural networks and decision trees) for protein interaction predictions. Unlike most of the existing methods which consider only domain pairs (one domain from one protein) and assume that domain-domain interactions are independent of each other, the proposed methods are capable of exploring all possible interactions between domains and make predictions based on all the domains. Compared to maximum-likelihood estimation methods, our experimental results show that the proposed schemes can predict protein-protein interactions with higher specificity and sensitivity, while requiring less computation time. Furthermore, the decision tree-based model can be used to infer the interactions not only between two domains, but among multiple domains as well.

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1. INTRODUCTION

Proteins play an essential role in nearly all cell functions such as composing cellular structure, promoting chemical reactions, carrying messages from one cell to another, and acting as antibodies. The multiplicity of functions that proteins execute in most cellular processes and biochemical events is attributed to their interactions with other proteins. It is thus critical to understand protein-protein interactions (PPIs) involved in a pathway or a cellular process in order to better understand protein functions and the underlined biological processes. PPI information can also help predict the function of uncharacterized proteins based on the classification of known proteins within the PPI network. Furthermore, a complete PPI map may directly contribute to drug development as almost all drugs are directed against proteins.

The recent development of high throughput technologies has provided experimental tools to identify PPIs systematically. These methods include two-hybrid system [1], mass spectrometry [2], protein chips [3], immunoprecipitation [4], and gel-filtration chromatography [5]. Protein interactions can also be measured by biophysical methods such as analytical ultracentrifugation [6], calorimetry [7], and optical spectroscopy [8]. Among those experimental methods, the two-hybrid system is mature and accurate enough to be used for obtaining the full protein interaction networks of Saccharomyces cerevisiae [9, 10]. However, such techniques are tedious and labor-intensive. In addition, the number of possible protein interactions within one cell is enormous, a potentially limiting factor for experimental analyses. The need for faster and cheaper techniques has prompted extensive research in seeking complementary in silico methods that are capable of accurately predicting interactions.

A number of computational approaches for protein interaction discovery have been developed over the years. These methods differ by the information they used for protein interaction prediction. Some earlier methodologies focus on estimating the interaction sites by recognizing specific residue motifs [11] or by using features and properties related to interface topology, solvent accessible surface area, and hydrophobicity [12]. Some computational techniques are based on genomic sequence analysis, for example, analyzing correlated mutations in amino acid sequences between interacting proteins [13], searching for conservation of gene neighborhoods and gene order [14], using the gene fusion method or “Rosetta Stone” [15, 16], employing genomic context to infer functional protein interactions [17], and exploring the principle on similarity of phylogenetic trees for protein interaction prediction [18, 19]. Protein
structural information is also used to predict protein interactions. Lu et al. [20] employ multimeric threading algorithm to assign quaternary structures and to predict protein interactions. Several papers propose to predict protein interaction sites based on profiles of a residue and its neighbors [21–23]. Bock and Gough introduced a method to predict protein interactions based on the primary structure and associated physicochemical properties [24]. There are also methods that integrate interaction information from many different sources [25–27].

Recently, methods for predicting protein interactions using domain information are developed and preliminary results have demonstrated their feasibility [26, 28–34]. The domain-based methods are motivated by the fact that protein-protein interactions are the results of physical interactions between their domains. In [33, 35], homology searches and clustering of domain profile pairs are used for protein interaction prediction. Kim et al. introduced a statistical domain-based algorithm called “potentially interacting domain pair (PID)” [32]; it is similar to the association method except in how scores are calculated for all possible domain pairs. Deng et al. [28] propose a probabilistic approach that employs the maximum likelihood estimation (MLE). Ng et al. [36] infer domain interactions from data collected through three different sources: experimental protein interaction data, intermolecular relationship data from protein complexes, and predict data by Rosetta Stone. Those domain-based techniques only consider single-domain pair interactions; however, protein-protein interactions could be the result of multiple-domain pairs or groups of domains interacting with each other. Han et al. [29, 30] introduced a domain combination-based method, which considers all possible domain combinations as the basic units of a protein. The domain combination interaction probability is also based on the number of interacting protein pairs containing the domain combination pair and the number of domain combinations in each protein. Thus, the method still suffers from a general limitation of the association method, that is, ignoring other domain-domain interaction information between the protein pairs. It assumes that one domain combination is independent of another. Furthermore, the domain combination method is computationally more expensive than the MLE method because all possible domain combinations are considered instead of just single-domain pairs. For example, if a protein contains $m$ domains and the other contains $n$ domains, $(2^m - 1) \times (2^n - 1)$ possible domain combination pairs between the two proteins have to be considered instead of $m \times n$ single-domain pairs in the MLE method.

Even though there are a number of progresses made toward protein interaction prediction using current computational methods, they still have a limited range of applicability: the specificity and sensitivity are normally low. In this paper, we develop two domain-based statistical approaches to predict protein-protein interactions. In the proposed methods, instead of considering single-domain pair as the basic unit of protein interactions, all the possible domain combinations will contribute to the protein interactions. In addition, the proposed models do not need to make the assumption that domain pairs are independent of each other. We compare the methods to the MLE method, better results (in terms of the specificity and sensitivity) are obtained. Furthermore, the decision tree-based model can be used to infer domain-domain interactions for each predicted protein pair.

The paper is organized into four sections. Section 2 introduces our predictive systems and methods. The experimental results are presented in Section 3. Finally, conclusions are drawn in Section 4.

2. SYSTEM AND METHODS

We formulate protein-protein interaction prediction problems as two-class classification problems: each protein pair is a sample belonging to either “interaction” class (the two proteins interact with each other) or “noninteraction” class (the two proteins do not interact with each other). Each protein pair is characterized by the domains of two proteins. In Section 2.1, we discuss how each sample is represented in terms of domains. In Section 2.2, we introduce a forward-pruning decision tree-based modeling, and in Section 2.3, we briefly discuss a neural network-based modeling.

2.1. Feature representation

Among all proteins in our data set, there are 4293 unique Pfam domains (for details, refer to Section 3.1). For ease of implementation, each domain is labeled with a number between 1 and 4293. Each protein is represented by a vector of 4293 binary numbers where each binary number is associated with the 4293 domains. For example, if a protein has a domain with label 5, then the 5th number of the feature vector is 1, otherwise 0. In order to represent a protein pair, two vectors of binary numbers corresponding to each protein are concatenated to form the final input feature vector (see Figure 1). The domain labels in the second protein are concatenated to form the final input feature vector. The domain labels in the second protein are concatenated to form the final input feature vector.

Most domain-based approaches consider how often a specific domain pair appears in the protein pairs that are interacted with each other. The hypothesis is that the more often the domain pairs appear in the interacted protein pairs, the more likely the two domains are interacted with each other. These methods, however, ignore the possible associations between different domain pairs (by assuming that all domain pairs are independent of each other). Furthermore, the traditional methods cannot handle cases when more than two domains are involved in protein interactions (which are highly possible). In our proposed models, the representation of each protein pair as a vector consisting of domains allows us to take all the possible combinations of domains into consideration. For example, in neural networks, each domain is associated with an input neuron and every domain (input) may contribute to the output of the neural networks differently, depending on the neuronal network weights.
2.2. Forward-pruning decision tree

Decision tree is one of the most popular classification methods. A decision tree is a tree consisting of two types of nodes: decision nodes and class nodes. A class node is a leaf node of a tree, which specifies a class. A decision node, also known as a nonleaf node, specifies a test to be carried out on a single attribute. An edge branches out from a decision node is associated with an attribute value.

Decision tree construction usually involves constructing and pruning. In the constructing phase, a decision tree is built level by level from a given training data set starting at the root. At each decision node, we need to select the best splitting attribute based on the measure called “goodness of split,” which assesses how well the attributes discriminate between classes. A number of measures were developed to select attributes for splitting. We use the information gain [19, 37] as the “goodness of split” measure, which is based on the classical formula from information theory. The information gain measures the theoretical information content of a code by \(\sum_i p_i \log(p_i)\), where \(p_i\) is the probability of the \(i\)th message.

Let \(D = \{X_1, X_2, \ldots, X_n\}\) represent the \(n\) training samples and let \(X_i = [x_{i1}^i, x_{i2}^i, \ldots, x_{im}^i, y_i]\) represent the \(i\)th sample with \(m\) attributes \(x\) belonging to the class \(y_i\). In our problem, \(y_i = 1\) represents for the “interaction” class and \(0\) for the “noninteraction” class. Assume that the numbers of samples in the “interaction” class and the “noninteraction” class are \(n_1\) and \(n_2\), respectively. The information needed to classify samples given only the decision class totals as a whole is

\[
H(C) = - (P(y = 0) \log P(y = 0) + P(y = 1) \log P(y = 1)),
\]

where \(P(y)\) is the class probability among all samples \((P(y = 1) = n_1/n\) and \(P(y = 0) = n_2/n)\). The information needed to classify samples, given knowledge of the attribute \(x_i\), is defined as

\[
H(C | x) = \sum_{i=1}^{2} P(x = x_i) H(C | x = x_i),
\]

where \(P(x = x_i)\) is the probability of the attribute \(x\) taking the value \(x_i\) (in our case, \(x\) can only take ones \((x_1 = 1)\) and zeros \((x_2 = 0)\)). The information needed given each attribute value, \(H(C \mid x = x_i)\), is defined by

\[
H(C \mid x = x_i) = - (P(y = 0 \mid x = x_i) \log P(y = 0 \mid x = x_i) - P(y = 1 \mid x = x_i) \log P(y = 1 \mid x = x_i)),
\]

where \(P(y = y_j \mid x = x_i)\) is the conditional probability of \(j\)th class given attribute value \(x_i\). Finally, the information gain (IG) measure for an attribute \(x\) can be calculated with (1) and (2):

\[
IG(x) = H(C) - H(C \mid x).
\]

During the tree construction phase, at each decision node, the attribute with the largest information gain and which has never been selected in the branch will be selected. The information gain for each attribute is calculated by (4) based on training samples classified at the decision node. After the attribute is selected, it splits the training samples according to the attribute values. Each decision node keeps splitting until all training samples at the node belong to the same decision class or no more attribute is left for splitting. The decision class associated with majority of the training samples at each leaf node is assigned as the prediction.

In order for the decision tree to work successfully and to avoid overfitting problems, branches with little statistical significance have to be pruned or removed. Traditionally, pruning methods begin with a full tree constructed from a set of training data, and remove tree branches in the bottom-up fashion. It has worked well for problems with a handful of attributes. However, for our problem, we have approximately 9000 attributes, and a full tree is expected to be extremely large. Therefore, pruning after building the full tree may not be a practical idea.

A forward-pruning technique that prunes during tree construction was implemented. It stops building the tree if certain conditions are met. First, we reserved 2/3 of the training data set as training and 1/3 as validation data set. The decision tree is then built with the training data set, and before each splitting attribute is selected, classification error for that particular decision node is calculated over the validation data set. Among all validation samples that go through the branch and reach the node for classification, the classification error is just the proportion of misclassified samples. If the error is less than or equal to a prespecified threshold, the node stops splitting and becomes a leaf node. This makes sure that branches with little statistical validity are not pursued.

2.3. Neural network

Artificial neural network is biologically inspired, and it solves problems by example mapping. A neural network learns from a collective behavior of the simple processing elements called neurons. The processing elements are connected by weighted connections. The weights on the connections contain the network knowledge. Each neuron performs limited
operations and works in parallel with other neurons to solve problems quickly.

Typically, a neural network has three layers: input, hidden, and output layers. The input layer is used to encode instance presented to the network for processing. The processing elements in the input layer are called input nodes, which may represent an attribute or feature value in the instance. In our application, each input neuron represents a domain. The hidden layer makes the network nonlinear through its hidden units. The output layer contains output units, which assign values to the input instance. The simple view of a neural network structure is depicted in Figure 2.

The nodes between layers are fully connected. For example, each hidden node is connected to all input nodes, and each output node is connected to all hidden nodes. There are no connections between nodes in the same layer. All connections point to the same direction from the input toward the output layer. The weights associated with each connection are real numbers in the range of [0, 1]. The connection weights are initialized to random small real numbers and are adjusted during network training. This structure can capture various combinations between domains, instead of only two domains at a time. Each domain will contribute to the network output, depending on the weights associated with the nodes. The network predicts a protein pair to be interacting if the output node value is larger than or equal to certain threshold. Normally, the threshold is set to be 0.5. In our experiment, we implement a multilayer feed-forward network using the delta rule.

3. EXPERIMENTAL RESULTS

3.1. Data sources

Protein-protein interaction data for the yeast organism were collected from the database of interacting proteins (DIP) [38], Deng et al. [28], and Schwikowski et al. [39]. The data set used by Deng et al. [28] is a combined interaction data experimentally obtained through two-hybrid assays on Saccharomyces cerevisiae by Uetz et al. [10] and Ito et al. [9]. Schwikowski et al. [39] gathered their data from yeast two-hybrid, biochemical, and genetic data.

We obtained 15409 interacting protein pairs for the yeast organism from DIP, 5719 pairs from Deng et al. [28], and 2238 pairs from Schwikowski et al. [39]. The data sets were then combined by removing the overlapping interaction pairs. Because domains are the basic units of protein interactions, proteins without domain information cannot provide any useful information for our prediction. Therefore, we excluded the pairs where at least one of the proteins has no domain information. To further reduce noises in our data, pairs with both proteins containing only one domain, which only occurred once among all proteins, were also excluded. Finally, we have 9834 protein interaction pairs among 3713 proteins, and it is separated evenly (4917 pairs each) into training and testing data sets. Negative samples are generated by randomly picking a pair of proteins. A protein pair is considered to be a negative sample if the pair does not exist in the interaction set. Total of 8000 negative samples were generated and also separated into two halves. The final training and testing data set both have 8917 samples, 4917 positive and 4000 negative samples.

The protein domain information was gathered from Pfam [40], a protein domain family database, which contains multiple sequence alignments of common domain families. Hidden Markov model profiles were used to find domains in new proteins. The Pfam database consists of two parts: Pfam-A and Pfam-B. Pfam-A is manually curated, and Pfam-B is automatically generated. Both Pfam-A and Pfam-B families are used here. In total, there are 4293 Pfam domains defined by the set of proteins.

3.2. Evaluation criteria

To evaluate the neural network and decision tree-based methods for predicting PPIs, we use both specificity and sensitivity. The specificity SP is defined as the percentage of matched noninteractions between the predicted set and the observed set over the total number of noninteractions. The sensitivity, denoted by SN, is defined as the percentage of matched interactions over the total number of observed interactions.

\[
SP = \frac{\# \text{ of true negative PPI}}{\# \text{ of observed negative PPI}} \quad \text{and} \quad SN = \frac{\# \text{ of true positive PPI}}{\# \text{ of observed positive PPI}}
\]

(5)

3.3. Predicting PPI

Training

In order to predict protein-protein interactions, our models need to be trained first with the training data set. In our neural network structure, 8586 input nodes are fed to the hidden layer. Values of the hidden neurons are then fed to the output layer with 1 output node. The number of hidden neurons and training cycles can greatly impact how well a network is trained; therefore, in order to make appropriate choices on the parameters, we have tested the network with various numbers of hidden neurons and training cycles. The training errors were plotted against different number of training cycles for different number of hidden neurons (see Figure 3). As shown in Figure 3, networks with different number of hidden neurons all converge. Most of them actually converge after 1000 training cycles. The difference is in the starting training errors and how fast a convergence is
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reached. Based on the results, we have chosen 50 hidden neurons and 2000 training cycles.

A decision tree is constructed over a set of training data with satisfied threshold for forward-pruning set to 0.01. Then the tree is used to classify samples in the test data set.

The results of our methods are compared with the maximum likelihood estimation (MLE) results \[ 14 \]. The MLE method is trained with false positive rate \( fp = 1.0 \times 10^{-5} \) and false negative rate \( fn = 0.85 \) over our training data.

**Test results**

To classify a new protein pair as either interacting or noninteracting, the pair is first converted to a feature vector as described in Section 2.1 and then used as input in the modes. Prediction is made based on the result generated by the models. For neural network, the output is a real number between 0 and 1. If the output is greater than or equal to a certain threshold (0.5), then the pair is said to be interacting. This threshold can be changed to produce an ROC curve for the neural network. In our decision tree model, the classification decision is either 0 for noninteracting or 1 for interacting. The decision tree ROC curve is constructed by tree chopping. When a decision tree is chopped to a certain height, its prediction accuracy in terms of specificity and sensitivity would change and consequently producing a point on the ROC curve.

Figure 4 compares the ROC curves of the three methods: neural network, decision tree, and MLE. As shown in Figure 4, both the neural network and decision tree outperform the MLE method. Table 1 shows the prediction results that each of the three methods can achieve over the test data set. From Table 1, we can see that with comparable sensitivities fixed at approximately the same level 78%, both of our methods can achieve above 60% in specificity, and MLE can only achieve specificity of 37.53%. Clearly from Figure 4, the neural network gives the best performance, and decision tree performs almost as well. Note that the training data set for decision tree is only 2/3 of the training data set used for the neural network and MLE. Therefore, with a full training data set, accuracy of decision tree is expected to be better.

**Computational speed**

Among the three methods, neural network-based system is the fastest, as the neural network’s running time mainly depends on the number of input nodes, hidden neurons, output nodes, and training cycles. Number of output nodes can be ignored in complexity calculation since it is one. Although the number of input nodes is extremely large, there are many zeros that do not involve in any calculation. Only the ones are considered by the network, so this is reduced to a quite small number mostly between 1 and 20. Therefore, the neural network’s computation time mainly depends on the number of hidden neurons and training cycles. The decision tree runs much slower than the neural network, however, is much faster than MLE methods. MLE takes the longest to run because it calculates interaction probability for all possible protein pairs, domain pairs, and protein pairs to be observed as interacting. For each type of pairs, there are several million possibilities, and it performs the calculations for a number of iterations. When running the models on an Intel Dual Xeon 2.6 GHz computer, it took approximately 20 minutes to train the neural network and several hours for the decision tree model. A single MLE iteration took several hours to execute, and many iterations are required for the likelihood function to converge.

The running time for training the models may not be a big issue if the number of training samples (protein pairs) is not very large. Once the models are trained, the running times for tests are very fast. However, as more and more interacted protein pairs are discovered, the running time for
MLE methods may become an issue, as it may take several months to train the model.

### 3.4. Inferring domain-domain interactions

With the decision tree method, we can also infer domain interactions. For each correctly predicted protein-protein interaction pairs, we can derive domains involved in the decision process by looking at the branch the protein pair took to reach the prediction. The branch of the decision tree contains all domains from both proteins that contribute to the correct classification. Among those involved domains, domains of two proteins with value of 1 indicate their existence in the protein pairs. Thus, these domains are interacted with each other. Apparently, we can retrieve more than two domains from each branch. This is attractive, as in some protein-protein interactions it is highly possible that more than two domains interact. Most of the existing methods can only identify domain pairs.

Table 2 lists some of the domain interacting pairs identified by the decision tree. We also found that those domain pairs are considered to be interacting pairs with a high confidence by the InterDom, a database of putative interacting domains, developed by Ng et al. [34]. For example, SH3 (PF00018) and Pkinase (PF00069) are derived from a protein-protein interaction only involving single-domain proteins. A protein is considered as a single-domain protein if it has only one domain and the domain accounts for at least 50% of the protein length. The domain interactions derived from single-domain protein interactions are usually considered to be highly likely. The domain SH3 is also found to interact with Pkinase_Tyr by Pfam [40]. The Pfam domain-domain interactions are determined by mapping Pfam domains onto the PDB structures, and interaction bonds are then identified. Pkinase and Pkinase_Tyr are both members of the protein kinase superfamily clan.

For each protein-protein interaction pair, there may be more than two domains involved. Using the decision tree, some domain combinations that could have involved in interaction are also identified. A domain combination is defined as two or more domains functioning as a whole during interaction. We list some of the identified domain combinations in Table 3. As an example, domains PF00172 (Znclus) and PB043568 (row 5) are discovered to bind together and interact as a whole unit with the PF00183 (HSP90) domain in another protein. We found that the PF00172 and PB043568 domains are the only domains existed in protein HAP1. The HAP1 protein is discovered to form biochemically distinctive higher-order complex with the HSP82 protein in the absence of heme [3]. The HSP82 protein contains two domains, which are HSP90 (PF00183) and HATPase_C (PF02518). Neither one of the two domains is identified to...
form an interacting domain pair with Znclus (PF00172) by iPfam [23]; therefore, our hypothesis formed from the prediction results is that the Znclus domain (PF00172) forms a domain combination with PB043568 and interacts with the HSP90 (PF00183) as a whole.

4. CONCLUSION

Proteins perform biological functions by interacting with other molecules. It is hypothesized that proteins interact with each other through specific intermolecular interactions that are localized to specific structural domains within each protein. Often, protein domains are structurally conserved among different families of proteins. Thus, understanding protein interactions at the domain level gives detailed functional insights onto proteins. Most of the existing domain-based computational approaches for predicting protein interaction assume that domain pairs are independent of each other and consider the interactions between two domains only. In this paper, we develop decision tree and neural network-based models to predict protein-protein interactions. These systems are capable of utilizing all the possible interactions between domains. For example, in the neural network-based method, all domains will contribute to the prediction of protein-protein interactions with different weights (e.g., the weights for domains that are not included in the protein pairs may be zeros). We compared our results with the maximum likelihood estimation method. The experimental results have shown that both methods can predict protein-protein interactions with higher specificity and sensitivity than the MLE method. Computationally, the MLE method needs extensive computation time and runs much slower than our methods. In addition, the decision tree method is particularly useful because domain-domain interactions can be inferred from the domains involved in predicting protein interactions, especially, this method allows for discovering interactions of domain combinations.

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