**Struct2Net: a web service to predict protein–protein interactions using a structure-based approach**

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Struct2Net: a web service to predict protein–protein interactions using a structure-based approach

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ABSTRACT

Struct2Net is a web server for predicting interactions between arbitrary protein pairs using a structure-based approach. Prediction of protein–protein interactions (PPIs) is a central area of interest and successful prediction would provide leads for experiments and drug design; however, the experimental coverage of the PPI interactome remains inadequate. We believe that Struct2Net is the first community-wide resource to provide structure-based PPI predictions that go beyond homology modeling. Also, most web-resources for predicting PPIs currently rely on functional genomic data (e.g. GO annotation, gene expression, cellular localization, etc.). Our structure-based approach is independent of such methods and only requires the sequence information of the proteins being queried. The web service allows multiple querying options, aimed at maximizing flexibility. For the most commonly studied organisms (fly, human and yeast), predictions have been pre-computed and can be retrieved almost instantaneously. For proteins from other species, users have the option of getting a quick-but-approximate result (using orthology over pre-computed results) or having a full-blown computation performed. The web service is freely available at http://struct2net.csail.mit.edu.

INTRODUCTION

Systems biology research is like solving a jigsaw puzzle: the goal is to figure out how the various parts (i.e. genes and proteins within the cell) interact and work together. The interactome of an organism is then analogous to the puzzle’s key: it describes the network of all the protein–protein interactions (PPIs) in a cell. As such, identifying all the protein-protein interactions for an organism is of great value, akin to sequencing its genome. Despite the use of high-throughput techniques in discovering PPIs, however, the coverage of experimentally determined PPI data remains poor (Table 1). Such low coverage is partly because the set of possible PPIs to be verified is so large (100 million for a species with 10 000 genes) that any exhaustive experimental verification will take a long time, even with high-throughput techniques. Indeed, the rate of PPI discovery has slowed down in recent years (Figure 1). Furthermore, the experimental approaches have limitations of their own. For example, tandem affinity purification experiments have historically had difficulty identifying transient interactions, while yeast two-hybrid experiments may produce false positives due to promiscuous proteins (1); recently, statistical methods have been proposed to improve confidence in the output of these experiments (2,3).

The paucity of interactome coverage has motivated significant research interest in methods for supplementing experimentally determined PPI data with interactions inferred or predicted from other sources. A wide variety of methods have been proposed. One approach is to use interologs, which are basically PPIs mapped from another species to the target species (4,5). The key problem there is to correctly map homologs across species (6,7). Another approach is to use functional genomic data and leverage the observation that a pair of interacting proteins is also likely to have similar GO annotations, occupy the same cellular sub-compartments, or correspond to genes with similar expression profiles (8,9). Consequently, many researchers have described machine learning-based approaches to predict PPI data from functional genomic data such as gene expression, cellular localization and GO annotation.

Predictions from many of these approaches have been aggregated into a number of databases/web services offering predicted PPIs. The STRING database (10)...
The use of structure-based approaches to predict interactions has been previously proposed. Aloy and Russell (16) suggested the use of structure-based approaches to predicting PPIs. Lu et al. (17) constructed statistical potential functions to evaluate potential PPIs and later described MultiProspector, a structure-based prediction algorithm (18). In a previous paper, we proposed a prediction algorithm (also used by Struct2Net). Our algorithm builds upon previous work like MultiProspector, by combining a threading approach for template alignment with a novel machine learning approach to estimate a confidence score for the interaction. In our previous proof-of-concept paper, we discussed how Struct2Net’s results compare favorably to related work (19).

Unfortunately, the progress made in prediction has not yet translated into comprehensive community resources. Aloy and Russell (20) have described InterPreTS, a web-server to predict PPIs for a given protein, using a homology modeling approach. We have already mentioned Lu et al.’s MultiProspector tool which also predicts PPIs (17). More recently, Fukuhara and Kawabata have described HOMCOS (21,22) a web-server that performs a similar task by homology modeling. MODBase is a database of homology models for protein complexes that have high sequence similarity to known structures (23). ADAN is a specialized database for prediction of PPIs mediated by linear motifs and utilizes position-specific matrices to assess putative interactions (24).

We believe that Struct2Net offers a significant advantage over such homology modeling approaches. Successful use of homology modeling requires relatively high sequence similarity between the query and template protein pairs. In contrast, we use a threading-based approach which widens the range of proteins for which predictions can be made. The use of threading also offers us improved performance: Fukuhara et al. (22) have reported that HOMCOS achieves a recall of 80% with a precision of about 10%; in comparison, Struct2Net achieves a recall of 80% with a precision of 30% [here, recall = (true positives)/(true positives + false negatives) and precision = (true positives)/(true positives + false positives)].

The Struct2Net approach can also be contrasted with methods that model PPIs based on domain-domain interactions. These approaches argue that the structural basis of protein interaction can be traced to the presence of interacting domains. A domain can be represented simply by its sequence motif or as a structure-fragment. Given a set of known PPIs, one can infer the set of domain pairings that are presumably the underlying cause of interaction. In principle, these pairs can then be used to make prediction for unannotated protein pairs. There has been a significant amount of work on analyzing PPIs using such domain interactions. Some researchers focus solely on the sequence signature of the domains, proposing methods to predict PPIs using these sequence domains (25,26). In previous work, we have discussed how such sequence-domain-based prediction can be combined with our approach in a machine-learning framework (19). We also described some results that suggest that Struct2Net’s predictive ability compares well with the sequence-domain approaches.

Table 1. Availability of experimental PPI data for major eukaryotic organisms

<table>
<thead>
<tr>
<th>Organism</th>
<th>Number of interactions</th>
<th>Percentage of proteins with at least one interaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouse</td>
<td>1486</td>
<td>6.0</td>
</tr>
<tr>
<td>Human</td>
<td>26,640</td>
<td>41.8</td>
</tr>
<tr>
<td>Worm</td>
<td>4559</td>
<td>14.5</td>
</tr>
<tr>
<td>Fly</td>
<td>22,740</td>
<td>52.7</td>
</tr>
<tr>
<td>Yeast</td>
<td>48,901</td>
<td>93.5</td>
</tr>
</tbody>
</table>

The statistics here and in Figure 1 have been computed using data from BioGRID (v 2.0.53) (35). Data based on phenotypic suppression/enhancement and synthetic interaction was excluded as these experiments do not provide evidence of a direct physical interaction between proteins.

Figure 1. Rate of discovery of new eukaryotic PPI data has slowed.

combines experimental datasets (e.g. KEGG, BioGRID, HPRD) with computational predictions based on co-expression, interologs and text-mining, etc. The entries in this database correspond to functional interactions, and may not always be directly interpretable as PPIs. Another database, IntAct (11), focuses more on inferring interactions from expert curation of data from literature. Other public services include DOMINO (12), InterDom (13) and I2D (14). However, all of these databases suffer from a common selection bias: often, the proteins that have been selected for PPI experiments are usually genes/proteins that have received some attention before and, as such, are also more likely to have functional genomic data.

In this article, we describe Struct2Net, a web service for predicting PPIs using a structure-based approach. Our method predicts interactions by threading each pair of protein sequences onto potential structures in the Protein Data Bank (PDB) (15). Struct2Net provides PPI predictions that are independent of all the non-structure-based approaches and may thus be combined with any of them. Another key advantage of our web server is that, apart from the PDB data, the prediction algorithm only requires protein sequence data as input. It can thus be applied to proteins for which no functional data is available provided there is a suitable PDB structural template available.

The use of structure-based approaches to predict interaction has been previously proposed. Aloy and Russell (16) suggested the use of structure-based approaches to predicting PPIs. Lu et al. (17) constructed statistical potential functions to evaluate potential PPIs and later described MultiProspector, a structure-based prediction algorithm (18). In a previous paper, we proposed a prediction algorithm (also used by Struct2Net). Our algorithm builds upon previous work like MultiProspector, by combining a threading approach for template alignment with a novel machine learning approach to estimate a confidence score for the interaction. In our previous proof-of-concept paper, we discussed how Struct2Net’s results compare favorably to related work (19).

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Other researchers have aimed to understand these domains from a structural perspective. Prieto and Las Rivas (27) have reviewed publicly available databases that facilitate analysis of domain-based PPIs: 3did (28), SNAPPI-DB (29), iPfam (30), PIBASE (31) and PSIBase (32). While our approach has some parallels with these approaches, our goal is significantly different. The domain interaction databases are essentially repositories of known structural data, analyzed specifically from a PPI perspective. Prediction, which is our core goal, is usually out of the scope of these approaches. In the ‘Methods Overview’ section below, we suggest how Struct2Net could take advantage of some of these databases.

METHODS OVERVIEW

The guiding intuition behind our prediction approach is that if a potential interaction is sufficiently favorable from a thermodynamics perspective, it is likely to be true. We provide a brief description of the algorithm here. For more details, see Singh et al. (19), which describes a proof-of-concept implementation of the algorithm.

Our approach proceeds in two broad stages. Given a pair of protein sequences, the first stage predicts the most likely structure of the complex formed by the two proteins and produces a vector of scores that quantitatively represent the thermodynamic suitability of this structure. For this task, we start by analyzing the PDB to construct a database of complex-structure templates; then we thread the two sequences jointly through the various templates in this database and identify the best fitting template. Our threading algorithm formulates the threading problem as an integer linear program (ILP) and uses branch-and-bound techniques to efficiently find the solution. The ideas in this algorithm, when applied to a single-protein threading context in the RAPTOR program, have performed well at various blind tests and competitions (33,34). To speed up prediction, we ran PSI-BLAST (35) before running our threading algorithm. If some templates in our database appear in the list of PSI-BLAST top hits (E-value < 10^-3), we simply thread the sequence pair to these templates instead of the whole template database. This speedup procedure does not lose accuracy since PSI-BLAST is good at close homolog detection.

We now briefly describe how the database of complex templates was constructed. We begin by using a simple geometric criterion to determine if two protein chains form a complex. This provides an unbiased and objective way of characterizing an interaction. Given two protein chains in the same PDB entry, we first calculate the distance between two (non-hydrogen) atoms from these two chains. We assume that there is an interaction between two residues of different chains if there is at least one pair of atoms from these two residues with distance <3.5 Å. If there are at least 10 interacting residue pairs between two chains A and B, we say these two chains form a complex. To avoid redundancy, we enforce the constraint that any two templates in the database share <70% sequence identity. Following this procedure, our database currently contains 10,111 dimers. While our template database (and the web server’s predictions) are currently built at the chain level, we intend to explore the incorporation of domain–domain interactions (from databases like SNAPPI, 3did, PSIBase, PIBASE, etc.) into it. This may help enlarge the database’s coverage.

The second stage of our prediction approach evaluates the likelihood of the interaction based on the predicted structure. We compute various energy scores that evaluate the structure (e.g. the quality of the interfacial region, the quality of fit for the individual proteins). Given these, we use logistic regression to predict whether an interaction will occur. Let \( y_i \) be an indicator variable representing protein interaction, i.e. \( y_i = 1 \) if the protein pair \( i \) interacts and 0 otherwise. Let \( x_i = \{x_i^1, x_i^2, \ldots \} \) be the vector of scores we use for prediction. We fit the following model:

\[
\log \frac{P(y_i = 1|x_i)}{P(y_i = 0|x_i)} = \alpha + \beta_1 x_i^1 + \beta_2 x_i^2 + \cdots
\]

where \( \alpha, \beta_1, \beta_2, \) etc. are parameters to be learned from data. To train this model, we constructed positive and negative training sets. Obviously, the choice of these sets can have a substantial impact on the prediction algorithm’s quality.

We have developed criteria for constructing these datasets. The exact criteria and a discussion about the rationale behind them are available at the Struct2Net website. Briefly, we require that the positive examples either come from a small set of trustworthy protocols or from low-throughput experiments; or roughly correspond to co-clustered protein pairs in the PPI network. We chose BioGRID (36) as our data-source, but other multi-species genome-wide databases [e.g. MINT (37) or APID (38)] could also be used. For negative examples, we require that the two proteins either be disconnected in the PPI network or be at least 3 hops away from each other. Using these criteria, we had a training set of 62,519 pairs and a test set of 15,635 pairs (with a positive:negative ratio of 1:6 approximately, in both sets). We believe that these datasets provide good evidence of validation. Our construction of the negative dataset was motivated by similar approaches in literature (8). For positive datasets, we believe that our approach identifies true PPIs with better confidence than an alternative approach that would select repeatedly observed PPIs (across multiple experiments). Our scheme emphasizes protocols and studies with low error-rates. In contrast, many high-throughput protocols (e.g. yeast two-hybrid) have systematic biases which may manifest as repeated false positives, even across multiple experiments.

In addition to the energy scores from the first stage, we aimed to enhance the model’s predictive power by adding extra terms to it. These included interaction terms, non-linear functions of the energy scores, as well as normalized scores (e.g. interfacial energy normalized by the average of the two proteins’ sequence length). We then used the Akaike information criterion (AIC) to select the model with the best trade-off of higher
We constructed a training set and test set of positive sequences at a time. In addition, with orthology-based approximation, users can specify just one protein identifier or FASTA sequence; in that case, all the interactions involving that protein will be returned.

Predictions are retrieved almost instantaneously when querying by ids. When querying by protein sequence and with orthology-based approximation selected, typical run-times are within 20 s. Full-blown computations finish within 45 mins, given query and subject sequences. Because of the potential for long run-times (e.g. if the server is overloaded), we encourage the user to supply an email address to which a job id and a link to the progress page are sent upon submission. Alternatively, users can check the progress of a submitted job by entering a job id in the ‘Fetch Job’ webpage. Upon completion of a job, an email with a link to the results page will also be sent.

For pre-computed predictions in \textit{S. cerevisiae}, \textit{D. melanogaster} and \textit{H. sapiens}, the output for each query protein sequence consists of a list of all predicted interactions along with their confidence scores (Figure 3D). Struct2Net interactively links each gene hit to various sequence databases along with associated GO annotations and aliases. Results are also cross-referenced with BioGrid in the case where experimental data is available for a predicted interaction. For predictions in other organisms using the Struct2Net algorithm, the output for each sequence pair contains details on the best-fit complex templates used during the computation including sequence alignments, alignment scores, their associated z-scores and an interfacial energy calculated between the sequence pair (Figure 3D). In addition, an overall confidence score is provided for each potential interaction. The confidence score ranges from 0 to 1, with 0 indicating minimum confidence and 1 indicating maximum confidence. In the ‘About’ page of the website, we discuss threshold choices that would allow a user to achieve a desired level of specificity in the output or a desired number of interactions above the threshold. For batch queries, results are separated by each pair of protein sequences.

For users interested in performing large-scale database analysis and classification, bulk download of predictions for \textit{S. cerevisiae}, \textit{D. melanogaster} and \textit{H. sapiens} is also available. We have further made available a script on the Download page that facilitates the integration of Struct2Net’s predictions with other tools. In the future, we plan to update our template database every 3 months. Every 6 months, we will update our pre-computed predictions using the latest template database.

In Table 2, we provide an example of our algorithm’s results on a set of protein pairs often used as test cases. For comparison, we have also displayed the results of HOMCOS and InterPretS for these pairs. MultiProspector no longer seems to be publicly available, and we could not include its results. The test cases we have chosen are the same as chosen by Fukuhara et al. for evaluating HOMCOS (22). As can be seen, for pairs that are thought to be interacting (Table 2), the final scores from Struct2Net are, on average, significantly higher than for non-interacting pairs (Table 2). Furthermore, normalizing the difference between the average interacting and non-interacting scores for each method by the

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{sensitivity_specificity.png}
\caption{Sensitivity versus specificity. The prediction algorithm can achieve 60% sensitivity while maintaining 75% specificity as measured on the test set. Here, sensitivity = (true positives)/(true positives + false negatives) and specificity = (true negatives)/(true negatives + false positives). We constructed a training set and test set of positive and negative examples from yeast and fly, using criteria we have developed to identify high-confidence positive and negative examples of PPIs (see the website FAQ for details). After training the logistic regression model on the training set, its performance was measured on the test set.}
\end{figure}
Figure 3. Web interface and output of Struct2Net. (A and B) Web server entry page. (C) A query option for either a quick-but-approximate approach (using orthology over pre-computed predictions from yeast, fly and human) or a full-blown computation using the Struct2Net algorithm. (D) Example of an output page when choosing to thread pairs of sequences onto all templates. Confidence scores for a potential interaction are displayed along with associated template-sequence alignments and threading details.
We chose sets of interacting and non-interacting protein pairs; these pairs are taken from an analysis by HOMCOS authors (22). Low confidence templates are indicated as ‘none’. For comparison, the scores from HOMCOS and InterPretS for these pairs are also shown. Struct2Net provides a coverage methods is able to capture alignments more in the ‘twilight zone’ by matching query sequences to structural templates (19). Furthermore, it has been shown that localized threading using interface profiles can further improve coverage and accuracy (39,40). While Struct2Net can be used for validation purposes (e.g. to double-check entries in BioGRID), its coverage limitations may at the present time make it better suited to be an exploratory tool, especially for unannotated proteins where only sequence information is available, or to be used in conjunction with low-confidence experimental data.

**CONCLUSIONS**

Although high-throughput biochemical approaches for discovering PPIs have proven very successful, the current experimental coverage of the interactome remains inadequate and would benefit from computational tools. The Struct2Net web server allows the user to easily query for high-probability structure-based interactions as a potentially high-quality, high-coverage data source for large-scale integrative approaches to interactome construction. The predicted interactions also include a numeric score, allowing users to further filter the data. To the best of our knowledge, this web server is the first of its kind and will be of considerable value to systems biologists interested in PPIs, partly because of the effort we have put into identifying high-confidence positive and negative examples of PPIs as inputs to machine learning algorithms and the extensive computational effort involved in making each prediction. A strength of this web service is its ongoing integration of up-to-date structural templates for improving its predictions. Struct2Net’s predictions may be used on their own or as one of the

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<th>Uniprot IDs</th>
<th>Templates</th>
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<th>Zseqcon</th>
<th>Zcon</th>
<th>Best Z-score</th>
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<td>1g6S5K</td>
<td>P22141, P30656</td>
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<td>Q70439, O83834</td>
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<td>SNTT3NHN</td>
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<tr>
<td>Average</td>
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<td>-3.57</td>
<td>2.87</td>
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</table>

**We chose sets of interacting and non-interacting protein pairs; these pairs are taken from an analysis by HOMCOS authors (22). Low confidence templates are indicated as ‘none’. For comparison, the scores from HOMCOS and InterPretS for these pairs are also shown.**

**Table 2. Struct2Net results for set of interacting and non-interacting protein pairs**
inputs into a computational framework that combines them with other sources (e.g. low-quality experimental data or predictions from functional genomic data). For example, Jensen et al. (10), Qi et al. (8) and Srinivasan et al. (9) have described some general approaches for combining various predictors of PPI data. Struct2Net’s predicted interaction scores can easily be integrated into such models.

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