

Identifying protein complexes from heterogeneous biological data

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ABSTRACT

With the increasing availability of diverse biological information for proteins, integration of heterogeneous data becomes more useful for many problems in proteomics, such as annotating protein functions, predicting novel proteinprotein interactions and so on. In this paper, we present an integrative approach called InteHC (Integrative Hierarchical Clustering) to identify protein complexes from multiple data sources. Although integrating multiple sources could effectively improve the coverage of current insufficient protein interactome (the false negative issue), it could also introduce potential false-positive interactions that could hurt the performance of protein complex prediction. Our proposed InteHC method can effectively address these issues to facilitate accurate protein complex prediction and it is summarized into the following three steps. First, for each individual source/feature, InteHC computes the matrices to store the affinity scores between a protein pair that indicate their propensity to interact or co-complex relationship. Second, InteHC computes a final score matrix, which is the weighted sum of affinity scores from individual sources. In particular, the weights indicating the reliability of individual sources are learned from a supervised model (i.e., a linear ranking SVM). Finally, a hierarchical clustering algorithm is performed on the final score matrix to generate clusters as predicted protein complexes. In our experiments, we compared the results collected by our hierarchical clustering on each individual feature with those predicted by InteHC on the combined matrix. We observed that integration of heterogeneous data significantly benefits the identification of protein complexes. Moreover, a comprehensive comparison demonstrates that InteHC performs much better than 14 state-of-the-art approaches. All the experimental data and results can be downloaded from http://www.ntu.edu.sg/home/zhengjie/data/InteHC.

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INTRODUCTION

Protein complexes are of great importance for understanding cellular organization and functions. For example, RNA-induced silencing complex (RISC complex)¹ plays a fundamental role in gene regulation by micro RNAs (miRNA) and in defence against viral infections by incorporating one strand of a small interfering RNA (siRNA) or miRNA. Another example is RNA polymerase II complex,² which transcribes genetic information into messages for ribosomes to produce proteins. Although protein complexes are crucial for many cellular processes in cell and molecular biology, they are still largely obtained through small-scale experimental techniques, which are time-consuming and tedious. In addition, many important protein complexes have not been detected by current wet-lab experiments. Therefore, computational methods for predicting protein complexes are highly desired.

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As a large amount of binary protein–protein interaction (PPI) data are now available, it becomes more prevalent to detect protein complexes in PPI networks where nodes are proteins and edges are protein interactions. There is an observation that protein complexes generally correspond to dense subgraphs or cliques in PPI networks.³ On the basis of this observation, a number of computational methods are proposed for identifying protein complexes, such as MCODE,⁴ MCL,⁵ DPClus,⁶ PCP,⁷ IPCA,⁸ COACH,⁹ CMC,¹⁰ and so on.

As two large-scale tandem affinity purification with mass spectrometry (TAP-MS) data were released^{11,12} in 2006, another batch of algorithms^{11–18} were proposed to detect protein complexes from these TAP-MS data. These methods can be classified into the following two categories. The methods in the first category compute the affinity scores between proteins (e.g., the probability of two proteins to be co-complex members) and convert the TAP-MS data to a PPI network based on the calculated affinity scores. Traditional graph clustering methods are then applied to detect protein complexes in the obtained PPI networks. The second category is to model TAP-MS data as bipartite graphs and then detect dense bipartite subgraphs as protein complexes.^{14,17}

Given the fact that PPI data are inherently noisy (e.g., false positives and false negatives), several methods had integrated other non-PPI sources to assess the reliability of PPI data for more accurate detection of protein complexes. For example, the above mentioned PCP⁷ and CMC¹⁰ algorithms utilized the topological weights of interactions as their reliability for detecting protein complexes. DECAFF¹⁹ and STM²⁰ first exploited functional information of proteins (e.g., Gene Ontology annotations) to assess the reliability of PPI data and then detected protein complexes from the refined PPI data. Meanwhile, MATISSE²¹ integrated gene expression data with PPI data to increase the confidence of interactions for the same purpose. More details for the above computational prediction of protein complexes can be found in this survey.²² However, each of the above integration methods generally combines a single data source (e.g., gene expression profiles or functional annotations) with PPI data to refine the quality of the PPI data. Currently, a highly diverse collection of sources for proteins is available, e.g., binary PPI data, gene expression profiles, functional Gene Ontology (GO) terms, TAP-MS data, and so on. Therefore, it is highly motivated to integrate these multiple heterogeneous sources for predicting protein complexes.

Although there are many studies on integrating multiple data sources to predict PPI,^{23–27} integrative methods for predicting complexes are still rather limited. To the best of our knowledge, there are two such existing works as follows. Xia *et al.*²⁶ integrated multiple data sources to create a database named IntNetDB for more confident PPI data and then predicted protein complexes by using



Figure 1

The overall framework of our proposed InteHC method.

MCODE algorithm on IntNetDB. However, they did not predict PPI or protein complexes in the model organism yeast. Another method named CMBI²⁸ integrated PPI data, gene expression profiles and gene essentiality data to predict protein complexes.

To address the above issue, we proposed an approach called InteHC (Integrative Hierarchical Clustering) to predict protein complexes by integrating multiple data sources. The overall framework of InteHC is shown in Figure 1. First, we use individual sources/features to compute the score matrices that store various protein affinities between all the pair-wise proteins. Second, we construct a final score matrix, which is the weighted sum of the matrices from individual features. In particular, these weights, indicating the reliability of each individual data sources, are learned by a supervised model (i.e., a linear ranking SVM) on a known set of positive and negative protein-protein interactions. This final score matrix not only integrates all the individual sources to address the false-negative issue but also takes the reliability of each individual sources into consideration to address the potential false positive issue. Finally, a hierarchical clustering algorithm is then applied to identify protein complexes from the final score matrix. We have conducted comprehensive experiments to evaluate our predicted complexes based on several metrics, such as recall, coverage rate, and accuracy. The comparison with other methods shows that the integration of heterogeneous sources significantly improves the coverage and accuracy for protein complex prediction.

MATERIALS AND METHODS

In this section, we first introduce various data sources used in this study. And then we provide detailed description of our proposed InteHC method as well as the evaluation metrics for predicted protein complexes.

Data source

Our InteHC integrated four data sources for identifying protein complexes, namely, PPI data, Gene Ontology data, gene expression profiles, and AP-MS data. DIP database for PPI data was downloaded from Ref. 29. GO data were downloaded from Ref. 30, while gene expression data were downloaded from Ref. 31. AP-MS data were downloaded from Refs. 11 and ¹². In summary, DIP data consist of 4930 proteins and 17,262 interactions and AP-MS data consist of 6498 purifications involving 2996 bait proteins and 5405 prey proteins.

In addition, 13,424 positive protein–protein interactions for training the linear ranking SVM [please refer to Eq. (4) in Section "Integration of Evidences"] were downloaded from Ref. 32. All these positive interactions are originally collected from the BioGrid database³³ and they are (1) detected by more than three experiments, or (2) in DIP-core,²⁹ or (3) are high-quality interactions in Krogan *et al*'s¹² purification data or Gavin *et al*'s¹¹ data. Meanwhile, we have also generated equal-size negative protein interactions if they do not occur in the positive interaction set. All the aforementioned data used in our experiments can also be downloaded from http:// www.ntu.edu.sg/home/zhengjie/data/InteHC.

Individual data sources for protein affinities

Estimation of the affinity scores between proteins is a crucial step for protein complex prediction.^{11,18,34,35} In the following, we introduce various protein affinities from heterogeneous data sources.

FSweight: A topological weighting in PPI networks

FSweight³⁶ was proposed to estimate the functional similarity between proteins based on their topological properties (i.e., common neighborhood) in PPI networks. In particular, two proteins with more neighbors in common are more likely to share similar functions. In this paper, we used the following simplified variant for FSweight as follows:

$$S_{FS}(p,q) = \frac{2 \times |N_p \cap Nq|}{|N_p - Nq| + 2 \times |N_p \cap Nq| + 1} \times \frac{2 \times |N_p \cap N_q|}{|N_q \cap Nq| + 2 \times |N_p \cap Nq| + 1}$$
(1)

where N_p includes direct neighbors of the protein p as well as p itself and $N_p \cap N_q$ includes the common neighbors between p and q. Thereafter, FS is short for FSweight in PPI networks.

Gene expression profiles

Given a protein pair (p, q), the proteins' propensity to interact can be measured by using the Pearson Correlation Coefficient between their encoded genes' expression profiles G_p and G_q as follows,

$$S_{GE}(p,q) = \frac{\left|\sum_{i=1}^{n} (p_i - \overline{p})(q_i - \overline{q})\right|}{\sqrt{\sum_{i=1}^{n} (p_i - \overline{p})^2} \sqrt{\sum_{i=1}^{n} (q_i - \overline{q})^2}} \quad (2)$$

where *n* is the number of time points for the expression profiles and p_i is the *i*th expression value of protein *p*'s expression profiles. \overline{p} is the average expression value across different time points.

Protein function profiles

Given two proteins p and q annotated by GO terms $\{g_{11}, \ldots, g_{1m}\}$ and $\{g_{21}, \ldots, g_{2n}\}$ respectively, their functional similarity as defined in Ref. 38, $S_{GO}(p, q)$, is calculated as follows.

$$S_{GO}(p,q) = \frac{\sum_{i=1}^{m} \max_{1 \le j \le n} sim(g_{1i}, g_{2j}) + \sum_{i=1}^{n} \max_{1 \le j \le m} sim(g_{1i}, g_{2j})}{m+n}$$
(3)

where $sim(g_{1i}, g_{2j})$ is the semantic similarity between GO terms g_{1i} and g_{2j} defined in Ref. 37. As we know, GO has three sub-ontologies, namely, biological process (BP), molecular function (MF), and cellular component (CC). BP sub-ontology contains the most number of GO terms and is the most informative,^{30,38} and thus we utilize

GO terms in BP sub-ontology to calculate the functional similarities between the pair-wise proteins in this article.

TAP-MS data profiles

With two large-scale TAP-MS data released, ^{11,12} additional computational methods have been proposed to assess the protein affinities based on the purification

tions. In

records, e.g., socio-affinity (SA),¹¹ purification enrichment (PE),³⁴ dice coefficient (DC),³⁵ and C2S scores.¹⁸ These methods generally assume that protein pairs, which occur more frequently in the same purifications (e.g., baitprey and prey-prey relationships), tend to have higher affinity scores. As C2S is the most recently developed scoring method, we finally used the normalized C2S scores $(S_{TAP}(p,q)) = \frac{C25(p,q)-min}{max-min}$, where max and min are the maximal and minimal C2S scores, respectively) to quantify the protein affinities from the TAP-MS data.

Integration of evidences

In the aforementioned subsection, we have introduced how to calculate the protein affinity scores based on four heterogeneous data sources. Now we are ready to integrate four score matrices together. Generally, a typical approach to integrating these affinity scores into a final score matrix is the linear weighted sum. This leads us to address the next question, "How to assign appropriate weights to the respective data sources given that they have different reliability?"

We first collected a set of known positive and negative protein-protein interactions.

Classification models can be learned from these known positive and negative PPIs to predict novel PPIs and assess protein affinities.³² In particular, each interaction can be represented as a feature vector $x_i = (s_{i1}, s_{i2}, \ldots, s_{ik})$ where s_{il} $(1 \leq l \leq k$, where k is the number of data sources and k = 4 in this work) is the affinity score from the l^{th} data source (i.e., S_{FS} , S_{GE} , S_{GO} and S_{TAP}). In addition, a linear ranking support vector machine (SVM)^{39,40} $y = w \cdot x + b$ can be learned from known positive and negative PPIs in Equation (4) by maximizing the area under ROC curve (AUC),

$$\min_{w} \frac{1}{2} ||w^{2}|| + c \sum_{i,j} l(w; (x_{i}, y_{i}, x_{j}, y_{j}))$$
s.t. $l(w; (x_{i}, y_{i}, x_{j}, y_{j})) = I_{y_{i} \neq y_{j}}$

$$\times \max(0, 1 - y_{i}w(x_{i} - x_{j}))$$
(4)

where y_i is the label of the interaction x_i (i.e., $y_i = 1$ means that x_i is a positive interaction and -1 otherwise), $I_{v_i \neq v_i}$ is the indicator function and *c* is a parameter.

Now let us focus on the loss function in Equation (4). Given that two interactions x_i and x_j have different labels, we assume that x_i is positive (i.e., $y_i = 1$) and x_j is negative (i.e., $y_i = -1$) without loss of generality. In this case, it is obvious that ranking x_i before x_i (i.e., $w \cdot x_i > w \cdot x_i$) will have a larger loss than ranking x_i before x_i . In other words, the linear ranking SVM in Eq. (4) will try to rank a positive interaction x_i before a negative interaction x_i to reduce the loss and maximize the AUC. With the weight vector $w = (w_1, \dots, w_k)$ learned from the known positive and negative PPIs, we can calculate the final affinity score for a protein pair (p, q), S(p, q) in Equation (5).

$$(x_j, y_j))$$

scores are more likely to be true protein-protein interac-
tions. In addition, the affinity scores in the matrix
$$S$$
 in
Eq. (5) are able to estimate the quality of protein inter-
actions more accurately than those from individual data

Obviously, protein pairs with higher final affinity

sources, because they combine multiple data sources and

also take the reliability of each source into account.

(5)

 $S(p, q) = \sum_{i=1}^{4} w_i \times S_{E_i}(p, q), \text{ where}$

 $E_i \in \{FS, GE, GO, TAP\}$

Hierarchical clustering for protein complexes

The hierarchical clustering algorithm is applied to detect protein complexes on the final score matrix in Eq. (5). First, it considers all singleton proteins as initial clusters. Second, it iteratively merges two clusters with the highest similarity in each iteration. The algorithm terminates when the quality of the detected complexes in the merging process has become maximal. The detailed procedure for the hierarchical clustering is illustrated in Algorithm 1. Given two clusters c_i and c_j , their similarity, $sim(c_i, c_i)$, is the average affinity score of all

Algorithm 1

Hierarchical Clustering for Protein Complexes Input: L, the set of proteins in a given organism (e.g., yeast); S, affinity scores between proteins in L. Output: C, the set of predicted protein complexes.

- 1: $C = \{\{p\} | p \in L\}; //$ Initialization
- 2: while(QualityFunction is maximal)
- 3: $(c_i^*, c_j^*) = \arg \max_{c_i, c_j} sim(c_i, c_j) //$ Finding two most similar clusters
- $c_{merge} = c_i^* \cup c_j^* //$ Merging two clusters 4:
- $C = C + \{c_{merge}\} \{c_i^*\} \{c_i^*\}$ // Removing two 5: original clusters
- 6: for each $c_k \in C$ 7: $sim(c_k, c_{merge}) = \frac{|c_i^*| \times sim(c_k, c_i^*) + |c_j^*| \times sim(c_k, c_j^*)}{|c_i^*| \times |c_j^*|} //$ Updating the similarity scores
- 8: end for
- 9: end while

the protein pairs between these two clusters as shown in Eq. (6). Next, we will introduce the quality function in Line 2.

$$sim(c_i, c_j) = \frac{1}{|c_i| \times |c_j|} \sum_{p \in c_i, q \in c_j} S(p, q)$$
(6)

Given a cluster c_i , we define its confidence score as the average affinity score of all possible protein pairs within shown in Eq. (7). Given a clustering it as

 $C = \{c_1, \dots, c_n\}$, we proposed the following two functions to measure the quality of this clustering *C* based on the affinity scores for protein pairs within these clusters. The first quality function $Q_1(C)$ in Eq. (8) is the average confidence score for all the clusters while the second quality function $Q_2(C)$ in Eq. (9) is the average affinity score for all the protein pairs within the clusters.

$$Conf(c_i) = \frac{\sum_{p,q \in c_i} S(p,q)}{|c_i| \times (|c_i| - 1)}$$
(7)

$$Q_1(C) = \frac{\sum_{i=1}^{n} Conf(c_i)}{n}$$
(8)

$$Q_2(C) = \frac{\sum_{i=1}^{n} \sum_{p,q \in c_i} S(p,q)}{\sum_{i=1}^{n} |c_i| \times (|c_i| - 1)}$$
(9)

In general, $Q_2(C)$ will be dominated by those large clusters as they contain many more protein pairs than those small clusters. Therefore, we can upgrade $Q_2(C)$ to the third quality function $Q_3(C)$ in Eq. (10) by dividing a factor (i.e., the square root of the cluster size, $\sqrt{|c_i|}$) in both denominator and numerator to eliminate the bias from those large clusters (this factor has also been utilized to eliminate the bias of cluster size in a previous study⁴¹). As such, we have three functions to evaluate the quality of a set of clusters.

$$Q_{3}(C) = \frac{\sum_{i=1}^{n} \frac{1}{\sqrt{c_{i}}} \sum_{p,q \in c_{i}} S(p,q)}{\sum_{i=1}^{n} \sqrt{|c_{i}|} \times (|c_{i}|-1)}$$
(10)

In fact, the hierarchical clustering here does not stop, i.e., it runs from the start with all individual proteins as clusters to the end with all the proteins as a whole cluster. During this process, the values for a given quality function can be monitored at each iteration. Finally, we output the clusters in a certain iteration where the given quality function achieves the highest value.

Evaluation metrics

We will utilize the sensitivity (Sn, i.e., the coverage rate) and positive predictive value (PPV) to evaluate the quality of predicted protein complexes. In particular, the Sensitivity and revised PPV between a benchmark complex b_i and a predicted complex c_j are defined in Eq. (11).^{9,42}. Accuracy is the geometric mean of sensitivity and PPV.

$$Sn = \frac{\sum_{i} \max_{j} T_{i,j}}{\sum_{i} |b_{i}|}, \quad PPV = \frac{\sum_{j} \max_{i} T_{i,j}}{\sum_{j} |\bigcup_{i} (b_{i} \cap c_{j})|} \quad (11)$$

and Accuracy = $\sqrt{Sn \times PPV}$

where $T_{i,j}$ is the number of proteins shared by b_i and c_j , i.e., $|b_i \cap c_j|$. We also use another metric, Recall, to evalu-

ate the predicted complexes. Given a predicted complex c and a real complex b, their neighborhood affinity score, NA(c, b) in Eq. (12), can be used to determine how well they match with each other. We consider them to be matching if $NA(c, b) \ge \omega$ (ω is a parameter which is normally set as 0.2 in previous studies^{4,6,9,19} and we also set it as 0.2 in our experiments). Recall is defined in Eq. (13) as the fraction of benchmark complexes that are matched by at least one predicted complex. In Eq. (13), B is the set of benchmark complexes. Here, we used CYC2008 catalog⁴³ as benchmark protein complexes for calculating both Accuracy and Recall.

$$NA(c, b) = \frac{|c \cap b|^2}{|c| \times |b|} , \qquad (12)$$

$$\text{Recall} = \frac{|\{b|b \in B, \exists c \in C, NA(c, b) \ge \omega\}|}{|B|}$$
(13)

With the Accuracy and Recall defined earlier, their average and harmonic average (*HAverage*) in Eq. (14) are utilized to evaluate the overall quality of predicted complexes. Now we have a comprehensive set of evaluation metrics for the quality of predicted complexes. For example, the sensitivity in Eq. (11) shows how many proteins in benchmark complexes are covered by the predicted complexes and thus measures protein-level coverage of predicted complexes. Meanwhile, Recall measures complex-level coverage of predicted complexes.

$$Average = \frac{Accuracy + Recall}{2} ,$$
$$HAverage = \frac{2 \times Accuracy \times Recall}{Accuracy + Recall} .$$
(14)

RESULTS

In this section, we demonstrate a comprehensive comparison between InteHC and various state-of-the-art approaches for predicting protein complexes. Moreover, we also introduce the experimental setting for a common parameter involved in ranking SVM as well as our choice for quality functions.

Linear ranking SVM

SVM^{*perf*³⁹} is utilized in this article to solve the optimization problem in Eq. (4). The parameter c in Eq. (4) is the trade-off between training error and margin. In general, the classifier (i.e., SVM) becomes more and more over-fitting and the generalization error increases as cbecomes larger. Therefore, an appropriate c should not be too large while maintaining a good ROC area (AUC).

To select an appropriate value for *c*, we performed five-folder cross-validation on 26,848 positive and negative protein interactions. Figure 2 shows the average



Figure 2 The average ROC area (AUC) of SVM as the parameter *c* increases.

AUC of our trained SVM using different values for c. We can observe that the AUC becomes more and more stable as c increases. We finally set c as 15 in our experiments as it satisfies the above two requirements—c is relatively small and its AUC (0.879) is also good. After learning, the weights for four data sources (FS, GO, GE, and TAP) in Eq. (5) are 8.49, 2.25, 0.84, and 2.89 respectively. On the basis of these learned weights for different biological sources, we are able to compute the final score matrix using Eq. (5).

Quality functions for a clustering

Once we get the final score matrix, the hierarchical clustering will run on it and output the predicted clusters as protein complexes when a specific quality function for the clustering is maximized. As such, the hierarchical clustering may predict different sets of protein complexes when we utilize different quality functions. Next, we will show the Recall and Accuracy of protein complexes predicted by our proposed quality functions.

Note that the aforementioned three quality functions for clusters are based solely on the properties of these clusters themselves (e.g., the affinity scores of protein pairs within clusters and the cluster size). Meanwhile, we can have another quality function for clusters, which is the sum of Recall and Accuracy collected by mapping them to known protein complexes (e.g., CYC2008 complexes). This quality function will output the optimal set of complexes which achieves higher sum of Recall and Accuracy. We thus denote it as Q^* and utilize it to verify the goodness of the quality functions Q_1 , Q_2 and Q_3 . As shown in Figure 3, it is obvious that the clusters gener-



Figure 3

The Recall and Accuracy of the complexes predicted by InteHC using various quality functions.

ated by the quality function Q_3 achieve higher Recall and Accuracy than those by Q_1 and Q_2 . Q_3 is also much closer to the optimal solution Q^* than Q_1 and Q_2 . This indicates that the quality of a set of clusters is well evaluated by considering the cluster size in a reasonable manner in our Equation (10). Thereafter, we will only show the results generated by Q_3 for comparisons.

The benefit of data integration

To demonstrate the power of data integration for identifying protein complexes, we will show the performance of hierarchical clustering algorithm using each individual data sources, as well as the combined set of all features in Figure 4.



Figure 4

The Recall and Accuracy of complexes predicted by the hierarchical clustering using individual data sources.

For the results using single data sources, we can observe that PPI network with FSweight (FS) covers the most benchmark complexes and achieves the highest Recall (0.708), demonstrating that traditional PPI networks are indeed informative and reliable for protein complex detection. GO can also achieve a good Recall (0.679). Meanwhile, complexes generated from TAP data have the highest sensitivity (0.679) and accuracy (0.752), indicating that TAP data are also a high quality source for predicting protein complexes.^{11,12}. However, it is obvious that FS has a low Accuracy while TAP has a relatively low Recall as shown in Figure 4, indicating that using individual sources alone will not produce very good results.

In Figure 4, it is demonstrated that data integration achieves a higher overall performance than using individual data sources, illustrating that our proposed InteHC can effectively integrate multiple sources for protein complex prediction. As mentioned above, FS has a high Recall and a low Accuracy while TAP has a low Recall and a high Accuracy. Here, InteHC manages to overcome the limitations of individual sources. In particular, InteHC has a comparable Recall with FS and has a significantly higher Accuracy than FS (0.769 vs. 0.627). Similarly, it has much higher Recall than TAP (0.711 vs. 0.625).

For completeness of our evaluation, we have 15 combinations with four data sources available (FS, GE, GO, and TAP). The performance of hierarchical clustering algorithm using all these 15 combinations is shown in Supporting Information Figure S1. For example, the combinations FS+GO and FS+GO+TAP also achieve high overall performance. This indicates once again that data integration indeed benefits the accurate prediction of protein complexes.

Comparison with methods on PPI networks

In this subsection, we compare InteHC with eight existing state-of-the-art approaches that detect protein complexes from PPI networks (e.g., the DIP data in our experiments), which include MCODE,⁴ MCL,⁵ DPClus,⁶ IPCA,⁸ DECAFF,¹⁹ COACH,⁹ HC-PIN,⁴⁴ and Pro-Rank.^{45,46} We also show the results of our hierarchical clustering algorithm on DIP data using FSweight, denoted as InteHC-FS.

As shown in Table I, InteHC has the highest coverage in both protein-complex level (i.e., Recall) and proteinlevel (i.e., sensitivity). Therefore, from the perspective of covering the known protein complexes, InteHC achieves the highest performance against all these methods designed for PPI networks. InteHC also achieves an Accuracy 76.9%, which is 11.5% higher than the second best method (DPClus, 65.4%). In addition, InteHC-FS achieves a high Recall while a low sensitivity, indicating that InteHC effectively improve the coverage of proteins

Table I

Comparisons between InteHC and Various Existing Methods on PPI Data

Methods	No. of complexes	Recall	Sn	PPV	Accuracy
MCODE	182	0.240	0.403	0.624	0.501
MCL	1116	0.586	0.541	0.763	0.643
DPClus	1140	0.642	0.511	0.836	0.654
IPCA	1242	0.539	0.501	0.766	0.620
DECAFF	2190	0.525	0.454	0.795	0.601
COACH	746	0.527	0.545	0.698	0.617
HC-PIN	99	0.169	0.573	0.347	0.446
HC-wPIN	147	0.289	0.516	0.673	0.589
ProRank	110	0.162	0.199	0.897	0.422
InteHC-FS	1307	0.708	0.434	0.906	0.627
InteHC	860	0.711	0.701	0.845	0.769

in known protein complexes by integrating other data sources.

Note that HC-PIN⁴⁴ is also a hierarchical clustering framework for detecting protein complexes. In Table I, HC-PIN and HC-wPIN represent the complexes predicted from unweighted and weighted DIP data, respectively. Here, FSweight³⁶ is utilized to compute the weights for protein interactions in HC-wPIN. We can observe that HC-PIN predicts a small number of complexes and achieves low Recall and Accuracy on both unweighted and weighted DIP data (similarly, a recent method ProRank^{45,46} also generates a small number of complexes). However, we must point out that HC-PIN has many good merits. For example, it has a low computational complexity. In addition, HC-PIN has a high fraction of predicted complexes (e.g., 86 of 147 complexes predicted by HC-wPIN) that can match with known complexes. ProRank is also similar to HC-PIN with 58 of 110 complexes matching with known complexes.

Comparison with methods on TAP-MS data

Here, we compare InteHC with five existing methods proposed for TAP-MS data, including C2S,¹⁸ CACHET,¹⁷ BT,¹³ Pu,¹⁶ and Hart.¹⁵ Similar to InteHC-FS in Table I, InteHC-TAP in Table II represents the set

Table II

Comparisons between InteHC and Various Existing Methods on TAP Data

Methods	No. of complexes	Recall	Sn	PPV	Accuracy
C2S CACHET	1035 449	0.630 0.515	0.676	0.847	0.757 0.665
B I Pu Hart	409 400 390	0.598 0.591 0.593	0.629 0.691	0.848 0.789 0.863	0.730 0.738 0.725
InteHC-TAP InteHC	974 860	0.623 0.711	0.679 0.701	0.833 0.845	0.752 0.769



Figure 5

Two examples of known protein complexes.

of complexes predicted by our hierarchical algorithm on TAP data.

As shown in Table II, InteHC achieves the highest Recall, sensitivity, and Accuracy. For example, InteHC has a Recall 71.1%, which is 8.1% higher than the second best method (C2S, 63.0%). Also, InteHC-TAP and C2S achieve very similar results. In fact, InteHC-TAP and C2S are regarded as operating on the same score matrix and have different forms of stop criteria for the hierarchical clustering. Meanwhile, C2S¹⁸ was demonstrated to have an effective stop criteria for the hierarchical clustering (however, it cannot be used in the framework of our InteHC) and outperformed existing methods including BT,¹³ Pu,¹⁶ and Hart.¹⁵ Therefore, we can confidently claim that our quality function in Eq. (10) also provides us with an effective stop condition for the hierarchical clustering.

Comparison with other integrative methods

Xia *et al.*²⁶ constructed a database named IntNetDB for PPI data predicted by integrating multiple data sources. They then predicted protein complexes by using MCODE algorithm on IntNetDB. However, they did not predict PPI or protein complexes for the model organism yeast. Therefore, we are not able to compare MCODE on IntNetDB with InteHC.

Another method named CMBI²⁸ was recently proposed to detect protein complexes by integrating multiple biological resources including PPI data, gene expression profiles and essential protein information. Essential proteins are first selected as seeds. Proteins, which have high topological similarity and gene expression correlation with those seeds, will be included to form protein complexes. In this subsection, we will briefly compare the results between InteHC and CMBI.

Using the data described in Ref. 28, CMBI predicted 760 protein complexes. 160 of 408 benchmark complexes will be covered, that is, CMBI has a Recall 0.392. The sensitivity, PPV, and Accuracy of CMBI are 0.508, 0.482, and 0.495, respectively. Using the same PPI and gene expression data, InteHC predicted 1469 protein complexes and achieves a Recall 0.720, sensitivity 0.464, PPV 0.900, and Accuracy 0.646. Thus, InteHC performs significantly better than CMBI in terms of these measures. As shown in Supp. Info. Table S1, different data sources have different importance for estimating protein affinities and predicting protein complexes. For example, the PPI data (FS scores) have the highest importance based on the leave-one-out AUC while gene expression profiles have the lowest. Correspondingly, our supervised model [i.e., the ranking SVM in Eq. (4)] assigns each data source with different weights, e.g., FS scores have the highest weight 8.49 while the gene expression profiles have the lowest 0.84.47 However, CMBI treats equally the topological weights (ECC, edge clustering coefficient) and gene expression correlations and thus overrates the importance of the gene expression profiles. This may be one of the reasons why InteHC performs better than CMBI.

Protein complexes more accurately detected by InteHC

In previous sections, we demonstrated that InteHC outperformed 14 methods in terms of various evaluation measures (e.g., Recall and Accuracy). Next, we introduce two example protein complexes that are more accurately detected by InteHC.

In Figure 5(A), the CCR4-NOT complex⁴⁸ in CYC2008 catalog has nine proteins and it is involved in

	CCI	R4-NOT complex (9 protein	s)	TFIID complex (15 proteins)			
Methods	NA score	Predicted size	Overlap	NA score	Predicted size	Overlap	
MCODE	0.790	9	8	0.152	11	5	
MCL	0.397	7	5	0.3	8	6	
DPClus	0.9	10	9	0.491	11	9	
IPCA	0.9	10	9	0.467	7	7	
DECAFF	1	9	9	0.474	9	8	
COACH	0.818	11	9	0.533	8	8	
HC-PIN	0.0316	285	9	0.0395	285	13	
HC-wPIN	0.692	13	9	0.436	22	12	
ProRank	0.037	3	1	0.267	4	4	
C2S	0.557	5	5	0.29	23	10	
CACHET	0.557	5	5	0.738	13	12	
BT	0.667	6	6	0.384	25	12	
Pu	0.571	7	6	0.369	26	12	
Hart	0.397	7	5	0.336	24	11	
CMBI	1	9	9	0.672	12	11	
InteHC-FS	0.711	10	8	0.333	5	5	
InteHC-GO	0.3333	3	3	0.15	4	3	
InteHC-GE	0.056	2	1	0.0333	2	1	
InteHC-TAP	0.556	5	5	0.29	23	10	
InteHC	1	9	9	0.8	12	12	

Table III								
The Mapped	Complexes	for CCR4	-NOT an	d TFIID	Complexes	Predicted	by Various	Methods

several aspects of mRNA metabolism, including repression and activation of mRNA initiation, control of mRNA elongation, deadenylation, and degradation. The complex (ID: 17) predicted by InteHC also has nine proteins and it can exactly match with the CCR4-NOT complex, that is., covering all the nine proteins in the CCR4-NOT complex. In Figure5(B), the known transcription factor TFIID complex⁴⁹ in CYC2008 catalogue contains 15 proteins, while the complex (ID: 186) predicted by InteHC has 12 proteins. All these 12 proteins (black circles) are involved in the benchmark complex while three remaining proteins (white circles) are not covered by the complex 186. More examples can be found in the Supporting Information.

From the set of complexes predicted by a specific approach (e.g., InteHC), we may pick up a complex which has the highest similarity score [i.e., NA score in Eq. (12)] to a given benchmark complex and denote it as the mapped complex for this benchmark complex. For instance, the complex 186 predicted by InteHC is the mapped complex for the TFIID complex. Table III shows the mapped complexes for CCR4-NOT complex and TFIID complex predicted by various methods. Take the TFIID complex for example, its mapped complex predicted by InteHC (i.e., the complex 186) covers 12 of its 15 proteins and has the highest NA score 0.8, while its mapped complexes predicted by CACHET, COACH, DPClus, and IPCA have NA scores 0.738, 0.533, 0.491, and 0.471, respectively. Meanwhile, the complex 17 predicted by InteHC is exactly the benchmark CCR4-NOT complex and has the optimal NA score (i.e., 1.0). Two example complexes more accurately detected by InteHC

show that a proper integration of multiple data sources indeed improves the computational prediction of protein complexes.

CONCLUSIONS AND DISCUSSIONS

In this article, we have proposed an integrative approach (InteHC, Integrative Hierarchical Clustering) for identifying protein complexes from heterogeneous sources, including PPI data, gene expression profiles, GO terms, and TAP-MS data. For each individual sources/ features, we calculated the affinities between proteins to show their propensity to interact. Subsequently, a supervised model (i.e., the ranking SVM) was utilized to learn the weight for each feature. The weighted sum of affinity scores from individual features resulted in a final score matrix. A hierarchical clustering algorithm on this final score matrix will generate a set of non-overlapping clusters as predicted protein complexes. In particular, we proposed a novel quality function [i.e., Q3 in Eq. (10)] for capturing the high-quality protein complexes. Experimental comparisons have shown that InteHC performs much better than 14 existing approaches in terms of several evaluation metrics, e.g., InteHC significantly improves the coverage for known protein complexes in both a protein-level (Recall) and complex-level (sensitivity). In addition, our InteHC is a flexible and generic framework to integrate multiple data sources for predicting protein complexes. It allows us to include new data sources by simple matrix operations to achieve even better results.

As discussed in our previous study,¹⁸ we plan to design new data-integration mechanisms for identifying protein complexes. For this purpose, we already managed to integrate multiple TAP-MS datasets in Ref. 18 and various heterogeneous datasets in this article. These techniques are characterized as raw dataset-level integration. In future work, we will focus on another kind of integration-result-level integration. A specific method for detecting protein complexes working on a specific dataset will predict a set of protein complexes. We can thus collect many sets of complexes by applying various methods on multiple datasets. We may expect to achieve higher prediction accuracy by analyzing and processing those integrated complexes. Furthermore, we plan to set up a platform or web-server to demonstrate all the above integration techniques for identifying protein complexes.

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