### Methods

**Predicting Protein Complex Membership Using Probabilistic Network Reliability**

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Evidence for specific protein–protein interactions is increasingly available from both small- and large-scale studies, and can be viewed as a network. It has previously been noted that errors are frequent among large-scale studies, and that error frequency depends on the large-scale method used. Despite knowledge of the error-prone nature of interaction evidence, edges (connections) in this network are typically viewed as either present or absent. However, use of a probabilistic network that considers quantity and quality of supporting evidence should improve inference derived from protein networks. Here we demonstrate inference of membership in a partially known protein complex by using a probabilistic network model and an algorithm previously used to evaluate reliability in communication networks.

[Supplemental material is available online at www.genome.org. Software called Complexpander, which predicts new members of partially known protein complexes, is available at the authors’ Web site http://llama.med.harvard.edu/Software.html.]

In the past few years, much effort has gone into developing and applying methods for discovering the complete set of protein–protein interactions in an organism—the interactome. The results of both large- and small-scale studies have been collected into databases (Hodges et al. 1999; Mewes et al. 2002; Xenarios et al. 2002; Bader et al. 2003), and it is apparent that the number of interactions derived from small-scale methods, such as affinity chromatography and cosedimentation, is now low compared with those derived from high-throughput techniques such as large-scale yeast two-hybrid (Y2H) screens (Uetz et al. 2000; Ito et al. 2001) and affinity purification with mass spectrometric identification (APMS) methods (Gavin et al. 2002; Ho et al. 2002).

Although these large-scale methods offer vast improvements in efficiency for identifying protein interactions, they are also prone to higher error rates than are conventional small-scale studies, and some large-scale methods are more reliable than others (Mrowka et al. 2001; Deane et al. 2002; Edwards et al. 2002; von Mering et al. 2002; Deng et al. 2003; Sprinzak et al. 2003). Notwithstanding high error rates, it seems likely that high-throughput studies will continue to be the largest source of information about protein–protein interactions.

Analyses of protein interaction evidence have generally treated interaction evidence as binary. By this, we mean that interaction data are often represented as a network in which edges are either present or absent based on the existence of supporting evidence. Conclusions are then drawn from this network without accounting for the quality or quantity of evidence supporting each interaction. This binary approach has been used previously in predicting protein function (Marcotte et al. 1999; Schwikowski et al. 2000) in predicting co-complexed proteins (Bader and Hogue 2003) and protein interaction networks (Marcotte et al. 1999; Schwikowski et al. 2003). We expected that information about quality and quantity of evidence lost in constructing a binary network could prove useful in improving these predictions.

We used a probabilistic model for protein interaction networks that accounts for quality of supporting evidence for each interaction. In this model, each pair of proteins with supporting evidence of interaction was assigned a weight, representing the probability that the two proteins interact directly and stably. This binary representation has been used with some success to predict protein complexes (Jansen et al. 2002; Bader and Hogue 2003; Bu et al. 2003; Rives and Galitski 2003) and protein function (Marcotte et al. 1999; Schwikowski et al. 2000). Given a collection of error-prone protein interaction data, how might we obtain a list of candidate proteins, ranked by probability of membership in a partially known protein complex?
We applied an algorithm that uses this probabilistic network model to generate a list of candidate proteins ranked according to the probability that each “connects” to one or more proteins in a query list—the core protein set. Here a protein is said to connect to the core if there is at least one path of direct and stable interactions between it and one of the core proteins. The algorithm we developed uses a large collection of networks, in which each edge is a binary interaction with existence that is determined by a Bernoulli trial using the probability from the corresponding edge in the probabilistic network. The probability that a given candidate protein connects to the core set is estimated as the fraction of sampled networks that contains a path connecting the candidate to the core. A probabilistic network and two sampled binary networks are illustrated in Figure 1.

Software called Complexpander, which implements this algorithm, was developed, allowing a user to submit a core set of proteins and obtain additional candidate members of this core complex using an evaluation of 10,000 sampled networks. In addition to returning a list of candidate proteins ranked by probability of connecting to the core complex, the software returns a graph representing the interaction neighborhood of the core set of proteins. Visualization of such graphs using Pajek (Batagelj and Mrvar 1998) is shown in Figure 2 for four example queries, each using a protein complex from the MIPS catalog (Mewes et al. 2002) as the “core” protein set. For each query, we used the MIPS complex catalog database as of June 20, 2003.

Results of a query with the SAGA complex (according to MIPS) are shown in Figure 2A. The top-ranked three candidate proteins returned (most reliably connected to core) were the TATA-binding protein associated factors Taf19, Taf145, and Taf40. Although none of these proteins are currently thought to be members of the SAGA complex, they are each members of the TFIID and the TBP-associated TAF[II] complexes, each of which has substantial overlap with the SAGA complex (Hodges et al. 1999). The fourth-ranked protein is Sgf29, a known member of the SAGA complex (Sanders et al. 2002), despite not being listed in MIPS. Other known members of the SAGA complex that are not listed in MIPS were also retrieved in the top 50: Sgf73 (Sanders et al. 2002), ranked 12th; Hfi1 (Grant et al. 1998), ranked 31st; and Ubp8 (Sanders et al. 2002), ranked 39th.

Results of a query with the NOT (negative on TATA) complex, as defined in the MIPS catalog, is shown in Figure 2B. The NOT complex was originally identified as a global negative regulator of transcription (Collart and Struhl 1994; Oberholzer and Collart 1998). Our query retrieved Ccr4 (ranked first) and Pop2 (ranked third), both known members of this complex—now more commonly called the Ccr4-NOT complex (Tucker et al. 2002)—that were not listed as NOT members in MIPS.

Results of a query with the replication factor C (RFC) complex, as defined by MIPS, are shown in Figure 2C. RFC is a “clamp loader” for the Proliferating Cell Nuclear Antigen (PCNA) replication processivity factor (Cullmann et al. 1995; Zhang et al. 1999) and is normally formed from proteins Rfc1-5 (the query set). Ctf18, ranked second, is an Rfc1-like protein that has been suggested to complex with Rfc2-5 (Hanna et al. 2001). Elg1, ranked third, participates in an alternate RFC with Rfc2-5 that has a role in DNA damage repair (Bellouai et al. 2003). Rad24, ranked 12th, is also known to form an alternative complex with Rfc2-5 (Majka and Burgers 2003).

As a final example, the results of a query with the Arp2/Arp3 complex, integrally involved in actin polymerization, are shown in Figure 2D. The top-ranked candidate protein was Arc40, a known member of the Arp2/Arp3 complex not listed in MIPS (Winter et al. 1999).

To rigorously assess the performance of our software in predicting new complex members, we examined a validation set of 27 complexes from the MIPS database of yeast protein complexes (see Methods). For each complex in the validation set, we left each protein out in turn and used the remaining proteins as the core set in a trial query for new complex members. The fraction of trial queries in which the target protein was found above a threshold rank R was assessed.

We wished to compare the performance of our “probabilistic network” (ProNet) method to an alternative method that uses a binary protein interaction network, in which an edge between two proteins exists if there is any evidence supporting their interaction. As no algorithm for ranking candidate members in a partially known protein complex was known to us (with one exception discussed below), we developed the “shortest path with evidence” (SPE) method. In the SPE method, the ranking of candidate protein members is simply by length of the shortest path through the binary network to one of the core proteins, with ties broken randomly.

The results for both ProNet and SPE methods are shown in Figure 3. In this graph, the fraction of correctly returned proteins above threshold rank R is plotted as a function of R. Because only one protein was left out of the known complex for each query, the lowest fraction of correctly returned proteins that any algorithm could hope to achieve is 1/R. As shown in Figure 3, the ProNet method outperformed the SPE method at every choice of threshold rank.

Despite the success of our approach, there are several issues that might be addressed in the future to improve identification of new protein complex members. It should be noted that in our methodology for predicting protein complex membership, we assume that complex membership is transitive across edges, that is, if protein A and B are co-complexed and B and C are co-complexed, then A and C are co-complexed. This is not necessarily the case, because there can be alternative complexes in which either A or C (but not both) are bound to B. However, it is not clear how or if such information can be extracted from currently available protein interaction information, and our model did not account for potential intransitivity of complex membership.

In developing our probabilistic network model, we made the simplifying assumption that the types of evidence we used may be treated as independent. This is an oversimplification, because biases inherent in one of the Y2H methods are likely to be present in the other. A similar statement applies to the two APMS methods. In addition, we assume that a given method treats all
classes of protein interactions identically; in fact, a particular method may have reporting biases, for example, against interactions involving membrane proteins. Despite these potential weaknesses, we have shown that our approach outperforms an alternative binary network-based approach. Improved models of dependence between interaction evidence types, and the use of additional evidence types (e.g., Jansen et al. 2003), should improve the performance of the network reliability approach described here.

During preparation of this manuscript, we became aware of a Markov random field (MRF) approach that predicts new members of partially known functional categories or protein complexes (Letovsky and Kasif 2003). This approach, as implemented, was unsuitable for our task because it does not accommodate multiple evidence types with varying reliability and more importantly because it requires that probability models be retrained for each query complex. A small or moderately sized complex provides few positive examples for training, and a query of one protein would provide none. However, an MRF approach with an alternative scheme for learning probability models is promising and merits further study.

In summary, we applied a network reliability approach previously applied in the field of communications theory to a probabilistic protein network model in order to predict new members of protein complexes. Furthermore, we showed that our probabilistic network approach was better than an alternative approach based on binary protein interactions.

### METHODS

The probability that a given candidate protein is in the same protein complex with a known core set of proteins may be expressed (to first approximation) as the probability that there exists a path of direct and stable protein interactions between that candidate and some member of the known complex. How can we estimate this probability? This problem is analogous to one previously considered in the theory of communications networks: the two-terminal network reliability problem (Colbourn 1987). In this case, one wishes to assess the probability that some path of functioning wires connects two terminals at any given time, given a graph of connections, each weighted by the probability that the corresponding wire is functioning at any given time. Although an exact solution to the two-terminal network reliability problem has been shown to be NP-hard (Valiant 1979), reliability can be approximated using Monte Carlo simulation and other approaches (Karger 1999). Here, we apply a Monte Carlo approach to the two-terminal network reliability problem to protein interaction networks.

A prerequisite to this approach is the existence of a weighted graph model of the protein interaction network, in which nodes
represents proteins and edges represent protein pairs that might stably interact. Edge weights represent the posterior probability that the two proteins interact stably and directly given available evidence. Assigning probability of interaction to protein pairs has been previously described (Edwards et al. 2002), and we follow a somewhat similar approach in constructing a probabilistic network model.

We first assembled a database of yeast protein interactions. Our interactome database consisted of the following four evidence types: (1 and 2) two high-throughput studies based on Y2H methods (Uetz et al. 2000; Ito et al. 2001); and (3 and 4) two high-throughput studies based on APMS methods (Gavin et al. 2002; Ho et al. 2002). For one of the Y2H studies (Ito et al. 2001), we used only interactions for which three or more “hits” have been observed (Ito et al. 2001); for both APMS studies, we only considered bait-to-prey interactions because these are thought to be more reliable (Bader and Hogue 2002).

The posterior probability \( P^* \) that a pair of proteins interact directly and stably (i.e., physically contact one another and are contained within the same protein complex) can be calculated according to Bayes’ rule:

\[
P^* = p(y = 1 | z) = \frac{\prod_{i=1}^{T} \left[ p(z_i | y = 1) \cdot p(y = 1) \right]}{\sum_{y=0,1} \left[ \prod_{i=1}^{T} \left[ p(z_i | y = 0) \cdot p(y = 0) \right] \right]}
\]

Here \( y \) is an indicator for interaction (\( y = 1 \) if the pair of proteins interacts directly and stably, and \( y = 0 \) otherwise), \( z \) is a vector of indicator variables for the presence or absence of evidence of each available type, and \( T \) is the number of evidence types. For example, \( z = (0,0,1,0) \) represents a pair of proteins for which evidence of interaction is derived only from the APMS method of Gavin et al. (2002). The equation above makes the naive Bayes assumption that different evidence types are independent of one another given the truth about stable interaction (discussed further below).

Unfortunately, there is no widely accepted and precise definition of what constitutes a stable interaction, or in fact what constitutes a protein complex. Furthermore, there exists no unbiased reference set of protein pairs for which the truth or falsehood of stable and direct interaction is known from which we might extract \( p(y) \), the prior probability of interaction, and \( p(z_i | y) \), likelihoods corresponding to each evidence type \( i \). Below, we make a reasonable estimate of \( p(y) \) and determine values for \( p(z_i | y) \) that optimize performance of our algorithm according to a training set of protein complexes. No protein in the training set of complexes is used subsequently for testing purposes.

We model the prior probability that two arbitrary proteins stably interact, \( p(y = 1) \), according to \( p(y = 1) = p(c = 1 | d = 1) = p(d = 1 | c = 1) \cdot p(d = 1) \), where \( d \) is an indicator variable for whether the two proteins interact directly, and \( c \) is an indicator variable for whether the two proteins are co-complexed. To calculate \( p(d = 1) \), we divide a previous estimate of the number of protein interactions in yeast (von Mering et al. 2002) by the number of possible \( (2 \times 10^7) \). We calculate the probability \( p(c = 1 | d = 1) \) by determining what fraction of those interactions supported by conventional studies (small-scale, non–high-throughput studies) are members of the same complex according to the MIPS complex catalog (Mewes et al. 2002). This resulted in a prior \( p(y = 1) \) probability of \( 7 \times 10^{-3} \), with \( p(y = 0) \) calculated according to \( p(y = 0) = 1 - p(y = 1) \).

The likelihood \( p(z_i | y) \) for each evidence type \( i \) is written in terms of prior probability \( p(y = 1) \), evidence type-specific error rate \( E_i = p(y = 0 | z_i = 1) \), and the fraction of gene pairs \( F_i = p(z_i = 1) \) with supporting evidence of type \( i \) as follows:

\[
p(z_i = 1 | y = 1) = \frac{(1 - E_i) \cdot F_i}{p(y = 1)},
\]

\[
p(z_i = 1 | y = 0) = \frac{E_i \cdot F_i}{p(y = 0)}.
\]

and

\[
p(z_i = 0 | y = 0) = 1 - p(z_i = 1 | y = 0).
\]

Each \( F_i \) value is calculated directly from a count of the number of edges supported by each evidence type, and each \( E_i \) is estimated using an optimization procedure described below at the end of the Methods section.

With estimates of \( p(y) \) and \( p(z_i | y) \) in hand, we can calculate \( P^* \) for each edge. To reduce subsequent computation costs, we set edge weights to zero for those edges with no supporting evidence of interaction. From this probabilistic network model, we sample many binary networks by using a Monte Carlo approach. Each sampled network represents a possible state of the actual network of stably interacting proteins. Sampled networks are obtained according to a Bernoulli trial on each edge, with probability of success equal to edge weight. Figure 1 illustrates a probabilistic network and two sampled binary networks. We sampled 10,000 networks for all results described here.

We count the fraction \( C_i \) of sampled binary networks in which a path exists from a candidate protein \( i \) to any protein in the core complex. The fraction \( C_i \) is our estimate of the probability that the protein \( i \) is a member of the core complex. Determining the existence of a path between two nodes in a sampled network reduces to a breadth-first search through this sparse network. To reduce computation, we restrict breadth-first searches to a maximum depth. The maximum depth is an adjustable parameter for which we used a default value of four. This was established by leaving each protein, in turn, out of a test set of 12 complexes and finding that in all cases a depth of four was sufficient to find a path back from the given protein to its parent complex, if such a path existed at all in the network. A user-defined core set of proteins is given as input, and a list of candidate proteins ranked by \( C_i \) is returned as output. Candidate proteins above a user-defined rank threshold \( R \) are displayed graphically along with all edges with nonzero weight. An example of a query with members of the SAGA complex is shown in Figure 2.

We used an optimization procedure to estimate the error rate vector \( E \) used above. Although estimates of error rate for some of these evidence types are available (Mrowka et al. 2001; Deane et al. 2002; Edwards et al. 2002; von Mering et al. 2002; Deng et al. 2003; Sprinzak et al. 2003), the error rates of each evidence type in predicting direct and stable interaction have not previously been estimated. We therefore sought to find an estimate for the \( E \) vector that optimized performance of our algorithm, searching the space of possible values using a steepest ascent method. As an objective function for optimization, we used area under the success rate versus rank curves. Success rate versus rank curves are calculated by using a cross-validation pro-
cEDURE in which each protein in a set of complexes is left out in turn, a ranked list of candidate proteins is obtained, and the success rate for candidate proteins ranked at or below R is as-
sessed (examples in Fig. 3).

To optimize our choice of error rate vector entries, we used a training set of seven protein complexes. These were chosen at

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