Discovering regulatory and signalling circuits in molecular interaction networks

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\textbf{ABSTRACT}

\textbf{Motivation:} In model organisms such as yeast, large databases of protein–protein and protein-DNA interactions have become an extremely important resource for the study of protein function, evolution, and gene regulatory dynamics. In this paper we demonstrate that by integrating these interactions with widely-available mRNA expression data, it is possible to generate concrete hypotheses for the underlying mechanisms governing the observed changes in gene expression. To perform this integration systematically and at large scale, we introduce an approach for screening a molecular interaction network to identify active subnetworks, i.e., connected regions of the network that show significant changes in expression over particular subsets of conditions. The method we present here combines a rigorous statistical measure for scoring subnetworks with a search algorithm for identifying subnetworks with high score.

\textbf{Results:} We evaluated our procedure on a small network of 332 genes and 362 interactions and a large network of 4160 genes containing all 7462 protein–protein and protein-DNA interactions in the yeast public databases. In the case of the small network, we identified five significant subnetworks that covered 41 out of 77 (53\%) of all significant changes in expression. Both network analyses returned several top-scoring subnetworks with good correspondence to known regulatory mechanisms in the literature. These results demonstrate how large-scale genomic approaches may be used to uncover signalling and regulatory pathways in a systematic, integrative fashion.

\textbf{Availability:} The methods presented in this paper are implemented in the Cytoscape software package which is available to the academic community at http://www.cytoscape.org.

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\textbf{Keywords:} molecular interactions; gene expression; data integration; simulated annealing; Monte carlo methods.

\textbf{INTRODUCTION}

Expression profiling and large-scale proteomics have revolutionized biology by generating vast amounts of data about cell state. Genes with significant changes in expression have immediate and widespread interest as markers for diseases, stages of development, and a variety of other cellular phenotypes (Altman and Raychaudhuri, 2001). Genes with correlated expression changes over many conditions are likely to be involved in similar functions or cellular processes; these genes often also share DNA sequence elements, providing evidence that they are regulated by common transcription factors. Analytical methods such as gene expression clustering (Eisen \textit{et al.}, 1998; Tamayo \textit{et al.}, 1999), significance testing (Kerr and Churchill, 2001; Rocke and Durbin, 2001; Ideker \textit{et al.}, 2000), and sequence motif identification (Pilpel \textit{et al.}, 2001) have been indispensable for enabling these discoveries and summarizing the data at each step. By performing these analyses, we hope ultimately to answer questions about the underlying molecular mechanism: \textit{What are the signalling and regulatory interactions in control of the observed gene expression changes? How is this control exerted?}

For model organisms such as yeast, new technologies and data sets are making it possible to address these questions more directly than ever before. For example, systematic two-hybrid screens and co-immunoprecipitation experiments are populating the public databases with thousands of protein–protein interactions and complexes (Uetz \textit{et al.}, 2000; Gavin \textit{et al.}, 2002). Other ongoing projects are defining large numbers of protein→DNA interactions (Ren \textit{et al.}, 2000), and protein microarrays are making it possible to map interactions between proteins and drugs, hormones, and other small molecules...
(Zhu et al., 2001). These molecular interactions provide a convenient framework for understanding changes in gene expression and for integrating a wide variety of global state measurements.

Along these lines, in recent work we used a molecular interaction network to analyze changes in expression observed over 20 perturbations to the yeast galactose-utilization (GAL) pathway (Ideker et al., 2001). To construct the network (shown in Figure 1), we screened a database of publicly available protein–protein and protein→DNA interactions to select 362 interactions linking genes that were differentially-expressed under one or more perturbations. We found that pairs of genes linked by molecular interactions in this network were more likely to have correlated expression profiles than genes chosen at random, and we reported particular interactions for which this pairwise correlation was strong. However, the general task of associating gene expression changes with higher-order groups of interactions, such as make up signalling and regulatory pathways in the cell, was not discussed.

To address this problem, we now introduce a general method for searching the network to find ‘active subnetworks’, i.e., connected sets of genes with unexpectedly high levels of differential expression1. When expression levels have been observed over multiple conditions, we also wish to determine which conditions significantly affect gene expression in each active subnetwork. In order to achieve these goals, we implement a statistical scoring system which captures the amount of gene expression in each active subnetwork. We then describe a search algorithm, based on simulated annealing, for identifying change in a given subnetwork. We also wish to determine which conditions significantly affect gene expression in each active subnetwork. In order to achieve these goals, we implement a statistical scoring system which captures the amount of gene expression in each active subnetwork. We then describe a search algorithm, based on simulated annealing, for identifying change in a given subnetwork. We then describe a search algorithm, based on simulated annealing, for identifying change in a given subnetwork. We then describe a search algorithm, based on simulated annealing, for identifying change in a given subnetwork.

To begin by assessing the significance of differential expression analysis tool that robustly models error would suffice, so long to a $z$-score $z_i = \Phi^{-1}(1 - p_i)$, where $\Phi^{-1}$ is the inverse normal CDF. Thus in random data, $p$-values are distributed uniformly from 0 to 1 and $z$-scores follow a standard normal, with smaller $p$-values corresponding to larger $z$-scores.

To produce an aggregate $z$-score $z_A$ for an entire subnetwork $A$ of $k$ genes, we sum the $z_i$ over all genes in the subnetwork:

$$z_A = \frac{1}{\sqrt{k}} \sum_{i \in A} z_i$$ (1)

Subnetworks of all sizes are comparable under this scoring system: if the $z_i$ are independently drawn from a standard normal distribution, $z_A$ will also be distributed according to a standard normal, independent of $k$. A high $z_A$ indicates a biologically active subnetwork.

**Calibrating $z$ against the background distribution**

In order to properly capture the connection between expression and network topology, we must determine whether the score $z_A$ of a subnetwork is higher than expected relative to a random set of genes (drawn from the same expression data but independently of the network). We randomly sample gene sets of size $k$ using a Monte Carlo approach, compute their scores $z_A$, then use these to derive estimates for the score mean $\mu_k$ and standard deviation $\sigma_k$ for each $k$. Because we expect the means and standard deviations to be a smooth function of $k$, we can reduce noise in the Monte Carlo estimates using a sliding window average. Using these estimates, the corrected subnetwork score $s_A$ is:

$$s_A = \frac{(z_A - \mu_k)}{\sigma_k}$$ (2)

Using this correction, the scores $s_A$ of randomized subnets are guaranteed to have $\mu = 0$ and $\sigma = 1$. An overview of this scoring process is shown in Figure 2a.

**Scoring over multiple conditions**

Our scoring system may be extended to accommodate gene expression changes measured over multiple conditions as shown in Figure 2b. In this case, we start with a matrix of $p$-values (genes versus conditions) and corresponding $z$-scores. Given a subnetwork $A$, we use eqn. (1) to produce $m$ different aggregate scores $(z_{A1}, z_{A2}, \ldots, z_{Am})$, one for each condition. These are then sorted from highest to lowest $(z_{A(1)}, \ldots, z_{A(j)}, \ldots, z_{A(m)})$; we compute the significance $r_{A(j)}$ of the $j$th highest score using a binomial

1 This is true because the variance of a sum is the sum of the variances for independent random variables.

Available at http://www.systemsbiology.org/VERAandSAM. Any gene expression analysis tool that robustly models error would suffice, so long as its output can be converted to $p$-values.
Discovering pathways in interaction networks

Fig. 1. Performance on a small molecular interaction network. Nodes represent genes, an edge directed from one node to another signifies that the protein encoded by the first gene can influence the transcription of the second by DNA binding (protein→DNA), and an undirected edge between nodes signifies that the corresponding proteins can physically interact. Z-scores (blue scale) indicate the likelihood of differential expression of each gene in a GAL80 knockout experiment. Z-scores were used to search for active subnetworks using our simulated annealing method; the five top-scoring subnets are shown. For further information on this network, including gene labels, see Figure 4 in (Ideker et al., 2001).

order statistic (Kendall et al., 1987) as follows: Let $P_z = 1 - \Phi(z_{A(j)})$ represent the probability that any single condition has a $z$-score above $z_{A(j)}$. Then:

$$p_{A(j)} = \sum_{h=j}^{m} \binom{m}{h} (P_z)^h (1 - P_z)^{m-h}$$

(3)

This summation gives the probability that at least $j$ of the $m$ conditions had scores above $z_{A(j)}$, which is equivalent to the $p$-value for $z_{A(j)}$ as the $j$th largest of a standard normal sample. We use the inverse CDF $r_{A(j)} = \Phi^{-1} (1 - p_{A(j)})$ to convert back into a standard normal $z$-score, which is now adjusted for rank; the maximum of these is the subnet’s new score, $r_A^{max}$:

$$r_A^{max} = \max_j (r_{A(j)})$$

(4)
We consider the subnetwork to be ‘active’ under conditions ranked 1 through $j$. As in the single condition case, we must also calibrate the score against the background distribution: first, a Monte Carlo technique is used to estimate means $\mu_k$ and standard deviations $\sigma_k$ for $r_A^{\text{max}}$ computed from random gene sets of size $k = 4$ (depicted by the histogram) producing the score of $-1.0$ by eqn. (2). Panel [b] shows the extended procedure for scoring a subnet under multiple conditions. Aggregate $z$ scores for each condition (i) are sorted (ii) and adjusted for rank (iii). As in [a], the maximum score of 4.2 is then corrected to 2.2 using the background score distribution (iv).

**Searching for high-scoring subnetworks via simulated annealing**

The above methods allow us to score a given subnetwork, but we must also find the highest-scoring subnetwork(s) in a full network of molecular interactions. Because the problem of finding the maximal-scoring connected subgraph is NP-hard$^5$, we implement an approach based on simulated annealing (Kirkpatrick et al., 1983). In practice, this approach is not guaranteed to find the maximal score overall; however, all high-scoring subnetworks are of strong biological interest regardless of whether they are strictly maximal$^1$.

Throughout the following algorithm, we associate an ‘active/inactive’ state with each node. $G_w$ denotes the ‘working’ subgraph of $G$ induced by the active nodes. At each iteration $i$, we define $s_i$ as the score ($s_A$ from above) of the highest-scoring component in $G_w$.

**Input:** A graph $G = (V, E)$ of molecular interactions, a number $N$ of iterations, and a temperature function $T_i$ which decreases geometrically from $T_{\text{start}}$ to $T_{\text{end}}$.

**Output:** A subgraph $G_w$ of $G$.

1. Initialize $G_w$ by setting each $v \in V$ to active/inactive with probability $\frac{1}{2}$.
2. FOR $i = 1$ to $N$ DO
3. Randomly pick a node $v \in V$ and toggle its state;
4. Compute the score $s_i$ for the working subgraph $G_w$;
5. IF ($s_i > s_{i-1}$), keep $v$ toggled;
6. ELSE keep $v$ toggled with probability $p = e^{(s_i - s_{i-1})/T_i}$
7. Output $G_w$ and its highest-scoring component $A$.

Finally, we ‘quench’ at temperature $T_i = 0$ until all adjoining possibilities have been explored and the score has reached a local maximum. Upon termination of annealing, the subnetwork $A$ represents a signalling or regulatory circuit of high biological interest$^*$. 

**Heuristics for improved annealing**

We have extended our annealing approach to search for $M$ subnetworks simultaneously (where $M$ is a user-definable parameter). Maintaining multiple components can dramatically improve the efficiency of annealing, because toggling the state of a node may cause a number of components with low score to merge into a single high-scoring component$^{10}$.

An additional heuristic increases the efficiency of annealing in networks with many ‘hubs’, i.e., nodes of high degree. Simulated annealing tends to perform poorly in such networks, because adding a hub to $G_w$ immediately creates a large component incorporating all nodes adjacent to the hub. Unless all adjacent nodes with low $z$-score are inactive, the resulting component will also have low score regardless of the contribution of the hub node itself. We have addressed this problem by a straightforward modification to step (3) of the algorithm: when adding a node of degree greater than a user-definable parameter $d_{\text{min}}$, simultaneously remove all neighbours that are not in the top-scoring component.

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$^5$Given suitable values for annealing parameters $T_i$ and $N$, the score of the final solution is guaranteed to be the global maximum (Landy and Mess, 1986). However, these values are generally unknown and can be impractically large.

$^*This search is robust with respect to false-positive interactions, because adding an interaction never disrupts an existing subnetwork. With respect to false negatives, it is possible for a missing interaction to prevent the formation of an otherwise high-scoring subnetwork; however, the remaining pieces of the subnetwork will still score well.

$^1$Full details are provided in the supplementary web materials.
RESULTS
Small network with a single perturbation
We searched for active subnetworks in the network of 362 protein–protein and protein→DNA interactions from our previous galactose utilization study (Ideker et al., 2001). This search was performed against gene expression data. We converted these significance values to z-scores, shown superimposed on the network in Figure 1.

Fig. 3. Score and temperature versus number of iterations. Simulated annealing was performed for the network and expression profile shown in Fig. 1, with parameters $N = 1 \times 10^5$, $T_{\text{start}} = 1$, $M = 5$, $d_{\text{min}} = 100$. Annealing temperature (right vertical axis; solid grey trace) decreases geometrically over consecutive iterations according to the set schedule. By the end of the run, scores for each of the five top-scoring subnetworks have increased to a local maximum (left vertical axis; solid black trace = top score, dashed = sum of all five).

Fig. 4. Distribution of subnetwork scores in actual and randomized data. Score distributions are shown for 200 simulated-annealing runs on actual (a) versus scrambled (b) data, for each of the top-five scoring subnetworks (vertical axis). Different initial conditions were used for each run, with identical network, expression data, and annealing parameters to those described for Figure 1. Greyscale intensity is proportional to the number of runs achieving a particular score; the bounding lines indicate the maximum and minimum values of each distribution. Scrambled data were generated by shuffling the mapping between genes and z-scores, thus preserving the score distribution over all genes but removing any correlation between gene expression and network location.

We ran our annealing algorithm with parameters ($N = 100,000; T_{\text{start}} = 1; T_{\text{end}} = 0.01; M = 5; d_{\text{min}} = 100$). Figure 1 shows the nodes involved in each of the five top-scoring subnetworks; Figure 3 tracks the increases in subnetwork score over the course of the run. Scores of the five subnetworks were (7.7, 3.1, 2.8, 2.5, 2.3); their corresponding sizes were (43, 5, 5, 3, 1). To verify that these subnetworks contained higher-than-expected levels of differential expression, we compared their scores to annealing runs performed on randomized expression data sets. As shown in Figure 4, the top score was greater than and non-overlapping with top scores from randomized data.

Taken together, the five subnetworks contained 41 out of 77 genes in the network with significant changes in expression ($p < 10^{-5}; z > 4.27$). In each case, the subnetwork gives clues as to how gene expression is transmitted from one gene to another. For instance, the top-scoring subnetwork included GAL80, the gene that was knocked out to produce the observed expression changes. It neighbours GAL4 (immediately right of GAL80 in Figure 1), a hub with protein→DNA interactions to seven other genes in the subnetwork. Therefore, one hypothesis is that GAL80 influences expression of these genes by a path through GAL4. The subnetworks contain many examples of genes with low z-score that were required to connect together several high-scoring genes. For example, subnetwork 3 (lower left corner of Figure 1) consists of four genes connected to a central transcription factor through protein→DNA interactions. As is typical for regulatory networks, the transcription factor shows only modest expression change compared to the four genes it regulates. Our search identifies this subnetwork because its total level of significance remains relatively high.

‡‡As described below in the Discussion, this hypothesis is well supported by the literature (Lohr et al., 1995).
Large network with multiple perturbations

Having characterized our methods on a small network, we wished to explore their performance on larger networks containing nearly all catalogued molecular interactions in yeast. We also wished to investigate how searching for active subnetworks was impacted by multiple conditions. To construct a large molecular interaction network for yeast, we included all 7145 protein–protein interactions in the BIND database (Bader et al., 2001) and all 317 protein→DNA interactions present in TRANSFAC (Wingender et al., 2001) as of September 2001. To find active subnetworks, we screened this network against a complete data set of 20 mRNA expression profiles gathered in response to different perturbations to genes in the GAL pathway (Ideker et al., 2001) (the GAL80 deletion was one of these). As before, all measurements were converted into z-scores representing the likelihood of differential expression for each gene and perturbation.

For this larger network, we optimized the performance of our annealing algorithm by evaluating a range of annealing parameters over multiple runs. As shown in Table 1, increasing $N$ from $10^3$ to $10^7$ led to a more than five-fold increase in top score [rows a]; tracking fewer than 10 subnetworks negatively impacted score [b]; the optimal value for $d_{min}$ was approximately 100 [c]; and temperature did not have a dramatic effect for the parameter sets we surveyed [d]. Using the near-optimal parameters ($N = 1 \times 10^7$, $T_{start} = 2$, $T_{end} = 0.01$, $M = 20$, $d_{min} = 100$), we identified seven subnetworks whose scores were significantly higher than expected in randomized data (by an analysis similar to that shown in Figure 4 for the small network). These subnetworks are shown in Figure 5, while Figure 6 shows the particular conditions used to maximize the score of each subnetwork, i.e., the conditions under which each subnetwork was active.

These results demonstrate that our approach can produce active subnetworks that are extremely large—340 genes in the case of the highest-scoring subnetwork! Although this entire structure is statistically significant, it is cumbersome to inspect visually or for use in formulating biological hypotheses. However, we may apply our search algorithm recursively to identify substructures that are particularly significant, i.e., scoring higher than would be expected at random within the 340-node subnetwork. By reducing the population of genes to those in the subnetwork, we change the distribution used to compute the correction factors $\mu_k$ and $\sigma_k$ in eqn. (2) and thus the scores $s_A$. Here, a second simulated annealing run performed on subnetwork 1 against all 20 perturbation conditions ($N = 1 \times 10^6$, $T_{start} = 1$, $M = 5$, $d_{min} = 1000$) identified five substructures of sizes (28, 13, 3, 7, 9); these are labelled #1a through #1e in Figures 5 and 6.

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DISCUSSION

Subnetworks are consistent with known regulatory circuits

Many of the subnetworks resulting from our analyses have striking overlap with well-known regulatory and signalling pathways described in the yeast literature. For example, the path $GAL3\rightarrow GAL80\rightarrow GAL4\rightarrow GAL1,7,10$ contained in subnetwork 1b (Figures 5 and 6b) forms the core of the known galactose-induction circuit (Lohr et al., 1995). Briefly, GAL4p is a transcription factor that induces the expression of the enzymes $GAL1,7,10$, though protein→DNA interactions, but in the absence of galactose, GAL80p inhibits this activity through a protein–protein interaction. In the presence of galactose, GAL3p associates with GAL80p via another protein–protein interaction, allowing GAL4p to transcribe the enzymes at a high level.

As a second example, subnetwork 1d contains interactions known to regulate genes involved in both the mating response (e.g., $STE2$, $MFA2$, $BAR1$) and cell cycle arrest ($CLB1$, $FAR1$). However, even without expert knowledge from yeast biology, visual inspection of these subnetworks provides us with ready hypotheses for why their genes are differentially expressed. On the strength of these known cases, it will be extremely interesting to examine subnet-
Fig. 5. Performance on a large interaction network using multiple conditions. The seven highest-scoring subnetworks screened from the large network are shown in decreasing order of score. Subnet 1, containing 340 nodes, was subjected to a second application of simulated annealing to produce five smaller subnets 1a–1e. Subnetworks are represented as in Figure 1, but with protein → DNA interactions denoted by yellow edges and protein–protein interactions denoted by blue edges.

Fig. 6. Perturbations affecting each subnetwork. Panel [a]: Red blocks indicate the particular perturbations (columns) maximizing the score of each subnetwork (rows) shown in Fig. 5. Panel [b]: Changes in expression level are shown using a red/green colorimetric scale for the genes and perturbations associated with subnet #1b. Note that inversely correlated genes (e.g., GAL1 and HXT6) may appear in the same subnet, and that genes without large changes in expression (e.g., GAL4) are nevertheless included because they connect several genes with dramatic changes (e.g., GAL10 and GAL7).

works containing genes whose functions and/or regulatory mechanisms are not already well understood.

Subnetworks versus gene-expression clusters
It is instructive to compare existing methods for clustering genes by expression, e.g., (Eisen et al., 1998; Tamayo et al., 1999), to those introduced here. Although all of these methods form groups of genes (clusters versus subnets), our method differs from established clustering techniques in at least four major ways. First, and most importantly, our approach groups genes subject to the constraints of the molecular interaction network. A striking consequence of this constraint is that subnetworks may contain genes without large expression changes so long as they are required to connect other, differentially expressed genes. Second, subnetworks are scored over only a subset of conditions; thus, genes are not required to be co-regulated over all conditions in order to group together. Third, while most clustering methods group genes by both magnitude and direction of change, we consider only the significance of change. Thus, we may connect strongly repressed...
induced genes to build subnets that represent complete signalling or regulatory pathways (see Figure 6b). Finally, our method leaves some genes unaffiliated with any subnet, unlike typical clustering methods which assign every gene to a distinct cluster.

**Future work**

In the near future, the most pressing task is to investigate our identified subnetworks in the laboratory. Because large interaction networks are suspected to contain many false-positives, an initial experiment would be to verify that the interactions in each subnet are reproducible and present under the subnet’s particular set of conditions. We also wish to investigate a number of extensions to our approach, including:

- annotating each interaction with its directionality and/or the specific conditions and compartments in which it has been observed, constraining subnetworks to plausible causal chains.
- accommodating new types of interaction data such as interactions between proteins and small molecules, and new types of biological state measurements such as changes in protein abundances, protein modifications, or concentrations of intracellular metabolites.
- potentially correcting for local network topology. In random expression data, some subnets are much more likely than others to have the highest score; it may be desirable to correct for this bias by computing a prior probability over the population of subnetworks.

As the core algorithms are developed further, we expect this approach to have immense impact in elucidating the underlying molecular mechanisms of a variety of organisms. Ultimately, we envision that biologists will use this approach to have immense impact in elucidating entire spectrum of drugs and human diseases.

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**REFERENCES**


