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Using Bayesian Networks to Analyze Expression Data

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ABSTRACT

DNA hybridization arrays simultaneously measure the expression level for thousands of genes. These measurements provide a "snapshot" of transcription levels within the cell. A major challenge in computational biology is to uncover, from such measurements, gene/protein interactions and key biological features of cellular systems. In this paper, we propose a new framework for discovering interactions between genes based on multiple expression measurements. This framework builds on the use of *Bayesian networks* for representing statistical dependencies. A Bayesian network is a graph-based model of joint multivariate probability distributions that captures properties of conditional independence between variables. Such models are attractive for their ability to describe complex stochastic processes and because they provide a clear methodology for learning from (noisy) observations. We start by showing how Bayesian networks can describe interactions between genes. We then describe a method for recovering gene interactions from microarray data using tools for learning Bayesian networks. Finally, we demonstrate this method on the *S. cerevisiae* cell-cycle measurements of Spellman *et al.* (1998).

Key words: gene expression, microarrays, Bayesian methods.

1. INTRODUCTION

Accentral goal of molecular biology is to understand the regulation of protein synthesis and its reactions to external and internal signals. All the cells in an organism carry the same genomic data, yet their protein makeup can be drastically different both temporally and spatially, due to regulation. Protein synthesis is regulated by many mechanisms at its different stages. These include mechanisms for controlling transcription initiation, RNA splicing, mRNA transport, translation initiation, post-translational modifications, and degradation of mRNA/protein. One of the main junctions at which regulation occurs is mRNA transcription. A major role in this machinery is played by proteins themselves that bind to regulatory regions along the DNA, greatly affecting the transcription of the genes they regulate.

In recent years, technical breakthroughs in spotting hybridization probes and advances in genome sequencing efforts lead to development of *DNA microarrays*, which consist of many species of probes, either oligonucleotides or cDNA, that are immobilized in a predefined organization to a solid phase. By using

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DNA microarrays, researchers are now able to measure the abundance of thousands of mRNA targets simultaneously (DeRisi *et al.*, 1997; Lockhart *et al.*, 1996; Wen *et al.*, 1998). Unlike classical experiments, where the expression levels of only a few genes were reported, DNA microarray experiments can measure *all* the genes of an organism, providing a "genomic" viewpoint on gene expression. As a consequence, this technology facilitates new experimental approaches for understanding gene expression and regulation (Iyer *et al.*, 1999; Spellman *et al.*, 1998).

Early microarray experiments examined few samples and mainly focused on differential display across tissues or conditions of interest. The design of recent experiments focuses on performing a larger number of microarray assays ranging in size from a dozen to a few hundreds of samples. In the near future, data sets containing thousands of samples will become available. Such experiments collect enormous amounts of data, which clearly reflect many aspects of the underlying biological processes. An important challenge is to develop methodologies that are both statistically sound and computationally tractable for analyzing such data sets and inferring biological interactions from them.

Most of the analysis tools currently used are based on *clustering* algorithms. These algorithms attempt to locate groups of genes that have similar expression patterns over a set of experiments (Alon *et al.*, 1999; Ben-Dor *et al.*, 1999; Eisen *et al.*, 1999; Michaels *et al.*, 1998; Spellman *et al.*, 1998). Such analysis has proven to be useful in discovering genes that are co-regulated and/or have similar function. A more ambitious goal for analysis is to reveal the structure of the transcriptional regulation process (Akutsu, 1998; Chen *et al.*, 1999; Somogyi *et al.*, 1996; Weaver *et al.*, 1999). This is clearly a hard problem. The current data is extremely noisy. Moreover, mRNA expression data alone only gives a partial picture that does not reflect key events, such as translation and protein (in)activation. Finally, the amount of samples, even in the largest experiments in the foreseeable future, does not provide enough information to construct a fully detailed model with high statistical significance.

In this paper, we introduce a new approach for analyzing gene expression patterns that uncovers properties of the transcriptional program by examining statistical properties of *dependence* and *conditional independence* in the data. We base our approach on the well-studied statistical tool of *Bayesian networks* (Pearl, 1988). These networks represent the dependence structure between multiple interacting quantities (e.g., expression levels of different genes). Our approach, probabilistic in nature, is capable of handling noise and estimating the confidence in the different features of the network. We are, therefore, able to focus on interactions whose signal in the data is strong.

Bayesian networks are a promising tool for analyzing gene expression patterns. First, they are particularly useful for describing processes composed of *locally* interacting components; that is, the value of each component *directly* depends on the values of a relatively small number of components. Second, statistical foundations for learning Bayesian networks from observations, and computational algorithms to do so, are well understood and have been used successfully in many applications. Finally, Bayesian networks provide models of causal influence: Although Bayesian networks are mathematically defined strictly in terms of probabilities and conditional independence statements, a connection can be made between this characterization and the notion of *direct causal influence*. (Heckerman *et al.*, 1999; Pearl and Verma, 1991; Spirtes *et al.*, 1993). Although this connection depends on several assumptions that do not necessarily hold in gene expression data, the conclusions of Bayesian network analysis might be indicative of some causal connections in the data.

The remainder of this paper is organized as follows. In Section 2, we review key concepts of Bayesian networks, learning them from observations and using them to infer causality. In Section 3, we describe how Bayesian networks can be applied to model interactions among genes and discuss the technical issues that are posed by this type of data. In Section 4, we apply our approach to the gene-expression data of Spellman *et al.* (1998), analyzing the statistical significance of the results and their biological plausibility. Finally, in Section 5, we conclude with a discussion of related approaches and future work.

2. BAYESIAN NETWORKS

2.1. Representing distributions with Bayesian networks

Consider a finite set $\mathcal{X} = \{X_1, \dots, X_n\}$ of random variables where each variable X_i may take on a value x_i from the domain $Val(X_i)$. In this paper, we use capital letters, such as X, Y, Z, for variable names and lowercase letters, such as x, y, z, to denote specific values taken by those variables. Sets of variables are denoted by boldface capital letters X, Y, Z, and assignments of values to the variables in these sets

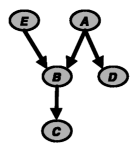


FIG. 1. An example of a simple Bayesian network structure. This network structure implies several conditional independence statements: I(A; E), $I(B; D \mid A, E)$, $I(C; A, D, E \mid B)$, $I(D; B, C, E \mid A)$, and I(E; A, D). The network structure also implies that the joint distribution has the product form: P(A, B, C, D, E) = P(A)P(B|A, E)P(C|B)P(D|A)P(E).

are denoted by boldface lowercase letters \mathbf{x} , \mathbf{y} , \mathbf{z} . We denote $I(X; Y \mid Z)$ to mean X is independent of Y conditioned on Z: P(X|Y,Z) = P(X|Z).

A Bayesian network is a representation of a joint probability distribution. This representation consists of two components. The first component, G, is a directed acyclic graph (DAG) whose vertices correspond to the random variables X_1, \ldots, X_n . The second component, θ describes a conditional distribution for each variable, given its parents in G. Together, these two components specify a unique distribution on X_1, \ldots, X_n .

The graph G represents conditional independence assumptions that allow the joint distribution to be decomposed, economizing on the number of parameters. The graph G encodes the *Markov Assumption*:

(*) Each variable X_i is independent of its nondescendants, given its parents in G.

By applying the chain rule of probabilities and properties of conditional independencies, any joint distribution that satisfies (*) can be decomposed into the *product form*

$$P(X_1,\ldots,X_n)=\prod_{i=1}^n P(X_i|\mathbf{Pa}^G(X_i)), \qquad (1)$$

where $\mathbf{Pa}^G(X_i)$ is the set of parents of X_i in G. Figure 1 shows an example of a graph G and lists the Markov independencies it encodes and the product form they imply.

A graph G specifies a product form as in (1). To fully specify a joint distribution, we also need to specify each of the conditional probabilities in the product form. The second part of the Bayesian network describes these conditional distributions, $P(X_i|\mathbf{Pa}^G(X_i))$ for each variable X_i . We denote the parameters that specify these distributions by θ .

In specifying these conditional distributions, we can choose from several representations. In this paper, we focus on two of the most commonly used representations. For the following discussion, suppose that the parents of a variable X are $\{U_1, \ldots, U_k\}$. The choice of representation depends on the type of variables we are dealing with:

- **Discrete variables.** If each of X and U_1, \ldots, U_k takes discrete values from a finite set, then we can represent $P(X \mid U_1, \ldots, U_k)$ as a table that specifies the probability of values for X for each joint assignment to U_1, \ldots, U_k . Thus, for example, if all the variables are binary valued, the table will specify 2^k distributions.
 - This is a general representation which can describe any discrete conditional distribution. Thus, we do not loose expressiveness by using this representation. This flexibility comes at a price: The number of free parameters is exponential in the number of parents.
- Continuous variables. Unlike the case of discrete variables, when the variable X and its parents U_1, \ldots, U_k are real valued, there is no representation that can represent all possible densities. A natural choice for multivariate continuous distributions is the use of Gaussian distributions. These can be represented in a Bayesian network by using *linear Gaussian* conditional densities. In this representation, the conditional density of X given its parents is given by:

$$P(X \mid u_1, \ldots, u_k) \sim N(a_0 + \sum_i a_i \cdot u_i, \sigma^2).$$

That is, X is normally distributed around a mean that depends *linearly* on the values of its parents. The variance of this normal distribution is independent of the parents' values. If all the variables in a network have linear Gaussian conditional distributions, then the joint distribution is a multivariate Gaussian (Lauritzen and Wermuth, 1989).

• **Hybrid networks**. When the network contains a mixture of discrete and continuous variables, we need to consider how to represent a conditional distribution for a continuous variable with discrete parents and for a discrete variable with continuous parents. In this paper, we disallow the latter case. When a continuous variable *X* has discrete parents, we use *conditional Gaussian* distributions (Lauritzen and Wermuth, 1989) in which, for each joint assignment to the discrete parents of *X*, we represent a linear Gaussian distribution of *X* given its continuous parents.

Given a Bayesian network, we might want to answer many types of questions that involve the joint probability (e.g., what is the probability of X = x given observation of some of the other variables?) or independencies in the domain (e.g., are X and Y independent once we observe Z?). The literature contains a suite of algorithms that can answer such queries efficiently by exploiting the explicit representation of structure (Jensen, 1996; Pearl, 1988)).

2.2. Equivalence classes of Bayesian networks

A Bayesian network structure G implies a set of independence assumptions in addition to (*). Let Ind(G) be the set of independence statements (of the form X is independent of Y given \mathbb{Z}) that hold in all distributions satisfying these Markov assumptions.

More than one graph can imply exactly the same set of independencies. For example, consider graphs over two variables X and Y. The graphs $X \to Y$ and $X \leftarrow Y$ both imply the same set of independencies (i.e., $Ind(G) = \emptyset$). Two graphs G and G' are *equivalent* if Ind(G) = Ind(G'). That is, both graphs are alternative ways of describing the same set of independencies.

This notion of equivalence is crucial, since when we examine observations from a distribution, we cannot distinguish between equivalent graphs. Pearl and Verma (1991) show that we can characterize *equivalence classes* of graphs using a simple representation. In particular, these results establish that equivalent graphs have the same underlying undirected graph but might disagree on the direction of some of the arcs.

Theorem 2.1 (Pearl and Verma, 1991). Two DAGs are equivalent if and only if they have the same underlying undirected graph and the same v-structures (i.e., converging directed edges into the same node, such that $a \to b \leftarrow c$, and there is no edge between a and c).

Moreover, an equivalence class of network structures can be uniquely represented by a *partially directed graph* (PDAG), where a directed edge $X \to Y$ denotes that all members of the equivalence class contain the arc $X \to Y$; an undirected edge X - Y denotes that some members of the class contain the arc $X \to Y$, while others contain the arc $Y \to X$. Given a DAG G, the PDAG representation of its equivalence class can be constructed efficiently (Chickering, 1995).

2.3. Learning Bayesian networks

The problem of learning a Bayesian network can be stated as follows. Given a training set $D = \{\mathbf{x}^1, \dots, \mathbf{x}^N\}$ of independent instances of \mathcal{X} , find a network $B = \langle G, \Theta \rangle$ that best matches D. More precisely, we search for an equivalence class of networks that best matches D.

The theory of learning networks from data has been examined extensively over the last decade. We only briefly describe the high-level details here. We refer the interested reader to Heckerman (1998) for a recent tutorial on the subject.

The common approach to this problem is to introduce a statistically motivated scoring function that evaluates each network with respect to the training data and to search for the optimal network according

¹To be more precise, under the common assumptions in learning networks, which we also make in this paper, one cannot distinguish between equivalent graphs. If we make stronger assumptions, for example, by restricting the form of the conditional probability distributions we can learn, we might have a preference for one equivalent network over another.

to this score. One method for deriving a score is based on Bayesian considerations (see Cooper and Herskovits (1992) and Heckerman *et al.* (1995) for complete description). In this score, we evaluate the posterior probability of a graph given the data:

$$S(G : D) = \log P(G \mid D)$$
$$= \log P(D \mid G) + \log P(G) + C$$

where C is a constant independent of G and

$$P(D \mid G) = \int P(D \mid G, \Theta) P(\Theta \mid G) d\Theta$$

is the marginal likelihood which averages the probability of the data over all possible parameter assignments to G. The particular choice of priors P(G) and $P(\Theta \mid G)$ for each G determines the exact Bayesian score. Under mild assumptions on the prior probabilities, this scoring rule is asymptotically consistent: Given a sufficiently large number of samples, graph structures that exactly capture all dependencies in the distribution, will receive, with high probability, a higher score than all other graphs (see, for example, Friedman and Yakhini (1996)). This means, that given a sufficiently large number of instances, learning procedures can pinpoint the exact network structure up to the correct equivalence class.

In this work, we use a prior described by Heckerman and Geiger (1995) for hybrid networks of multinomial distributions and conditional Gaussian distributions. (This prior combines earlier works on priors for multinomial networks (Buntine, 1991; Cooper and Herskovits, 1992; Heckerman *et al.*, 1995) and for Gaussian networks (Geiger and Heckerman, 1994).) We refer the reader to Heckerman and Geiger (1995) and Heckerman (1998) for details on these priors.

In the analysis of gene expression data, we use a small number of samples. Therefore, care should be taken in choosing the prior. Without going into details, each prior from the family of priors described by Heckerman and Geiger can be specified by two parameters. The first is a *prior network*, which reflects our prior belief on the joint distribution of the variables in the domain, and the second is an *effective sample size* parameter, which reflects how strong is our belief in the prior network. Intuitively, setting the effective sample size to K is equivalent to having seen K samples from the distribution defined by the prior network. In our experiments, we choose the prior network to be one where all the random variables are independent of each other. In the prior network, discrete random variables are uniformly distributed, while continuous random variables have an a priori normal distribution. We set the equivalent sample size 5 in a rather arbitrary manner. This choice of the prior network is to ensure that we are not (explicitly) biasing the learning procedure to any particular edge. In addition, as we show below, the results are reasonably insensitive to this exact magnitude of the equivalent sample size.

In this work we assume *complete data*, that is, a data set in which each instance contains the values of all the variables in the network. When the data is complete and the prior satisfies the conditions specified by Heckerman and Geiger, then the posterior score satisfies several properties. First, the score is *structure equivalent*, i.e., if G and G' are equivalent graphs they are guaranteed to have the same posterior score. Second, the score is *decomposable*. That is, the score can be rewritten as the sum

$$S(G:D) = \sum_{i} ScoreContribution(X_i, \mathbf{Pa}^G(X_i):D),$$

where the contribution of every variable X_i to the total network score depends only on the values of X_i and $\mathbf{Pa}^G(X_i)$ in the training instances. Finally, these local contributions for each variable can be computed using a closed form equation (again, see Heckerman and Geiger (1995) for details).

Once the prior is specified and the data is given, learning amounts to finding the structure G that maximizes the score. This problem is known to be NP-hard (Chickering, 1996). Thus we resort to heuristic search. The decomposition of the score is crucial for this optimization problem. A *local* search procedure that changes one arc at each move can efficiently evaluate the gains made by adding, removing, or reversing a single arc. An example of such a procedure is a greedy hill-climbing algorithm that at each step performs the local change that results in the maximal gain, until it reaches a local maximum. Although this procedure does not necessarily find a global maximum, it does perform well in practice. Examples of other search methods that advance using one-arc changes include beam-search, stochastic hill-climbing, and simulated annealing.

2.4. Learning causal patterns

Recall that a Bayesian network is a model of dependencies between multiple measurements. However, we are also interested in modeling the mechanism that generated these dependencies. Thus, we want to model the flow of causality in the system of interest (e.g., gene transcription in the gene expression domain). A *causal network* is a model of such causal processes. Having a causal interpretation facilitates predicting the effect of an *intervention* in the domain: setting the value of a variable in such a way that the manipulation itself does not affect the other variables.

While at first glance there seems to be no direct connection between probability distributions and causality, causal interpretations for Bayesian networks have been proposed (Pearl and Verma, 1991; Pearl, 2000). A causal network is mathematically represented similarly to a Bayesian network, a DAG where each node represents a random variable along with a local probability model for each node. However, causal networks have a stricter interpretation of the meaning of edges: the parents of a variable are its immediate causes.

A causal network models not only the distribution of the observations, but also the effects of *interventions*. If X causes Y, then manipulating the value of X affects the value of Y. On the other hand, if Y is a cause of X, then manipulating X will not affect Y. Thus, although $X \to Y$ and $X \leftarrow Y$ are equivalent Bayesian networks, they are not equivalent causal networks.

A causal network can be interpreted as a Bayesian network when we are willing to make the *Causal Markov Assumption*: given the values of a variable's immediate causes, it is independent of its earlier causes. When the casual Markov assumption holds, the causal network satisfies the Markov independencies of the corresponding Bayesian network. For example, this assumption is a natural one in models of genetic pedigrees: once we know the genetic makeup of the individual's parents, the genetic makeup of her ancestors is not informative about her own genetic makeup.

The central issue is: When can we learn a causal network from observations? This issue received a thorough treatment in the literature (Heckerman *et al.*, 1999; Pearl and Verma, 1991; Spirtes *et al.*, 1993; Spirtes *et al.*, 1999). We briefly review the relevant results for our needs here. For a more detailed treatment of the topic, we refer the reader to Pearl (2000) and to Cooper and Glymour (1999).

First it is important to distinguish between an *observation* (a passive measurement of our domain, i.e., a sample from \mathcal{X}) and an *intervention* (setting the values of some variables using forces outside the causal model, e.g., gene knockout or over-expression). It is well known that interventions are an important tool for inferring causality. What is surprising is that some causal relations can be inferred from observations alone.

To learn about causality, we need to make several assumptions. The first assumption is a modeling assumption: we assume that the (unknown) causal structure of the domain satisfies the Causal Markov Assumption. Thus, we assume that causal networks can provide a reasonable model of the domain. Some of the results in the literature require a stronger version of this assumption, namely, that causal networks can provide a perfect description of the domain (that is, an independence property holds in the domain if and only if it is implied by the model). The second assumption is that there are no *latent* or hidden variables that affect several of the observable variables. We discuss relaxations of this assumption below.

If we make these two assumptions, then we essentially assume that one of the possible DAGs over the domain variables is the "true" causal network. However, as discussed above, from observations alone, we cannot distinguish between causal networks that specify the same independence properties, i.e., belong to the same equivalence class (see Section 2.2). Thus, at best we can hope to learn a description of the equivalence class that contains the true model. In other words, we will learn a PDAG description of this equivalence class.

Once we identify such a PDAG, we are still uncertain about the true causal structure in the domain. However, we can draw some causal conclusions. For example, if there is a directed path from X to Y in the PDAG, then X is a causal ancestor of Y in all the networks that could have generated this PDAG including the "true" causal model. Moreover, by using Theorem 2.1, we can predict what aspects of a proposed model would be detectable based on observations alone.

When data is sparse, we cannot identify a unique PDAG as a model of the data. In such a situation, we can use the posterior over PDAGs to represent *posterior* probabilities over causal statements. In a sense the posterior probability of "X causes Y" is the sum of the posterior probability of all PDAGs in which this statement holds. (See Heckerman *et al.* (1999) for more details on this Bayesian approach.) The situation is somewhat more complex when we have a combination of observations and results of different

interventions. From such data we might be able to distinguish between equivalent structures. Cooper and Yoo (1999) show how to extend the Bayesian approach of Heckerman *et al.* (1999) for learning from such mixed data.

A possible pitfall in learning causal structure is the presence of latent variables. In such a situation the observations that X and Y depend on each other probabilistically might be explained by the existence of an unobserved common cause. When we consider only two variables we cannot distinguish this hypothesis from the hypotheses "X causes Y" or "Y causes X". However, a more careful analysis shows that one can characterize all networks with latent variables that can result in the same set of independencies over the observed variables. Such equivalence classes of networks can be represented by a structure called a partial ancestral graphs (PAGs) (Spirtes et al., 1999). As can be expected, the set of causal conclusions we can make when we allow latent variables is smaller than the set of causal conclusions when we do not allow them. Nonetheless, causal relations often can be recovered even in this case.

The situation is more complicated when we do not have enough data to identify a single PAGs. As in the case of PDAGs, we might want to compute posterior scores for PAGs. However, unlike for PDAGs the question of scoring a PAG (which consists of many models with different number of latent variables) remains an open question.

3. ANALYZING EXPRESSION DATA

In this section we describe our approach to analyzing gene expression data using Bayesian network learning techniques.

3.1. Outline of the proposed approach

We start with our modeling assumptions and the type of conclusions we expect to find. Our aim is to understand a particular *system* (a cell or an organism and its environment). At each point in time, the system is in some *state*. For example, the state of a cell can be defined in terms of the concentration of proteins and metabolites in the various compartments, the amount of external ligands that bind to receptors on the cell's membrane, the concentration of different mRNA molecules in the cytoplasm, etc.

The cell (or other biological systems) consists of many interacting components that affect each other in some consistent fashion. Thus, if we consider random sampling of the system, some states are more probable. The likelihood of a state can be specified by the joint probability distribution on each of the cells components.

Our aim is to estimate such a probability distribution and understand its structural features. Of course, the state of a system can be infinitely complex. Thus, we resort to a partial view and focus on some of the components. Measurements of attributes from these components are random variables that represent some aspect of the system's state. In this paper, we are dealing mainly with random variables that denote the mRNA expression level of specific genes. However, we can also consider other random variables that denote other aspects of the system state, such as the phase of the system in the the cell-cycle process. Other examples include measurements of experimental conditions, temporal indicators (i.e., the time/stage that the sample was taken from), background variables (e.g., which clinical procedure was used to get a biopsy sample), and exogenous cellular conditions.

Our aim is to model the system as a joint distribution over a collection of random variables that describe system states. If we had such a model, we could answer a wide range of queries about the system. For example, does the expression level of a particular gene depend on the experimental condition? Is this dependence direct or indirect? If it is indirect, which genes mediate the dependency? Not having a model at hand, we want to learn one from the available data and use it to answer questions about the system.

In order to learn such a model from expression data, we need to deal with several important issues that arise when learning in the gene expression domain. These involve statistical aspects of interpreting the results, algorithmic complexity issues in learning from the data, and the choice of local probability models.

Most of the difficulties in learning from expression data revolve around the following central point: Contrary to most situations where one attempts to learn models (and in particular Bayesian networks), expression data involves transcript levels of thousands of genes while current data sets contain at most a few dozen samples. This raises problems in both computational complexity and the statistical significance of the resulting networks. On the positive side, genetic regulation networks are believed to be sparse,

i.e., given a gene, it is assumed that no more than a few dozen genes directly affect its transcription. Bayesian networks are especially suited for learning in such sparse domains.

3.2. Representing partial models

When learning models with many variables, small data sets are not sufficiently informative to significantly determine that a single model is the "right" one. Instead, many different networks should be considered as reasonable explanations of the given data. From a Bayesian perspective, we say that the posterior probability over models is not dominated by a single model (or an equivalence class of models).²

One potential approach to deal with this problem is to find all the networks that receive high posterior score. Such an approach is outlined by Madigan and Raftery (1994). Unfortunately, due to the combinatoric aspect of networks the set of "high posterior" networks can be huge (i.e., exponential in the number of variables). Thus, in a domain such as gene expression with many variables and diffused posterior, we cannot hope to explicitly list all the networks that are plausible given the data.

Our solution is as follows. We attempt to identify properties of the network that might be of interest. For example, are X and Y "close" neighbors in the network. We call such properties *features*. We then try to estimate the posterior probability of features given the data. More precisely, a feature f is an indicator function that receives a network structure G as an argument and returns 1 if the structure (or the associated PDAG) satisfies the feature and 0 otherwise. The posterior probability of a feature is

$$P(f(G) \mid D) = \sum_{G} f(G)P(G \mid D). \tag{2}$$

Of course, exact computation of such a posterior probability is as hard as processing all networks with high posteriors. However, as we shall see below, we can estimate these posteriors by finding representative networks. Since each feature is a binary attribute, this estimation is fairly robust even from a small set of networks (assuming that they are an unbiased sample from the posterior).

Before we examine the issue of estimating the posterior of such features, we briefly discuss two classes of features involving pairs of variables. While at this point we handle only pairwise features, it is clear that this type of analysis is not restricted to them, and in the future we are planning to examine more complex features.

The first type of feature is *Markov relations*: Is Y in the *Markov blanket* of X? The Markov blanket of X is the minimal set of variables that shield X from the rest of the variables in the model. More precisely, X given its Markov blanket is independent from the remaining variables in the network. It is easy to check that this relation is symmetric: Y is in X's Markov blanket if and only if there is either an edge between them or both are parents of another variable (Pearl, 1988). In the context of gene expression analysis, a Markov relation indicates that the two genes are related in some joint biological interaction or process. Note that two variables in a Markov relation are directly linked in the sense that no variable in the model mediates the dependence between them. It remains possible that an unobserved variable (e.g., protein activation) is an intermediate in their interaction.

The second type of feature is *order relations*: Is X an ancestor of Y in all the networks of a given equivalence class? That is, does the given PDAG contain a path from X to Y in which all the edges are directed? This type of feature does not involve only a close neighborhood, but rather captures a global property. Recall that under the assumptions discussed in Section 2.4, learning that X is an ancestor of Y would imply that X is a cause of Y. However, these assumptions are quite strong (in particular the assumption of no latent common causes) and thus do not necessarily hold in the context of expression data. Thus, we view such a relation as an indication, rather than evidence, that X might be a causal ancestor of Y.

3.3. Estimating statistical confidence in features

We now face the following problem: To what extent does the data support a given feature? More precisely, we want to estimate the posterior of features as defined in (2). Ideally, we would like to sample networks from the posterior and use the sampled networks to estimate this quantity. Unfortunately, sampling from the posterior is a hard problem. The general approach to this problem is to build a *Markov Chain Monte*

²This observation is not unique to Bayesian network models. It equally well applies to other models that are learned from gene expression data, such as clustering models.

Carlo (MCMC) sampling procedure (see Madigan and York, 1995, and Gilks et al., 1996, for a general introduction to MCMC sampling). However, it is not clear how these methods scale up for large domains.

Although recent developments in MCMC methods for Bayesian networks, such as Friedman and Koller (2000), show promise for scaling up, we choose here to use an alternative method as a "poor man's" version of Bayesian analysis. An effective, and relatively simple, approach for estimating confidence is the *bootstrap* method (Efron and Tibshirani, 1993). The main idea behind the bootstrap is simple. We generate "perturbed" versions of our original data set and learn from them. In this way we collect many networks, all of which are fairly reasonable models of the data. These networks reflect the effect of small perturbations to the data on the learning process.

In our context, we use the bootstrap as follows:

- For i = 1...m.
 Construct a data set D_i by sampling, with replacement, N instances from D.
 Apply the learning procedure on D_i to induce a network structure G_i.
- For each feature f of interest calculate

$$\operatorname{conf}(f) = \frac{1}{m} \sum_{i=1}^{m} f(G_i)$$

where f(G) is 1 if f is a feature in G, and 0 otherwise.

See Friedman, Goldszmidt, and Wyner (1999) for more details, as well as for large-scale simulation experiments with this method. These simulation experiments show that features induced with high confidence are rarely false positives, even in cases where the data sets are small compared to the system being learned. This bootstrap procedure appears especially robust for the Markov and order features described in Section 3.2. In addition, simulation studies by Friedman and Koller (2000) show that the confidence values computed by the bootstrap, although not equal to the Bayesian posterior, correlate well with estimates of the Bayesian posterior for features.

3.4. Efficient learning algorithms

In Section 2.3, we formulated learning Bayesian network structure as an optimization problem in the space of directed acyclic graphs. The number of such graphs is superexponential in the number of variables. As we consider hundreds of variables, we must deal with an extremely large search space. Therefore, we need to use (and develop) efficient search algorithms.

To facilitate efficient learning, we need to be able to focus the attention of the search procedure on relevant regions of the search space, giving rise to the *Sparse Candidate* algorithm (Friedman, Nachman, and Pe'er, 1999). The main idea of this technique is that we can identify a relatively small number of *candidate* parents for each gene based on simple local statistics (such as correlation). We then restrict our search to networks in which only the candidate parents of a variable can be its parents, resulting in a much smaller search space in which we can hope to find a good structure quickly.

A possible pitfall of this approach is that early choices can result in an overly restricted search space. To avoid this problem, we devised an iterative algorithm that adapts the candidate sets during search. At each iteration n, for each variable X_i , the algorithm chooses the set $C_i^n = \{Y_1, \ldots, Y_k\}$ of variables which are the most promising candidate parents for X_i . We then search for G_n , a high scoring network in which $\mathbf{Pa}^{G_n}(X_i) \subseteq C_i^n$. (Ideally, we would like to find the highest-scoring network given the constraints, but since we are using a heuristic search, we do not have such a guarantee.) The network found is then used to guide the selection of better candidate sets for the next iteration. We ensure that the score of G_n monotonically improves in each iteration by requiring $\mathbf{Pa}^{G_{n-1}}(X_i) \subseteq C_i^n$. The algorithm continues until there is no change in the candidate sets.

We briefly outline our method for choosing C_i^n . We assign to each variable X_j some score of relevance to X_i , choosing variables with the highest score. The question is then how to measure the relevance of potential parent X_j to X_i . Friedman *et al.* (1999) examine several measures of relevance. Based on their experiments, one of the most successful measures is simply the improvement in the score of X_i if we add X_j as an additional parent. More precisely, we calculate

This quantity measures how much the inclusion of an edge from X_j to X_i can improve the score associated with X_i . We then choose the new candidate set to contain the previous parent set $\mathbf{Pa}^{G_{n-1}}(X_i)$ and the variables that are most informative given this set of parents.

We refer the reader to Friedman, Nachman, and Pe'er (1999) for more details on the algorithm and its complexity, as well as empirical results comparing its performance to traditional search techniques.

3.5. Local probability models

In order to specify a Bayesian network model, we still need to choose the type of the local probability models we learn. In the current work, we consider two approaches:

- **Multinomial model.** In this model we treat each variable as discrete and learn a multinomial distribution that describes the probability of each possible state of the child variable given the state of its parents.
- Linear Gaussian model. In this model we learn a linear regression model for the child variable given its parents.

These models were chosen since their posterior can be efficiently calculated in closed form.

To apply the multinomial model we need to discretize the gene expression values. We choose to discretize these values into three categories: underexpressed (-1), normal (0), and overexpressed 1, depending on whether the expression rate is significantly lower than, similar to, or greater than control, respectively. The control expression level of a gene either can be determined experimentally (as in the methods of DeRisi. $et\ al.$ (1997)) or it can be set as the average expression level of the gene across experiments. We discretize by setting a threshold to the ratio between measured expression and control. In our experiments, we choose a threshold value of 0.5 in logarithmic (base 2) scale. Thus, values with ratio to control lower than $2^{-0.5}$ are considered underexpressed, and values higher than $2^{0.5}$ are considered overexpressed.

Each of these two models has benefits and drawbacks. On one hand, it is clear that by discretizing the measured expression levels we are loosing information. The linear-Gaussian model does not suffer from the information loss caused by discretization. On the other hand, the linear-Gaussian model can only detect dependencies that are close to linear. In particular, it is not likely to discover combinatorial effects (e.g., a gene is over-expressed only if several genes are jointly over-expressed, but not if at least one of them is not under-expressed). The multinomial model is more flexible and can capture such dependencies.

4. APPLICATION TO CELL CYCLE EXPRESSION PATTERNS

We applied our approach to the data of Spellman *et al.* (1998). This data set contains 76 gene expression measurements of the mRNA levels of 6177 *S. cerevisiae* ORFs. These experiments measure six time series under different cell cycle synchronization methods. Spellman *et al.* (1998) identified 800 genes whose expression varied over the different cell-cycle stages.

In learning from this data, we treat each measurement as an independent sample from a distribution and do not take into account the temporal aspect of the measurement. Since it is clear that the cell cycle process is of a temporal nature, we compensate by introducing an additional variable denoting the cell cycle phase. This variable is forced to be a root in all the networks learned. Its presence allows one to model dependency of expression levels on the current cell cycle phase.³

We used the Sparse Candidate algorithm with a 200-fold bootstrap in the learning process. We performed two experiments, one with the discrete multinomial distribution, the other with the linear Gaussian distribution. The learned features show that we can recover intricate structure even from such small data sets. It is important to note that our learning algorithm uses no prior biological knowledge nor constraints. All learned networks and relations are based solely on the information conveyed in the measurements themselves. These results are available at our WWW site: http://www.cs.huji.ac.il/labs/compbio/expression. Figure 2 illustrates the graphical display of some results from this analysis.

³We note that we can learn temporal models using a Bayesian network that includes gene expression values in two (or more) consecutive time points (Friedman *et al.*, 1998). This raises the number of variables in the model. We are currently perusing this issue.

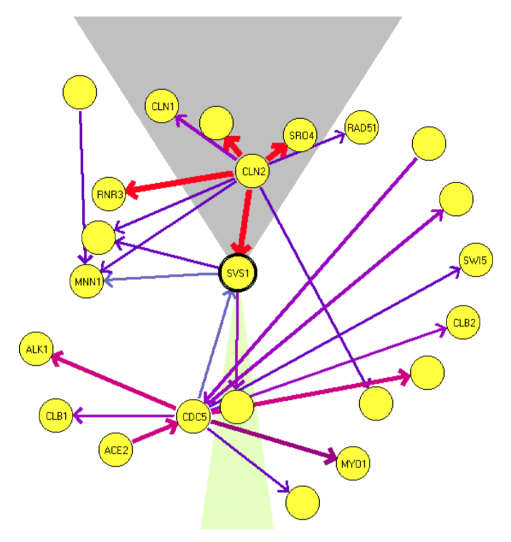
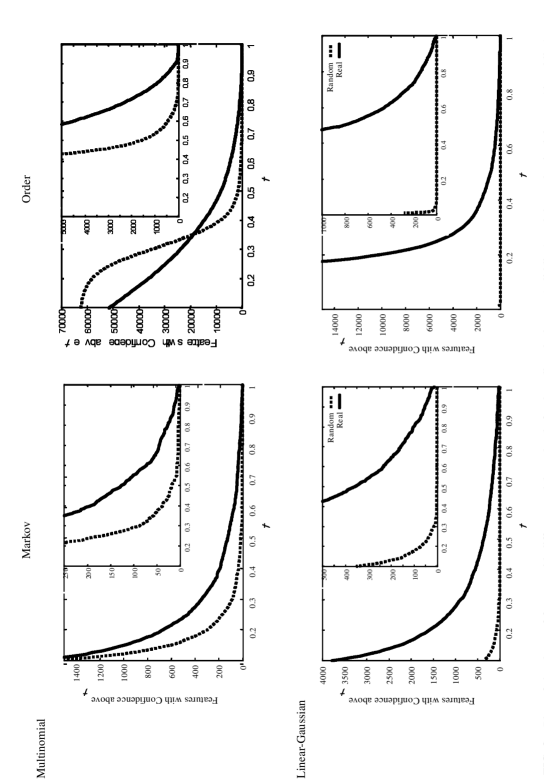


FIG. 2. An example of the graphical display of Markov features. This graph shows a "local map" for the gene SVS1. The width (and color) of edges corresponds to the computed confidence level. An edge is directed if there is a sufficiently high confidence in the order between the genes connected by the edge. This local map shows that CLN2 separates SVS1 from several other genes. Although there is a strong connection between CLN2 to all these genes, there are no other edges connecting them. This indicates that, with high confidence, these genes are conditionally independent given the expression level of CLN2.

4.1. Robustness analysis

We performed a number of tests to analyze the statistical significance and robustness of our procedure. Some of these tests were carried out on a smaller data set with 250 genes for computational reasons.

To test the credibility of our confidence assessment, we created a random data set by randomly permuting the order of the experiments independently for each gene. Thus for each gene the order was random, but the composition of the series remained unchanged. In such a data set, genes are independent of each other, and thus we do not expect to find "real" features. As expected, both order and Markov relations in the random data set have significantly lower confidence. We compare the distribution of confidence estimates between the original data set and the randomized set in Figure 3. Clearly, the distribution of confidence estimates in the original data set have a longer and heavier tail in the high confidence region. In the linear-Gaussian model we see that random data does not generate any feature with confidence above 0.3. The multinomial model is more expressive and thus susceptible to over-fitting. For this model, we see a smaller gap between the two distributions. Nonetheless, randomized data does not generate any feature with confidence above 0.8, which leads us to believe that most features that are learned in the original data set with such confidence are not artifacts of the bootstrap estimation.



x-value. The graphs on the left column show Markov features, and the ones on the right column show Order features. The top row describes features found using the multinomial model, and the bottom row describes features found by the linear-Gaussian model. Inset in each graph is a plot of the tail The x-axis denotes the conpdence threshold, and the y-axis denotes the number of features with conpdence equal or higher than the corresponding FIG. 3. Plots of abundance of features with different conbdence levels for the cell cycle data set (solid line), and the randomized data set (dotted line). of the distribution.

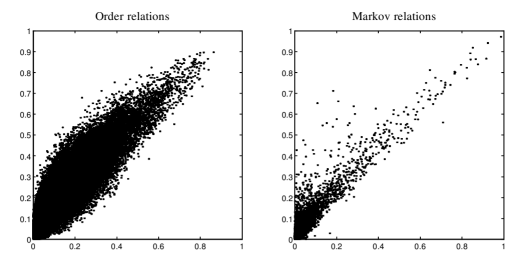


FIG. 4. Comparison of confidence levels obtained in two data sets differing in the number of genes, on the multinomial experiment. Each relation is shown as a point, with the x-coordinate being its confidence in the the 250 genes data set and the y-coordinate the confidence in the 800 genes data set. The left figure shows order relation features, and the right figure shows Markov relation features.

To test the robustness of our procedure for extracting dominant genes, we performed a simulation study. We created a data set by sampling 80 samples from one of the networks learned from the original data. We then applied the bootstrap procedure on this data set. We counted the number of descendents of each node in the synthetic network and ordered the variables according to this count. Of the 10 top dominant genes learned from the bootstrap experiment, 9 were among the top 30 in the real ordering.

Since the analysis was not performed on the whole *S. cerevisiae* genome, we also tested the robustness of our analysis to the addition of more genes, comparing the confidence of the learned features between the 800-gene data set and a smaller 250-gene data set that contains genes appearing in eight major clusters described by Spellman *et al.* (1998). Figure 4 compares feature confidence in the analysis of the two data sets for the multinomial model. As we can see, there is a strong correlation between confidence levels of the features between the two data sets. The comparison for the linear-Gaussian model gives similar results.

A crucial choice for the multinomial experiment is the threshold level used for discretization of the expression levels. It is clear that by setting a different threshold we would get different discrete expression patterns. Thus, it is important to test the robustness and sensitivity of the high-confidence features to the choice of this threshold. This was tested by repeating the experiments using different thresholds. The comparison of how the change of threshold affects the confidence of features show a definite linear tendency in the confidence estimates of features between the different discretization thresholds (graphs not shown). Obviously, this linear correlation gets weaker for larger threshold differences. We also note that order relations are much more robust to changes in the threshold than are Markov relations.

A valid criticism of our discretization method is that it penalizes genes whose natural range of variation is small: since we use a fixed threshold, we would not detect changes in such genes. A possible way to avoid this problem is to *normalize* the expression of genes in the data. That is, we rescale the expression level of each gene so that the relative expression level has the same mean and variance for all genes. We note that analysis methods that use *Pearson correlation* to compare genes, such as those of Ben-Dor *et al.* (1999) and Eisen *et al.* (1999), implicitly perform such a normalization. When we discretize a normalized data set, we are essentially rescaling the discretization factor differently for each gene, depending on its variance in the data. We tried this approach with several discretization levels, and got results comparable

⁴An undesired effect of such a normalization is the amplification of measurement noise. If a gene has fixed expression levels across samples, we expect the variance in measured expression levels to be noise either in the experimental conditions or the measurements. When we normalize the expression levels of genes, we lose the distinction between such noise and true (i.e., significant) changes in expression levels. In the Spellman *et al.* data set we can safely assume this effect will not be too grave, since we only focus on genes that display significant changes across experiments.

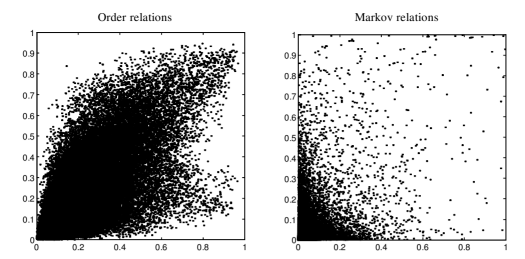


FIG. 5. Comparison of confidence levels between the multinomial experiment and the linear-Gaussian experiment. Each relation is shown as a point, with the *x*-coordinate being its confidence in the multinomial experiment, and the *y*-coordinate its confidence in the linear-Gaussian experiment. The left figure shows order relation features, and the right figure shows Markov relation features.

to our original discretization method. The 20 top Markov relations highlighted by this method were a bit different, but interesting and biologically sensible in their own right. The order relations were again more robust to the change of methods and discretization thresholds. A possible reason is that order relations depend on the network structure in a global manner and thus can remain intact even after many local changes to the structure. The Markov relation, being a local one, is more easily disrupted. Since the graphs learned are extremely sparse, each discretization method "highlights" different signals in the data, which are reflected in the Markov relations learned.

A similar picture arises when we compare the results of the multinomial experiment to those of the linear-Gaussian experiment (Figure 5). In this case, there is virtually no correlation between the Markov relations found by the two methods, while the order relations show some correlation. This supports our assumption that the two methods highlight different types of connections between genes.

Finally, we consider the effect of the choice of prior on the learned features. It is important to ensure that the learned features are not simply artifacts of the chosen prior. To test this, we repeated the multinomial experiment with different values of K, the effective sample size, and compared the learned confidence levels to those learned with the default value used for K, which was 5. This was done using the 250-gene data set and discretization level of 0.5. The results of these comparisons are shown in Figure 6. As can be seen, the confidence levels obtained with a K value of 1 correlate very well with those obtained with the default K, while when setting K to 20 the correlation is weaker. This suggests that both 1 and 5 are low enough values compared to the data set size of 76, making the prior's affect on the results weak. An effective sample size of 20 is high enough to make the prior's effect noticeable. Another aspect of the prior is the prior network used. In all the experiments reported here, we used the empty network with uniform distribution parameters as the prior network. As our prior is noninformative, keeping down its effect is desired. It is expected that once we use more informative priors (by incorporating biological knowledge, for example) and stronger effective sample sizes, the obtained results will be biased more toward our prior beliefs.

In summary, although many of the results we report below (especially order relations) are stable across the different experiments discussed in the previous paragraph, it is clear that our analysis is sensitive to the choice of local model and, in the case of the multinomial model, to the discretization method. It is probably less sensitive to the choice of prior, as long as the effective sample size is small compared to the data set size. In all the methods we tried, our analysis found interesting relationships in the data. Thus, one challenge is to find alternative methods that can recover all these relationships in one analysis. We are currently working on learning networks with semi-parametric density models (Friedman and Nachman, 2000; Hoffman and Tresp, 1996) that would circumvent the need for discretization on one hand and allow nonlinear dependency relations on the other.

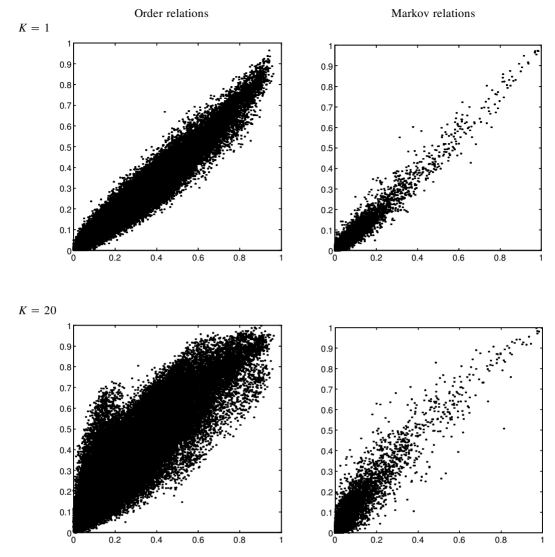


FIG. 6. Comparison of confidence levels between runs using different parameter priors. The difference in the priors is in the effective sample size, K. Each relation is shown as a point, with the x-coordinate being its confidence in a run with K = 5, and the y-coordinate its confidence in a run with K = 1 (top row) and K = 20 (bottom row). The left figure shows order relation features, and the right figure shows Markov relation features. All runs are on the 250 gene set, using discretization with threshold level 0.5.

4.2. Biological analysis

We believe that the results of this analysis can be indicative of biological phenomena in the data. This is confirmed by our ability to predict sensible relations between genes of known function. We now examine several consequences that we have learned from the data. We consider, in turn, the order relations and Markov relations found by our analysis.

4.2.1. Order relations. The most striking feature of the high-confidence order relations, is the existence of dominant genes. Out of all 800 genes, only a few seem to dominate the order (i.e., appear before many genes). The intuition is that these genes are indicative of potential causal sources of the cell-cycle process. Let $C_o(X, Y)$ denote the confidence in X being an ancestor of Y. We define the dominance score of X as $\sum_{Y,C_o(X,Y)>t} C_o(X,Y)^k$, using the constant k for rewarding high confidence features and the threshold t to discard low confidence ones. These dominant genes are extremely robust to parameter selection for both t and k, the discretization cutoff of Section 3.5, and the local probability model used. A list of the highest-scoring dominating genes for both experiments appears in Table 1.

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TABLE 1	LICT OF DOMINANT	CENEC IN THE	Ordering Relations ¹
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	Score in experiment			
Gene/ORF	Multinomial	Gaussian	Notes	
MCD1	550	525	Mitotic Chromosome Determinant, null mutant is inviable	
MSH6	292	508	Required for mismatch repair in mitosis and meiosis	
CSI2	444	497	Cell wall maintenance, chitin synthesis	
CLN2	497	454	Role in cell cycle START, null mutant exhibits G1 arrest	
YLR183C	551	448	Contains forkheaded associated domain, thus possibly nuclear	
RFA2	456	423	Involved in nucleotide excision repair, null mutant is inviable	
RSR1	352	395	GTP-binding protein of the RAS family involved in bud site selection	
CDC45	_	394	Required for initiation of chromosomal replication, null mutant lethal	
RAD53	60	383	Cell cycle control, checkpoint function, null mutant lethal	
CDC5	209	353	Cell cycle control, required for exit from mitosis, null mutant lethal	
POL30	376	321	Required for DNA replication and repair, null mutant is inviable	
YOX1	400	291	Homeodomain protein	
SRO4	463	239	Involved in cellular polarization during budding	
CLN1	324	_	Role in cell cycle START, null mutant exhibits G1 arrest	
YBR089W	298	_		

¹Included are the top 10 dominant genes for each experiment.

Inspection of the list of dominant genes reveals quite a few interesting features. Among them are genes directly involved in initiation of the cell cycle and its control. For example, CLN1, CLN2, CDC5, and RAD53 whose functional relation has been established (Cvrckova and Nasmyth, 1993; Drebot *et al.*, 1993). The genes MCD1, RFA2, CDC45, RAD53, CDC5, and POL30 were found to be essential (Guacci *et al.*, 1997). These are clearly key genes in essential cell functions. Some of them are components of prereplication complexes(CDC45,POL30). Others (like RFA2,POL30, and MSH6) are involved in DNA repair. It is known that DNA repair is associated with transcription initiation, and DNA areas which are more active in transcription are also repaired more frequently (McGregor, 1999; Tornaletti and Hanawalk, 1999). Furthermore, a cell cycle control mechanism causes an abort when the DNA has been improperly replicated (Eisen and Lucchesi, 1998).

Most of the dominant genes encode nuclear proteins, and some of the unknown genes are also potentially nuclear (e.g., YLR183C contains a forkhead-associated domain which is found almost entirely among nuclear proteins). A few nonnuclear dominant genes are localized in the cytoplasm membrane (SRO4 and RSR1). These are involved in the budding and sporulation processes which have an important role in the cell cycle. RSR1 belongs to the RAS family of proteins, which are known as initiators of signal transduction cascades in the cell.

4.2.2. Markov relations. We begin with an analysis of the Markov relations in the multinomial experiment. Inspection of the top Markov relations reveals that most are functionally related. A list of the top scoring relations can be found in Table 2. Among these, all those involving two known genes make sense biologically. When one of the ORFs is unknown, careful searches using Psi-Blast (Altschul et al., 1997), Pfam (Sonnhammer et al., 1998), and Protomap (Yona et al., 1998) can reveal firm homologies to proteins functionally related to the other gene in the pair. For example YHR143W, which is paired to the endochitinase CTS1, is related to EGT2—a cell-wall-maintenance protein. Several of the unknown pairs are physically adjacent on the chromosome and thus presumably regulated by the same mechanism (see Blumenthal (1998)), although special care should be taken for pairs whose chromosomal location overlap on complementary strands, since in these cases we might see an artifact resulting from cross-hybridization. Such an analysis raises the number of biologically sensible pairs to nineteen out of the twenty top relations.

Some interesting Markov relations were found that are beyond the limitations of clustering techniques. Among the high-confidence Markov relations, one can find examples of conditional independence, i.e., a group of highly correlated genes whose correlation can be explained within our network structure. One such example involves the genes CLN2,RNR3,SVS1,SRO4, and RAD51. Their expression is correlated, and in Spellman *et al.* (1998) they all appear in the same cluster. In our network CLN2 is with high confidence a parent of each of the other 4 genes, while no links are found between them (see Figure 2). This agrees with biological knowledge: CLN2 is a central and early cell cycle control, while there is

Confidence	Gene 1	Gene 2	Notes
1.0	YKL163W-PIR3	YKL164C-PIR1	Close locality on chromosome
0.985	PRY2	YKR012C	Close locality on chromosome
0.985	MCD1	MSH6	Both bind to DNA during mitosis
0.98	PHO11	PHO12	Both nearly identical acid phosphatases
0.975	HHT1	HTB1	Both are histones
0.97	HTB2	HTA1	Both are histones
0.94	YNL057W	YNL058C	Close locality on chromosome
0.94	YHR143W	CTS1	Homolog to EGT2 cell wall control, both involved in cytokinesis
0.92	YOR263C	YOR264W	Close locality on chromosome
0.91	YGR086	SIC1	Homolog to mammalian nuclear ran protein, both involved in nuclear function
0.9	FAR1	ASH1	Both part of a mating type switch, expression uncorrelated
0.89	CLN2	SVS1	Function of SVS1 unknown
0.88	YDR033W	NCE2	Homolog to transmembrame proteins suggest both involved in protein secretion
0.86	STE2	MFA2	A mating factor and receptor
0.85	HHF1	HHF2	Both are histones
0.85	MET10	ECM17	Both are sulfite reductases
0.85	CDC9	RAD27	Both participate in Okazaki fragment processing

TABLE 2. LIST OF TOP MARKOV RELATIONS, MULTINOMIAL EXPERIMENT

no clear biological relationship between the others. Some of the other Markov relations are intercluster pairing genes with low correlation in their expression. One such regulatory link is FAR1–ASH1: both proteins are known to participate in a mating-type switch. The correlation of their expression patterns is low and Spellman *et al.* (1998) cluster them into different clusters. When looking further down the list for pairs whose Markov relation confidence is high relative to their correlation, interesting pairs surface, for example, SAG1 and MF-ALPHA-1, a match between the factor that induces the mating process and an essential protein that participates in the mating process. Another match is LAC1 and YNL300W. LAC1 is a GPI transport protein and YNL300W is most likely modified by GPI (based on sequence homology).

The Markov relations from the Gaussian experiment are summarized in Table 3. Since the Gaussian model focuses on highly correlated genes, most of the high-scoring genes are tightly correlated. When we checked the DNA sequence of pairs of physically adjacent genes at the top of Table 3, we found that there is significant overlap. This suggests that these correlations are spurious and due to *cross hybridization*. Thus, we ignore the relations with the highest score. However, in spite of this technical problem, few of the pairs with a confidence > 0.8 can be discarded as biologically false.

Some of the relations are robust and also appear in the multinomial experiment (e.g., STE2–MFA2, CST1–YHR143W). Most interesting are the genes linked through regulation. These include: SHM2 which converts glycine that regulates GCV2 and DIP5 which transports glutamate which regulates ARO9. Some pairs participate in the same metabolic process, such as: CTS1–YHR143 and SRO4–YOL007C, all which participate in cell wall regulation. Other interesting high-confidence (> 0.9) examples are: OLE1–FAA4 linked through fatty acid metabolism, STE2–AGA2 linked through the mating process, and KIP3–MSB1, both playing a role in polarity establishment.

5. DISCUSSION AND FUTURE WORK

In this paper we presented a new approach for analyzing gene expression data that builds on the theory and algorithms for learning Bayesian networks. We described how to apply these techniques to gene expression data. The approach builds on two techniques that were motivated by the challenges posed by this domain: a novel search algorithm (Friedman, Nachman, and Pe'er, 1999) and an approach for estimating statistical confidence (Friedman, Goldszmidt, and Wyner, 1999). We applied our methods to the real expression data of Spellman *et al.* (1998). Although, we did not use any prior knowledge, we managed to extract many biologically plausible conclusions from this analysis.

TABLE 3	LIST OF	TOP MARKOV	RELATIONS,	Gaussian	Experiment ¹	
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Confidence	Gene 1	Gene 2	Notes
1.0	YOR263C	YOR264W	Close locality on chromosome
1.0	CDC46	YOR066W	YOR066W is totally unknown
1.0	CDC45	SPH1	No suggestion for immediate link
1.0	SHM2	GCV2	SHM2 interconverts glycine, GCV2 is regulated by glycine
1.0	MET3	ECM17	MET3 required to convert sulfate to sulfide, ECM17 sulfite reductase
1.0	YJL194W-CDC6	YJL195C	Close locality on chromosome
1.0	YGR151C	YGR152C	Close locality on chromosome
1.0	YGR151C	YGR152C-RSR1	Close locality on chromosome
1.0	STE2	MFA2	A mating factor and receptor
1.0	YDL037C	YDL039C	Both homologs to mucin proteins
1.0	YCL040W-GLK1	WCL042C	Close locality on chromosome
1.0	HTA1	HTA2	Two physically linked histones
0.99	HHF2	HHT2	both histones
0.99	YHR143W	CTS1	Homolog to EGT2 cell wall control, both involved in cytokinesis
0.99	ARO9	DIP5	DIP5 transports glutamate which regulates ARO9
0.975	SRO4	YOL007C	Both proteins are involved in cell wall regulation at the plasma membrane

¹The table skips over 5 additional pairs with which close locality.

Our approach is quite different from the clustering approach used by Alon *et al.* (1999), Ben-Dor *et al.* (1999), Eisen *et al.* (1999), Michaels *et al.* (1998), and Spellman *et al.* (1998) in that it attempts to learn a much richer structure from the data. Our methods are capable of discovering causal relationships, interactions between genes other than positive correlation, and finer intracluster structure. We are currently developing hybrid approaches that combine our methods with clustering algorithms to learn models over "clustered" genes.

The biological motivation of our approach is similar to the work on inducing *genetic networks* from data (Akutsu *et al.*, 1998; Chen *et al.*, 1999; Somogyi *et al.*, 1996; Weaver *et al.*, 1999). There are two key differences: First, the models we learn have probabilistic semantics. This better fits the stochastic nature of both the biological processes and noisy experiments. Second, our focus is on extracting features that are pronounced in the data, in contrast to current genetic network approaches that attempt to find a single model that explains the data.

We emphasize that the work described here represents a preliminary step in a longer-term project. As we have seen above, there are several points that require accurate statistical tools and more efficient algorithms. Moreover, the exact biological conclusions one can draw from this analysis are still not well understood. Nonetheless, we view the results described in Section 4 as definitely encouraging.

We are currently working on improving methods for expression analysis by expanding the framework described in this work. Promising directions for such extensions are: (a) developing the theory for learning local probability models that are suitable for the type of interactions that appear in expression data; (b) improving the theory and algorithms for estimating confidence levels; (c) incorporating biological knowledge (such as possible regulatory regions) as prior knowledge to the analysis; (d) improving our search heuristics; (e) learning temporal models, such as *Dynamic Bayesian Networks* (Friedman *et al.*, 1998), from temporal expression data, and (f) developing methods that discover *hidden variables* (e.g., protein activation).

Finally, one of the most exciting longer-term prospects of this line of research is discovering causal patterns from gene expression data. We plan to build on and extend the theory for learning causal relations from data and apply it to gene expression. The theory of causal networks allows learning from both observational and *interventional* data, where the experiment intervenes with some causal mechanisms of the observed system. In the gene expression context, we can model knockout/over-expressed mutants as such interventions. Thus, we can design methods that deal with mixed forms of data in a principled manner (see Cooper and Yoo (1999) for recent work in this direction). In addition, this theory can provide

tools for *experimental design*, that is, understanding which interventions are deemed most informative to determining the causal structure in the underlying system.

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