

Mining Functional Modules in Genetic Networks with Decomposable Graphical Models

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Short title: Mining functional genetic modules

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Abstract:

In recent years graphical models have become an increasingly important tool for the structural analysis of genome-wide expression profiles at the systems level. Here we present a new graphical modelling technique, which is based on decomposable graphical models, and apply it to a set of gene expression profiles from acute lymphoblastic leukemia (ALL). The new method explains probabilistic dependencies of expression levels in terms of the concerted action of underlying genetic functional modules, which are represented as so-called “cliques” in the graph. In addition, the method uses continuous-valued (instead of discretized) expression levels, and makes no particular assumption about their probability distribution. We show that the method successfully groups members of known functional modules to cliques. Our method allows the evaluation of the importance of genes for global cellular functions based on both link count and the clique membership count.

1. Introduction

With the development of genome-wide expression measurements by DNA microarrays (Brown and Botstein, 1999) and related techniques, it became possible to gain a systems level view of the instantaneous state of the genetic regulatory network controlling most of the cellular processes. An increasing amount of modelling work focuses on the extraction of functional relationships from the generated database which consists of high-dimensional gene expression measurements (for reviews, see Berrar et al., 2002; Stetter et al., 2003, 2004). The ultimate goal is to gain a better quantitative understanding of the biological processes which emerge from the dense interplay of the genes, the transcription and the proteome at the level of large subcellular systems or even whole cells. One prominent recent class of approaches adopts graphical modelling techniques to efficiently learn statistical relationships between the expression levels of many genes. From these relationships one can then try to infer the underlying biological functional links (Friedman et al., 2000; Hartemink et al., 2001; Imoto et al., 2002; Berrar et al., 2002; Dejori and Stetter, 2003; Segal et al., 2003.; Dejori et al., 2004). The underlying hypothesis is that a more quantitative understanding of the structure and function of the genetic regulatory network will be the basis for entirely new and systematic approaches to produce highly efficient disease markers, for drug target finding and drug discovery, for tissue engineering and many other fields.

In this paper we address two specific issues of relevance to the systems-level modelling of microarray data using graphical models. The first issue is concerned with the fact, that gene expression levels are often discretized during preprocessing. It is clearly unsatisfactory to learn graphical models on continuous variables by discretization, since discretization might mask statistical relationships in the data. In our approach we are able to model arbitrary non-Gaussian continuous densities by employing *kernel density estimators* to model the statistical distributions.

The second issue is concerned with the modularity, which is considered an important feature of biological organization (Hartwell et al., 1999). For example, genetic and protein networks often show scale-free topologies (Jeong et al., 2000), which implies that they decompose into sets of densely connected gene clusters (Watts and Strogatz, 1998). One reason for this cliquishness might be that many proteins consist of several subunits, encoded by different genes, and become only functional when the corresponding genes are all expressed. Similarly, proteins can form larger assemblies and carry out their function only in a particular assembly. Finally, proteins can also be part of reaction cascades, which sub-serve a common task. In summary, gene products are often naturally linked to *functional modules*. Hence, expressing genes usually means expressing groups of genes necessary to manufacture a functional module. Our analysis is tailored to account for this modular structure: The basis of our approach is a *decomposable graphical model*, which is capable to efficiently learn the graphical structure of continuous data. The decomposable model can be represented as a *clique tree*, where each clique represents a group of genes (the nodes of the clique), which are fully connected. This dense statistical dependency structure is thought to reflect the dense biological link of a functional module causing the gene expression patterns. By its structure, the decomposable model explains the statistical structure of the data in terms of interacting functional modules (represented by cliques), instead of only the functionally linked genes (the nodes). Hence, our approach has the potential to truthfully reflect the modular structure of gene expression.

We proceed by first giving a brief introduction to the decomposable model approach in section 2, and apply this technique to estimate the structure of functional modules in genetic regulatory networks involved in the pathogenesis of childhood acute lymphoblastic leukaemia (ALL, Yeoh et al., 2002).

2. Materials and Methods

We apply decomposable graphical models, which – by their inherently clique-like structure - mirror the modular architecture inherent to genetic networks in a natural way. Since our knowledge about the structure of underlying gene-gene relationships is incomplete, we learn this structure from data. Decomposable graphical models and structural learning therein have been described in detail elsewhere (Pearl, 1988; Lauritzen, 1996; Hofmann and Tresp, 1998; Cowell et al., 1999; Schwaighofer et al., 2004). Hence, because the algorithm we use here has been introduced in another related study (Schwaighofer et al., 2004), we will in the following provide only a brief summarizing description of it.

2.1 Decomposable Models

A graphical model (or, probabilistic network) describes a family of joint probability distributions in form of a graph (see figure 2a, b for illustrative examples of joint probability densities). The nodes $\mathbf{x} = (x_1, \dots, x_n)$ in the graph represent random variables and an edge between two nodes represents direct statistical dependencies (figure 1c). In the following we only consider graphs with undirected edges. Let E be the set of all edges in the graph. The absence of an edge between variables x_i and x_j implies that both variables are statistically independent, conditioned on all other random variables in the domain, i.e. that there is no direct dependency between the variables except the ones which are mediated by the remaining nodes in the network. In our application each variable x_i represents the expression level of one gene and edges mark statistical dependencies between the corresponding pairs of gene expression levels. The set of nodes \mathbf{x} and the set of edges E form the graph structure G .

Figure 1 about here

In the following we will only consider a particular subclass of undirected graphical models, so-called *decomposable models*. They have the particular property that each cycle of four nodes has a *cord*, i.e. an edge linking two non-adjacent nodes in the cycle. Practically speaking, in a decomposable model the variables can be grouped into overlapping subsets of fully linked nodes. Any graphical model can be transformed into a decomposable model by adding a sufficient number of edges. For inference and for the calculation of the likelihood function we need the notion of a *clique*, which is defined as a maximal subgroup of nodes, which are mutually fully connected. A *clique tree* of a decomposable model is a particular tree in which the cliques form the nodes. Each edge in the tree represents a *separator* which contains the nodes common to the cliques linked by the edge. Let Σ be the set of all separators. Figure 1c shows an example of a small undirected graph for a decomposable model of 7 random variables, together with one equivalent clique tree in Figure 1d.

As their most important property, decomposable models allow the joint probability density to be written in terms of marginal densities of the random variables contained in a clique (cf figure 1a for the definition of joint and marginal probability densities). More precisely, the joint probability density in a decomposable model factorizes: it can be written as a product of simpler probability densities of the form

$$p(\mathbf{x}) = \frac{\prod_{C \in \mathcal{K}} p(\mathbf{x}_C)}{\prod_{S \in \Sigma} p(\mathbf{x}_S)}, \quad (1)$$

where \mathbf{x}_C denotes the subset of variables x_i (i.e., the components of \mathbf{x}) that form clique C and \mathbf{x}_S denotes the subset of variables x_j that form separator S . The major advantage of this formulation is that the marginal densities over cliques and separators are usually much lower-dimensional than the full joint probability density of all involved variables.

The goal of graphical modelling is to identify and describe significant statistical dependencies in a finite data set $D = (\mathbf{x}^1, \dots, \mathbf{x}^m)$ where here each data sample \mathbf{x}^m represents one DNA microarray measurement. For this purpose one needs to tackle two problems: (i), find the structure of a clique tree, which is suitable for describing the dependency structure (structure learning), and (ii) find suitable descriptions for the marginal densities $p(\mathbf{x}_C)$ and $p(\mathbf{x}_S)$ within this structure. In practice, these problems are solved simultaneously.

2.2 Nonparametric Density Estimation in Decomposable Models

In our work we use a nonparametric probability density estimate - namely a *kernel density estimator* - for modelling the marginal clique and separator densities. A kernel density estimator consists of the superposition of m kernel functions g centered at each data point:

$$p(\mathbf{x} | D, \Theta) = \frac{1}{m} \sum_{i=1}^m g(\mathbf{x}; \mathbf{x}^i, \Theta). \quad (2)$$

We use a Gaussian kernel with diagonal covariance matrix:

$$g(\mathbf{x}; \mathbf{x}^i, \Theta) = \frac{1}{(2\pi)^{n/2} |\text{diag } \Theta|^{1/2}} \exp\left(-\frac{1}{2} (\mathbf{x} - \mathbf{x}^i)^T (\text{diag } \Theta)^{-1} (\mathbf{x} - \mathbf{x}^i)\right). \quad (3)$$

In eq. (3), Θ denotes the vector of variances along the n dimensions of the data space. It is worth noting that although the kernel function is Gaussian, the nonparametric probability density estimate eq. (2) is a superposition of Gaussians and can in principle (i.e., in the limit of many data points) describe general probability density functions. This is an important extension to earlier approaches, which were restricted to Gaussian densities: the parametric form of the probability densities of microarray experiment data is not known but it is likely to be non-Gaussian. Other researchers discretized the data prior to graphical modelling which is also problematic since discretization might mask statistical dependencies (Friedman et al., 2000, Dejori et al., 2003).

In our approach the parameters Θ are fitted once for a fully connected model using a leave-one-out procedure. The densities for the low-dimensional marginal densities $p(\mathbf{x}_C)$ and $p(\mathbf{x}_S)$ can be calculated by marginalizing the joint model. Thus, we can write

$$p(\mathbf{x}) = \frac{\prod_{C \in \mathcal{K}} p(\mathbf{x}_C | D_C, \hat{\Theta}_C)}{\prod_{S \in \Sigma} p(\mathbf{x}_S | D_S, \hat{\Theta}_S)}, \quad (4)$$

where the vectors in D_C , Θ_C and D_S , Θ_S contain only the dimensions (or nodes) which belong to clique C or separator S , respectively. The hat denotes the fitted values of a variable.

This procedure has two important advantages:

- (i) The density estimate is consistent. In other words, calculating a marginal probability density with different sequences of marginalization always yields the same result.
- (ii) The probability density models Θ do not need to be re-fitted during the structural learning procedure. This greatly reduces the computational complexity.

2.3 Learning Decomposable Models

Structural learning of decomposable models requires (i) a criterion to score the quality of a given decomposable model structure for describing the data and (ii) an efficient search strategy which allows to successively generate new graph structures, while staying within the class of decomposable models. In our approach, the structure of a decomposable model is evaluated using the *predictive assessment criterion*, namely 5-fold cross-validation. For this, the data set is divided into disjunct subsets D^k , $k=1, \dots, 5$. The model is learned using 80 percent $D \setminus D^k$ of the available data ($D \setminus D^k$ denotes all data points in D except the ones in D^k), and is evaluated based on the log-likelihood of the remaining data points D^k , which had not been used for training. The log-likelihood represents a measure of how likely the observed data could have been generated by the model, or in other words, how good the model describes the test dataset. This procedure is repeated 5 times and the final score consists of the average of the 5 test set log-likelihoods. Cross validation enforces the model to explain previously unseen data. This procedure gives a high score to models that have learned the underlying statistical structure and largely ignored the effect of fluctuations in the finite data set.

Since the joint probability density factorizes as given by eq. (4), the log-likelihood becomes

$$L(T) = \sum_{j=1}^A L(C_j) - \sum_{i=1}^B L(S_B)$$

(5)
with

$$L(C) = \sum_{k=1}^5 \sum_{\mathbf{x}_C \in D_C^k} \log p(\mathbf{x}_C | D_C] D_C^k, \hat{\Theta}_C)$$

(6)

$$L(S) = \sum_{k=1}^5 \sum_{\mathbf{x}_S \in D_S^k} \log p(\mathbf{x}_S | D_S] D_S^k, \hat{\Theta}_S)$$

(7)

and where A is the number of cliques and B is the number of separators.

The log likelihood eq. (5) is used to score the goodness of a model (model score). It is maximized successively by modifying the structure of the decomposable graphical model. We use greedy forward search, where, starting from an initially empty graph, edges are successively added sequentially, subject to two conditions: (i) An edge may be added only if the resulting new graphical model is still decomposable, and (ii) if, among all edges allowed according to (i) the addition of the edge leads to the highest increase in the total score eq. (5).

Our check for decomposability is based on a *cordiality check procedure* introduced in (Ibarra, 2000), and is described in detail in (Schwaighofer et al., 2004).

2.4 Estimating the Confidence of Decomposable model Features

Structure learning in graphical models is unstable, which means that the learned structure can be sensitive to small modifications of the data. To distinguish between stable and unstable structure we used a 20-fold *bootstrap approach* (Friedman et al., 1999; Steck and Jaakkola, 2004). In the bootstrap, we generate $Q=20$ perturbed versions of the original data set, by sampling with replacement. For a data set D with m examples, each of these perturbed data sets D^i also contains m examples, which are drawn at random (with replacement) from D . We learn a structure, as outlined in the previous sections, on each bootstrap data set D^i , $i=1, \dots, Q$, and obtain an estimated structure G^i of the graphical model. The confidence of a particular edge between nodes u and v can be estimated as the fraction of structures G^i where this edge is present.

For each of the perturbed data sets D^i , from the ALL data set described below, we use the structure obtained when the graph contained 3500 edges. In the analysis, only edges that have a confidence of 90% or above, i.e., edges that were found in at least 18 out of the 20 replications, were considered. This thresholding by edge confidence may lead to a non-decomposable resulting model. To again obtain a decomposable model, some of the lowest confidence edges were pruned until decomposability was reached.

2.5 ALL Microarray Data Set and Data Preprocessing

DNA microarray measurements reflect the expression level of thousands of genes in a cell by measuring the mRNA concentration for each gene simultaneously (Brown and Botstein, 1999). The data set analyzed here consists of measurements of 12,000 probes from 327 patients suffering from one of 7 different subtypes of pediatric acute lymphoblastic leukemia (ALL) (Yeoh et al., 2002).

ALL is a heterogeneous disease. It appears in various subtypes which differ markedly in their response to medical treatment. Leukemic ALL-cells are related to bone marrow cells, which are destined to either become T-lymphocytes (T-lineage) or B-lymphocytes (B-lineage). There is one homogeneous T-lineage-related ALL (15 % of cases), referred to as 'T-ALL', the pathogenesis of which is not yet well-understood, and several subtypes of B-lineage related ALL (85 % of cases), which can be retraced to specific genetic lesions. (E2A-PBX1, hyperdip > 50, BCR-ABL, TEL-AML1, MLL, 'novel'). The goal of the study of Yeoh and coworkers was to use expression profiling for identifying each of the known prognostically and therapeutically relevant disease subtypes and to assign patients to one subtype, thereby identifying patients who are at high risk for failing conventional therapeutic approaches. This was done by hierarchical clustering. Here we re-analyze the same data set using decomposable model learning in order to identify disease relevant genes, and group together such genes to functional modules in a data-driven way.

Out of the 12,000 measured gene probes, we selected those that best define the individual subtypes using Chi-square testing according to Yeoh et al. (2002): The 40 most discriminative probes for each of the 7 disease subtypes were selected, resulting in 280 gene probes. Out of those, 9 probes appear in more than one cluster but only once in our final data set, such that 271 probes remain included in the data set. Hence, each microarray measurement forms a $n=271$ -dimensional data vector \mathbf{x} , and the complete data matrix D contains $m=327$ vectors. In this data set, 239 genes were measured by

one probe, 13 genes by two probes each, and two genes by three probes. The probes are in multiple copies partly to overcome problems with alternative splicing or to test for variability within a microarray measurement. Here we use the redundancy to benchmark the ability of the decomposable model to detect functionally linked genes: Every gene is by definition strongly linked to itself, and therefore repeated measurements of a single gene by duplicate or multiple probes should be grouped together within a clique by the model.

Figure 2 about here

3. Results

In this section we apply decomposable model analysis to the ALL microarray data set in order to assign functional roles and obtain functional groups of genes related to this disease.

3.1 Analysis of the Dependency Graph

First we analyze the graph structure G , which was learned from the data according to the procedure described in the methods section. The graph structure is rather large and complex, thus we will only present a detailed description of some particularly interesting parts. Figure 2 shows a section of the obtained ALL graph structure. Most edges in the graph connect genes belonging to the same ALL subtype. Hence, the result correctly reflects the co-activation of genes grouped to a cluster, but in addition imposes a substructure within each cluster. Moreover, it can be seen that many genes are linked to other genes by only one or two edges, but a few of them show a high number of links. In other words, the number of edges from or to a certain gene, referred to as its degree, varies strongly between different genes.

One possible interpretation of the graph, which has been suggested for Bayesian networks (Friedman et al., 2000, Dejori et al., 2004), is to interpret each edge as an underlying biological (e.g., transcriptional regulatory) relationship between genes. According to this view, the biological relationship indirectly causes the statistical dependency of the gene expression levels, which is then reflected by the edge of the graph. In light of this hypothesis, genes with a high degree interact – via their product – with many other genes or gene products. Hence it can be argued that a gene with a high degree is crucial for the coordinated action of many other genes, and therefore is important for the correct operation of critical cellular life processes. Conversely, any damage on a high degree gene is likely to have a deeper impact on cellular processes at a systems level, than damaging a low-degree gene. Following this rationale, we suggest high-degree genes as critically involved in pathogenic mechanisms and to be candidates for drug targets.

Since the structure is learned from leukaemia data, the genes with high degree should be important for leukaemogenesis or for tumour development in general. In figure 2, gene *PSMD10* (Affymetrix- ID 37350_at, at the bottom of the figure) is found to be linked to a high number of other genes, and is therefore predicted by the model as important for the stability of cellular function. *PSMD10* is a regulatory subunit of the 26S proteasome, a protein complex which -in agreement with the model topology - degrades a large family of proteins that are marked to be destroyed, and thus helps regulating the protein turnover in eukaryotic cells. Hence, it is known to be crucial for normal cellular function. Conversely, a malfunction of *PSMD10* is known to result in a defective regulation of a large number of intracellular proteins that govern cell division, tumour growth, and

tumour survival, and which are functionally altered in cancer cells. Indeed, recent work has shown (Adams, 2002), that the PSMD10 pathway is often subject to cancer-related deregulation and can underlie processes such as oncogenic transformation or tumour progression.

Table 1 about here

Table 1 summarizes the 4 genes with the highest degree over the whole graph, together with their annotation. All highly connected genes are either known to be genes with an oncogenic characteristic or known to be involved in critical biological processes while being altered during oncogenesis (Jordanova et al., 2003).

3.2 Analysis of Functional Modules

In a second step of analysis, we consider individual cliques of the learned decomposable model as functional modules. Table 2 summarizes all cliques of size 3 and higher obtained from the ALL data set. Horizontal lines separate different cliques. A first observation concerns multiply occurring gene probes. In the given data set, 13 genes are measured by two probes and 2 by three probes on the chip. We noticed that each pair or triple of probes is grouped into the same clique. This can also be seen in part from table 2, where all multiple probes contained are highlighted in light grey. This observation forms an important benchmark test for the model, because each gene has by definition the maximum functional link to itself, and therefore multiple measurements of one gene should always be grouped into a clique. The fact that all 15 multiple measurements have been correctly assigned to 15 cliques demonstrates that the model can successfully detect functional cliques with high robustness.

Table 2 about here

We next focused on genes known to be subunits of a common functional complex. From annotation data we found three functional complexes with more than one member present in the data set: (i) The major histocompatibility complex class 2 (MHC II) (five members, one measured twice), the p26 proteasome (two members), and the T3 complex (two members). Table 3 lists these genes for three complexes. Interpreting cliques as functional modules, the genes for each of these complexes should also be grouped into a common clique. It was observed that the members of each of the complexes were always put into one clique, or into adjacent cliques, by the decomposable model. Adjacent positions were defined as being linked by a single separator in the decomposable model. Moreover, when the genes were assigned to adjacent cliques instead of a single clique, which occurred only for the MHC II group, the separator was very strong (the cliques contained 4 members, three of which formed the separator). This can also be seen from cliques 3, 4 and 5 from top in table 2.

Functionally related genes are often located on nearby places on the chromosome. This is true for the members of the MHC class II functional module (cf. cliques 3 - 5 from top, column 3, in table 2). Therefore we next considered the locations of genes grouped into individual cliques. In 17 out of 23 cliques in table 2 (not counting the multiple probes) there are regularities in the gene locations. Either all genes were located nearby on the same chromosome (cliques 3, 5, 17, 18, 23, 24), in part located nearby (cliques 4, 9, 10 and 11), or showed some regularity of the location. For example, cliques 1, 2, 12, 14, 15 and 16 always contain genes at the X chromosome locations near p22 and p28. The widespread presence of regularities in gene-loci within cliques indicates that genes are grouped in a meaningful way by decomposable models.

If we identify a clique as a functionally linked group of genes (gene products), which subserves a certain common task, genes which belong to many cliques can be predicted to be of more central (or common) purpose than genes belonging to few or only one clique. Therefore, we next ranked genes by the number of cliques they contribute to. Table 1 summarizes the four genes with the highest number of clique memberships. As can be seen, both the identity of the genes and their ranking coincide with the ranking obtained from the degree distribution. However the absolute number of the rank differs slightly. In summary, both the degree and clique membership count draw a consistent picture of the functional importance of genes.

Table 3 about here

Motivated by this observation, we re-analyzed the three functional modules provided in table 2, in terms of their fine structure as ranked by the clique membership count. Genes listed in the upper part of table 3 are part of the MHC II complex. Class II molecules are composed of two polypeptide chains, α and β chains. The MHC II molecules themselves are highly polymorphic (meaning that there are many different variants of these genes within the population), forming different MHC II variants for different antigens. Yet, *HLA-DRA* itself is monomorph, thus it is present in almost each of the MHC modules. This is reflected in the high number of cliques in which *HLA-DRA* is involved, in comparison to the lower clique counts of the polymorphic components (table 3). Also, whereas some cliques link *HLA-DRA* to genes outside the MHC II, each clique containing a MHC II member also contains *HLA-DRA* itself.

Table 3 also contains one example of a doubly measured gene, *HLA-DPB1*. It can be seen that in both cases the clique count of this gene is low but not identical, (clique counts 3 and 1). This variability can serve as a measure for the confidence in the clique count. Although there is some variability seen between both counts, the *HLA-DPB1* counts are much smaller than the clique count found for *HLA-DRA*. Hence, the difference in clique counts between *HLA-DRA* and *HLA-DPB1* is likely to be significant.

Genes listed in the second part of table 3 are subunits of the 26S proteasome complex. Whereas *PSMD10* is present in many cliques the other subunit, *PSMC1*, is present only in one clique namely with *PSMD10*. This can be evidence for a more dominant role of *PSMD10* in protein degradation than of *PSMC1*. In the learned structure, *PSMD10* connects most cliques that contain genes which are altered in ALL subtype hyperdiploidy, predicting a dominant role of *PSMD10* in the hyperdiploidy subtype. This prediction seems reasonable, since the 26S proteasome is involved in general protein degradation and is therefore likely to be hyperactive in response to excess protein production by hyperdiploidy.

The third part of table 2 contains the two subunits of the data set which contribute to the T-cell antigen receptor complex. *CD3D* forms the δ subunit of the T3 complex, and has been identified as the gene which discriminates between ALL of T and of B-lineage (Yeoh et al., 2002). Hence, *CD3D* seems to play a central role in ALL disease mechanisms. At the same time, *CD3D* is assigned to be a member in many (12) cliques by our analysis, pointing towards a central function as well. In contrast, the gene for the ϵ subunit of the T3 complex, *CD3E*, appears only in one clique: Correspondingly, the decomposable model assigns a less dominant role of *CD3E* in ALL.

4. Discussion

We presented a novel approach towards a systems level analysis of concerted cellular mechanisms, and applied the model to a set of genome-wide expression profiles from ALL patients. The approach is based on a graphical modelling technique called

decomposable model, which puts particular emphasis on the modular way, in which bio-molecules act together to accomplish a certain task, and on the continuous yet noisy nature of the data to be analyzed. A decomposable model tries to explain the statistics in a data set by the action of mutually linked functional modules, so-called cliques. Decomposable models with continuous variables have significant advantages for this application domain:

(i) Previous approaches (e.g., Friedman et al., 2000, Pe'er et al., 2001) have mostly concentrated on learning discrete valued models from such data. Hence, one needs to first discretize the continuous-valued expression level. This is a crucial and quite delicate pre-processing step that needs to be conducted carefully (Friedman et al., 2000). In contrast, the approach adopted here accounts in a natural way for the continuous nature of the measurements, and for their unknown and probably non-Gaussian joint probability distribution.

(ii) Molecular networks often show a “small-world” topology (Jeong et al., 2000), in which the network is decomposable into smaller groups of densely connected clusters (Watts and Strogatz, 1998). This finding might render decomposable models with their intrinsic modular or clique-like structure particularly suitable for describing genetic networks.

(iii) Functional modules are considered to be a critical level of biological organization (Hartwell et al., 1999). One example are modules in transcriptional regulation. Transcription factors work by binding to DNA-motifs and affecting the rate of transcription. Many binding sites occur in spatial and functional clusters called enhancers, promoter elements, or regulatory modules. Thus, the promoter regions suggest a hierarchical or modular style of the transcription complex. Two further examples of molecular modules are subunits of multimeric proteins, where the subunits are coded by separate genes, or protein groups which associate into larger structures termed macromolecular assemblies. In the latter two cases, the genes for the different subunits or the genes that code for proteins of the same macromolecular assembly are functionally grouped to a module. Finally, gene products can also form a functional module by carrying out a certain cellular function in a concerted way, but without being physically grouped to a molecular assembly. The inherent modular structure of a decomposable model imposes a strong drive for it towards explaining the data in terms of densely linked gene groups. By this, decomposable models should be able to detect with particularly high sensitivity the signature of a concerted action of gene modules in the data. In light of this rationale, cliques are likely to contain *functionally highly correlated* genes, as opposed to gene clusters (Eisen et al., 1998, Yeoh et al., 2002), where genes are grouped together by mere co-expression. Hence, as opposed to clustering, the learned structure also reveals some information about the possible statistical relationship of genes within a cluster.

A decomposable model approach preserves some of advantages of the related systems-level modelling techniques by Bayesian networks (Friedman et al., 2000; Dejori et al., 2003, 2004), namely (i) it takes into account the systemic nature of many biological processes, which arise from the interactions of many genes rather than from actions of an individual gene, and (ii) it accounts for the statistical and noisy nature of the data by adopting a probabilistic approach. A difference is that Bayesian networks allow a causal interpretation whereas decomposable models are restricted to identifying strongly coupling sets of genes. The two main advantages of our approach are that (i) it directly works with the continuous expression data and does not depend on preprocessing by discretization or assumptions like Gaussianity of the data. (ii) The approach is tailored to account for the modular nature of biological molecular life processes, which frequently involve the collective action of protein subunits, protein assemblies, and other functional modules.

An apparent restriction of a decomposable graphical model, namely its special structure as a set of linked cliques, turns out to be its strength: Decomposable models are particularly sensitive to the signature of functional modules in the data, because they are designed to explain all the statistics in terms of interacting cliques of genes. In applying the model to ALL data, genes known to encode subunits of known complexes were correctly inked into individual or closely linked adjacent cliques, demonstrating a high sensitivity for detecting functional modules. If subunits were not grouped into a single clique but in adjacent cliques, the link connecting them was very strong. This means that for such adjacent cliques only one or few edges in the graph were missing to render them a single clique, which might be a consequence of statistical fluctuations due the limited number and the noisiness of the data set. Based on the clique and the link structure learned from the data, it became possible to formulate two new scores for ranking genes according to their putative importance in cellular processes: the number of links to a gene and the number of cliques it participates in.

One important ingredient of cellular processes is their dynamical nature, which is linked with molecular reaction constants (Stetter et al., 2003). Unfortunately, due to the vast complexity of these dynamics and the small amount of data available the dynamics of molecular networks at a large scale have rarely been investigated so far. At present, dynamical considerations can be at most applied to small and experimentally very well-characterized subsystems. Decomposable models might be able to simplify a dynamic analysis by suggesting small (i.e., low-dimensional) tightly coupled functional modules. These modules might be natural breakpoints for the separation of scales, for example by assuming an adiabatic approximation within a functional module and by explicitly modelling only the time constants for interactions between modules.

Decomposable models are not restricted to analyzing the transcriptome of a cell and its changes under various pathological conditions. As cellular life processes are strongly affected and even dominated by a enormous multitude of protein-protein interactions, the technique presented here will be suitable to analyse whole proteome measurements from cells in laboratories of the near future, putting emphasis in the level of interaction between proteins. Of similar importance is the extraction of the modularity of these interactions, where small functional modules will be continuously grouped together to accomplish more and more complex tasks, up to whole cellular genetic programs. In light of this view, genome wide and proteome wide modular analysis and related techniques might form a key ingredient of modern functional genomics and proteomics.

Acknowledgments: M. Dejori and A. Schwaighofer gratefully acknowledge support through an Ernst v Siemens scholarship.

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Figure captions:

Figure 1: Schematic illustration of decomposable graphical models. **(a)** Sketch of a two-dimensional joint probability density $p(x_1, x_2)$. The marginal densities $p(x_1)$ and $p(x_2)$ are obtained by integrating over the respective other variable. In this example, the conditional probability density $p(x_2/x_1)$ differs from the marginal density, reflecting a statistical dependency between x_1 and x_2 . **(b)** Same as **(a)** but for statistically independent variables x_1 and x_2 . Here conditional and marginal probability densities coincide and the joint probability density factorizes. **(c)** Graph structure of a simple decomposable graphical model with 7 nodes. Each node i stands for a variable x_i , and each edge reflects a direct statistical dependency. **(d)** A join tree equivalent to the graph structure of **(c)**. Each node of the tree stands for a clique of fully connected variables, and each edge reflects a set of variables common to adjacent cliques: their separator.

Figure 2: A part of the decomposable model structure learned from the ALL data set, visualized as graph structure G . The highly connected gene *PSMD10* (Affymetrix- ID 37350_at, at the bottom of the figure) is thought to be involved in cellular deregulations that lead to oncogenesis.

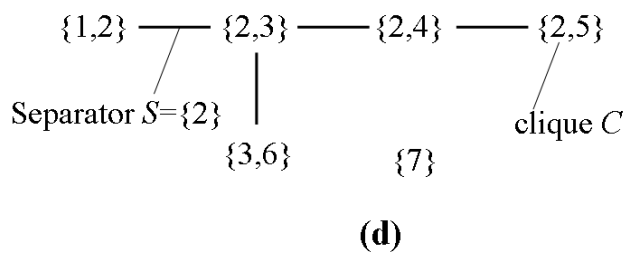
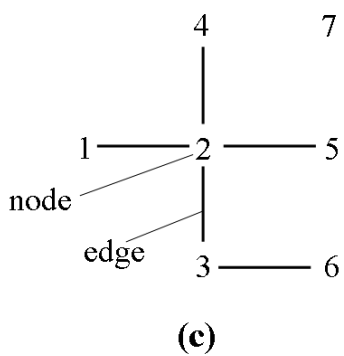
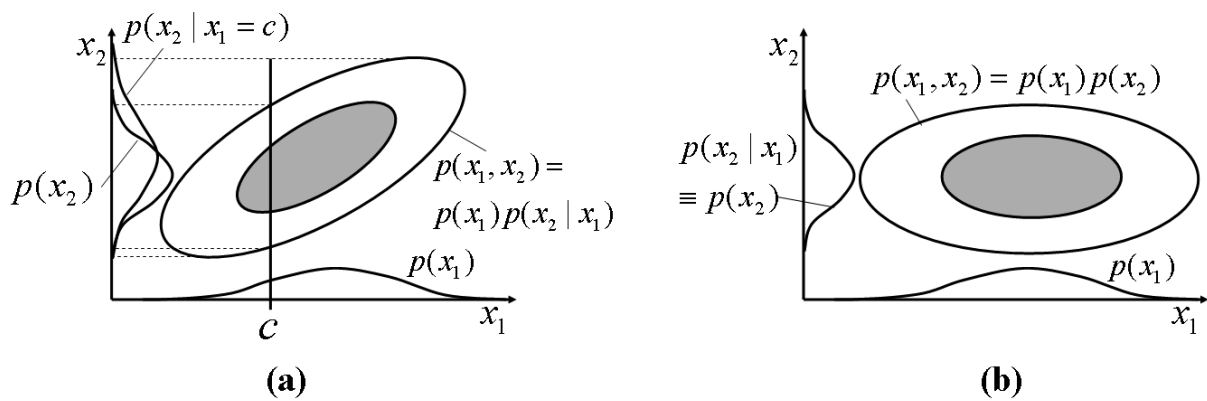


Figure 1 of Dejori et al., top

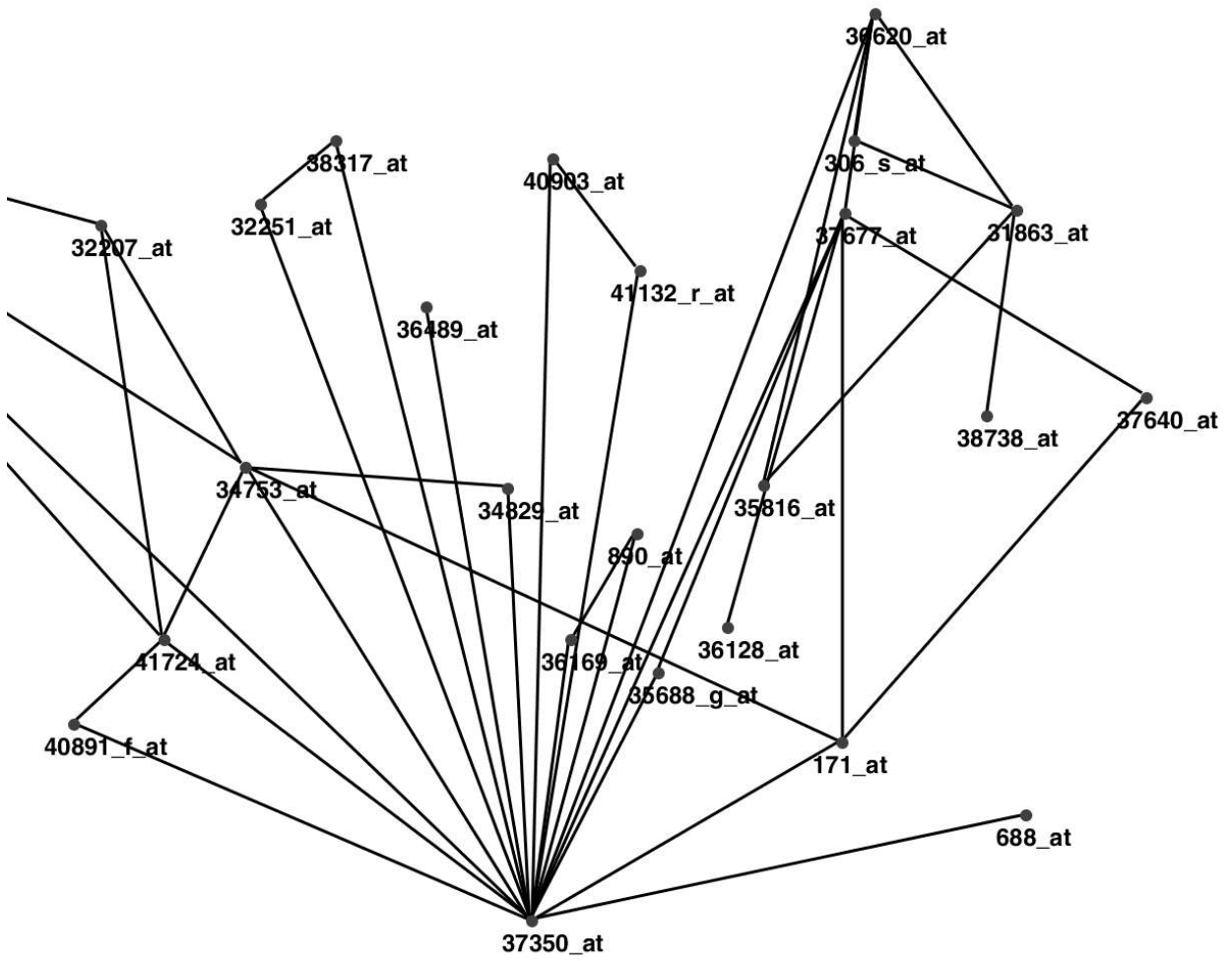


Figure 2 of Dejori et al., top

Table 1 : Genes in the ALL data set, ranked by the number of connections

Gene	Affymetrix ID	Degree	no. cliques	Putative function
<i>PSMD10</i>	37350_at	17	12	Proteasome, protein degradation
<i>HLA-DRA</i>	37039_at	13	8	immune response, antigene presentation
<i>SCML2</i>	38518_at	9	6	embryogenesis, transcription factor
<i>POU2AF1</i>	36239_at	7	6	transcription cofactor, antipathogene response

Table 1 of Dejori et al.

Table 2: List of all cliques of size 3 and higher

Gene	Affym.-ID	Location	Putative function
BCAP31	41724_at	Chr:Xq28	accessory protein BAP31
MPP1	32207_at	Chr:Xq28	membrane protein, palmitoylated 1, 55kDa
SCML2	38518_at	Chr:Xp22	sex comb on midleg- like 2 (Drosophila)
SYBL1	34753_at	Chr:Xq28	synaptobrevin- like 1
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
BCAP31	41724_at	Chr:Xq28	accessory protein BAP31
SCML2	38518_at	Chr:Xp22	sex comb on midleg- like 2 (Drosophila)
SYBL1	34753_at	Chr:Xq28	synaptobrevin- like 1
HLA- DPB1	38095_i_at	Chr:6p21.3	major histocompatibility complex, class II, DP beta 1
HLA- DPA1	38833_at	Chr:6p21.3	major histocompatibility complex, class II, DP alpha 1
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
HLA- DMA	37344_at	Chr:6p21.3	major histocompatibility complex, class II, DM alpha
CD74	35016_at	Chr:5q32	CD74 antigen (invariant polypeptide of MHC II)
HLA- DPB1	38095_i_at	Chr:6p21.3	major histocompatibility complex, class II, DP beta 1
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
HLA- DMA	37344_at	Chr:6p21.3	major histocompatibility complex, class II, DM alpha
HLA- DPB1	38095_i_at	Chr:6p21.3	major histocompatibility complex, class II, DP beta 1
HLA- DPB1	38096_f_at	Chr:6p21.3	major histocompatibility complex, class II, DP beta 1
HLA- DPA1	38833_at	Chr:6p21.3	major histocompatibility complex, class II, DP alpha 1
HLA- DMA	37344_at	Chr:6p21.3	major histocompatibility complex, class II, DM alpha
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
CD3D	38319_at	Chr:11q23	CD3D antigen, delta polypeptide (TiT3 complex)
CD19	1096_g_at	Chr:16p11.2	CD19 antigen
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
CD79A	38017_at	Chr:19q13.2	CD79A antigen (immunoglobulin- associated alpha)
POU2AF1	36239_at	Chr:11q23.1	POU domain, class 2, associating factor 1
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
CD3D	38319_at	Chr:11q23	CD3D antigen, delta polypeptide (TiT3 complex)
BLNK	38242_at	Chr:10q23.2	B- cell linker
HLA- DRA	37039_at	Chr:6p21.3	major histocompatibility complex, class II, DR alpha
TCL1A	39318_at	Chr:14q32.1	T- cell leukemia/lymphoma 1A
CD24	266_s_at	Chr:6q21	CD24 antigen (small cell lung carcinoma cl. 4 antigen)
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
VBP1	171_at	Chr:Xq28	von Hippel- Lindau binding protein 1
PGK1	37677_at	Chr:Xq13	phosphoglycerate kinase 1
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
SOD1	36620_at	Chr:21q22.11	superoxide dismutase 1, soluble (ALS 1 (adult))
PGK1	37677_at	Chr:Xq13	phosphoglycerate kinase 1
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
VBP1	171_at	Chr:Xq28	von Hippel- Lindau binding protein 1
SYBL1	34753_at	Chr:Xq28	synaptobrevin- like 1
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
PGK1	37677_at	Chr:Xq13	phosphoglycerate kinase 1
---	35688_g_at	---	---
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
HNRPH2	41132_r_at	Chr:Xq22	heterogeneous nuclear ribonucleoprotein H2 (H')
ATP6IP2	40903_at	Chr:Xq21	ATPase, H+ transp., lysosomal interacting protein 2
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
BCAP31	41724_at	Chr:Xq28	accessory protein BAP31
DXS9879E	40891_f_at	Chr:Xq28	DNA segment on X (unique) 9879 expr. sequence
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
SYBL1	34753_at	Chr:Xq28	synaptobrevin- like 1
DKC1	34829_at	Chr:Xq28	dyskeratosis congenita 1, dyskerin
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
UBE2A	890_at	Chr:Xq24- 25	ubiquitin- conjugating enzyme E2A (RAD6 homolog)
NDUFA1	36169_at	Chr:Xq24	NADH dehydrogenase 1 alpha subcomplex, 1, 7.5kDa
PSMD10	37350_at	Chr:Xq22.3	proteasome 26S subunit, non- ATPase, 10
TCEAL1	38317_at	Chr:Xq22.1	transcription elongation factor A (SII)- like 1
FLJ21174	32251_at	Chr:Xq22.1	hypothetical protein FLJ21174
PTPRM	31892_at	Chr:18p11.2	protein tyrosine phosphatase, receptor type, M
PTPRM	994_at	Chr:18p11.2	protein tyrosine phosphatase, receptor type, M
PTPRM	995_g_at	Chr:18p11.2	protein tyrosine phosphatase, receptor type, M
ITGA6	33410_at	Chr:2q31.1	integrin, alpha 6
ITGA6	41266_at	Chr:2q31.1	integrin, alpha 6
ITGA6	33411_g_at	Chr:2q31.1	integrin, alpha 6
BCR	1635_at	Chr:22q11.23	breakpoint cluster region
BCR	1636_g_at	Chr:22q11.23	breakpoint cluster region
ABL1	39730_at	Chr:9q34.1	v- abl AML viral oncogene homolog 1
CD44	2036_s_at	Chr:11p13	CD44 antigen (homing function, Indian blood group s.)
CD44	40493_at	Chr:11p13	CD44 antigen (homing function, Indian blood group s.)
---	1126_s_at	---	---
SOD1	36620_at	Chr:21q22.11	superoxide dismutase 1, soluble (ALS 1 (adult))
KIAA0179	31863_at	Chr:21q22.3	KIAA0179 protein
HMG1	306_s_at	Chr:21q22.2	high- mobility group nucleosome binding domain 1
SOD1	36620_at	Chr:21q22.11	superoxide dismutase 1, soluble (ALS 1 (adult))
KIAA0179	31863_at	Chr:21q22.3	KIAA0179 protein
CSTB	35816_at	Chr:21q22.3	cystatin B (stefin B)
VBP1	171_at	Chr:Xq28	von Hippel- Lindau binding protein 1
PGK1	37677_at	Chr:Xq13	phosphoglycerate kinase 1
HPRT1	37640_at	Chr:Xq26.1	hypoxanthine phosphoribosyltransferase 1
SCML2	38518_at	Chr:Xp22	sex comb on midleg- like 2 (Drosophila)
TNRC11	40998_at	Chr:Xq13	trinucleotide repeat containing 11 (THR- associated)
---	34374_g_at	---	---

Table 2 of Dejori et al.

Table 3 : Genes of three functional complexes, ranked by the number of cliques they belong to.

<i>Genes</i>	Affy.- ID	no. cliques	Putative function
<u>MHC, class II</u>			
<i>HLA- DRA</i>	37039_at	8	MHC II, DR α
<i>HLA- DMA</i>	37344_at	4	MHC II, DM α
<i>HLA- DPB1</i>	38095_i_at	3	MHC II, DP β 1
<i>HLA- DPA1</i>	38833_at	2	MHC II, DP α 1
<i>HLA- DPB1</i>	38096_f_at	1	MHC II, DP β 1
<i>HLA- DRB1</i>	41723_s_a t	1	MHC II, DR β 1
<u>Proteasome p26</u>			
<i>PSMD10</i>	37350_at	12	26S, non- ATPase regulatory subunit 10
<i>PSMC1</i>	688_at	1	26S, ATPase regulatory subunit 1
<u>T3 complex</u>			
<i>CD3D</i>	38319_at	12	T3 complex, δ polypeptide subunit
<i>CD3E</i>	36277_at	1	T3- cpmlex, ϵ polypeptide subunit

Table 3 of Dejori et al.