Discovering gene regulatory networks of multiple phenotypic groups using dynamic Bayesian networks

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Abstract

Dynamic Bayesian networks (DBNs) can be used for the discovery of gene regulatory networks (GRNs) from time series gene expression data. Here, we suggest a strategy for learning DBNs from gene expression data by employing a Bayesian approach that is scalable to large networks and is targeted at learning models with high predictive accuracy. Our framework can be used to learn DBNs for multiple groups of samples and highlight differences and similarities in their GRNs. We learn these DBN models based on different structural and parametric assumptions and select the optimal model based on the cross-validated predictive accuracy. We show in simulation studies that our approach is better equipped to prevent overfitting than techniques used in previous studies. We applied the proposed DBN-based approach to two time series transcriptomic datasets from the Gene Expression Omnibus database, each comprising data from distinct phenotypic groups of the same tissue type. In the first case, we used DBNs to characterize responders and non-responders to anti-cancer therapy. In the second case, we compared normal to tumor cells of colorectal tissue. The classification accuracy reached by the DBN-based classifier for both datasets was higher than reported previously. For the colorectal cancer dataset, our analysis suggested that GRNs for cancer and normal tissues have a lot of differences, which are most pronounced in the neighborhoods of oncogenes and known cancer tissue markers. The identified differences in gene networks of cancer and normal cells may be used for the discovery of targeted therapies.

Keywords: dynamic Bayesian networks, time series, gene expression, Bayesian learning, classification, MCMC

Introduction

Learning gene regulatory networks (GRNs) from gene expression data has been the focus of much research in the last decades [7, 10, 38, 62]. The precise knowledge of GRNs can help to understand the molecular mechanisms driving diseases and facilitate the search for targeted therapies [3, 32]. Multiple computational methods can be used to learn GRNs from observational data, including correlation analysis [20, 29, 34], Boolean networks [31, 36], Bayesian networks [5, 12, 59], differential equation models [60, 61] and machine learning approaches [19]. A recent benchmarking study [63] revealed no clear winner among different methods for GRN reconstruction, with different methods demonstrating advantageous in different settings.

A Bayesian network is a probabilistic graphical model representing dependencies between random variables via a directed acyclic graph (DAG). Due to its probabilistic nature, this model is well suited to describe noisy biological data. However, static Bayesian networks do not allow directed cycles, rendering it impossible for them to model feedback loops. The Dynamic Bayesian Network (DBN) model overcomes this problem by including dependencies between nodes at different time points and accommodating the possibility of cycles [28, 39, 50].

DBN models were used to learn biological networks [35], including GRNs [2, 6, 15, 30, 59, 64] and multi-omics networks [48]. Learning DBN structures from data is computationally challenging because the number of possible network topologies grows exponentially with the number of nodes. Some methods solve this issue by employing a greedy search [48, 59], others restrict the network topology by prohibiting instantaneous dependencies between genes or limiting the number of possible incoming edges per each node [18, 35, 57]. However, topological restrictions may potentially result in the discovery of suboptimal models [41].

Another limitation of most network learning methods lies in the assumption that all samples in the dataset represent the same GRN, however this assumption may be violated. For example, it has been shown experimentally that protein–protein interactions differ drastically.
between tumor and normal cell lines [23]. Hence the
discovery of context-specific GRNs can facilitate the dis-
covery of targeted therapies [56]. Only limited research
was devoted to learning DBNs from distinct but related
contexts [24, 42, 43]. However, none of the methods was
applied to networks with more than 40 nodes, and all
suggested approaches utilized limited DBN topologies
that assume no instantaneous dependencies between
genes.

The goal of this study was to create a scalable frame-
work for learning DBN models that provide high pre-
dictive accuracy and can be used for learning GRNs
for multiple subgroups of samples, defined, for exam-
ple, by molecular, histological or clinical phenotypes. We
employed a Bayesian approach [26] for learning DBNs
that is scalable to networks with hundreds of nodes and
implemented in the R-package BiDAG [52]. BiDAG was
previously used for context-specific learning of static
gene networks [27, 51]. This package allows selecting
from a wide range of network topologies, including prior
information from public gene interaction databases and
modeling gene interactions whose strength changes over
time. In addition, the Bayesian approach to structure
learning implemented in the package is well equipped
to prevent overfitting, a known problem occurring in the
analysis of high-dimensional biological data.

Apart from BiDAG, we found five R-packages for learn-
ing DBNs, namely G1DBN [28], dbnlearn [8], dbnR [44],
ebdbNet [45] and bnstruct [9]. Only dbnR and bnstruct
are able to learn DBNs with the same range of topologies
as BiDAG. However, these packages can only learn models
whose parameters are constant over time, while BiDAG
can be used for learning both constant and time-varying
models. We compared BiDAG with these tools in simu-
lation studies to determine which tool best reconstructs
network structures.

Apart from BiDAG, none of the mentioned DBN learn-
ing tools includes functions enabling classification. For
this reason, for classification comparison, we chose stan-
dard classification tools that cannot perform network
reconstruction. In addition, we compared our results
with the DBN-based classifier reported in [24] for the
same datasets, however the code of this classifier is not
available.

To demonstrate the applications of the described
approach, we identified time-series datasets in the Gene
Expression Omnibus (GEO) database [1], which included
gene expression data for at least two different phenotypic
groups of the same tissue and comprised at least 50
observations in each of two consecutive time slices.
We found two datasets (GSE5462 and GSE37182) that
satisfied these criteria. To inform model selection in
the absence of ground truth, we used a cross-validated
measure of predictive accuracy that was previously used
to perform DBN model selection [35, 48]. In addition, we
used cross-validated classification accuracy to assess the
different models’ ability to distinguish between the
analyzed phenotypic groups. Concerning applications,
the suggested framework helped to understand if the
phenotypic groups in each dataset could be better
represented by GRNs with the same structure (but not
parameters) or if gene regulation differs substantially so
that different structures more accurately represent the
analyzed subgroups. In addition, our analysis demon-
strated that the range of modeling possibilities offered
by BiDAG is helpful for the discovery of models that
reach the highest predictive accuracy, while the DBN-
based classifier demonstrated competitive classification
accuracy.

Methods and data
A DBN is a probabilistic graphical model for the joint
distribution of random variables \( X = (X_1, \ldots, X_t) \) observed at
time points \( t = 0, 1, \ldots, T \). The DBN model uses a directed
graph to encode a factorization of the joint distribution of
\( X \) along the time slices \( t = 0, \ldots, T \) (Figure 1A).
Here, we consider DBNs in which structures are identical for all
time slices. We also assume that variables in time slice
\( t \) can depend on other variables in time slice \( t \) and on
variables in time slice \( t - 1 \), i.e.

\[
P(X_t | X_{t-1}, \ldots, X_0) = P(X_t | X_{t-1}).
\]  

Such DBN models are referred to as first-order DBNs.
The joint probability distribution of a DBN with \( T + 1 \)
time slices is

\[
P(X_0, X_1, \ldots, X_T) = P(X_0) \prod_{t=1}^{T} P(X_t | X_{t-1}).
\]
With these assumptions, the unfolded structure of a DBN (Figure 1A) can be represented in a compact way with two DAGs \((G^0, G^-)\) which are referred to as initial structure and transition structure, respectively (Figure 1B). The initial structure contains only edges in the first time slice. The transition structure describes relationships between gene expression levels in all other time slices, \(t > 0\). The edges within one time slice are called intra-edges and edges between time slices are called inter-edges. In \(G^0\), only intra-edges are present, while \(G^-\) contains both intra and inter-edges. We are primarily interested in discovering the transition structure because it describes both instantaneous dependencies (represented by intra-edges) and dependencies between gene expression levels at different time points (represented by inter-edges).

Within each time slice \(t > 0\) the joint distribution of \(X_t, \ldots, X_n\) is factorized according to a Bayesian network model:

\[
P(X_t | X_{t-1}) = \prod_{i=1}^{n} P(X_t^i | \mathbf{Pa}_t^i),
\]

where \(\mathbf{Pa}_t^i\) denotes the set of parents of node \(X_t^i\) in time slices \(t\) and \(t-1\) in \(G^-\). For \(G^0\) the parent sets \(\mathbf{Pa}_0^i\) are used instead to factorize \(P(X_0^i)\).

To fully specify a DBN, we also need parameters \(\theta\) which describe probabilistic dependencies between each node \(X_t^i\) and its parents in a DBN structure. We assume that \(X_t^i\) are jointly normally distributed. This results in the distribution of each node \(X_t^i\) being a linear regression on its parents [14]:

\[
P(X_t^i | G^- , \theta^i) = \mathcal{N}\left(X_t^i | m_t^i + \sum_{t'=t-1}^{t} \sum_{j \in \mathbf{Pa}_t^i} \beta_{j,t'}X_{t'}^j , (\sigma_t^i)^2\right).
\]

For each time slice \(t\), we have the parameters \(\theta^i = (m_t^i , B_t^i , (\sigma_t^i)^2)\), where \(m_t^i\) is a vector of regression intercepts, \(B_t^i = (\beta_{j,t'}^i)^i\) a set of all regression coefficients and \((\sigma_t^i)^2\) a vector of variances. For \(G^0\), the sum over parents in the previous time slice is dropped. We consider two cases, namely stationary DBNs where parameters stay constant over time \(\theta^1 = \ldots = \theta^T =: \theta^-\) and time-varying DBNs, where \(\theta^1 , \ldots , \theta^T\) are generally different. The parameters \(\theta^0\) and \(\theta^-\) are different even for a stationary model. In a time-varying model, we assume time-varying parameters, while the structure \(G^-\) is assumed to be the same across time slices \(1, \ldots , T\).

We also consider a special case where the initial structure \(G^0\) is the same as the internal structure of the transition structure \(G^-\), i.e. for all nodes, all intra-slice edges in \(G^-\) are the same as these edges in \(G^0\).

For learning the DBN structure from observational data \(D\), we employ the Bayesian approach implemented in the R package BiDAG [26, 52], and use the BGe score for learning and sampling the structures of Bayesian networks [14, 25]. The BGe score of a graph \(S(G | D)\) is derived from its posterior probability that is proportional to its marginal likelihood and graph prior:

\[
P(G | D) \propto P(D | G)P(G) = : S(G | D)
\]

As was shown in [11, 14], when some technical assumptions are fulfilled, the score \(S(G | D)\) decomposes in terms, each depending on a single node and its parents (see details in Supplementary data):

\[
S(G | D) = P(D | G)P(G) = \prod_{i=1}^{n} S(X_t^i , \mathbf{Pa}_t^i | D) \tag{6}
\]

The BGe score assumes a normal-Wishart prior on parameters [14] that satisfies the assumptions required for score decomposition in Equation (6).

For DBNs, the dataset \(D\) consists of \(N\) observations from \(T + 1\) time slices. To learn a time-varying DBN, we divide \(D\) in \(T + 1\) parts and define the BGe score of a DBN structure as

\[
S(G | D) = \prod_{i=1}^{n} S(X_t^i , \mathbf{Pa}_t^i | D^0) \prod_{i=1}^{n} S(X_t^i , \mathbf{Pa}_t^i | D^\rightarrow). \tag{7}
\]

To perform structure learning for a stationary model we divide the data into two parts: \(D^0\) and \(D^\rightarrow\), where \(D^\rightarrow\) contains observations from all pairs of neighboring time slices. Equation (7) then simplifies to

\[
S(G | D) = \prod_{i=1}^{n} S(X_t^i , \mathbf{Pa}_t^i | D^0) \prod_{i=1}^{n} S(X_t^i , \mathbf{Pa}_t^i | D^\rightarrow). \tag{8}
\]

We use the iterative order Markov chain Monte Carlo (MCMC) scheme [26] to estimate the \(a\ posteriori\) (MAP) structures \(G^0\) and \(G^-\). In addition, we estimate consensus structures by averaging over a sample of graphs from the posterior distribution and composing consensus structures of edges whose posterior probability is higher that 0.9 [26, 52].

**Learning DBN models for phenotypic subgroups**

In the proposed framework, each sample \(D_m\) contains gene expression levels of one patient from all time points and is assigned to a phenotypic subgroup \(Z_m = k\), \(k \in (1, \ldots , K)\). In this work, we analyzed two datasets, each comprising gene expression from \(K = 2\) subgroups (Figure 2A), however the model can be extended to an arbitrary number of groups.

Since the analyzed subgroups of samples in each dataset are related, we propose considering two models:
one which assumes that DBN structures are subgroup-specific and the other one that represents all subgroups by a single DBN structure (Figure 2B). In the latter case, the differences between subgroups can be explained by differences in DBN parameters. From a biological perspective, it is interesting to understand to which extent the interaction networks of different subgroups, for example, defined by different phenotypes, differ from each other.

To inform the choice between different structural and parametric model assumptions, we suggest computing mean absolute error (MAE) as a measure of predictive accuracy, which was already used in previous applications of DBN to biological datasets [35, 48]. MAE reflects how well the model predicts the changes in gene expression levels in time.

To avoid overfitting, we estimated MAE using leave-one-out cross-validation (CV). In each CV run, we removed one sample $D_m$ from the data $D$ and used the remaining data $(D)_{-m}$ to learn DBN structure and parameters. After that, we plugged in the values $D_m^0$ containing gene expression levels of the test sample from the first time slice and predicted gene expression levels in all other time slices iteratively according to the learned model. Finally, we computed MAE for each node and time slice and averaged it across all genes, slices, and test samples.

In addition to the predictive accuracy, we measured cross-validated classification accuracy to evaluate how well the DBN-based classifier can discriminate between the analyzed subgroups.

Other DBN learning tools do not provide functions for classification. For this reason, we compared our DBN-based classifier against random forest and naive Bayes classifiers [17, 33]. We ran the CV 100 times for the random forest classifier to average out randomness in the results.

### Structural assumptions

Including prior biological knowledge can improve network learning [65], so we consider two different ways to include such knowledge: by penalizing the edges that cannot be found in public protein–protein interaction databases, such as, e.g. the STRING database [53] (Figure 1E) and by excluding these edges completely from the search space (Figure 1F). Penalization is implemented by imposing a nonuniform prior over structures:

$$P(G) \propto \prod_{i=1}^{n} \frac{1}{\prod_{j \in \mathcal{Pa}_i} \pi_{ij}},$$

where $\pi_{ij} = 1$ if the interaction between genes $X_i$ and $X_j$ can be found in the STRING database with a confidence level of at least 0.4, and $\pi_{ij} = 2$ otherwise.

Most DBN models and tools prohibit intra-edges. We do not assume the presence or absence of intra-edges by default. Instead, we include the model without intra-edges in the set of investigated models (Figure 2D) and compare its predictive accuracy with other models, including the model without any structural restrictions (Figure 2C).

### Work steps of the model

Our goal is to evaluate DBN models using all possible combinations of structural assumptions depicted in Figure 2B and Figure 2C.F. In addition, we consider models with time-varying and constant parameters for datasets where more than two time slices are present. For each combination of structural and parametric assumptions we perform the following CV procedure:
Leave-one-out CV procedure for DBNs.

Input:
Time series gene expression data $D$ and class membership assignments of each sample (row of $D$): $Z_1, ..., Z_N$
Blacklist matrix, $P = \{p_i\}$—penalization matrix, parameter assumptions, $p$—posterior probability threshold

Output
Cross-validated clustering accuracy of MAP and consensus models
Cross-validated MAE of MAP and consensus models

For $m = 1 : N$
1. Define training data $(D_m)_{m \neq m}$ and test sample $D_m$
2. Learn MAP and consensus structures and parameters
   Given $(D_m)_{m \neq m}, P, B$
   For $k = 1 : K$
   Learn MAP DBN structure $G_k$
   Obtain a sample of DBN structures from the posterior distribution $G_1, ..., G_k$
   Given $G_1, ..., G_k$ estimate consensus structure $Z_k$
   Compute class posteriors $P(D_m | k, G, \theta_d)$ and $P(D_m | k, G, \theta)$
   END For
3. Assign membership and compute MAE
   Assign class memberships given MAP and consensus models
   $\hat{y}_m = \text{argmax}_k P(Z_m = k | D_m, G, \theta)$
   $\overline{y}_m = \text{argmax}_k P(Z_m = k | D_m, Z, \theta)$
   Given $\hat{y}_m, \overline{y}_m, P, D_m$ compute MAE (MAP model)
   Compute clustering accuracy by comparing $\hat{y}, \overline{y}$ and $Z$
   END For
4. Compute clustering accuracy and global MAE of MAP and consensus models

Posterior probabilities of class memberships $Z_m$ of observations $D_m$ are computed as follows:

$$P(Z_m = k | D_m, G, \theta_d) = \frac{r_k P(D_m | G_k, \theta_d)}{\sum_{k' \neq k} r_k P(D_m | G_{k'}, \theta_{d_{k'}})}$$

(10)

where likelihoods $P(D_m | G_k, \theta_d)$ are computed according to the learned DBN structures and parameters:

$$P(D_m | G_k, \theta_d) = P(D_m^0 | G_k^0, \theta_d^0)^{\tau} \prod_{t=1}^{\tau} P(D_m^t | G_k^t, \theta_d^t)$$

(11)

and $r_k = \frac{1}{k}$ are estimated from the training data.

When the same structure for the analyzed subgroups is assumed, the graphs in step 2 need to be learned only once instead of $K$ times separately for each subgroup.

BiDAG package

The R-package BiDAG [52] implements a collection of MCMC methods that can be used for learning and sampling of static Bayesian network structures as well as DBNs. To implement the work steps of the model described in Section 2.3, we used the following functions:

- \text{iterativemcmc} implements a hybrid MCMC approach introduced in [26] and was used for MAP structure search
- \text{ordermcmc} was used for sampling from the posterior distribution
- \text{modep} was used for model averaging
- \text{scoreagainstDBN} was used to compute likelihoods from Equation (11)
- \text{compareDBNs} was used for model comparison

Data

We applied the described framework to two biological datasets, each containing time-series gene expression data of two phenotypic subgroups of the same tissue type (Section 4).

The dataset GSE5462 contains gene expression data of 116 biopsies from 58 breast cancer patients at two time points: pre-treatment and 10–14 days after treatment with letrozole [37]. We log2-transformed and normalized the raw data using robust multiarray averaging (RMA, R-package affy, [13]) for subsequent DBN analysis.

The second dataset, GSE37182, contains expression data of 172 biopsies from 15 colorectal cancer patients, totaling 88 normal tissue biopsies and 84 tumor tissue biopsies [40]. The samples were obtained during surgery and left at room temperature at four time points: 20 min ($t = 0$), 60 min ($t = 1$), 180 min ($t = 2$) and 360 min ($t = 3$). Afterwards, the samples were stored at $-80°C$ until RNA extraction. The data from the repository were already normalized separately within each group (tumor and non-tumor). To make samples between two conditions comparable, we used the package NormalizerDE [58] and performed median normalization.

Gene filtering

To select genes to be included in the DBNs we performed DGE analysis using the R package limma [46]. We considered genes as differentially expressed between conditions if their false discovery rate (FDR)-adjusted P-value was smaller than 0.05. We did not apply a log2-fold-change cutoff.

Simulation studies

We generated 50 two-step DBNs structures. For each DBN structure, we generated 30 training samples from four consecutive time slices and two test samples. We learned MAP and consensus structures corresponding to posterior thresholds of $p \in \{0.3, 0.5, 0.7, 0.9, 0.99\}$ using the Bayesian approach implemented in BiDAG ([52], Section 2.4). We also learned best-scoring structures using greedy hill climbing and the BIC score from the R-package bnlearn [49] with the limits on the number of parents of 3 and 5. For each limit, we also learned consensus structures based on bootstrap support levels of $p \in \{0.3, 0.5, 0.7, 0.9, 0.99\}$. Finally, we learned DBN structures using the R-packages dbnR and bnstruct.

dbnR implements the MMHC approach for DBNs [54].
The package bnstruct implements hill climbing as well, however it automatically discretizes continuous data.

We compared the learned structures with the ground truth using true positive rate (TPR), FDR and structural Hamming distance (SHD). SHD is defined as the number of edge additions, deletions and reversals needed to make the two graphs match [54].

**Results**

**Simulated data**

In the simulation studies (Section 2.7), we generated 50 random DBNs and data from these DBNs, followed by network reconstruction using available software packages. We explored the situation when the number of observations between neighboring time points is smaller than the number of nodes.

The MCMC approach reached the highest TPR, followed by hill climbing and MMHC (Figure 3A). However, hill climbing applied to discretized data showed the worst result discovering less than 40% of true positives. Such a poor performance likely demonstrates the effect of information loss due to data discretization. Notably, all best-scoring structures resulted in a high FDR. Structural overfitting of maximum score structures in the high-dimensional setting was previously demonstrated in [26] and Bayesian model averaging proved to be effective for decreasing the FDR.

Only bnlearn and BiDAG provide tools for model averaging. Hence we did not include other approaches in the comparison of consensus graphs. We observed that consensus structures contained fewer false-positive edges but also fewer true positives. SHD, which sums all differences (TP, FP and errors in directions of edges), was lower for consensus than for MAP structures (Figure 3B). The lowest SHD was achieved with the MCMC scheme at the posterior threshold of 0.99, demonstrating an advantage of using BiDAG for DBN structure learning.

The better performance of BiDAG comes at the cost of longer runtimes. BiDAG needed 14 min to find the MAP structure and perform the sampling from the posterior distribution. Hill climbing required 4.5 min, including 100 bootstrap runs needed to estimate consensus structures. Hill climbing applied to discretized data required the longest time of 68 min to learn the best scoring graph.

**Analysis of time-series gene expression data**

We applied the proposed approach to two transcriptomic datasets from the GEO repository (Section 2.5, Section 4): the colorectal cancer dataset GSE37182 and the breast cancer dataset GSE5462. For each dataset, we learned several DBN models (Sections 2.1–2.2) using the Bayesian approach and measured, via leave-one-out cross-validation (CV), how they perform with regard to predictive accuracy and classification accuracy (Section 2.3).

For the colorectal cancer dataset, we learned a DBN with time-varying parameters and compared it with a DBN assuming parameters that stay constant across all time slices \( t > 0 \). A time-varying DBN can describe the underlying process with higher precision. However, it can also lead to overfitting.

**Analysis of breast cancer time-series gene expression data**

The GSE5462 dataset contains gene expression measurements for two groups of breast cancer patients: responders and non-responders to treatment (Section 2.5). We selected the genes that were either differentially expressed between responders and non-responders or differentially expressed in post-treatment compared with pre-treatment samples (Section 2.6). In addition, we included all transcription factors of the identified genes found in the database Omnipath [55].

The best model learned by BiDAG yielded a higher classification accuracy than naive Bayes and random forest (Table 2). We further noted that all models in this work outperformed the highest classification accuracy of DBN models reported in [24] as well as the ML approaches that the authors used for comparison.

The lowest MAE was reached for DBNs learning the same DBN structure for both subgroups (Table 3). This finding aligns well with the differential gene expression and pathway enrichment analysis. Since out of 22283 genes, only 19 were differentially expressed, we can assume that the GRNs are very similar in responders and non-responders. However, the highest classification accuracy of 0.85 was reported for models that learned DBN structures independently for responders and non-responders (Table 2).

We chose the MAP model for the downstream analysis that learned the same DBN structure for responders and non-responders and blacklisted all non-STRING interactions. Even though the classification accuracy of this model was only the second highest, the lower MAE suggests that it better predicts the changes in post-treatment gene expression levels and hence is more appropriate for the analysis of gene expression dynamics.

Pathway enrichment analysis showed that no KEGG [21] pathway was enriched in the differentially expressed genes. However, when we assessed the set of all parent nodes of these genes in the estimated DBN structure (Supplementary data, Figure S1), three KEGG pathways (p53 signaling, cellular senescence and cell cycle) were enriched (FDR < 0.05). Thus, the DBN model connected genes found to be important for treatment response to

**Table 1. Cross-validated classification accuracy demonstrated by DBN-based and standard classification tools**

| Model                          | Accuracy | # genes |
|-------------------------------|----------|---------|
| BiDAG, different DBN structures | 0.85     | 125     |
| Naive Bayes                   | 0.83     | 125     |
| Random forest                 | 0.79     | 125     |
| BiDAG, same DBN structure     | 0.79     | 125     |
| DBN-based, Kourou et al. [24] | 0.71     | 39      |
| Various ML approaches Kourou et al. [24] | 0.58–0.66 | 39 |
Discovering GRNs using DBSs

Figure 3. Comparison of performance of DBN structure learning algorithms on simulated data. A total of 50 random two-step DBN structures were generated with \( n = 120 \) nodes and three parents on average for each node in the transition structure. The training datasets contained 30 samples from four consecutive time slices, the test datasets included two samples each. MCMC (blue, R-package BiDAG), hill climbing (HC, red and green, R-package bnlearn), MMHC (violet, R-package dbnR) and hill climbing applied to discretized data (HC.discr, yellow, R-package bnstruct) were used to learn the DBN structures and compare them with the ground truth using (A) the TPR and FDR and (B) SHD. The performance of the hill climbing was evaluated for two limits for the parent set size: \( \text{maxp} = 3 \) (red) and \( \text{maxp} = 5 \) (green). Consensus models for MCMC and hill climbing were learned using a range of posterior thresholds and bootstrap support levels of \((0.3, 0.3, 0.7, 0.9, 0.99)\).

Table 2. Ten DBN models with the lowest cross-validated MAE learned by BiDAG for the breast cancer dataset

| Model          | intra_edges | blacklist       | prior  | init_trans        | class_structures | MAE  |
|----------------|-------------|-----------------|--------|-------------------|------------------|------|
| MAP            | +           | non-STRING      | -      | sharing intra     | same             | 0.428|
| MAP            | +           | non-STRING      | -      | sharing intra     | different        | 0.436|
| MAP            | +           | non-STRING      | -      | no sharing        | different        | 0.437|
| consensus      | -           | non-STRING      | -      | sharing intra     | same             | 0.438|
| consensus      | +           | non-STRING      | -      | no sharing        | same             | 0.438|
| consensus      | +           | non-STRING      | -      | sharing intra     | different        | 0.448|
| consensus      | -           | non-STRING      | -      | no sharing        | different        | 0.448|
| MAP            | -           | non-STRING      | STRING | sharing intra     | same             | 0.450|
| consensus      | -           | STRING          | STRING | sharing intra     | same             | 0.452|

genes from major cancer-related pathways. Among these genes, the most connected node was CDK1 (Cyclin Dependent Kinase 1), which is a known target for treating breast cancer [22]. Interestingly, Cdk inhibitors are already approved for treating breast cancer as the first-line treatment in combination with letrozole (used in the analyzed dataset) [47] which confirms the discovered link.

Analysis of colorectal cancer time-series gene expression data

For the colorectal cancer dataset GSE37182, we performed the DGE analysis at three consecutive time points, using \( t = 0 \) as a reference (Section 2.6). The number of differentially expressed genes increased with time. In total, we identified 58 genes that were
differentially expressed over all time points in cancer and tumor biopsies.

We proceeded with the identification of transcription factors that may be involved in regulating the identified genes using the Omnipath database. We combined them with the first set of genes and used their union for the DBN analysis with BiDAG. We learned multiple DBN models using various structural and parametric assumptions (Sections 2.1–2.2) and performed cross-validation as described above to select the best model (Section 2.3). The classification accuracy was 100% for all models and higher than the accuracy of the best model reported in [24] (98.5%). The MAE was clearly the lowest for DBNs assuming time-varying parameters (Table 4) as none of the 10 best models assumed constant parameters. From a biological perspective, the time-varying model is also plausible. First, the time lags between the measurements were nonuniform. Second, the tissue was left at room temperature, and the process of degradation likely led to changes in the strengths of dependencies between genes. Among the time-varying models, the lowest MAE was reached for models where intra-edges were prohibited.

Finally, we observed that DBN models that learn structures for tumor and normal subgroups independently resulted in the lowest MAE. Consequently, for the downstream analysis, we selected a consensus DBN model which learns structures separately for normal and tumor samples and blacklists intra-slice edges. Despite being learned independently, the DBN models for cancer and normal subgroups shared 60% of edges. Such a high overlap suggests that a lot of underlying processes in cancer and normal cells can be described by the same dependencies between genes.

To highlight the differences and similarities between the analyzed phenotypic groups, we identified the nodes with the most similar and different interaction partners in networks representing tumor and normal subgroups. There were 18 nodes that had neighborhoods with empty intersections in two networks. Out of these, three genes (FOS, JUN, GADD45B) belong to the KEGG colorectal cancer pathway. Two genes from this set, namely FOSB and JUN, were identified and validated as markers for colorectal tumor tissue degradation [40] (Figure 4A). Out of 20 nodes with most similar neighborhoods 10 can be found of a generic transcription pathway (Figure 4B, Supplementary data, Figure S2, [4]).

**Discussion**

DBNs are powerful models for analyzing time-series gene expression data because they allow us to shed light on the GRNs that orchestrate molecular processes. Recently, a lot of research has focused on learning context-specific gene networks [23, 27, 42, 51]. In this work, we proposed a framework for learning DBNs for multiple phenotypic groups. This framework employs the Bayesian approach to structure learning of DBNs implemented in the R-package BiDAG and provides a broad range of modeling options, including various DBN topologies, constant and time-varying parameters, the inclusion of priors as well as Bayesian model averaging. We demonstrated in simulation studies that BiDAG outperforms other available tools for DBN structure learning.

We applied the proposed framework to two time-series gene expression datasets, each comprising data from two subgroups of samples. The GSE5462 dataset included gene expression data of breast tumor biopsies taken before and after treatment with letrozole. Our analysis suggested that GRNs do not differ substantially between responders and non-responders. However, the analysis of the learned DBN structure suggested that differences in the signaling pathways of the subgroups might lie at the phosphoproteome level since the kinase CDK1 appeared to be densely connected to a few genes that were differentially expressed between the responders and non-responders. Even with only a few differences detected at the gene expression level between the subgroups, the classification accuracy was higher than reported in the previous study [24].

For the colon cancer dataset, the best predictive accuracy was reached for the model assuming different DBN structures for tumor and normal samples. This finding indicates that GRNs differ considerably in normal and tumor cells and aligns well with experimental results obtained from the analysis of normal and tumor cell lines.
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Figure 4. Subnetworks of DBN transition structures discovered for the GSE37182 dataset. Structures of time-varying DBN models without intra-edges were learned independently for normal and tumor subgroups. The blue edges denote the edges which are specific to the normal model, red edges are specific to the tumor model. Black edges are present in both models. The solid lines correspond to edges between genes which were found as interactors in the STRING database. Genes from the colorectal cancer pathway (KEGG) are highlighted in orange. (A) Most differently connected genes (FOSB, JUN, FOS, GADD45B) in DBN transition structures of cancer and normal DBN models that are either enriched in the colorectal cancer signaling pathway or were previously validated as biomarkers of cancer and their parents in the learned DBN models. (B) Most similarly connected genes, which were also found on the generic transcription pathway [4] (highlighted in green).

Key Points
- The proposed strategy for learning GRNs of multiple phenotypic groups unifies the efficient method of DBN structure learning and the versatile approach to model selection, enabling the discovery of models with high predictive and classification accuracy.
- The efficient Bayesian approach to structure learning is better equipped to prevent overfitting than greedy hill climbing coupled with other conventional techniques.
- Application of the proposed method to the real transcriptomic data revealed differences and similarities between the regulatory networks of cancer and normal cells that aligned well with previous findings and can be used to facilitate the discovery of targeted therapies.

Supplementary data
Supplementary data are available online at https://academic.oup.com/bib.

Author contributions statement
P.S. and J.K. conceived the research project. N.B. supervised the research project. P.S. and J.K. designed and implemented the computational framework and conducted the analyses. P.S. and N.B. wrote the manuscript. N.B. and J.K. reviewed and edited the manuscript.

Data and code availability
The unprocessed data is available at the public GEO repository under identifiers GSE5462 and GSE37182.
The datasets can be accessed at https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE5462 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE37182

The reproducible code and the results are available at the GitHub repository https://github.com/cbg-ethz/DBNclass. The latest version of the BiDAG package including implemented updates is available at CRAN repository https://cran.r-project.org/web/packages/BiDAG

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