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Inferring Causal Relations from Multivariate Time Series: A Fast Method for Large-Scale Gene Expression Data

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Abstract—Various multivariate time series analysis techniques have been developed with the aim of inferring causal relations between time series. Previously, these techniques have proved their effectiveness on economic and neurophysiological data, which normally consist of hundreds of samples. However, in their applications to gene regulatory inference, the small sample size of gene expression time series poses an obstacle. In this paper, we describe some of the most commonly used multivariate inference techniques and show the potential challenge related to gene expression analysis. In response, we propose a directed partial correlation (DPC) algorithm as an efficient and effective solution to causal/regulatory relations inference on small sample gene expression data. Comparative evaluations on the existing techniques and the proposed method are presented. To draw reliable conclusions, a comprehensive benchmarking on data sets of various setups is essential. Three experiments are designed to assess these methods in a coherent manner. Detailed analysis of experimental results not only reveals good accuracy of the proposed DPC method in large-scale prediction, but also gives much insight into all methods under evaluation.

I. INTRODUCTION

One major step in recent genomic research is the advance of high-throughput microarray technique, which allows expression levels of all genes in the genome to be measured at particular time points. The resulting gene expression dynamics are important, since they directly reveal the active components within the cell over time, indicating gene regulatory relationships on the transcriptional level. However, they also pose a dimensionality problem in the subsequent data analysis, with the number of genes far exceeding the number of samples/time points [1]. The situation is quite the opposite to classical time series analysis, thus objective technique is needed for learning gene regulatory relationships from these data. Also, large number of genes/variables poses a challenge to interaction inference in a directed form, since there are twice as many possibilities as there are in an undirected network.

Many methods have been proposed for studying the interdependence/causality relationships between genes/variables. The study should ultimately lead to the reconstruction of gene regulatory networks and provide new insights into the functioning of the regulatory system. One of the most popular directed network inference methods, dynamic Bayesian networks (DBNs) [2], [3] has been applied in this area. DBNs are graphical models trained to maximise the joint probability of a set of observed data and their conditional dependencies. DBNs have been routinely applied to data, mainly long time series, to provide information about system dynamics. However, a major concern about DBNs is its inefficiency in large-scale prediction, i.e., with the presence of many variables.

Recently, a directed network inference approach, namely shrinkage vector autoregressive method (SVAR), was proposed by Rhein et al. [4] to circumvent the small sample problem. The basic procedure consists of first computing the shrinkage estimates of covariance matrices to obtain regression coefficients for fitting autoregressive models. Then, instead of using the regression coefficients directly, the corresponding partial correlation coefficients are statistically tested. Significant coefficients are selected using False Discovery Rate (FDR) [5] to be included into the reconstructed network.

Another recent advance in this area is the introduction of the concept of Granger causality (GC) [6], a statistical technique for causal inference well known in economics. Time series A is said to Granger cause time series B, if the forecast of B has incremental predictive power with the addition of A. The predictive power can be measured by the variances of residuals as a result of linear model fitting. Informally, the method measures the influence of one time series on another by checking if the prediction of the response can be improved by incorporating the knowledge of a predictor. Subsequently, for the application to gene expression data, one of the first attempts is a simple bivariate model that uses Granger causality to infer relationships between pairs of variables without taking into account other variables [7]. For
the purpose of our comparative experiments, we implemented a multivariate model in R, since the bivariate model could lead to false positive edges such as the ones in Fig. 1(c), compared with the true network (Fig. 1(a)).

All above three methods infer directed networks. One undirected but popular method, a shrinkage estimate method using graphical Gaussian models (GGMs), is proposed by Schäfer and Strimmer [8] to tackle the dimensionality problem. GGMs are undirected models which describe conditional dependence structure among variables. In essence, partial correlation is used as the mathematical foundation for establishing direct interactions among genes (Fig. 1(a)). When inferring the relationship between two temporal signals/gene expressions, the rest of the signals are also taken into account in the computation of partial correlation to discriminate between direct (Fig. 1(a)) and indirect (Fig. 1(b)) interactions. Significant coefficients of partial correlation can then be selected using FDR for the reconstruction of gene networks.

Although this method is fast and well suited for small sample data analysis [9], the inferred interactions are undirected. In an undirected network, the role that a gene plays in the regulatory activities is unknown. Therefore, based on partial correlation, we propose a directed approach specifically targeted at small-sample gene expression data. It is then compared with the existing directed inference methods as described above, DBNs, SV AR, and Granger causality, to demonstrate its effectiveness.

Although there are many comparative studies of interdependence inference techniques in the literature [10], [11], few of them are conducted on microarray data sets. For example, in a comparative study on inference algorithms for multivariate time series interaction [10], GC is reported to perform well for stationary times series data, but is sensitive to non-linearity. In contrast, our study focuses on the small sample problem in gene expression data analysis. This paper aims at illustrating the application of multivariate time series analysis in the reign of gene interaction inference. It also sheds light on the application of multivariate time series analysis in gene expression data analysis. This paper aims at illustrating the technical details for three representative methods, focusing on their abilities in analysing gene expression data. Next, the proposed algorithm is formulated. These technical details provide us strong foundation for the discussions later on experimental results. Based on the interpretation of experimental results, we hope to shed some light on the nature of inference techniques, their advantages and inherent problems.

A. Existing multivariate time series inference methods

1) Vector autoregressive models (VAR): Suppose \( Y = \{y_i|i = 1, 2, ..., n\} \) is a multivariate stationary time series consisting of \( n \) variables and \( t \) time points. A \( p \)-order vector autoregressive VAR(p) model specifies that the value of the \( i \)th variable at time point \( t, y_i(t) \), is a linear combination of a constant/mean value, the past of the multivariate time series, and a noise component

\[
Y(t) = B + A \sum_{u=1}^{p} Y(t-u) + \varepsilon(t). \tag{1}
\]

\( B \) is a constant matrix of size \( n \times t \). \( \varepsilon \) consists of vectors of residuals \( \{\varepsilon_i|i = 1..n\} \), each assumed to be zero mean noise with covariance matrix \( \Sigma_i \). \( A \) is the \( n \times n \) coefficient matrix representing the dynamic structure. When \( A \) is a constant matrix, this model assumes homogeneity across time. A special case of the \( p \)-order VAR process, the first-order autoregressive model (VAR(1)), is often considered when analysing microarray data for the sake of simplicity [3], [4]

\[
Y(t) = B + AY(t-1) + \varepsilon(t). \tag{2}
\]

2) Granger causality inference method (GC): We start with Granger causality method (GC) in the multivariate case. Let \( Y^- \) symbolise the past state of \( Y, Y^- = \{Y(u)|u = 1, ..., t-1\} \), and let \( y^-_i \) symbolise the past of variable \( y_i \). The Granger causality measure of prediction power of one variable \( y_i \) on the other variable \( y_j, i \neq j \), is defined by

\[
g_{y_i \rightarrow y_j} = \ln \left( \frac{\sigma_{y_i|Y^-}}{\sigma_{y_j|Y^-}} \right). \tag{3}
\]

Symbol “\(|\)” denotes operation “condition on” and symbol “\( /\)” denotes “without”. \( \sigma_{y_i|Y^-} \) is the variance of the residual \( \varepsilon(t) \) in the VAR(1) model for \( y_j \) conditioned on the past of all variables \( Y^- \). It is compared to \( \sigma_{y_j|Y^-} \), which is conditioned on the past of all variables but \( y_j \), \( Y^- \). GC directly measures the prediction power of \( y_i \) for \( y_j \), as a result of the reduction of prediction errors by incorporating \( y_i \) into the VAR(1) model for \( y_j \). In other words, if introducing \( y_i \) significantly reduces the variance of the prediction error of \( y_j \), then a variable \( y_i \) Granger causes the variable \( y_j \). Since it requires fitting autoregressive model with all variables and their past states, GC can only be applied to data satisfying: \( t > n(p + 1) \), indicating its limited potential in gene expression analysis.
3) Shrinkage VAR method (SVAR): Although the VAR model has been widely used in economics and neuroscience, it has its own limitations when small samples are encountered. An effective shrinkage estimation procedure was developed for learning the VAR models from small sample data [4]. The idea is that a shrinkage estimate can replace the covariance matrix for the joint matrix of both the present data and the lagged data, which then leads to the computation for regression coefficients. The covariance matrix would be otherwise ill-conditioned, given the large number of variables (2 × n) and short time series \( t, t \ll n \).

Let \( \Phi \) denote the joint matrix of the multivariate \( Y \)'s present state \( (Y^+ = \{Y(u)|u = 2, \ldots, t\}) \) and past state \( (Y^- = \{Y(u)|u = 1, \ldots, t - 1\}), \Phi = [Y^+Y^-] \). Assuming that the data has zero mean, an unbiased estimate of the covariance matrix for \( \Phi \) is

\[
\text{cov}(\Phi) = \frac{1}{t-1} [Y^+Y^-][Y^+Y^-] \tag{4}
\]

Note that this matrix contains the sub-matrices \( Y^{-}Y^{-} \) and \( Y^{-}Y^{+} \). Meanwhile, the ordinary least squares (OLS) estimation [12] for the regression coefficient \( A \) in the VAR(1) model (Eq.2) is:

\[
\hat{A}(1) = (Y^{-}Y^{-})^{-1}Y^{-}Y^{+}. \tag{5}
\]

Therefore, the shrinkage estimation of \( \text{cov}(\Phi) \) will lead to the estimated coefficient matrix \( \hat{A} \). Then the partial correlation coefficients \( q \) can be computed from \( \hat{A} \) and the FDR is used to select significant coefficients. With large number of variables, this method gave good result in the comparative simulation study using simulated autoregressive data in the original paper [4].

4) Dynamic Bayesian networks inference method (DBNs): DBNs implementations are usually designed for data with hundreds or thousands of samples. The limitation of microarray experimental costs prohibits most of the techniques from exploring small sample gene expression data. In this paper, we use the implementation of the R package G1DBN [3], which is based on a trivariate AR(1) model:

\[
\begin{align*}
Y(1) & \sim \mathcal{N}(\mu_1, \Sigma_1), \\
Y(t) & = B + AY(t - 1) + \varepsilon(t), \\
\varepsilon(t) & \sim \mathcal{N}(0, \sigma),
\end{align*} \tag{6}
\]

with predefined \( \mu_1, \Sigma_1, \) and \( \sigma \). This method measures the conditional dependence between two variables \( y_i, y_j \) by testing the null hypothesis \( H_0: \alpha_{ijk} = 0 \) on every third variable \( \{y_k | k \neq i, j\} \). Then, a score is assigned to the potential edge \( y_i \rightarrow y_j \) corresponding to the maximum \( p \)-values from the tests \( p_{\text{max}}(y_i \rightarrow y_j) \). This means the algorithm has a computational complexity of \( \mathcal{O}(n^3) \). The computation of this method may be too heavy for data with more than a hundred variables. Since the simulated data are generated following the characteristic assumption of small sample, we hope to cast light on this particular aspect.

III. PROPOSED DIRECTED PARTIAL CORRELATION INFERENCE METHOD (DPC)

The shrinkage estimate for partial correlation in [8] was formulated specifically for the inference from small sample gene expression data. Although partial correlation is undoubtedly fast in computation and suitable for small sample problem, it can only infer undirected networks. Another problem is, variable time lag cannot be taken into account as in a VAR(1) model. We introduce the notion of directed partial correlation (DPC) for fast inference of directed gene networks. The idea is similar to the idea behind Granger causality - a variable A has causal influence on another variable B, if the removal/addition of A has a large impact on the prediction of B. While GC measures this impact by comparing the residuals before and after adding A to the prediction of B, DPC measures it by examining the correlation coefficients.

A. Zero-order directed partial correlation DPC(0)

Directed partial correlation aims to investigate the effect of including a variable into the predictions of another gene, i.e. the change of partial correlations among other genes. Let \( Q_Y \) of size \( n \times n \) denote the partial correlation matrix for \( Y \). Each element \( q(i,j|Y) \) in \( Q_Y \) is the partial correlation between \( y_i \) and \( y_j \) given \( Y \), \( i = 1, \ldots, n, j = 1, \ldots, n, i \neq j \), i.e., the correlation between \( y_i \) and \( y_j \) after the linear effects of the rest of variables are removed. This can be formulated as \( q(i,j|Y) \).

Removal of linear effects from others means that resulting partial correlation indicates the direct relationship between two variables. Fig. 2(a) shows \( q(i,j|Y) \), \( k \neq i, j \), which denotes the partial correlation between \( y_i \) and \( y_k \) when effects from all others, including \( y_j \), are removed.

However, relationship indicated by \( q(i,j|Y) \) is undirected. To investigate the influence \( y_j \) has on \( y_i \), we propose the following. If we delete the variable \( y_j \) from \( Y \), the partial correlation between \( y_i \) and another variable \( y_k \), \( k \neq i, j \), is denoted as \( q(i,k|Y_{-j}) \) in the matrix \( Q_{Y_{-j}} \). As shown in Fig. 2(b), in the prediction of relationship between \( y_i \) and any other variable \( y_k \), \( k \neq i, j \), \( q(i,k|Y_{-j}) \) no longer remove the effect from \( y_j \), which means \( y_j \) no longer take part in the prediction.
of \( y_i \). Consequently, there are two groups of statistics related to the prediction of \( y_i \), each corresponding to coefficients before and after the removal of \( y_j \). To be more specific, the first group is the \( i \)th row in \( Q_y \) without the \( i \)th and \( j \)th element, \( g_1 = \{ q(i,j\mid Y), j \neq i \} \), shown in dark green in Fig. 2(a). The second group corresponding to the dark green elements in Fig. 2(b) is the \( i \)th row in \( Q_y \) without the \( i \)th element, \( g_2 = \{ q(i,j\mid Y/k), j \neq i, k \} \). Both groups have the length of \( n-2 \). The effect \( y_j \) has on the prediction of \( y_i \) is defined as:

\[
e^0_{y_i \rightarrow y_j} = t\text{-test} (g_1, g_2)
\]

for the joint matrix \( \Phi = [Y^+ Y^-] \), with \( Y^+ \) the present state \( Y^+ = \{ Y(u) \mid u \in 2...t \} \) and \( Y^- \) the past state \( Y^- = \{ Y(u) \mid u \in 1...t-1 \} \).

We use Student’s t-test on the two groups to see if there exists an effect on the prediction of other variables with the removal of variable \( y_j \). The null hypothesis is that there is no significant difference between the two groups, before and after the removal. Student’s t-test compare the sample means:

\[
\tau = \frac{\bar{g}_1 - \bar{g}_2}{\sqrt{(n-2)^{-1}(\sigma_{g1}^2 + \sigma_{g2}^2)}}
\]

where \( \sigma_{g1} \) stands for the standard deviation of \( g_1 \), the denominator of \( \tau \) is the standard error of the difference between two means, the degree of freedom for the test is \( 2n - 6 \).

In summary, we take advantage of the fact that in computing partial correlation between two variables, all effects from other variables need to be removed. In other words, \( y_j \) takes part in the predictions of \( y_i \) with all other variables \( y_k \). We measure \( y_j \)’s influence on \( y_i \) by comparing partial correlation coefficients related to \( y_i \) before and after the deletion of \( y_j \), since \( y_j \) does not take part in the prediction of \( y_i \) after the deletion.

B. First-order directed partial correlation DPC(1)

A key feature of the proposed DPC method is that it can be easily extended to include time lags. The correlation between the variables measured as a function of time lag is of interest because such a time lag may reflect a causal relationship. Let \( \Phi \) be the joint matrix of the present state and the past state of data, \( \Phi = [Y^+ Y^-] \). To compute the correlation matrix for \( \Phi \), we note that the covariance matrix of \( \Phi \) is ill-conditioned for small sample data and therefore not suitable. We use the shrinkage estimate method in Eq.4 to compute the partial correlation matrix \( Q_{Y}^{(1)} \) for \( \Phi \):

\[
Q_{Y}^{(1)} = \begin{bmatrix}
Q^{++} & Q^{+-} \\
Q^{*-} & Q^{--}
\end{bmatrix}
\]

Hence each element in the sub-matrix \( Q^{++} \), \( q^{(1)}(i,j) \) with \( i = 1...n, j = 1...n \), stands for the partial correlation between \( y_i \) and \( y_j \), when the effects of the present states of other variables and the past states of all variables are removed. If a variable \( y_j \) is deleted from the joint matrix \( \Phi \), the corresponding partial correlation matrix \( Q_{Y|\beta}^{(1)} \) has equivalent meaning as described in the zero-order model, i.e. the effect of \( y_j \) is not taken into account in the prediction of the other variables. The first-order directed partial correlation from \( y_j \) to \( y_i \) can be formulated as:

\[
e^{(1)}_{y_j \rightarrow y_i} = t\text{-test}\left( \{ q^{(1)}(i,j\mid Y) \mid j \neq i \}, \{ q^{(1)}(i,k\mid Y) \mid k \neq i, k \} \right).
\]

Note that although the partial correlation matrix is of size \( 2n \times 2n \), only the sub-matrix \( Q^{++} \) is used for computing \( e^{(1)} \). The probability of the directed interaction is indicated by the resultant \( p \)-values. Using FDR, adjusted \( p \)-values are selected in accordance to confidence levels, for example, 2% of FDR means accepting all tests with adjusted \( p \)-values < 0.02 as significant.

Conceptually, DPC tests the effect of one variable on the predictions of another by all the rest of variables at the same time, hence is able to monitor the dynamic process within reasonable computation time. It avoids linear model fitting, thus is more efficient and less constrained by the sample size.

IV. EXPERIMENTS

Previously, SVAR and DBNs were experimentally proved to be useful using simulated data from autoregressive models [3], [4]. These methods are based on the autoregressive model and their performance on other types of data is still not clear. When the data satisfy the model assumption, we can expect the corresponding technique to perform well. Therefore, an important question pertains to which assumption best describes gene expression data. In this section, we aimed at investigating the following question: how well the inference methods can meet the requirements of microarray data.

Since real expression data are generally noisy, they may not be fully reflective of the gene relationships and the ground truth is unknown, comparisons of performance are conducted
with synthetic data. SynTReN is well suited for testing module network algorithms [13]. By using topologies generated based on previously described source networks, SynTReN allows good approximation of the statistical properties of real biological networks.

Four multivariate time series inference algorithms as described above are evaluated in this experiment. Their ways of inferring the final network vary and each requires fine tuning for the parameters, which could be subjective for large-scale experiments (in the synthetic data experiment altogether 142 data sets are used). To eliminate any subjective element and enable a fair comparison, we decide to compare directly on their preliminary output, the score matrices. For clarity, the related symbols for each score matrix in the algorithms’ technical details are listed in Table II.

For the inferred score matrices, we compute their true positives (TP), false positives (FP), true negatives (TN), and false negatives (FN) given a threshold. This procedure was repeated 500 times for each test statistic and variance scenario, to obtain Receiver Operator Characteristic (ROC) curves [14], [15] for describing the dependence of true positive rate/sensitivity $TPR = TP/(TP + FN)$ and false positive rate/specificity $FPR = FN/(TN + FP)$. ROC curves provide a straightforward graphical representation of the performance of the algorithms, hence are especially useful in statistically principled comparisons. It avoids issues related to a chosen threshold, by using all possible thresholds. As a summary metric for ROC, the area under the ROC curve (AUC), as its name indicates, can measure the average accuracy of the prediction.

While AUC provides quantitative measurement on average performance for a method, maximum F-score [16] evaluates each method at its point of optimum. F-score is the harmonic mean of precision ($TP/(TP + FP)$) and recall ($TP/(TP + FN)$). As a composite measure, F-score challenges algorithms with higher specificity and benefits algorithms with higher sensitivity. Both of the metrics are used when appropriate in the following experiment section. Apart from these metrics, we also base our evaluation on the time consumed for the computation and the true positive rate at the point of 0.2 false positive rate, since usually a low false positive rate is preferred.

SynTReN produces synthetic transcriptional regulatory networks and the corresponding simulated microarray data sets, parameterized by the network topology, size of the network/number of genes, levels of biological, experimental, and input noise etc. Network topologies can be generated by selecting sub-networks from previously described biological networks or by using random graph models. The former method is used here to offer better approximation. Two different strategies to select a connected subgraph are implemented: neighbour addition (neighAdd) and cluster addition (clusterAdd). It was suggested that sub-network selection by cluster addition is preferable [17], since the resulting sub-network preserves features of scale-free networks such as having hubs. However, consider that during variable selection process one may not include all neighbours of a hub gene, sub-networks by neighbour addition may sometimes represent a more realistic situation in gene network analysis. Hence we use both strategies in simulated data generation.

After the topologies of the synthetic networks are sampled, transition functions can be determined and enzyme kinetic equations are selected for each gene and its regulators. Combined with external conditions that trigger the network, the expression levels of genes in each experiment are generated according to the activities of their regulators. The final data generated by SynTReN are quantiled to the range of [0, 1] where 0 indicates no expression and 1 indicates maximal expression. We normalize the data to the $log_2$ ratio by selecting one of the samples as the control.

We design three sets of experiments in order to assess the methods’ performance in a coherent manner. In the first and second experiment, data sets of small sample, varying network sizes and data sets of large network, varying sample sizes are generated. In the third experiment, sample size slightly larger than network size/gene number are generated. However in the first experiment, data sets are analysed by DPC, SVAR and DBNs but not GC, since GC requires sample size larger than network size ($n < t$).

The experiments are designed to give a thorough evaluation of the proposed algorithm, to observe its performance in different scenarios, to compare four algorithms in a coherent manner. Specifically, we note that settings in the first experiment are perhaps closest to realistic situation for microarray data analysis, and therefore results from the first experiment should receive careful consideration and interpretation.

A. Networks of various sizes and small sample size

In this experiment, we assess the influence of network size on the performance of inference algorithms, by fixing the sample size to a small number and vary the network size.
This means GC cannot be applied in this experiment, since it requires long time series \((t \gg n)\) to fit linear models. First, \textit{E.coli} sub-networks are selected by cluster addition as the network topologies. We generate altogether 45 data sets with 25 samples varied in network size from 10 nodes to 300 nodes, and 45 data sets with 15 samples varied in network size from 10 nodes to 300 nodes. The configurations in the network topology selection, sample size and network size are provided in Table I. In this table, parameter noise level refers to all the noise parameters in SynTReN: levels of biological, experimental, and input noise. They are set to the same value. The noise level is set relatively lower for the 15-sample data sets than the 25-sample data sets.

For each algorithms, the resulting AUC values for data set 1, 2 and 3 are averaged and plotted in Fig. 3(a), so are the inference results for data sets 4, 5, and 6 in Fig. 3(b). Because of the computation costs of DBNs, we only compute results for networks of size 10 \(\sim\) 100 for the 25-sample data and size 10 \(\sim\) 150 for the 15-sample data. The plots show the effect of the amount of available gene expression data on the inference results. Although SVAR and DBNs sometimes outperform DPC with smaller networks/less variables, when faced with increasing network size, their performance both drops rapidly when there are only small amount of data available. However, DPC’s overall performance dramatically improves from the beginning, i.e. for smaller networks. Then it stays the same regardless of the changes in network size.

Generally, the performance of most of the algorithms degrades as the number of genes increase. This conforms to current theory. In contrast, DPC outperforms others only when the network size is big enough. However, this is reasonable. The two-sample test DPC based on is effective only when there are big enough sample populations, which, in this scenario, are equivalent to the number of genes/variables. In summary, DPC shows superior performance in inferring large-scale gene networks, although for small networks it is sometimes outperformed by SVAR and DBNs.

### B. Networks of fixed size and various sample sizes

To assess the influence of sample size on the performance of the inference algorithms, we generate data sets for a fixed network size but of various sample sizes. Four \textit{E.coli} sub-networks selected by cluster addition of size 50, 50, 100, and 100 genes were chosen as the network topologies. For each of the two 50-gene networks, we generate 12 gene expression data sets with sample sizes varied from 60 to 1000. For each of the two 100-gene networks, we generate 8 data sets with sample sizes varied from 120 to 1000. Altogether 40 data sets are generated and used as input for the four algorithms. The resulting AUC values for the two 50-gene networks are averaged and plotted in Fig. 4(a) and the same for the resulting AUC values from two 100-gene networks plotted in Fig. 4(b).

All algorithms show different behaviours when faced with increased amount of data. In Fig. 4(a), the performance of all methods improve as the sample size grows at the beginning (60 \(\sim\) 180 samples), but then level off eventually. Compared with the AUC values for DBNs and DPC, the AUC values for SVAR and GC show larger variations. The best performer in this case when the network size is 50 genes is DBNs, followed by DPC(0).

However, this situation is reversed when the network size is increased to 100 genes. Performance of DBNs drops sharply, reflecting its sensitivity to the network size. Interestingly, the performance of DPC improves dramatically compared to that in the case of 50-genes. With bigger network/more variables, one would expect a decent in the performance of inference methods. DPC, in contrast, perform even better, which conform to the conclusion and interpretation for DPC in the first experiment.

With increasing number of samples, the improvement of performance is not as dramatic as one would expect in the case of 50-gene networks. A performance plateau was reached at 180 samples for most of the algorithms. However, the case with 100-gene networks are quite different in that general
The AUC values for four network inference algorithms on different data sets with noise level at 10% (a) fixed network size of 50 genes, sample size 60 ∼ 1000, (b) fixed network size of 100 genes, sample size 120 ∼ 1000.

Performance only starts to improve when more than 300 samples are available. An exception is DPC, which show fairly steady performance across all sample sizes. From this experiment we can observe that expecting dramatic improvement in the performance of inference algorithms by increasing sample size is unrealistic, especially for microarray data whose samples are costly. Again DPC outperforms other algorithms for larger networks.

C. Networks of typical sizes

In the third experiment, we investigate the situation when the network size is selected to be slightly smaller than the sample size. This is close to a scenario when a researcher chooses the number of genes to be included in the network according to the number of microarray samples available. Four data set configurations with 50 genes × 100 samples, 80 genes × 100 samples, 100 genes × 150 samples, 150 genes × 180 samples are considered. Three data sets are generated for each configuration. By assigning different random seeds to SynTReN to select random nodes as a starting point, all data sets are guaranteed to relate to different network topologies.

The noise level for all the noise parameters (e.g. biological noise and experiment noise) in SynTReN is set to 10%.

To evaluate the performance of the four algorithms, we first

Fig. 5. ROC curves for the comparisons of the four network inference algorithms using their score matrices, (a) synthetic network of 50 genes and 100 samples, (b) synthetic network of 80 genes and 100 samples, (c) synthetic network of 100 genes and 150 samples, (d) synthetic network of 150 genes and 180 samples.

TABLE II

<table>
<thead>
<tr>
<th>Method</th>
<th>DPC(1)</th>
<th>DPC(0)</th>
<th>SVAR</th>
<th>GC</th>
<th>DBNs</th>
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<tr>
<td>Score matrix</td>
<td>$e^{(1)}$</td>
<td>$e^{(0)}$</td>
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<td>r</td>
<td>$</td>
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<tr>
<td>Data size</td>
<td>Average AUC value</td>
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<td>Average true positive rate (false positive rate=0.2)</td>
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plot the resulting ROC curves in Fig. 5 individually for the four scenarios. In each plot, for each algorithm, its results on all three data sets are first plotted as the dotted curves, then their boxplots are used as the summary curves.

It is easy to observe, for SVAR, GC and DBNs, a descent in their performances as network expands. In contrast, DPC shows robustness to network size in this case, when sample size is reasonable in comparison to the network size.

Quantitative measurements of performance including AUC values, true positive rates, F scores and average computation time for each methods are provided in Table II, with best results bolded. For clarity, score matrices are the symbols corresponding to the technical details in the previous section. The average performance are reported as a result of using three data sets for each typical data size. AUC values are calculated from Fig. 5, and the average true positive rates are given for each algorithm when the false positive rate is 20%. Average consumed time is for a PC (Intel Pentium 4 2.80GHz). From this table, noticeable advantage to DPC can be seen in terms of both accuracy and efficiency, although SVAR is the most efficient.

V. Conclusions

This paper reviews some recent advances on multivariate time series inference in the field of gene expression data analysis and reports a new method, aiming at shedding light on future research. We performed thorough experiments to investigate the properties of the proposed method and other methods in comparison, although settings for the first experiment are closer to realistic situation for microarray data.

Superior performance of the proposed directed partial correlation (DPC) for large-scale network inference is observed throughout the experiments. Its excellent property of robustness to gene number/network size is made explicit in the first experiment, while other methods in comparison show fast descent in their performance. Moreover, there is no obvious effect of the sample size on the performance of inference algorithm, as it is shown in the second experiment. The marginal influence on the inference results by increasing samples to a realistic limit indicates that, exploring fundamental advancement in inference algorithms is perhaps the key to success in this field.

Finally, we note that a major difference between SVAR and DPC is, SVAR inspects the regression coefficients of the full linear model, while DPC takes advantage of the idea behind Granger causality and tests the effect on the removal of individual variables. Unlike the method using Granger causality, the computation cost for DPC is low and not substantially affected by network size. Following the success of DPC on synthetic data, we would like to test it on real biological data and this will be our new line of investigation in the near future.

REFERENCES