

A Practical Approach to Inferring Large Graphical Models from Sparse Microarray Data

Juliane Schäfer

Department of Statistics, University of Munich, Germany.

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Coauthor:

Korbinian Strimmer

Department of Statistics, University of Munich, Germany

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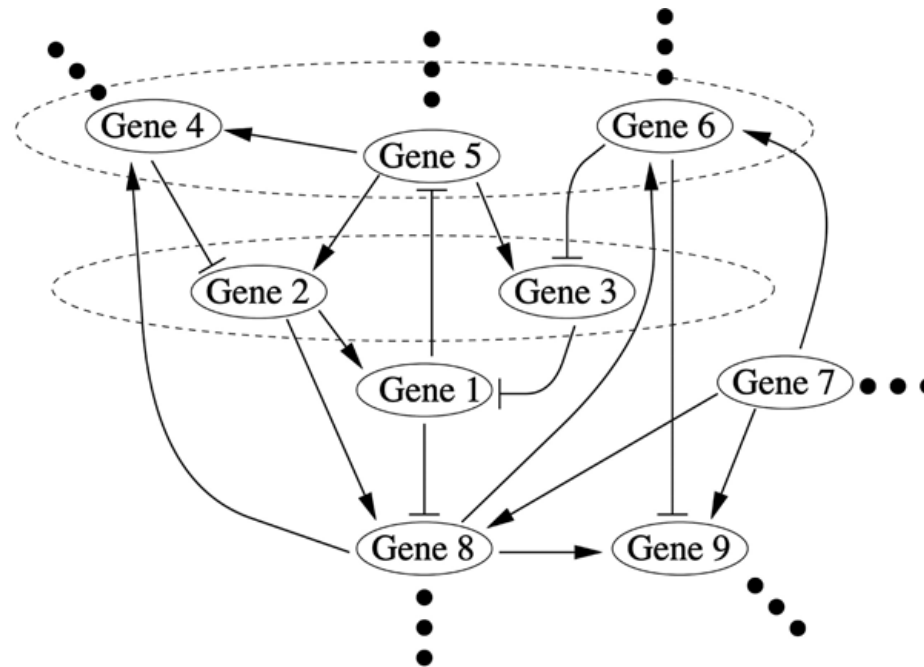
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Motivation: Gene regulatory networks

Cellular processes lead to complex dependency structure in gene expressions



Microarray experiment

Central dogma: DNA $\xrightarrow{\text{transcription}}$ mRNA $\xrightarrow{\text{translation}}$ protein

- explore transcript abundance, taken as a proxy for gene expression
- hybridization properties
- gene expression profile data: measurements under different conditions (certain points in time, treatments, tissues, etc.)

Reverse engineering problem

- Given a set of measurements (=multiple time series data), what can we deduce about the underlying network structure?

In particular:

Dimensionality problem: data feature space \gg sample size

- Challenging problem whose tractability is controversially discussed (e.g. Friedman et al. (2000) were the first to propose the use of Bayesian networks)
- What can we expect from available microarray data?

Graphical models

- Graphical models provide appropriate statistical framework:
 - association structure between multiple interacting quantities
 - distinguish between direct and indirect correlations
 - visualization in graph $G = (V, E)$
 - concept of conditional independence
- There are many different graphical models:
 - undirected vs. directed models
 - dynamic vs. static models

Some Definitions

Sample covariance matrix (with empirical mean $\hat{\mu}_i = \bar{y}_{\cdot i} = \frac{1}{N} \sum_{k=1}^N y_{ki}$)

$$\hat{\sigma}_{ij} = s_{ij} = \frac{1}{N} \sum_{k=1}^N (y_{ki} - \bar{y}_{\cdot i})(y_{kj} - \bar{y}_{\cdot j}) \quad (1 \leq i, j \leq G)$$

Empirical correlation coefficient matrix according to Bravais-Pearson

$$\hat{\rho}_{ij} = r_{ij} = \frac{s_{ij}}{\sqrt{s_{ii}s_{jj}}} \quad (1 \leq i, j \leq G)$$

Genetic Correlations

Possible reasons for high pairwise correlation coefficient:

- direct interaction
- indirect interaction
- regulation by common gene

Not accounting for intermediates can lead to considerably biased conclusions (pseudo correlations, hidden correlations)!

We are mainly interested in **direct** interactions.

Graphical Gaussian models

We focus in this talk on a very simple class of graphical models:

Undirected graphical Gaussian models (Dempster, 1972; Whittaker, 1990)

- Starting point:
 - correlation structure, neither direction nor causality
 - multivariate Normal distribution with parameters μ and Σ assumed
- Based on the following:
 - Conditional distribution of genes i and j , given all the rest of the genes, is bivariate normal
 - Partial correlations as opposed to simple correlations

Graphical Gaussian models: Technical Details

- Partial correlations $\Pi = (\pi_{ij})$ are computed from the inverse of the $(G \times G)$ correlation matrix $(\omega_{ij}) = \Omega = P^{-1}$, with $P = (\rho_{ij})$
- the following are equivalent
 1. $\omega_{ij} = 0$
 2. genes i and j **conditionally independent** given the remainder of the genes
 3. **partial correlation coefficient** $\pi_{ij} = \rho_{ij|\text{rest}} = \frac{-\omega_{ij}}{\sqrt{\omega_{ii}\omega_{jj}}} = 0$
- Significance tests based on deviance difference between successive models (i. e. large sample tests based on limiting χ^2 distribution)

Problems arising in application to microarray data

- **unstable** partial correlation estimators for $G > N$
- **multicollinearity**: (nearly) linear dependencies in the data
- **model selection**: N is small, hence needs to be based on exact tests

↪ Application of GGMs so far restricted to assess relationships between small number of genes (Waddell & Kishino, 2000) or clusters of genes (Toh & Horimoto, 2002)

↪ Problem of interpretability

Small sample GGM framework needed!

Trick 1: Use pseudoinverse to invert correlation matrix

- failure of standard definition for inverse of a matrix for singular matrices
- generalization using singular value decomposition: $A = U \Sigma V^T$
- Pseudoinverse (Moore Penrose inverse): $A^+ = V (\Sigma^T \Sigma)^{-1} \Sigma U^T$
- $\sum (A^+ A - I)^2$ minimized

This allows for computing partial correlations for $N < G$.

Trick 2: Use Bagging (Bootstrap aggregation)

General algorithm to improve estimates (Breiman 1996):

Step 1 Generate bootstrap sample y^{*b} with replacement from original data. Repeat process $b = 1, \dots, B$ times independently (e. g. $B = 1000$).

Step 2 Calculate for each bootstrap sample y^{*b} estimate $\hat{\theta}^{*b}$.

Step 3 Compute bootstrap mean

$$\frac{1}{B} \sum_{b=1}^B \hat{\theta}^{*b}$$

Small Sample Estimates of Partial Correlation

1. $\hat{\Pi}^1$: use pseudoinverse for inverting \hat{P} but do not perform bagging (= observed partial correlation).
2. $\hat{\Pi}^2$: use bagging to estimate correlation matrix P , then invert with pseudoinverse (= partial bagged correlation).
3. $\hat{\Pi}^3$: use bagging on estimate $\hat{\Pi}^1$, i. e. use pseudoinverse for inverting each bootstrap replicate estimate \hat{P}^{*b} (= bagged partial correlation).

Simulation study

To assess the statistical properties of the proposed procedures we need to perform a simulation study:

1. Generate random artificial network, i. e. true matrix of partial correlations Π
2. Compute corresponding matrix of correlations P
3. Simulate data from respective multivariate Normal distribution (with zero mean and variance one)
4. Estimate partial correlations $\hat{\Pi}^i$ from simulated data

Trick 3: Generating GGMs

Problem: true P must be **positive definite**, thus completely randomly chosen partial correlations do not necessarily correspond to valid graphical Gaussian model.

Solution:

1. generate random **diagonally dominant matrix**
2. standardize to obtain partial correlation matrix Π

—→ resulting model is guaranteed to be valid

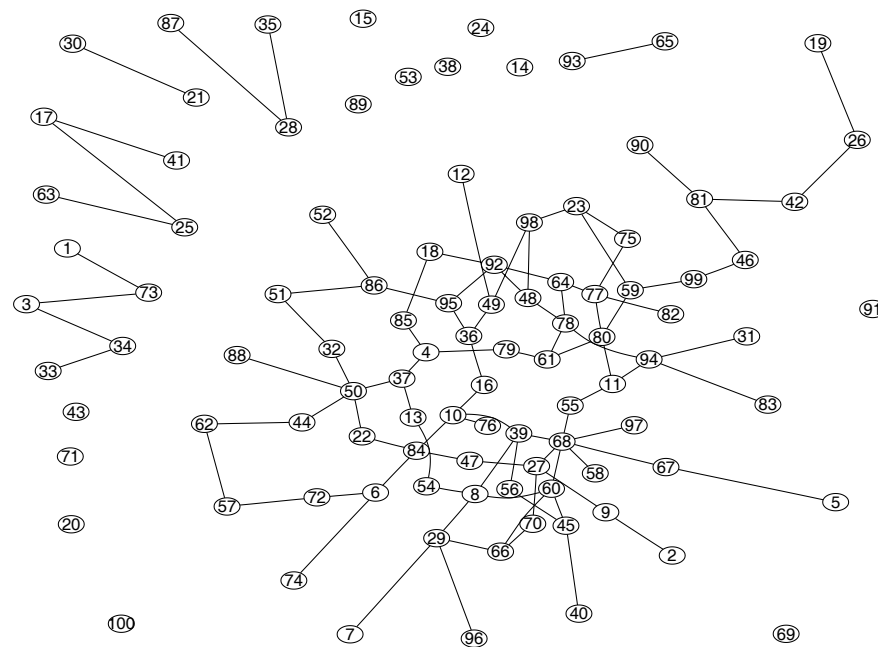
Evaluation of empirical mean squared error

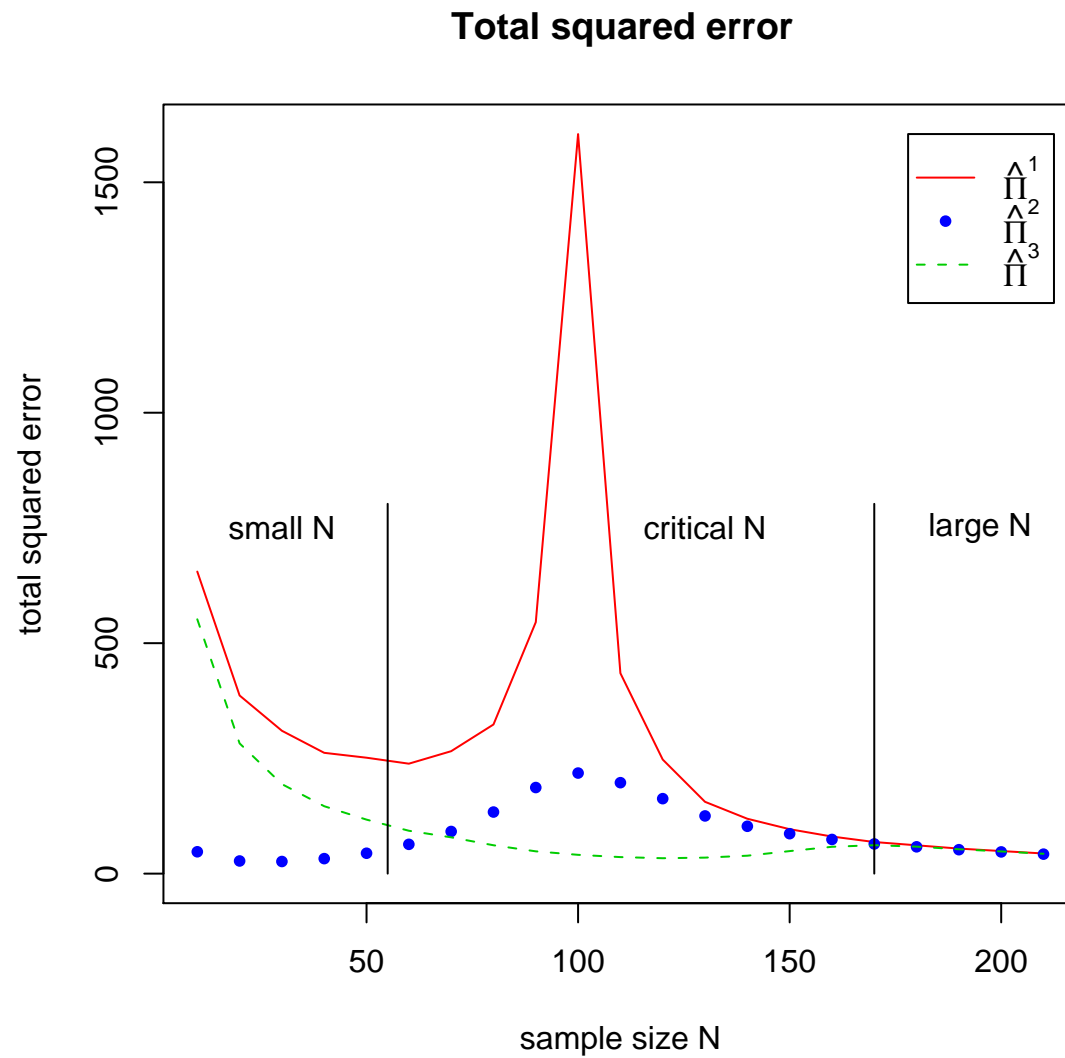
$$\sum_{1 \leq i < j \leq G} (\hat{\pi}_{ij}^k - \pi_{ij})^2 \quad (k = 1, 2, 3)$$

Example simulation setup:

- 100 nodes
- 2% non-zero partial correlations (biological networks are known to be sparse)
 - ↪ 99 true edges out of 4950 potential edges
- 1000 bootstrap replicates
- 50 simulation runs/sample size

Random network with 100 nodes and edge fraction 0.02





Peaking phenomenon

- From a statistical point of view: VERY surprising!
- estimates expected to improve with increasing sample size

But:

- well known in small-sample regression and classification problems (Raudys & Duin, 1998; Skurichina & Duin, 2002)

Comparison of Point Estimates

- extremely bad performance of observed partial correlation $\hat{\Pi}^1$ in critical region (sample size $N \approx$ feature size G)
- Partial bagged correlation $\hat{\Pi}^2$ performs well for very small sample sizes (reason: bagged *sample* correlation matrix positive definite)
- Bagged partial correlation estimate $\hat{\Pi}^3$ best in critical region $N \approx G$
- the three methods coincide for $N \gg G$ (note that this is where classical GGM theory applies)

Model selection

Determination of network topology

- try all potentially adequate graphical models and evaluate their goodness of fit
 \hookrightarrow impossible in realistic applications due to enormous effort
- textbook methods (e. g. stepwise selection based on significance tests that are asymptotic χ^2 -tests based on the deviance difference between successive models) are unreliable for small sample sizes

Alternative strategy used here:

multiple testing of all possible edges using exact correlation test

Null Distribution

Density under null hypothesis, i. e. $\rho = 0$, of Normal (partial) correlation coefficient (Hotelling 1953):

$$f_0(r) = (1 - r^2)^{(\kappa-r)/2} \frac{\Gamma(\frac{\kappa}{2})}{\pi^{\frac{1}{2}} \Gamma(\frac{\kappa-1}{2})} \quad (1)$$

where κ is the degree of freedom.

For $\rho = 0$ the degree of freedom is equal to the inverse of the variance, i.e. $\text{Var}(r) = \frac{1}{\kappa}$, and to sample size minus one ($\kappa = N - 1$).

For partial correlations: $\kappa = N - 1 - (G - 2) = N - G + 1$.

Negative for $N < G!!!$

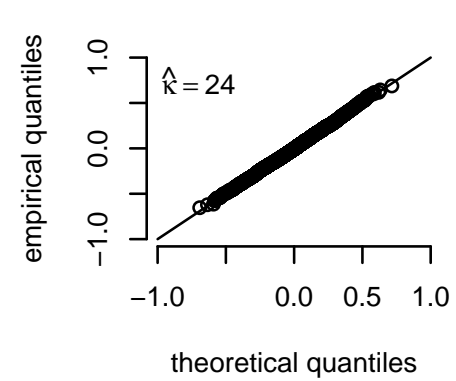
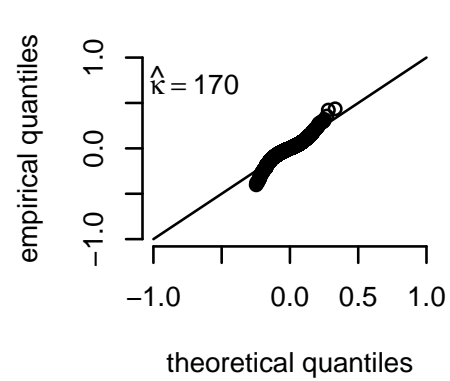
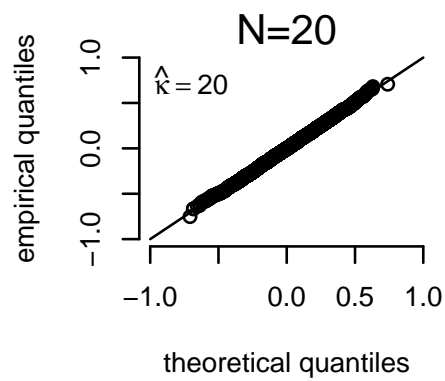
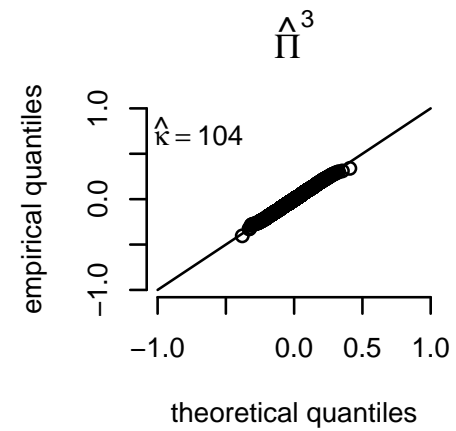
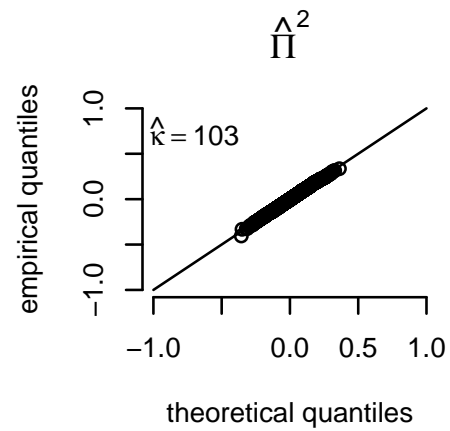
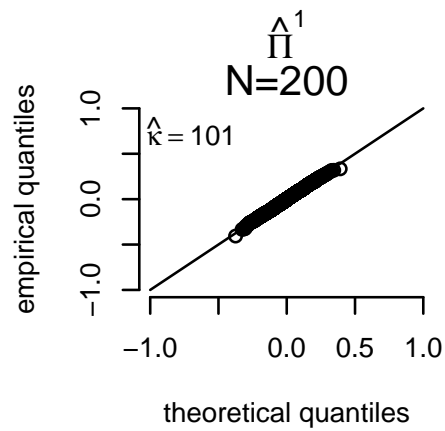
Model Validation

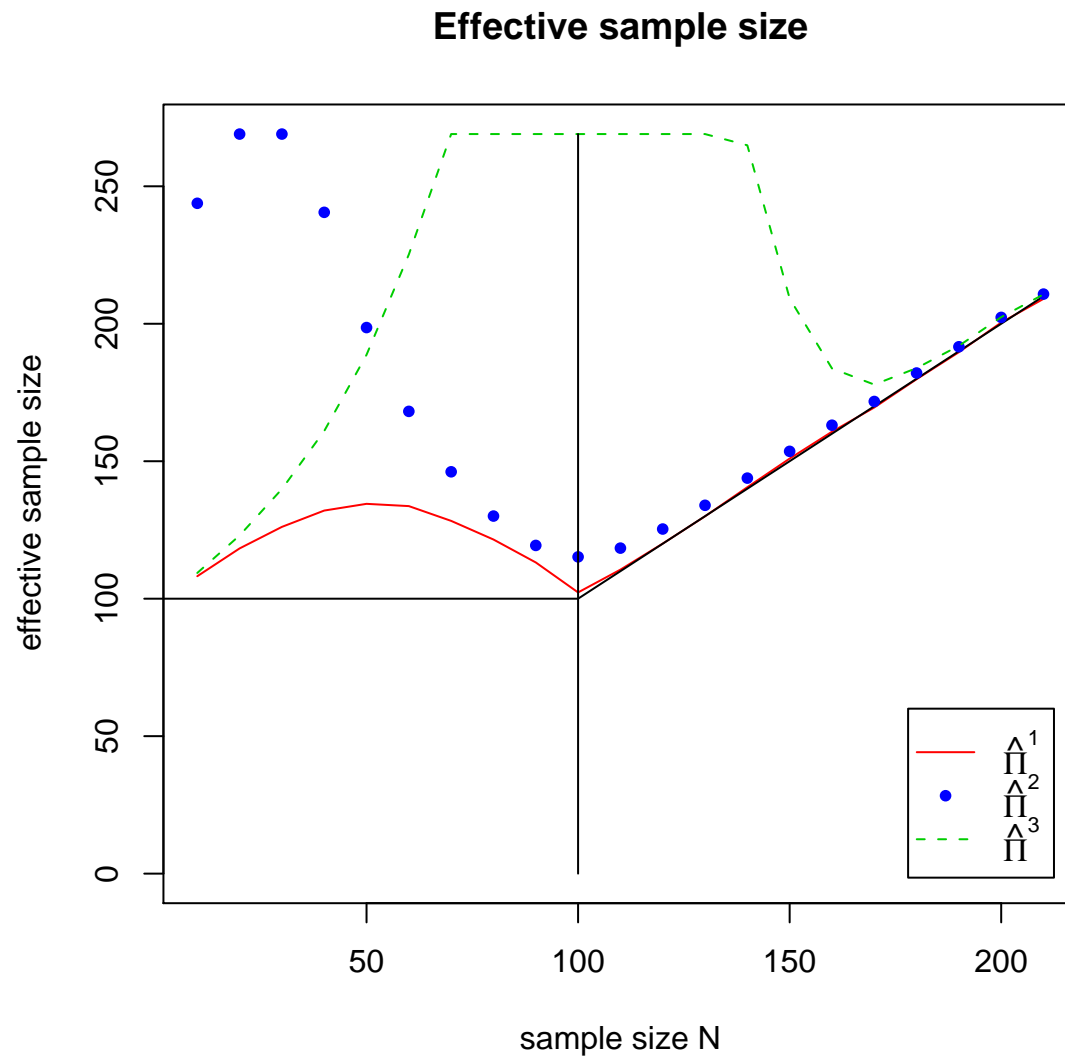
Do small sample estimates $\hat{\pi}_{ij}^1$, $\hat{\pi}_{ij}^2$, and $\hat{\pi}_{ij}^3$ of partial correlations under H_0 indeed follow this distribution?

Trick 4: Estimate degree of freedom κ adaptively (details later).

Next two slides:

- QQ plots of all three point estimates for large ($N=200$, top row) and small ($N=20$, bottom row) sample size. Data simulated assuming $G = 100$ and no edges at all in underlying graph.
- plot of effective sample size $N_{\text{eff}} = \hat{\kappa} + G - 1$





Results: Fit of Null-Model

- Empirical null distributions of estimates $\hat{\Pi}^i$ agree to a high degree with the theoretical distribution for the normal sample correlation.
- Estimated variance, degree of freedom and effective sample size differ among estimators and investigated region ($N \ll G, N \approx G, N \gg G$).
- Small total mean squared error and large effective sample size coincide

Inference of Edges

Trick 5: Exploit highly parallel structure of the problem and sparsity of biomolecular networks.

- Assume most edges to be zero.
- more specifically: observed partial correlations p across all edges follow mixture distribution:

$$f(p) = \eta_0 f_0(p; \kappa) + \eta_A f_A(p) \quad (2)$$

with $\eta_0 + \eta_A = 1$ and $\eta_0 \gg \eta_A$.

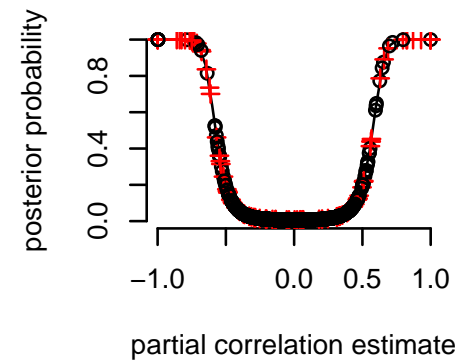
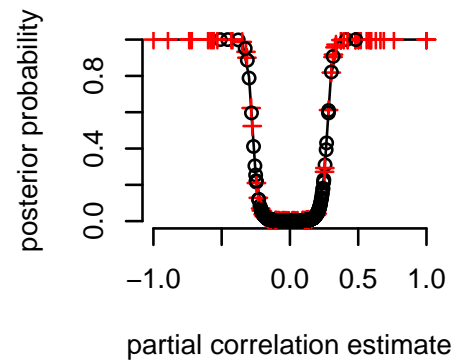
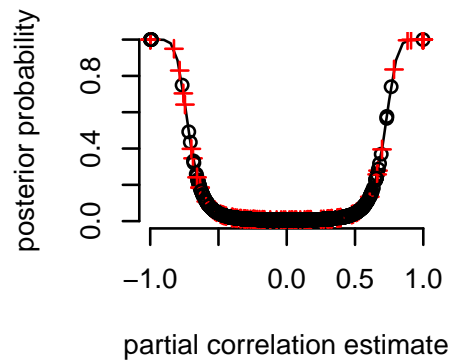
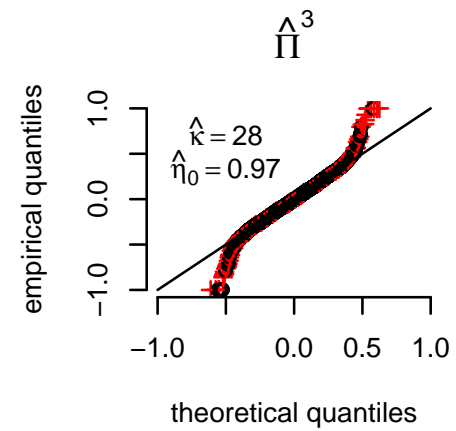
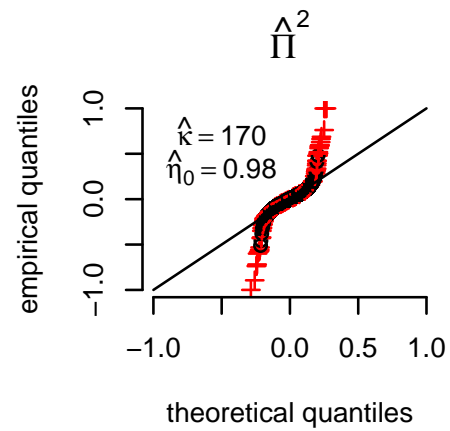
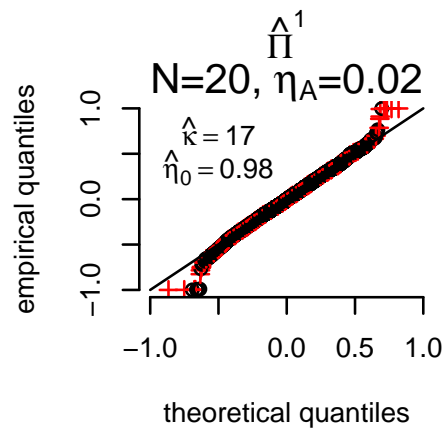
- alternative distribution f_A : uniform distribution from -1 to 1

Trick 5 in style of empirical Bayes methods for problems of differential expression (Sapir & Churchill, 2000; Efron *et al.*, 2001; Efron, 2003)

Fit of Mixture Distribution (next slide):

- QQ plots for all three estimates in small-sample example with $N = 20$, $G = 100$, and $\eta_A = 0.02$ (top row)
- supplementary: empirical posterior probability plots of an edge being present (bottom row)

$$\text{pr}(\text{non-zero edge}|\hat{p}) = \frac{\hat{\eta}_A f_A(\hat{p})}{f(\hat{p}; \hat{\kappa})} \quad (3)$$



Model Selection Using FDR Multiple Testing

False discovery rate criterion (Benjamini & Hochberg, 1995): control expected proportion of false positives

1. Set of ordered p -values $p_{(1)}, p_{(2)}, \dots, p_{(M)}$ corresponding to all potential edges $e_{(1)}, e_{(2)}, \dots, e_{(M)}$
2. Let i_Q be largest i with $p_{(i)} < \frac{i}{M} \frac{Q}{\eta_0}$
3. Reject null hypothesis of zero partial correlation for edges $e_{(1)}, e_{(2)}, \dots, e_{(i_Q)}$

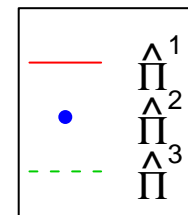
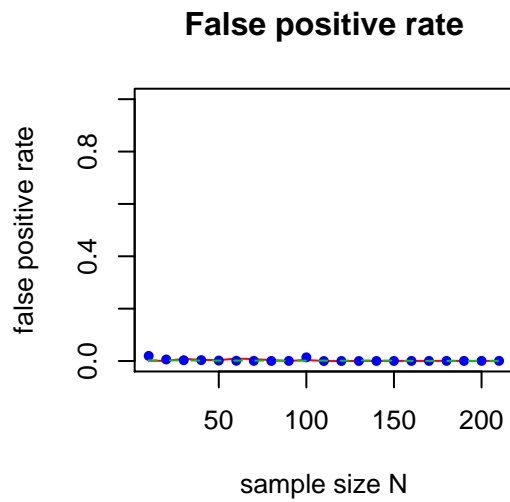
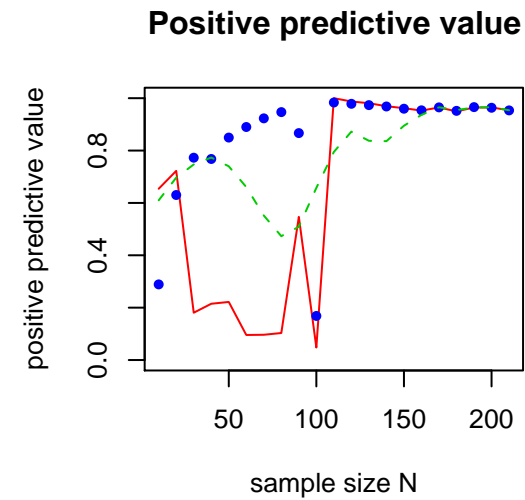
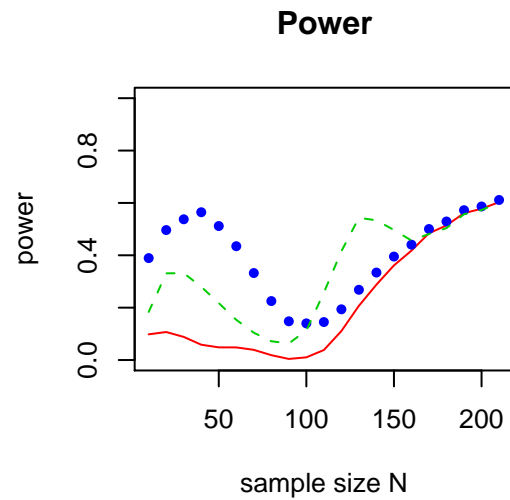
Approximation to proper model search!

Power analysis

Investigation of statistical properties of proposed model selection procedure for $\hat{\Pi}^1$, $\hat{\Pi}^2$, and $\hat{\Pi}^3$:

- FDR level $Q = 0.05$
- empirical power (sensitivity, true positive rate)
- empirical false positive rate (1-specificity)
- positive predictive value

Simulation setup: $G = 100$ and $\eta_A = 0.02$ with $N = 10, 20, \dots, 210$

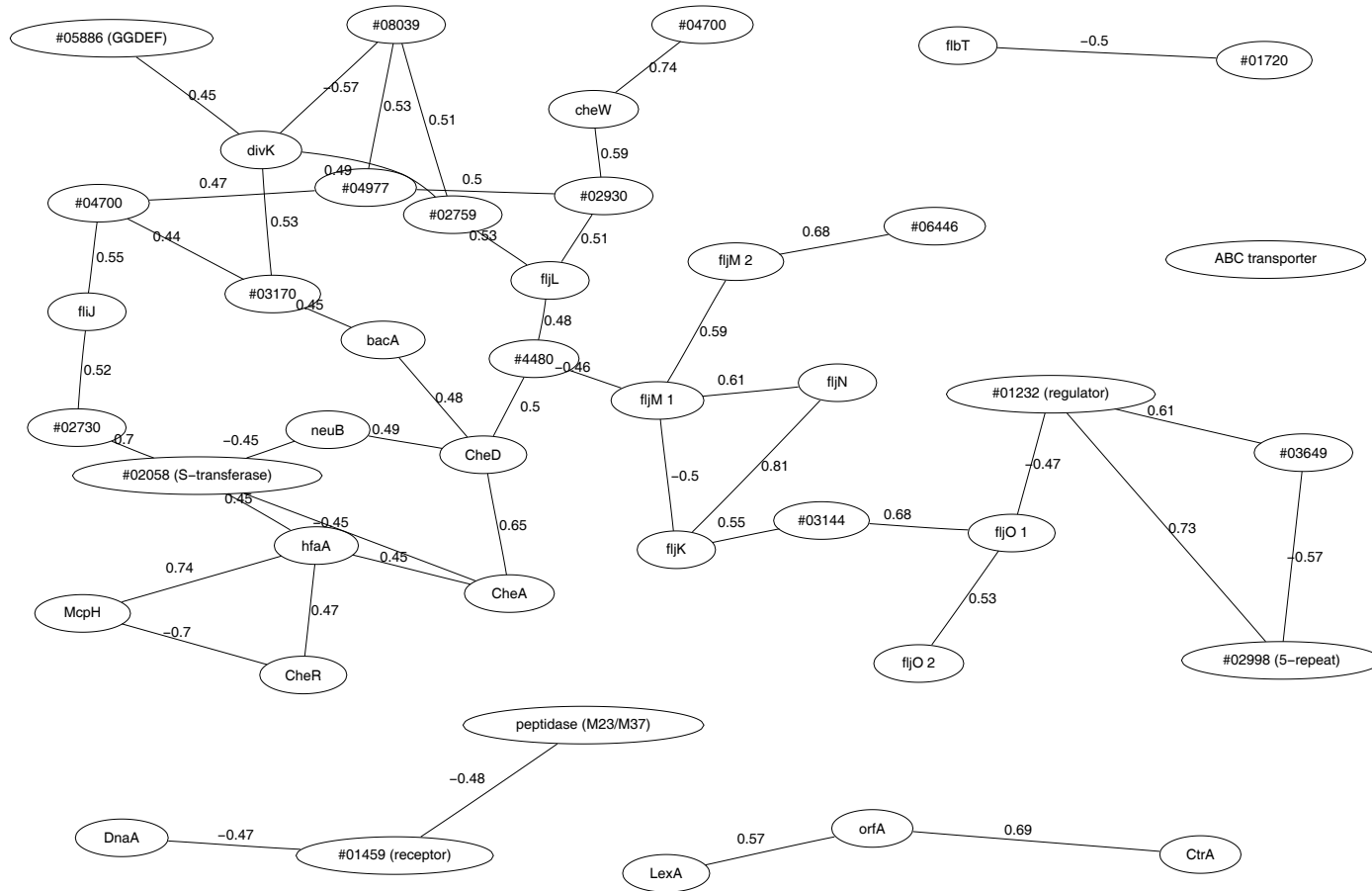


Summary: Recipe of Analysis

1. choose suitable point estimate of partial correlation
2. estimate degree of freedom κ of underlying null distribution
3. compute two-sided p -values and posterior probabilities, respectively, for all possible edges
4. apply multiple testing procedure using FDR criterion to determine graph topology (exploratory tool!)
5. visualize resulting network structure

Molecular Data

- cell cycle in *Caulobacter crescentus* (Laub *et al.*, 2000)
- 3062 genes and ORFs at 11 sampled time points
- reduced to 1444 (due to missing values) and further to 42 potentially interesting genes and ORFs (Wichert *et al.*, 2004)
- 47 significantly non-zero partial correlations



Discussion

We have presented a novel framework for inferring large GGMs from small-sample data sets such as microarray (time series) data sets.

Key Insights:

- we may employ bagging to obtain improved point estimates of partial correlation
- we can exploit the sparsity of the network to estimate the null distribution from the point estimate of the correlation matrix
- heuristic (but fast) model selection can be done via multiple testing (using frequentist FDR method or empirical Bayes)

Discussion ctd.

Advantages:

- in contrast to other applications of GGMs to micorarray data the analysis can take place on the gene level (interpretability)
- our simulation results suggest that sensible estimation of sparse graphical models is possible in the proposed graphical Gaussian modeling framework, even for small samples.
- the inference procedure is computationally efficient
- software will soon be made available in R (GeneTS version 2.0)

Discussion ctd.

Further points to consider:

- critical review of model assumptions (i.i.d., normality)
- though estimation of κ somehow accounts for longitudinal autocorrelation in the data, data should be treated as proper time series
- heuristic network search may be improved
- imperfect null distribution of $\hat{\Pi}^2$ may be modified to improve statistical testing for very small samples
- GGMs may serve as a starting point to build more sophisticated graphical models (Bayesian nets, dynamics etc).
- graphical model framework is suitable statistical approach to modeling, but inference and model selection remain challenging