

Pathway & Network Analysis of Omics Data: Introduction

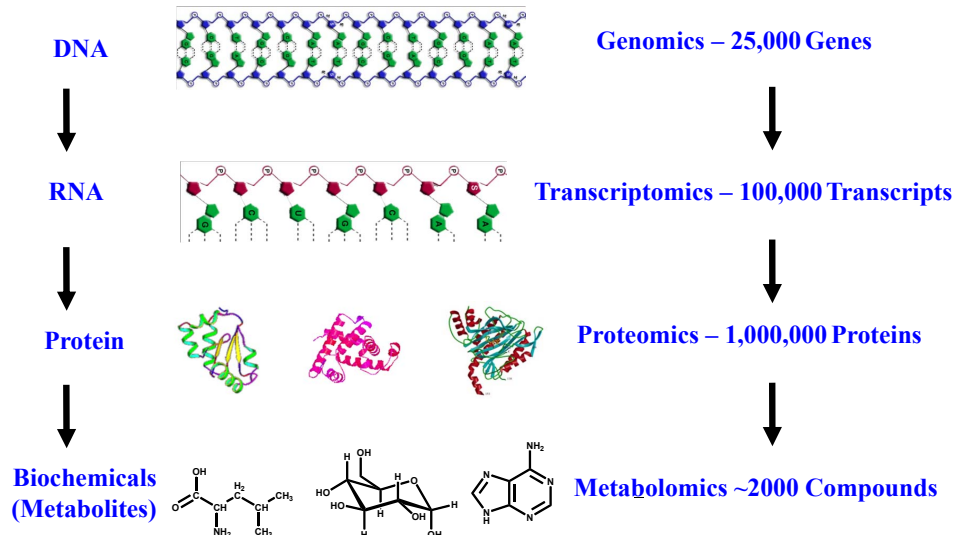
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Summer Institute for Statistical Genetics – 2020

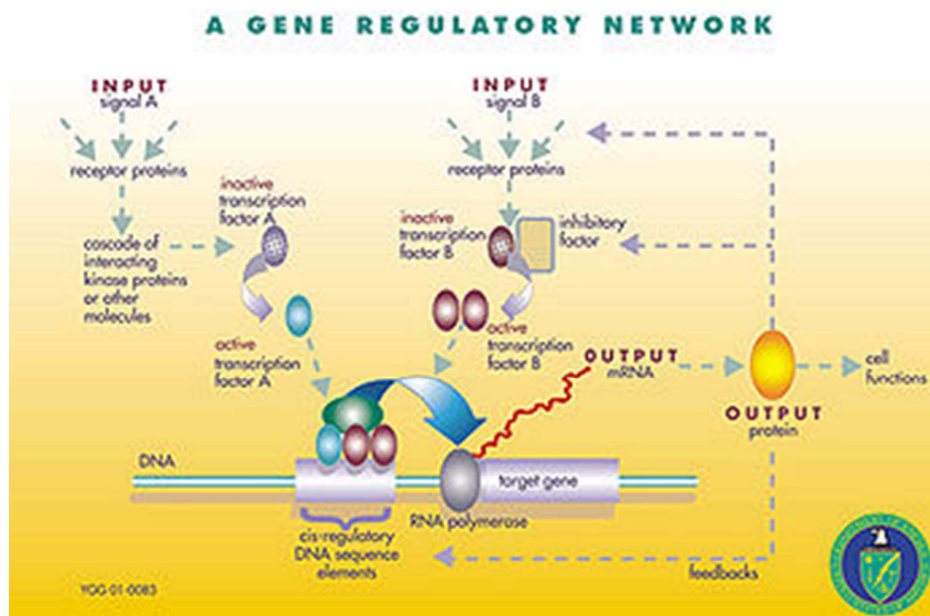
Why Study Networks?

- ▶ Components of biological systems (genes, proteins etc) interact with each other to carry out cell functions.
- ▶ Examples of such interactions include signaling, regulation and interactions between proteins.
- ▶ We cannot understand the function and behavior of biological systems by studying individual components ($2 + 2 \neq 4!$).
- ▶ Networks provide an efficient representation of complex interactions in cells, and a basis for mathematical/statistical models to study these systems.

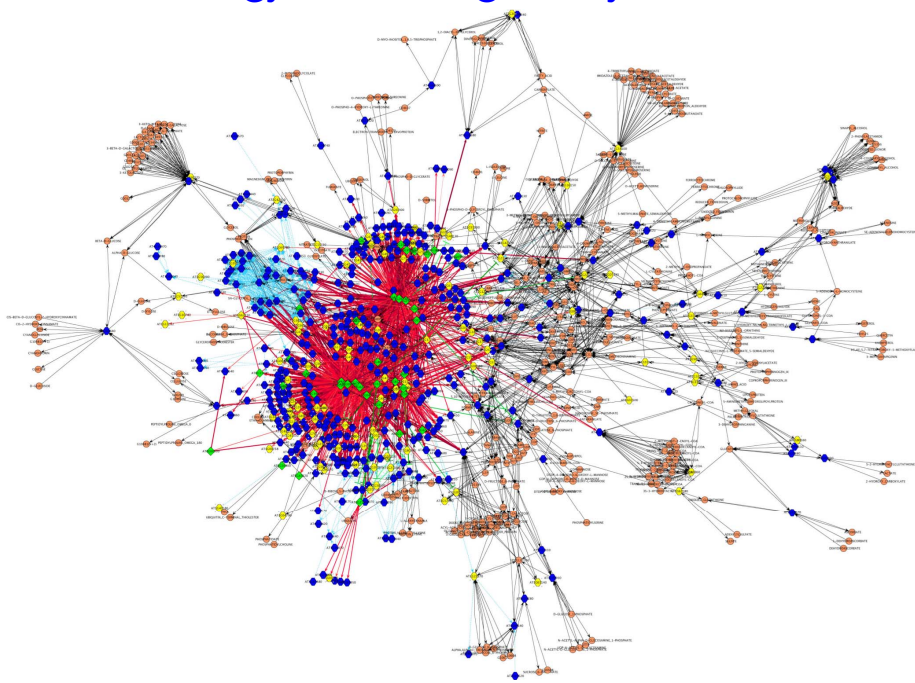
Central Dogma of Molecular Biology (Extended)



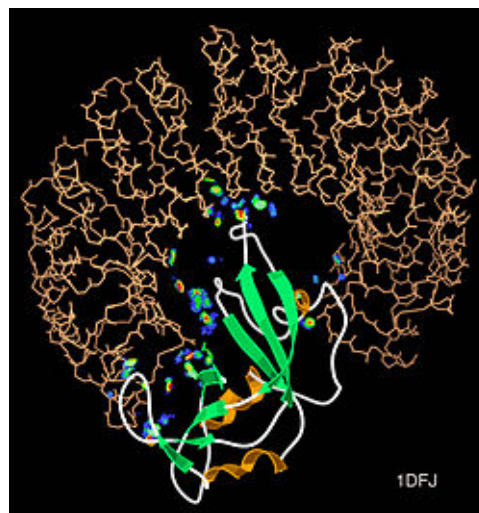
Networks in Biology: Gene Regulatory Interactions



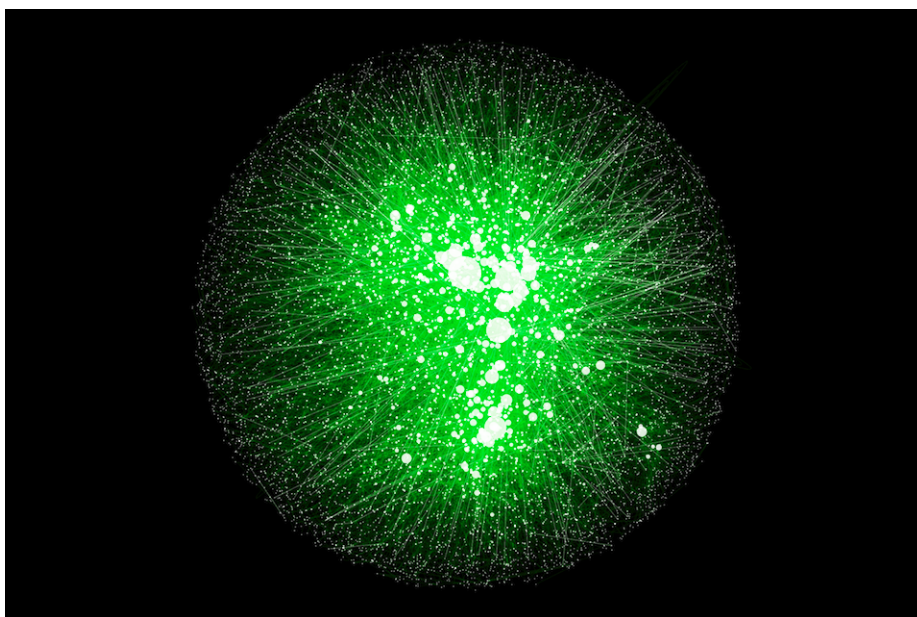
Networks in Biology: Gene Regulatory Networks



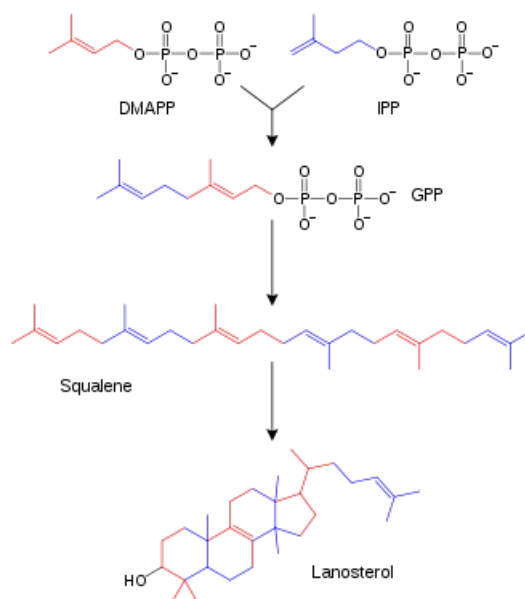
Networks in Biology: Protein-Protein Interaction



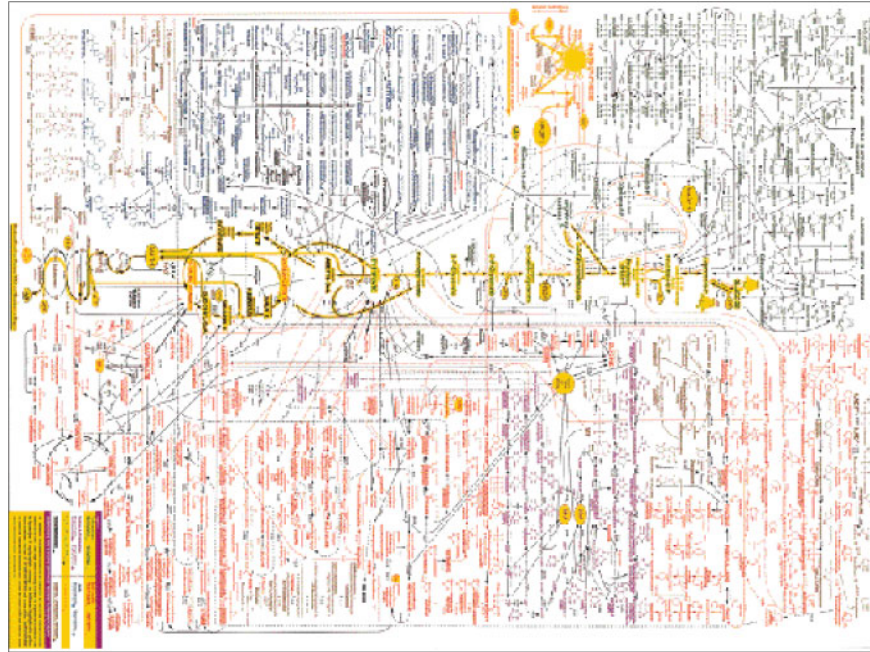
Networks in Biology: Protein-Protein Interactions (PPI)



Networks in Biology: Metabolic Reactions



Networks in Biology: Metabolic Pathways



But Do Networks Matter?

- ▶ They Do!
- ▶ Recent studies have linked changes in gene/protein networks with many human diseases.

Systems Biology and Emerging Technologies

Gene Networks and microRNAs Implicated in Aggressive Prostate Cancer

Liang Wang,¹ Hui Tang,² Venugopal Thayanithy,³ Subbaya Subramanian,³ Ann L. Oberg,² Julie M. Cunningham,¹ James R. Cerhan,² Clifford J. Steer,⁴ and Stephen N. Thibodeau¹

¹Departments of Laboratory Medicine and Pathology and ²Health Sciences Research, Mayo Clinic, Rochester, Minnesota; and Departments of ³Laboratory Medicine and Pathology, ⁴Medicine, and Genetics, Cell Biology, and Development, University of Minnesota, Minneapolis, Minnesota

But Do Networks Matter?

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Estrogen-Regulated Gene Networks in Human Breast Cancer Cells: Involvement of E2F1 in the Regulation of Cell Proliferation

Joshua D. Stender, Jonna Frasor, Barry Komm, Ken C. N. Chang, W. Lee Kraus, and Benita S. Katzenellenbogen

Departments of Biochemistry (J.D.S.) and Molecular and Integrative Physiology (J.F., B.S.K.), University of Illinois at Urbana-Champaign, Urbana, Illinois 61801-3704; Women's Health and Musculoskeletal Biology (B.K., K.C.N.C.), Wyeth Research, Collegeville, Pennsylvania 19426; and Department of Molecular Biology and Genetics (W.L.K.), Cornell University, Ithaca, New York 14853-4203

But Do Networks Matter?



Cancer Cell
Article

A Transcriptional Signature and Common Gene Networks Link Cancer with Lipid Metabolism and Diverse Human Diseases

Heather A. Hirsch,^{1,7} Dimitrios Iliopoulos,^{1,7} Amita Joshi,^{1,7} Yong Zhang,² Savina A. Jaeger,³ Martha Bulyk,^{3,4,5} Philip N. Tsichlis,⁶ X. Shirley Liu,² and Kevin Struhl^{1,*}

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But Do Networks Matter?

And, incorporating the knowledge of networks **improves our ability to find causes of complex diseases.**

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biology

REPORT

Network-based classification of breast cancer metastasis

Han-Yu Chuang^{1,5}, Eunjung Lee^{2,3,5}, Yu-Tsueng Liu⁴, Doheon Lee³ and Trey Ideker^{1,2,4,*}

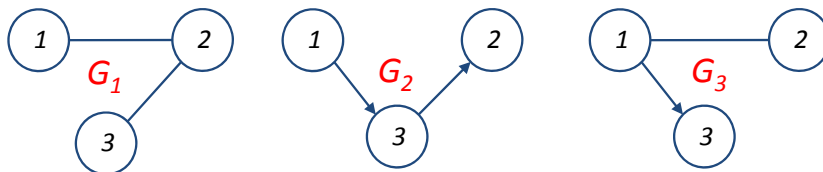
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Networks: A Short Primer

- ▶ A network is a collection of **nodes** V and **edges** E .
- ▶ We assume the network has p nodes, **corresponding to random variables** $X_1, \dots, X_p \equiv$ **biological measurements**.
- ▶ Edges can be **directed** $X \rightarrow Y$ or **undirected** $X - Y$.



- ▶ In all these example, the **node set** is $V = \{1, 2, 3\}$.
- ▶ The **edges** are:

$$E_1 = \{1 - 2, 2 - 3\}$$

$$E_2 = \{1 \rightarrow 3, 3 \rightarrow 2\}$$

$$E_3 = \{1 - 2, 1 \rightarrow 3\}$$

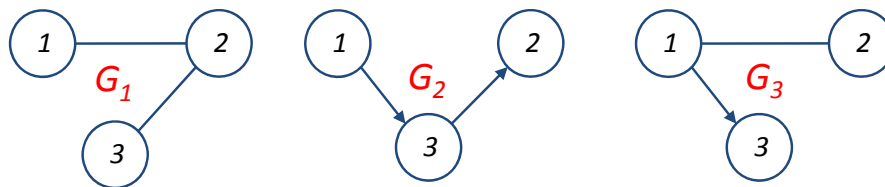
Networks: A Short Primer

- A convenient way to represent the **edges** of the network is to use an **adjacency matrix A**
- A **matrix** is a rectangular array of data (similar to a table)
- Values in each **entry** are shown by **indices of row and column**

$$A = \begin{bmatrix} . & x & . \\ . & . & . \\ . & . & . \end{bmatrix} \text{ Here, } x \text{ is in row 1 and column 2}$$

- Adjacency matrix is a **square** matrix, which has a **1** if there is **an edge** from a **node in one row** to a **node in another column**, and **0** otherwise
- For **undirected edges**, we add a **1** in both directions

Networks: A Short Primer



$$A = \begin{bmatrix} 0 & 1 & 0 \\ 1 & 0 & 1 \\ 0 & 1 & 0 \end{bmatrix} \quad A = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 0 & 1 & 0 \end{bmatrix} \quad A = \begin{bmatrix} 0 & 1 & 1 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}$$

What Do Edges in Biological Networks Mean?

- ▶ In **gene regulatory networks**, an edge from gene i to gene j often means that i **affects the expression of j** ; i.e. as i 's expression changes, we expect that expression of j to increase/decrease.
- ▶ In **protein-protein interaction networks**, an edge between proteins i and j often means that *the two proteins bind together and form a protein complex*. Therefore, we expect that these proteins are generated at similar rates.
- ▶ In **metabolic networks**, an edge between compound i and j often means that *the two compounds are involved in the same reaction*, meaning that they are generated at relative rates.
- ▶ Thus, edges represent some type of **association among genes, proteins or metabolites**, defined generally to include *linear or nonlinear* associations; more later....

Statistical Models for Biological Networks

- ▶ We use the framework of **graphical models**
- ▶ In this setting, **nodes correspond to “random variables”**
- ▶ In other words, each node of the network represents one of the variables in the study
 - ▶ In gene regulatory networks, **nodes \equiv genes**
 - ▶ In PPI networks, **nodes \equiv proteins**
 - ▶ In metabolic networks, **nodes \equiv metabolites**
- ▶ In practice, we observe n measurements of each of the variables (genes/proteins/ metabolites) for say different individuals, and want to determine which variables are connected, or use their connection for statistical analysis

Our Plan

We will cover the following topics

- ▶ Methods for **detecting signal on known networks**
 - ▶ Network analysis based on **centrality and clustering**
 - ▶ **Topology-based pathway enrichment analysis**
- ▶ Methods for **learning undirected networks**
 - ▶ Co-expression networks
 - ▶ ARACNE
 - ▶ Conditional independence graphs
 - ▶ Gaussian observations (glasso, etc)
 - ▶ Non-Gaussian and non-linear data (nonparanormal, etc)
- ▶ Methods for **learning directed networks**
 - ▶ Bayesian Networks (basic concepts, reconstruction algorithm)
 - ▶ Learning directed networks from time-course data (dynamic Bayesian networks)

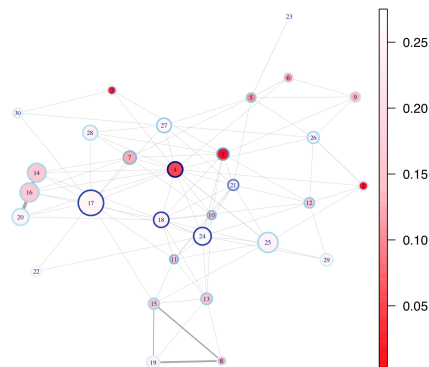
Pathway & Network Analysis of Omics Data: Analysis of Network-Structured Data

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Summer Institute for Statistical Genetics – 2020

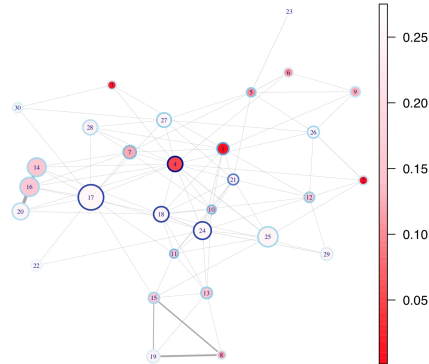
Introduction

Suppose we observe **activities of individual nodes** (genes, proteins, brain regions, etc) on a **network** (gene regulatory network, structural connectivity network, etc)



How can we identify the **important nodes**?
and what does this even mean?

Identifying Important Nodes



How can we identify the **important nodes**?

- ▶ We can select the **significant nodes** based on p-values, after adjusting for multiple comparisons (FDR, etc)
- ▶ But the signal is often weak for lots of tests
- ▶ If we believe the network is informative, it may make sense to **use the network to guide our selection**

Identifying Important Nodes

Possible strategies:

- ▶ Identify **individual nodes** associated with the outcome by incorporating the network (signal detection on network)
- ▶ Test if (pre-specified) **subnetworks** are associated with the outcome (**topology-based pathway enrichment analysis**)
- ▶ Identify **collections of (connected) nodes** that are associated with the outcome (*de-novo identification of enriched modules*)

Signal Detection on Networks

Signal Detection on Networks

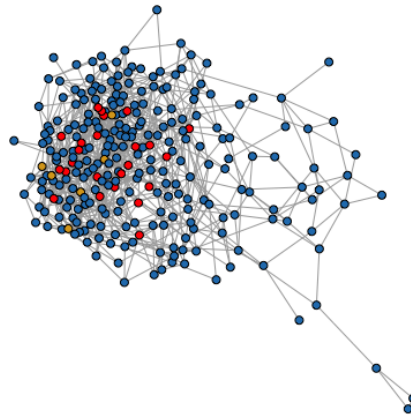
How can we identify the **important nodes in a network**?

The simplest option is to limit our search/testing to the **central nodes** in the network:

- ▶ Nodes connected to many other nodes, aka **hub nodes**
- ▶ Nodes that are **close to many other nodes** (**closeness**)
- ▶ Nodes that are **on many network paths** (**betweenness**)

Example: Functional Relevance of Hub Nodes

- Inferred genetic interaction network of cancer-related pathway in prostate cancer (data from TCGA)
- Hubs defined as nodes whose degrees are at the 75th percentile of the degree distribution



Other Measures of Centrality

- **Closeness**: Total distance of each node to other nodes:

$$cl_j = \left(\sum_{k \in V} d(j, k) \right)^{-1}$$

where $d(j, k)$ is the (shortest path) distance between j and k .

- **Betweenness**: The number of *paths* that go through a node:

$$bw_j = \sum_{i \neq j \neq k} \frac{\pi_{ik}(j)}{\pi_{ik}}$$

where $\pi_{ik}(j)$ is the number of paths between i and k that go through j , and π_{ik} is the total number of paths between them.

Identifying “Central” Nodes

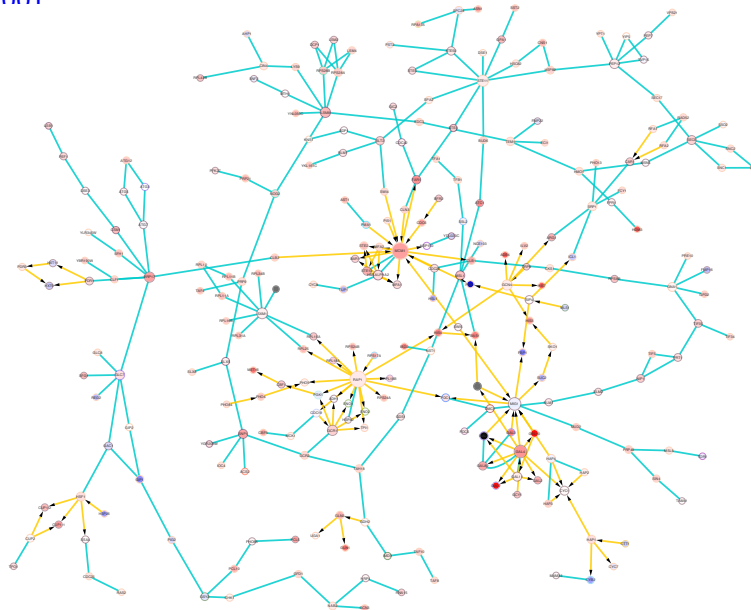
Calculating centrality measures using igraph:

- ▶ Hub nodes: `hub_score(graph)`
- ▶ Closeness: `closeness(graph, vids)`
 - ▶ use `estimate_closeness()` for larger networks
- ▶ Betweenness: `betweenness(graph, vids)`
 - ▶ use `estimate_betweenness()` for larger networks

Topology-Based Pathway Enrichment Analysis

Yeast GAL Pathway

Ideker et al, 2001



Topology-Based Pathway Enrichment Analysis

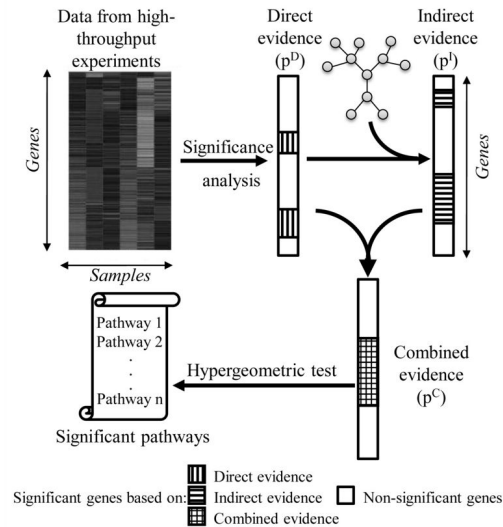
Test for **changes in activities of node** (genes, brain ROIs, etc) in **pre-specified subnetworks**, while **incorporating network information**

Two possible null hypotheses:

- ▶ **Competitive** null hypothesis: activity of each pathway is **compared with other pathways**, often using a **permutation test**
 - ▶ Assume few genes are differentially connected, and may be sensitive to the choice of gene sets
- ▶ **Self-contained** null hypothesis: activity of each pathway is compared **against the null** distribution
 - ▶ More rigorous, but may be sensitive to modeling assumptions (Goemen & Buhlmann (07), Ackermann & Strimmer (09))

PathNet¹

A simple topology-based pathway enrichment method:



PathNet: Details

- Each gene's p -value from differential expression is combined with p -values of its neighbors using **Fisher's methods**

$$SI_j = \sum_{k \in \text{ne}(j)} \left\{ -\log_{10} (p_k^D) \right\}.$$

- The indirect p -value, p^I is calculated from SI_j by permutation
- Direct (p_j^D) and indirect (p_j^I) p -values are then combined (p_j^C)
- The significance of p_j^C for genes in each pathway is assessed using a **hypergeometric test**
- Implemented in Bioconductor package PathNet

topologyGSA²

- ▶ topologyGSA (Gene Set Analysis Exploiting Pathway Topology) assumes that data are normally distributed:

$$X^1 \sim N(\mu^1, \Sigma^1), \quad X^2 \sim N(\mu^2, \Sigma^2)$$

- ▶ It obtains estimates of Σ^1 and Σ^2 based on the networks (think graphical lasso, but with **known nonzero entries**)
- ▶ It then **performs two tests**:
 - ▶ equality of covariance matrices: $H_0^c : \Sigma^1 = \Sigma^2$
 - ▶ equality of means $H_0^m : \mu^1 = \mu^2$ — it uses different methods depending on the result of H_0^c
- ▶ Implemented in R-package topologyGSA (also in graphite)

²Massa et al (2010)

Signaling Pathway Impact Analysis (SPIA)³

- ▶ Combines overrepresentation analysis (ORA) with measure of perturbation of a given pathway under a given condition
- ▶ A bootstrap procedure is used to assess the significance of the observed pathway perturbation (difficult to extend to comparison of > 2 conditions)
- ▶ Currently not applicable to all pathways (more later)
- ▶ Analyzes each pathway separately (ignores connections between pathways)
- ▶ Implemented in the Bioconductor package SPIA

³Tarca et al (2009)

The SPIA Methodology

SPIA combines two types of evidence

- (i) the **overrepresentation** of DE genes in a given pathway
 - measured by the p-value for the given number of DE genes

$$P_{NDE} = P(X \geq N_{DE} \mid H_0)$$

The SPIA Methodology

SPIA combines two types of evidence

- (ii) the **abnormal perturbation of the pathway**
 - the **perturbation for each gene** in the pathway is defined as

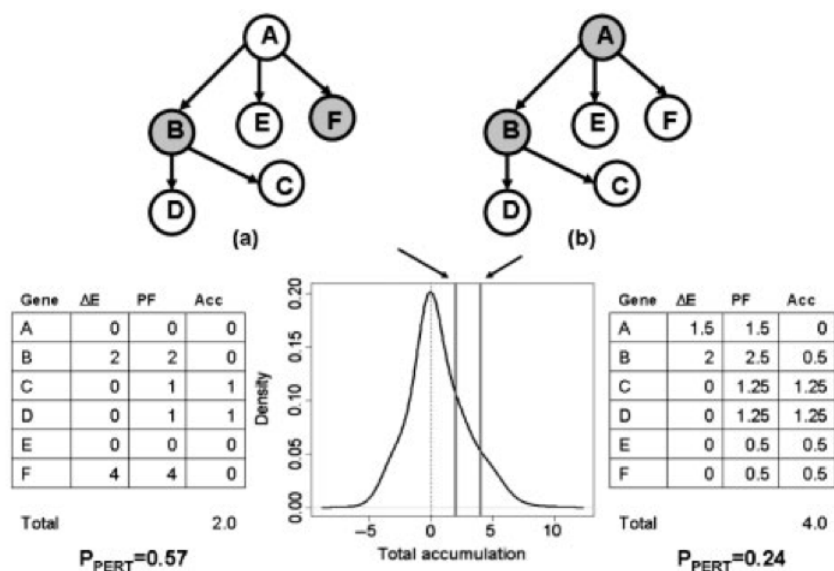
$$PF(g_i) = \Delta E(g_i) + \sum_{j=1}^p \beta_{ij} \frac{PF(g_j)}{N_{DS}(g_j)}$$

- $PF(g_i)$ is the **perturbation factor** of gene i (not known)
- β_{ij} is the **magnitude of effect** of gene j on gene i ; currently,
 $\beta_{ij} = 1$ if $j \rightarrow i$
- $\Delta E(g_i)$ is the **fold change** in expression of gene i
- $N_{DS}(g_j)$ is the **number of downstream** genes from gene j

The SPIA Methodology

- ▶ The **accumulated activity of each gene** can then be calculated as $ACC(g_i) = B \cdot (I - B)^{-1} \Delta E$
 - ▶ B is the **normalized matrix of β 's**: $B_{ij} = \beta_{ij} / N_{DS}(g_j)$
 - ▶ ΔE is the **vector of fold changes**
 - ▶ **Requires B to be invertible**; would not work otherwise
- ▶ The **total accumulated perturbation of the pathway** is then given by $t_A = \sum_i ACC(g_i)$
- ▶ The **p-value** for pathway perturbation is given by $P_{PERT} = P(T_A \geq t_A | H_0)$, which is calculated using a bootstrap approach

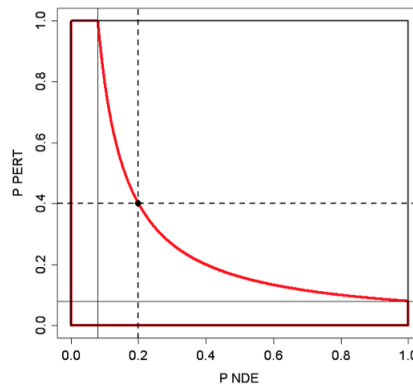
The SPIA Methodology



The SPIA Methodology

SPIA combines two types of evidence

- ▶ The **final p-value for each pathway** is calculated based on the p-values from parts (i) and (ii):
 - ▶ $P_G(i) = c_i - c_i \ln(c_i)$
 - ▶ $c_i = P_{NDE}(i)P_{PERT}(i)$



An Example in R: Data on Colorectal Cancer

```
data(colorectalcancer)

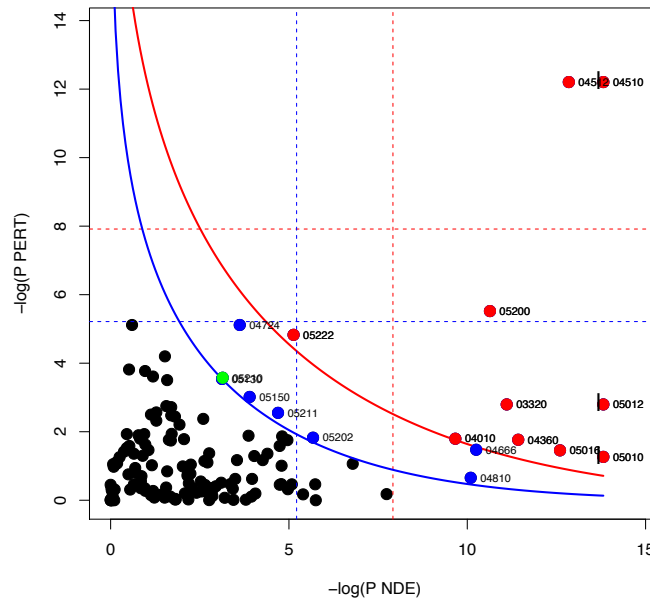
#pathway analysis using SPIA
#use nB=2000 or higher for more accurate results
#uses older version of KEGG signaling pathways graphs
res <- spia(de=DE_Colorectal, all=ALL_Colorectal, organism="hsa", beta=NULL,
            nB=2000, plots=FALSE, verbose=TRUE, combine="fisher")

#now combine pNDE and pPERT using the normal inversion method without
#running spia function again
res$pG=combfunc(res$pNDE,res$pPERT,combine="norminv")
res$pGFdr=p.adjust(res$pG,"fdr")
res$pGFWER=p.adjust(res$pG,"bonferroni")
plotP(res,threshold=0.05)

#highlight the colorectal cancer pathway in green
points(I(-log(pPERT))~I(-log(pNDE)),data=res[res$ID=="05210",],col="green",
      pch=19,cex=1.5)
```

The SPIA Methodology

SPIA two-way evidence plot



Network-Based Gene Set Analysis (NetGSA)⁴

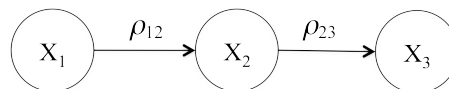
- Generalizes SPIA, to allow for more complex experiments & incorporate interactions among pathways
- Assesses the overall behavior of arbitrary subnetworks (pathways): **changes in gene expression & network structure**
- Uses **latent variables** to model the interaction between genes defined by the network
- Uses **mixed linear models** for inference in complex data
- Computationally challenging for large networks, unless pathways separately analyzed (similar to SPIA)

⁴S & M (2009, 2010); Ma, S & M (2016)

Problem Setup

- ▶ Gene (protein/metabolite) expression data for K experimental conditions and J_k time points
- ▶ Network information (partially) available in the form of a directed weighted graph $G = (V, E)$, with vertex set V corresponding to the genes/proteins/metabolites and edge set E capturing their associations
- ▶ Network edges can be directed $j \rightarrow k$ or undirected $j \leftrightarrow k$
- ▶ Edges defines the effect of nodes on their immediate neighbors; the weight associated with each edge corresponds to the value of partial correlation
- ▶ Represent the network by its adjacency matrix A : $A_{jk} \neq 0$ iff $k \rightarrow j$ & for undirected edges, $A_{jk} = A_{kj}$

The Latent Variable Model: Main Idea



$$X_1 = \gamma_1$$

$$X_2 = \rho_{12}X_1 + \gamma_2 = \rho_{12}\gamma_1 + \gamma_2$$

$$X_3 = \rho_{23}X_2 + \gamma_3 = \rho_{23}\rho_{12}\gamma_1 + \rho_{23}\gamma_2 + \gamma_3$$

Thus $X = \Lambda\gamma$ where

$$\Lambda = \begin{pmatrix} 1 & 0 & 0 \\ \rho_{12} & 1 & 0 \\ \rho_{12}\rho_{23} & \rho_{23} & 1 \end{pmatrix}$$

The Latent Variable Model

- ▶ Let Y be the i th sample in the expression data
- ▶ Let $Y = X + \varepsilon$, with **signal** X and **noise** $\varepsilon \sim N_p(0, \sigma_\varepsilon^2 I_p)$
- ▶ The **influence matrix** Λ measures the **propagated effect of genes on each other** through the network, and can be calculated based on the adjacency matrix A
- ▶ Using $X = \Lambda\gamma$, we get

$$Y = \Lambda\gamma + \varepsilon, \quad \Rightarrow \quad Y \sim N_p(\Lambda\mu, \sigma_\gamma^2 \Lambda\Lambda' + \sigma_\varepsilon^2 I_p)$$

where $\gamma \sim N_p(\mu, \sigma_\gamma^2 I_p)$ are **latent variables**

Mixed Linear Model Representation

Rearranging the expression matrix into np -vector Y , we can write

$$Y = \Psi\beta + \Pi\gamma + \varepsilon$$

where β and γ are fixed and random effect parameters and

$$\varepsilon \sim N_{np}(0, R(\theta_\varepsilon)), \quad \gamma \sim N_{np}(0, \sigma_\gamma^2 I_{np})$$

- **Temporal Correlation** incorporated through R

In general, the **design matrices**, Ψ and Π depend on the experimental settings (similar to ANOVA), and are **functions of Λ**

Estimation of MLM Parameters

MLE for β :

$$\hat{\beta} = (\Psi' \hat{W}^{-1} \Psi)^{-1} \Psi' \hat{W}^{-1} Y$$

where $W = \sigma_{\gamma}^2 \Pi \Pi' + R$.

$\hat{\beta}$ depends on estimates of σ_{γ}^2 and θ_{ε}^2 (estimated using **restricted maximum likelihood** (REML)).

Inference using MLM

- Let ℓ be a **contrast vector** (a linear combination of fixed effects), and consider the test:

$$H_0 : \ell\beta = 0 \quad \text{vs.} \quad H_1 : \ell\beta \neq 0$$

- Use t-test to test the significance of each hypothesis separately

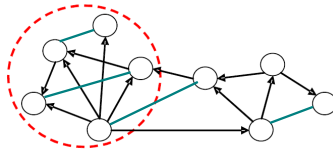
$$T = \frac{\ell\hat{\beta}}{\sqrt{\ell\hat{C}\ell'}}$$

where $C = (\Psi' W^{-1} \Psi)^{-1}$

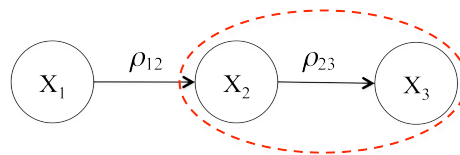
- Under the null hypothesis, T is approximately t -distributed with degrees of freedom that needs to be estimated

“Optimal” Choice of Contrast Vector

- An intuitive choice is the **indicator (membership) vector** for the pathway, **b**, but this only captures changes in mean
- Need to **de-couple the effect of subnetwork** from other nodes



“Optimal” Choice of Contrast Vector



$$\Lambda = \begin{pmatrix} 1 & 0 & 0 \\ \rho_{12} & 1 & 0 \\ \rho_{12}\rho_{23} & \rho_{23} & 1 \end{pmatrix}$$

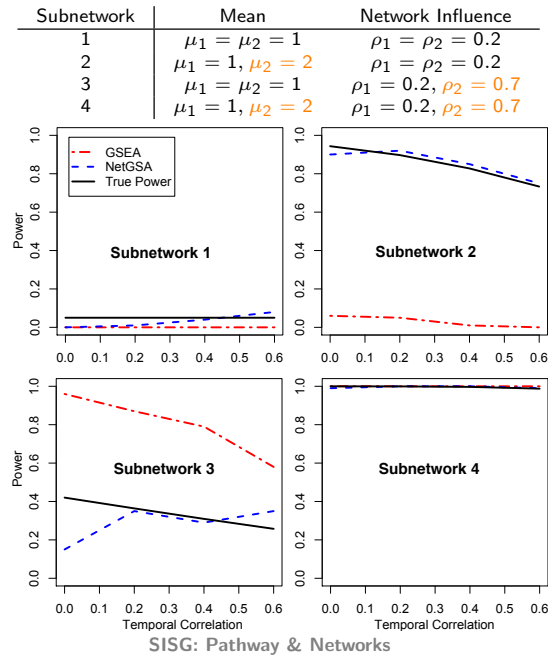
Consider the set, $b = (0, 1, 1)$; then

$$(b\Lambda) = (\rho_{12} + \rho_{12}\rho_{23}, 1 + \rho_{23}, 1)$$

On the other hand,

$$(b\Lambda \cdot b) = (0, 1 + \rho_{23}, 1)$$

Comparison in Simulated Data



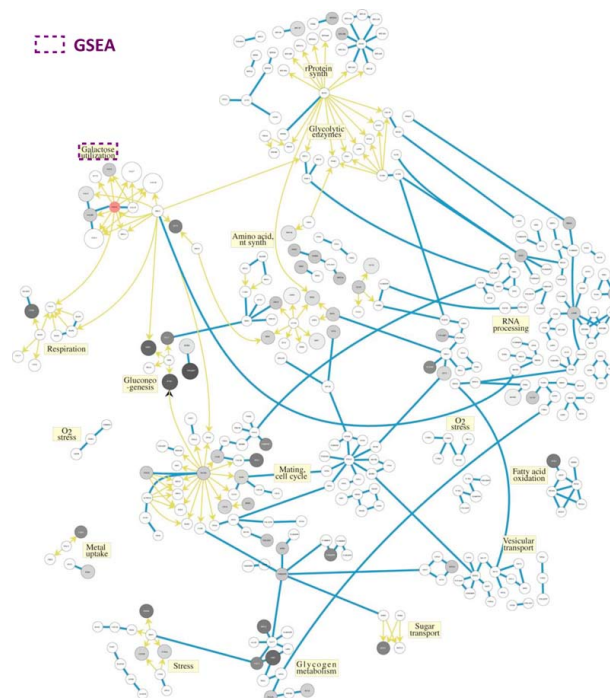
Yeast Galactose Utilization Pathway

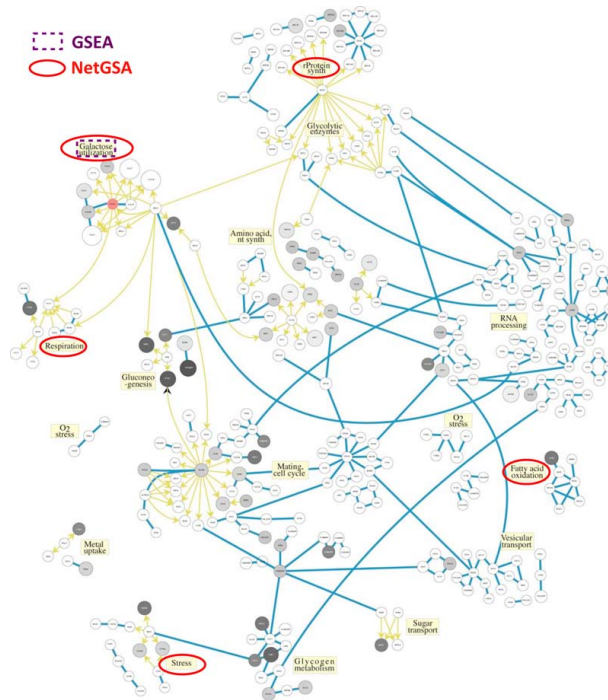
Ideker et al (2001) data on yeast Galactose Utilization Pathway

- Gene expression data for 2 experimental conditions: (gal+) and (gal-)
- Gene-gene and protein-gene interactions as well as association weights found from previous studies
- Q: which pathways respond to the change in growth medium?

Analysis of Yeast GAL Data

- Data:
 - gene expression data for 343 genes
 - 419 interactions found from previous studies and integration with protein expression (**association among genes also available**)
- Results:
 - **GSEA** finds *Galactose Utilization Pathway* significant
 - **NetGSA** finds several other pathways with biologically meaningful functions related to survival of yeast cells in gal–





Environmental Stress Response in Yeast

Gene expression data on Yeast Environmental Stress Response (ESR) (*Gasch et al.*, 2000)

- ▶ 3 combinations of experimental factor, heat shock and osmotic changes (sorbitol), over 3 time points
- ▶ **Temporal correlation**
- ▶ **Network correlation**
- ▶ **Q**: Which **pathways** indicate response to environmental stress
 - ▶ in different **experimental conditions**
 - ▶ over **time**

Yeast ESR Data

Gasch et al (2000)

► Gene Expression Data

Experiment	Obs. Time (after 33C)
Mild heat shock (29C to 33C), no sorbitol	5, 15, 30 min
Mild Heat Shock, 1M sorbitol at 29C & 33C	5, 15, 30 min
Mild Heat Shock, 1M sorbitol at 29C	5, 15, 30 min

► Network Data

- Use **YeastNet** (Lee et al., 2007) for gene-gene interactions (102,000 interactions among 5,900 yeast genes)
- Use independent experiments of Gasch et al. to **estimate weights**
- Pathways are defined using **GO** functions

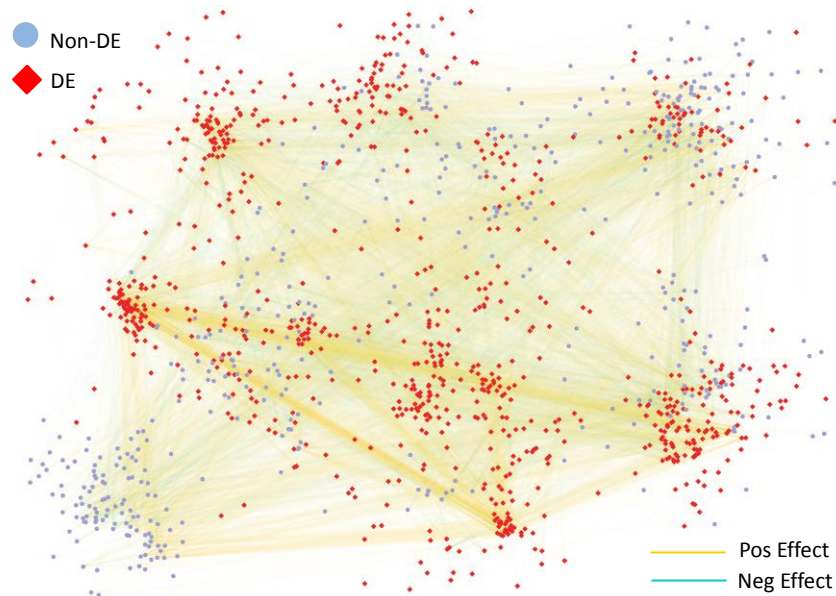
Model and Results

- Model: Let j and k be indices for **time** and levels of **sorbitol**

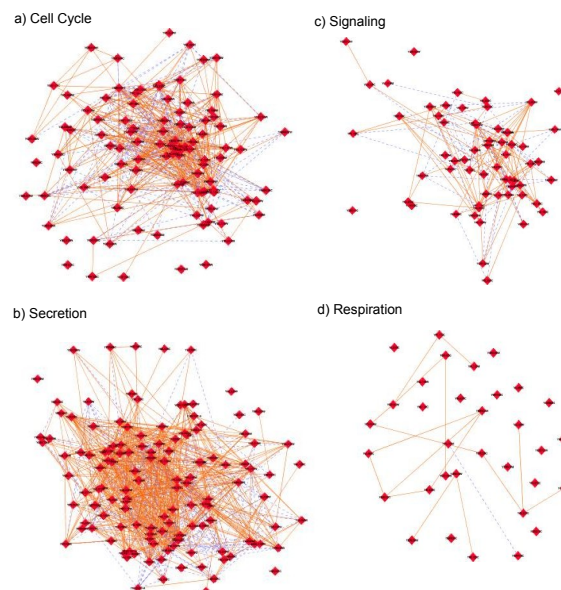
$$\mathbb{E} Y_{11} = \Lambda \mu, \quad \mathbb{E} Y_{jk} = \Lambda(\mu + \alpha_j + \delta_k) \quad j, k = 2, 3$$

- **Temporal correlation** is modeled directly via R (as $AR(1)$ process)
- Results:
 - ~ 3000 genes,
 - 47 pathways showed significant changes of expression
 - 24 pathways showed changes over **time**
 - 29 pathways showed changes in response to different **sorbitol** levels
 - 12 pathways showed **both** types of changes
 - Significant pathways overlap with the gene functions recognized by Gasch et al.

Yeast ESR Network

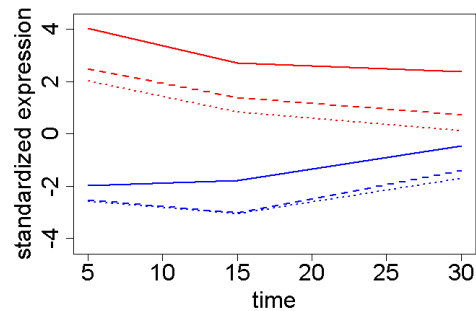


Significant subnetworks



Expression Profiles

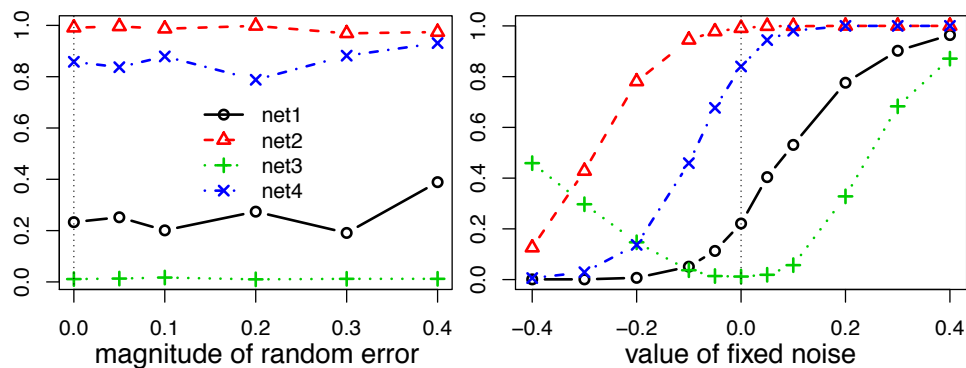
Average Standardized Expression Levels of Pathways



- **Induced** and **Suppressed** Pathways
- Can observe the **transient patterns of expressions** as predicted by *Gasch et al.*

Effect of Noise In Network Information

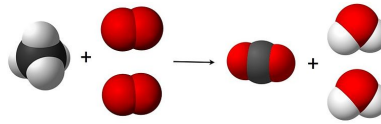
- Let \tilde{A} be observed network information, and A be the truth.
- It can be shown that, if $\|\tilde{A} - A\|$ is small then, NetGSA **still works** (is asymptotically most powerful unbiased test)



Metabolic Profiling in Bladder Cancer

Targeted metabolic profiling of bladder cancer (BCa) (*Putluri et al.*, 2012)

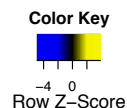
- ▶ 58 bladder cancer and adjacent benign samples
- ▶ Pathways information obtained from **KEGG**



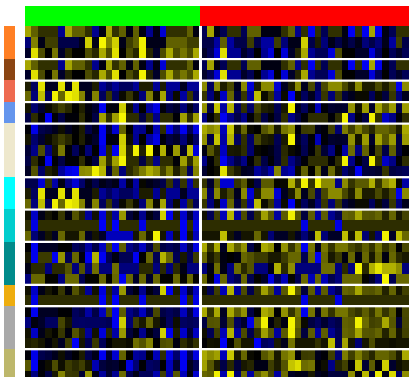
- ▶ Varying number of identified metabolites per pathway (3-15)
- ▶ **Q**: Which **pathways** show differential activity in BCa?

Metabolic Profiling in BCa

- ▶ 63 metabolites identified, mapped to 70 pathways
- ▶ 27 pathways with at least 3 members

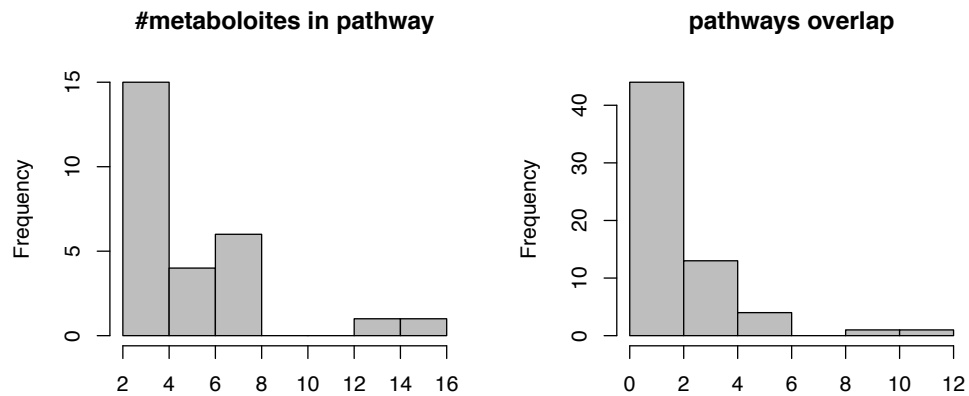


- Fatty acid biosynthesis
- Biosynthesis of unsaturated fatty acids
- Sulfur metabolism
- Lysine degradation
- Alkaloid biosynthesis II
- Methionine metabolism
- Valine, leucine and isoleucine biosynthesis
- Pyrimidine metabolism
- Valine, leucine and isoleucine degradation
- Pantothenate and CoA biosynthesis
- Phenylalanine, tyrosine and tryptophan biosynthesis



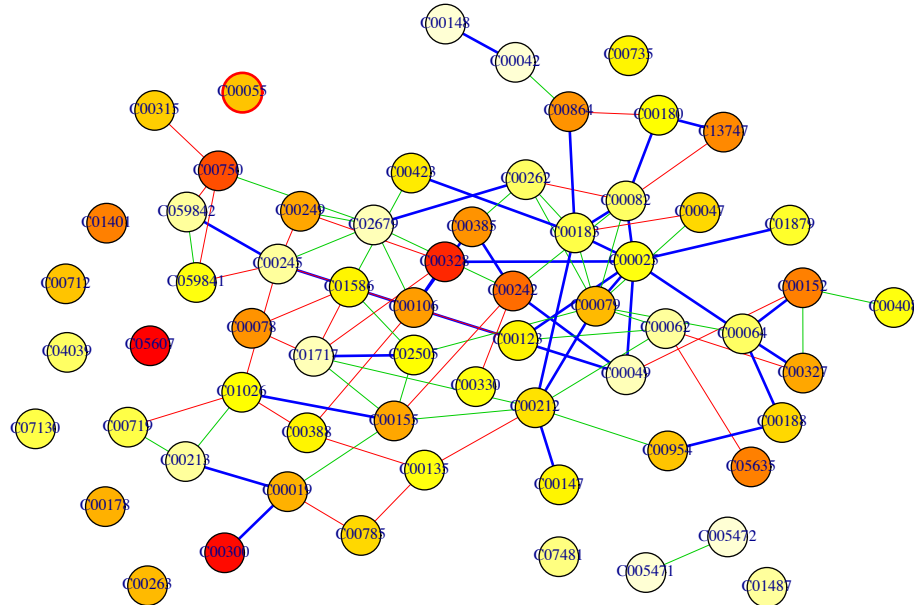
Metabolic Profiling in BCa

- Small pathway sizes & significant overlap among pathways



- Existing methods may not work well...

Metabolic Interaction Network



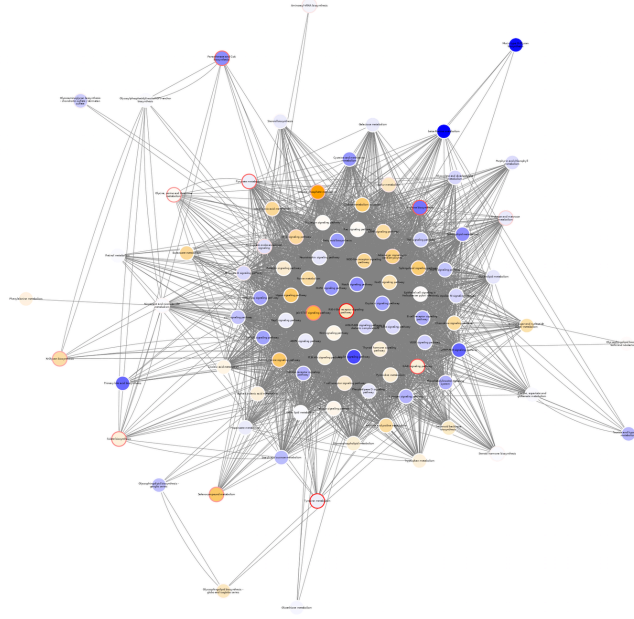
Significant Pathways

- ▶ **GSEA** does not identify any pathway as differential
- ▶ **GSA** identifies **Fatty Acid Biosynthesis** as differential
- ▶ **NetGSA** identifies another 7 pathways corresponding to role of **Amino Acid Metabolism** in BCa, similar to *Putluri et al* (2012)

R-Package netgsa

- ▶ Basic usage:
$$\text{NetGSA}(A, x, \text{group}, \text{pathways})$$
- ▶ A : List of $p \times p$ **weighted adjacency matrices** for each condition (e.g. normal vs cancer), to **capture changes in the network**
- ▶ pathways : a $K \times p$ 0-1 matrix of pathway membership:
 $\text{pathways}_{k,j} = 1$ if gene/.../metabolite j in pathway k
- ▶ **Output**: test statistics and p-values for each pathway
- ▶ The NetGSA function takes a weighted A as input. The package includes functions to **learn A for undirected networks** from a (partial) list of network edges

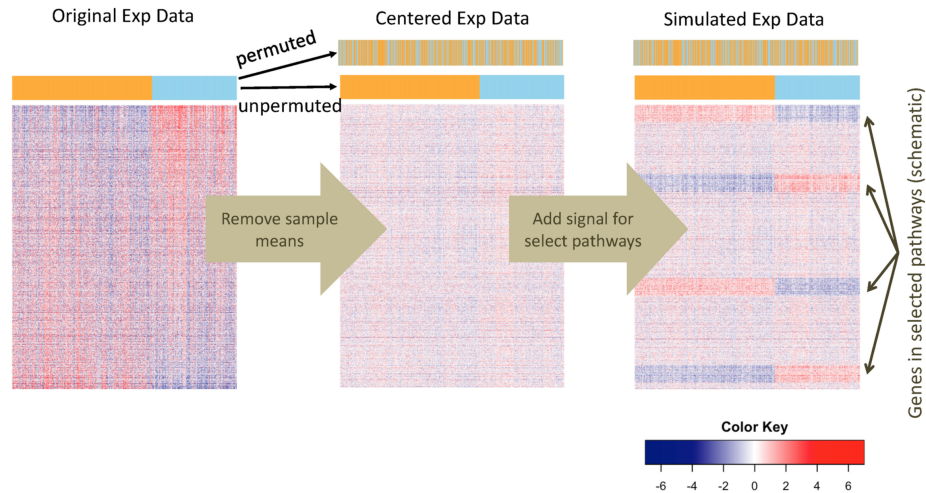
R-Package netgsa — coming soon...



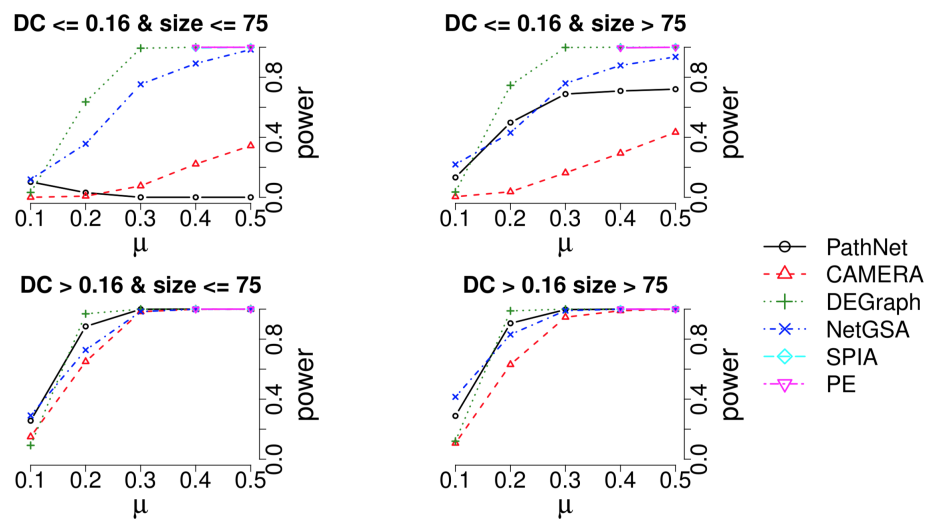
Comparison Using Synthetic Data (Ma, S., Michailidis, 2018)

- ▶ Comparison of topology-based pathway enrichment methods using two synthetic data sets
 - ▶ Gene expression data $p \approx 3000$
 - ▶ Metabolomics data $p \approx 100$
- ▶ *In silico* data sets with known signal:
 1. Remove the original signal, but **keep the correlation structure**
 2. **Perturb means in one condition** (differential expression) for nodes in selected pathways
 3. Also use sample permutation to **create data with equal correlation structure**

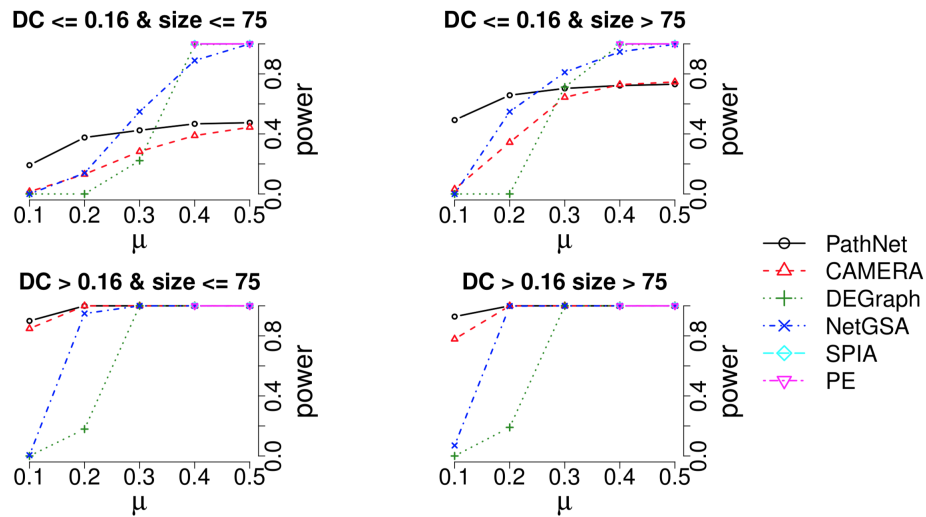
Comparison Using Synthetic Data



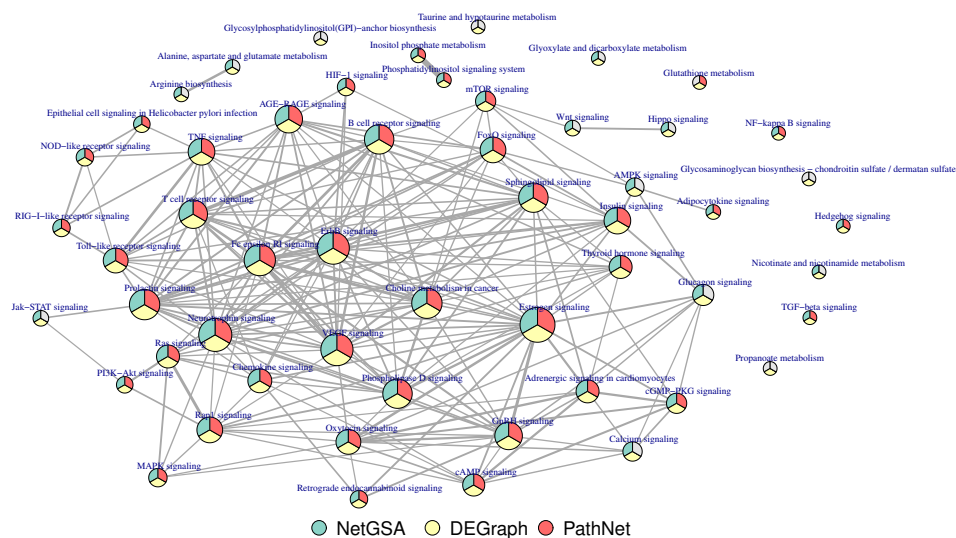
Results for Gene Expression Data — Equal Covariance



Results for Gene Expression Data — Diff Covariance

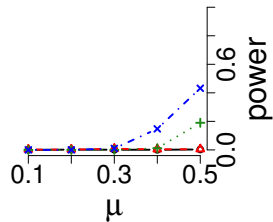


Results for Gene Expression Data

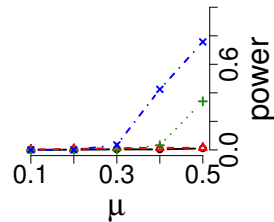


Results for Metabolomics Data — Equal Covariance

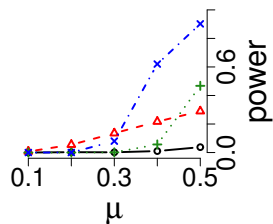
DC ≤ 0.5 & size ≤ 11



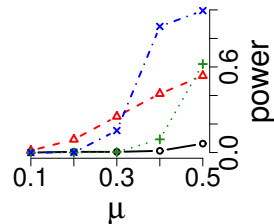
DC ≤ 0.5 & size > 11



DC > 0.5 & size ≤ 11



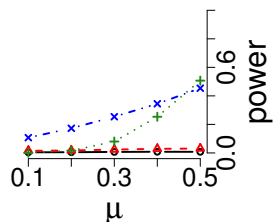
DC > 0.5 size > 11



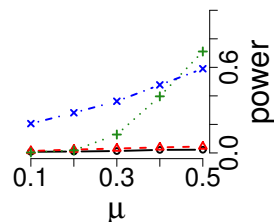
—○— PathNet
-△- CAMERA
-+- DEGraph
-x- NetGSA

Results for Metabolomics Data — Diff Covariance

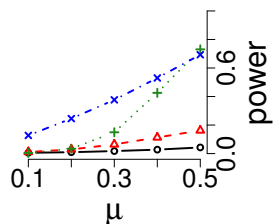
DC ≤ 0.5 & size ≤ 11



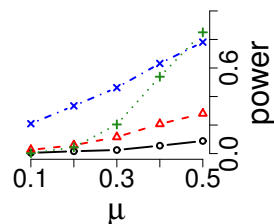
DC ≤ 0.5 & size > 11



DC > 0.5 & size ≤ 11

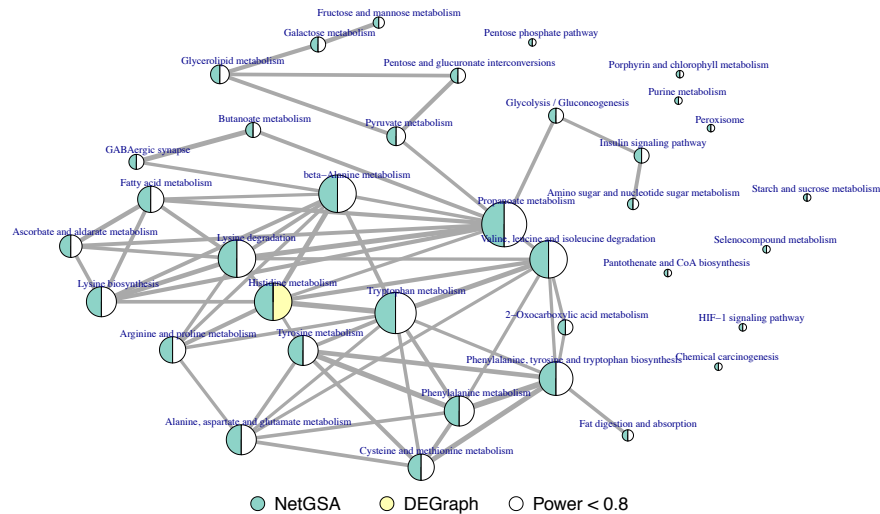


DC > 0.5 size > 11



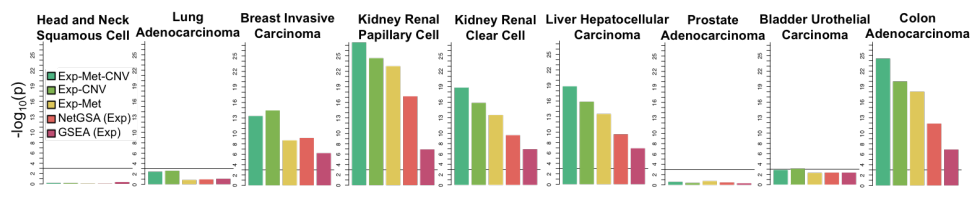
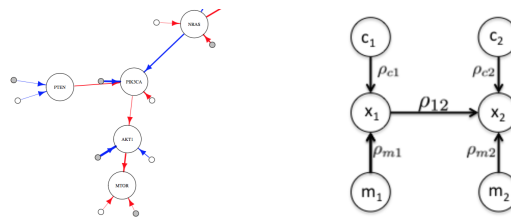
—○— PathNet
-△- CAMERA
-+- DEGraph
-x- NetGSA

Results for Metabolomics Data



Multi-Omics NetGSA

Pan-cancer integration of **expression**, **methylation** and **CNV** in **BRAF** (TCGA data)⁵



⁵Zhang et al (2018)

Identifying Enriched Modules in Networks

Identifying Enriched Modules in Networks

Two general strategies:

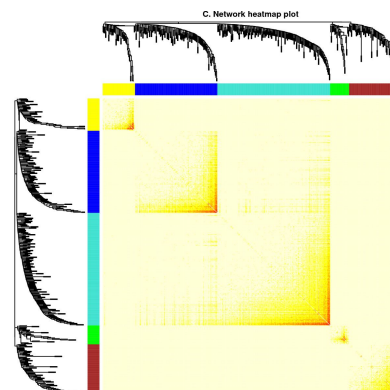
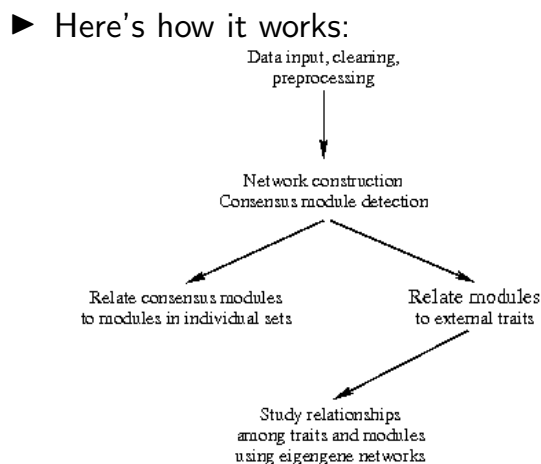
- ▶ Assess the significance of **data-driven modules** (WGCNA):
 1. Identify modules (network clustering, etc)
 2. Assess the significance of modules
- ▶ **Search** for enriched (connected) subnetworks (often using greedy search methods)
- ▶ Advantage: No need to rely on known pathways — especially useful when known pathways are not complete, etc
- ▶ Disadvantage: Interpretation may become challenging...

WGCNA⁶

- ▶ We previously talked about weighted gene co-expression (WGCNA), but for **estimating** networks
- ▶ However, WGCNA is also used for topology-based enrichment analysis, although in a different way than many other topology-based methods
- ▶ Here's how it works:
 1. Estimate the **co-expression network** (more in the next lecture)
 2. Find **modules** by **clustering** the nodes in the estimated network
 3. Summarize the expressions of genes in each module using PCA (**eigen-genes**)
 4. Test if the eigen-genes are associated with the outcome

⁶Horvath & Zhang (2005); Langfelder et al (2008)

WGCNA



Let's look at an example in R...

Walktrap⁷

- Searches for connected modules containing significant genes
 - Weights each edges based on the **significance of its corresponding nodes**

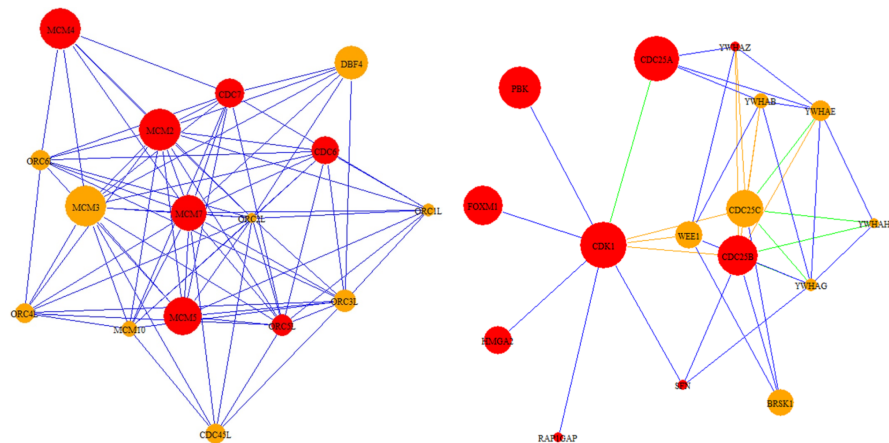
$$w_{ij} = (|FC_i| + |FC_j|)/2$$

- Connected significant modules are found through **community detection** using a **random walk** with transition probability

$$P_{ij} = \frac{w_{ij}}{\sum_j w_{ij}}$$

⁷Petrochilos et al (2013)

Identifying Cancer-Related Modules

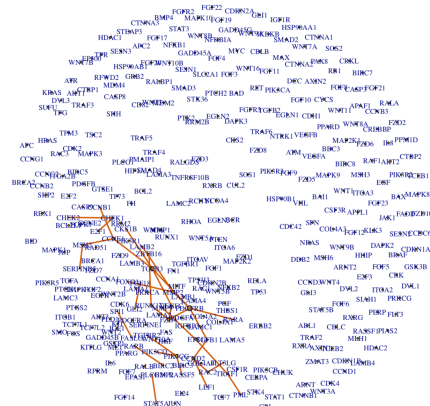
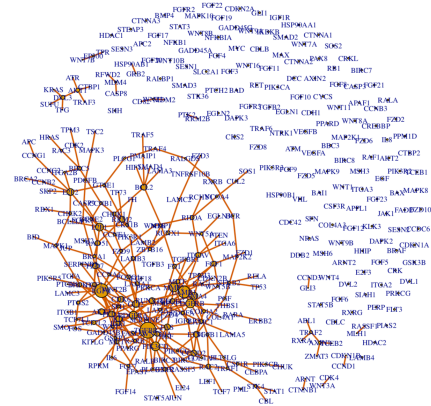


Summary

- Network-based methods (centrality-based, pathway topology, etc) rely on network information — helpful if correct network information avail
- What if network information is not available?

Summary

- Focus is shifting towards estimating changes in the structure of networks: **differential network biology**⁸



⁸Ideker & Krogan (2012); Shojaie (2020)

Pathway & Network Analysis of Omics Data: Learning Undirected Networks

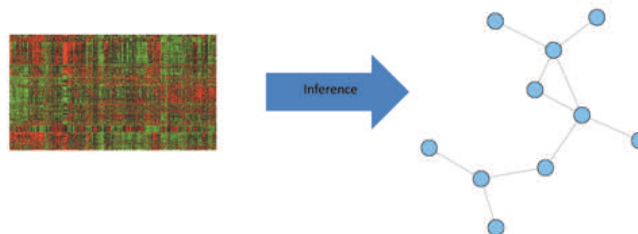
Ali Shojaie
Department of Biostatistics
University of Washington
faculty.washington.edu/ashojaie

Summer Institute for Statistical Genetics – 2020

Learning Undirected Networks

Learn network from data (**structure learning**):

- ▶ Data matrix: $X_{n \times p}$.
- ▶ Features correspond to the p nodes in the network.
- ▶ Goal: Learn edges between nodes \equiv learn the **statistical relationships** between features.



Why Do We Need Network Inference?

- ▶ Despite progress, our knowledge of interactions is limited.
- ▶ The entire genome is a vast landscape, and **experiments for discovering networks are very expensive**.
- ▶ From a statistical point of view, **network estimation is related to estimation of covariance matrices**, which has many independent applications in statistical inference and prediction (*more about this later*).
- ▶ Finally, and perhaps most importantly, **gene and protein networks are dynamic** and changes in these networks have been attributed to complex diseases.

Network Inference — An Overview

Two general classes of network inference methods:

- ▶ Methods based on **marginal measures of association**:
 - ▶ Co-expression Networks (based on linear measures of association)
 - ▶ Methods based on **mutual information** (can accommodate non-linear associations)
- ▶ Methods based on **conditional measures of association**:
 - ▶ Methods assuming (multivariate) normality (glasso, etc)
 - ▶ Generalizations to allow for nonlinear dependencies (nonparanormal, etc)

Graphical Models

Probabilistic Graphical Models ¹

Joint multivariate probability distribution where dependencies can be represented as a network.

Advantages:

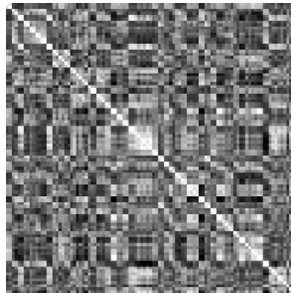
- ▶ Graphical models offer efficient factorized forms for joint distributions with easily interpretable dependencies.
 - ▶ **Conditional dependencies** denoted via an edge in network.
- ▶ Convenient visual representation.

¹A detailed technical introduction to these models is provided in *Graphical Models, Exponential Families, and Variational Inference*; Wainwright and Jordan (2008)

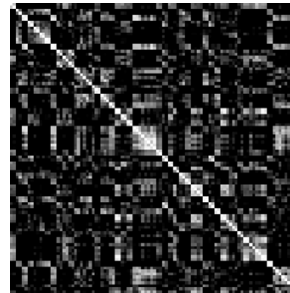
Marginal Association Networks

Correlation Networks (Association Networks)

- ▶ Simplest (and most-widely used!) method for estimating networks — key assumption:
large correlation \equiv presence of an edge
- ▶ Let $r(i, j)$ be correlation between X_i and X_j ; we claim an **edge between i and j if $|r(i, j)| > \tau$** .
 - ▶ τ : a user-specified threshold (**tuning parameter**).



Correlation matrix



Thresholded correlation matrix

Limitations of Correlation Networks

1. The estimation is highly dependent on the **choice of τ** .
2. Correlations capture **linear associations**, but **many real-world relationships are nonlinear**.
3. Large correlations can occur due to **confounding**.

Limitations of Correlation Networks

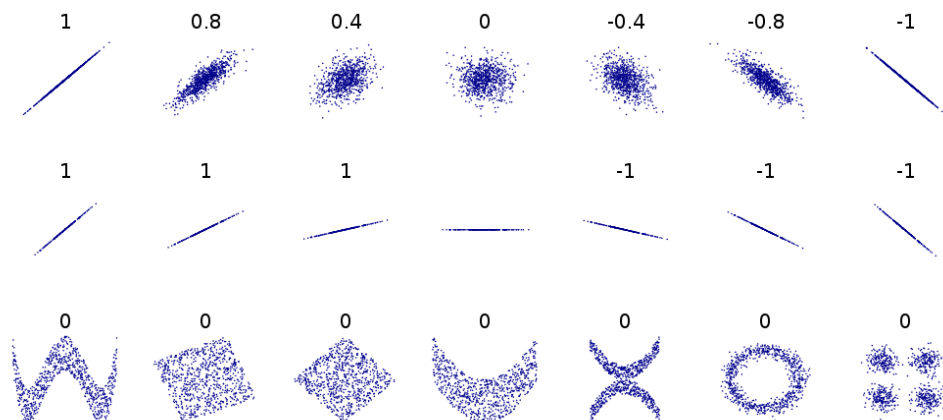
The estimation is highly dependent on the choice of τ .

- ▶ We can work with **weighted co-expression networks** (WGCNA)
- ▶ We can instead **test $H_0 : r_{xy} = 0$**
 - ▶ A commonly used test is based on the **Fisher transformation**

$$Z = \frac{1}{2} \ln \left(\frac{1+r}{1-r} \right) = \text{artanh}(r) \sim_{H_0} N \left(0, \frac{1}{\sqrt{n-3}} \right)$$

Limitations of Correlation Networks

Correlations capture **linear** associations, but **many real-world relationships are nonlinear**.



Limitations of Correlation Networks

Correlations capture **linear** associations, but **many real-world relationships are nonlinear**.

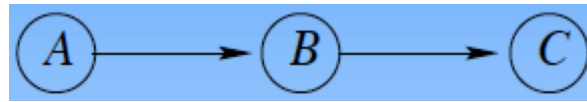
- ▶ We can use other measures of association, for instance, **Spearman correlation** or **Kendal's τ** .
 - ▶ These methods define the correlation between two variables, based on the **ranking** of observations, and not their exact values.
 - ▶ They can better capture non-linear associations.
- ▶ We can instead use **mutual information**; this has been used in many algorithms, e.g. ARACNE.

ARACNE: Algorithm for the Reconstruction of Accurate Cellular NEtworks²

1. Identifies statistically significant gene-gene co-regulation based on mutual information
2. It then eliminates indirect relationships in which two genes are co-regulated through one or more intermediates

²Margolin et al (2006)

Key Idea: Data Processing Inequality (DPI)



$$I(A, C) \leq \min[I(A, B), I(B, C)]$$

where

$$I(g_i, g_j) = \log P(g_i, g_j) / P(g_i)P(g_j)$$

- ▶ Look at every triplet and remove the weakest link
- ▶ Need to estimate marginal and joint (pairwise) probabilities (using Gaussian Kernel)

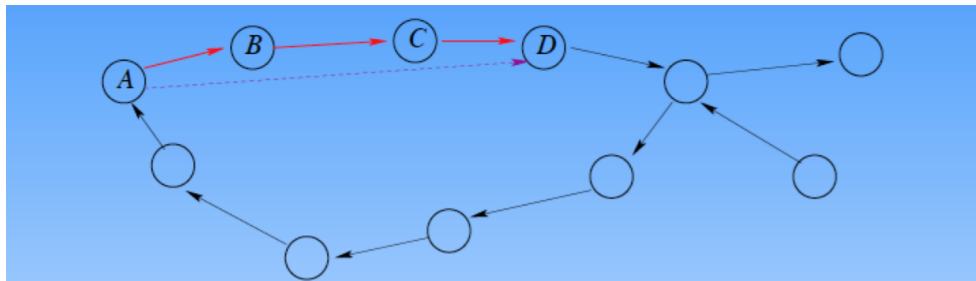
Algorithm Details

- ▶ The algorithm examines each gene triplet for which all pairwise MIs are greater than a cut-off and removes the edge with the smallest value based on DPI.
 - ▶ Each triplet is analyzed even if its edges have been selected for removal by prior DPI applications to other triplets.
 - ▶ The least of the three MIs can come from indirect interactions only, and checking against the DPI may identify **gene pairs that are not independent, but still do not interact.**

Rationale and Guarantees

- ▶ If MIs are estimated with no errors, then ARACNE reconstructs the underlying interaction network exactly, if the network is a tree and has only pairwise interactions.
- ▶ The maximum MI spanning tree is a subnetwork of the network built by ARACNE.

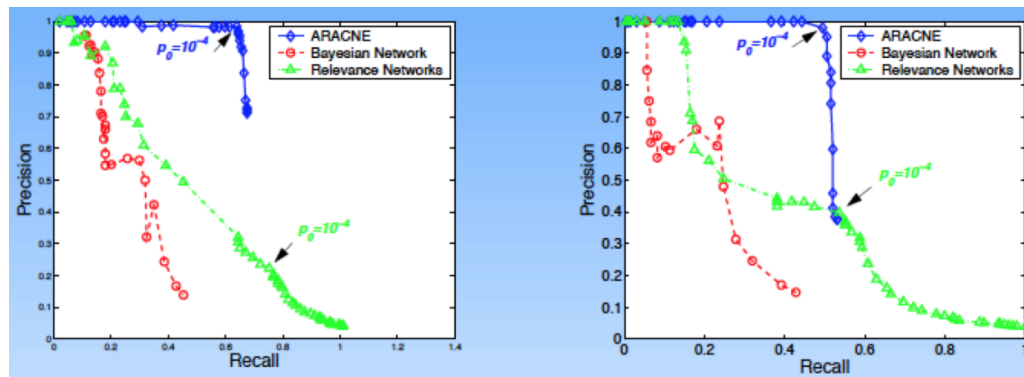
Rationale and Guarantees



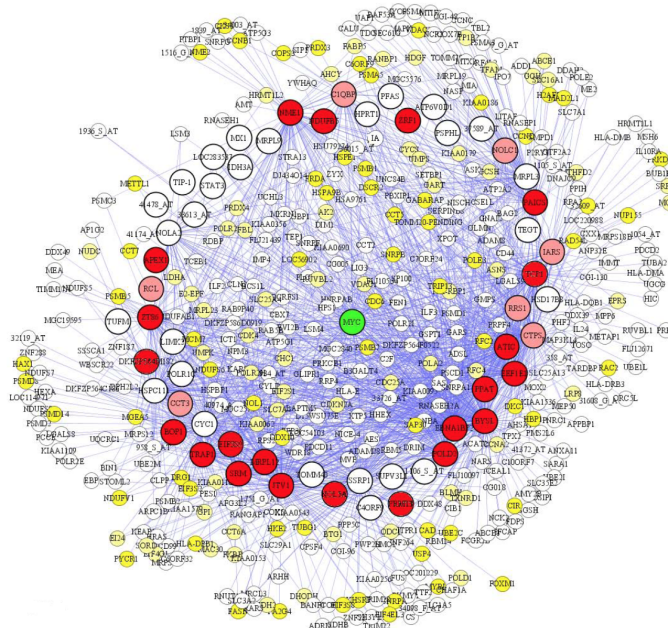
Theorem. Let π_{ik} be the set of nodes forming the shortest path in the network between nodes i and k . Then, if MIs can be estimated without errors, ARACNE reconstructs an interaction network without false positives edges, provided: (a) the network consists only of pairwise interactions, (b) for each $j \in \pi_{ik}$, $I_{ij} \geq I_{ik}$.

Further, ARACNE does not produce any false negatives, and the network reconstruction is exact iff (c) for each directly connected pair ij and for any other node k , we have $I_{ij} > \min[I_{ik}, I_{jk}]$.

Performance on Synthetic Data



Application: B-lymphocytes Expression Data



Application: B-lymphocytes Expression Data

- ▶ MYC (proto-oncogene) subnetwork (2063 genes)
- ▶ 29 of the 56 (51.8%) predicted first neighbors biochemically validated as targets of the MYC transcription factor.
- ▶ New candidate targets were identified, 12 experimentally validated.
 - ▶ 11 proved to be true targets.
- ▶ The candidate targets that have not been validated are possibly also correct.

Software

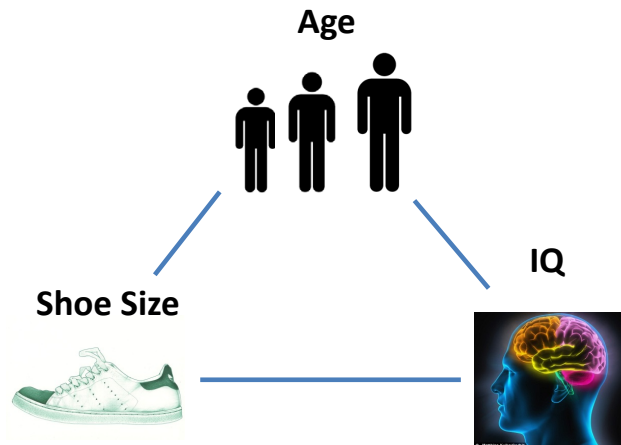
- ▶ Implemented in the R-package minet:

```
source("http://bioconductor.org/biocLite.R")
biocLite("minet")
```
- ▶ Main estimation function `aracne(mim, eps=0)`
 - ▶ `mim`: mutual information matrix

```
mim <- build.mim(syn.data, estimator="spearman")
```
 - ▶ `eps`: threshold for setting an edge to zero, prior to searching over triplets

Limitations of Correlation Networks

Large correlations can occur due to **confounding**.

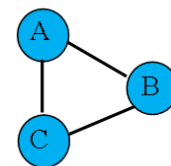


Markov Networks

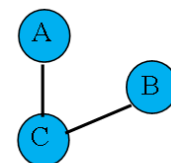
Markov Network

An *undirected graphical model* that characterizes **conditional dependence** (\equiv direct relationships).

- *Edge*: Two nodes are **conditionally dependent**.
- *No edge*: Two nodes are **conditionally independent**.
- Conditions on all other nodes.



$$A \perp B \mid C$$

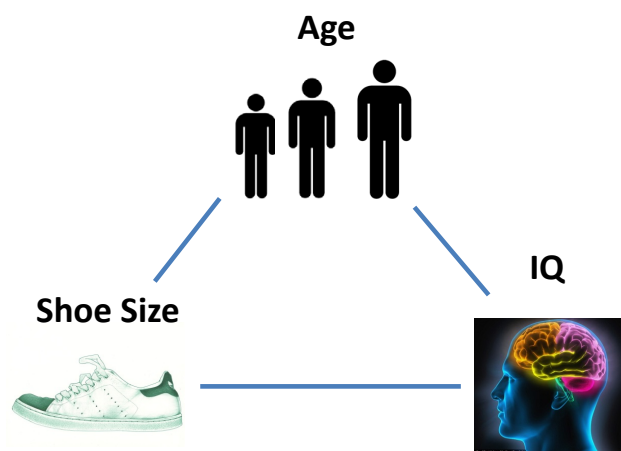


Markov Networks — Conditional Dependence

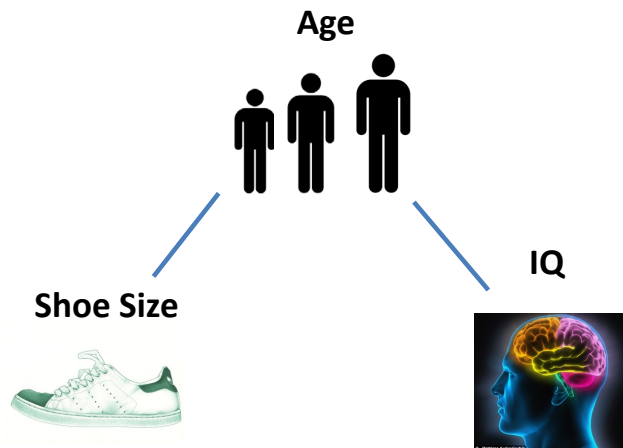
Regression Interpretation:

- ▶ Imagine trying to predict the observations in **Node A** (response) by the observations of all other nodes (predictors).
- ▶ **Node B** predictive of **Node A** (with all other nodes in model).
 - ▶ **A** is conditionally dependent on **B**.
 - ▶ Edge.
- ▶ Because of other nodes in model, **Node B** does not add any predictive value for **Node A**.
 - ▶ **A** is conditionally independent of **B**.
 - ▶ No Edge.

Markov Networks — Conditional Dependence



Markov Networks — Conditional Dependence



Conditional Dependence.

Markov Networks — Conditional Dependence

How can we learn conditional dependencies?

- ▶ A and B are conditionally independent given C if

$$P(A, B \mid C) = P(A \mid C)P(B \mid C)$$

- ▶ Generally difficult (need to estimate multivariate densities).
- ▶ Alternatively, can use nonparametric approaches, e.g. **conditional mutual information**, but not easy in high dimensions.
- ▶ Often resort to models, or simple measures, such as **partial correlations**...

Partial Correlation

- ▶ Partial correlation measures the correlation between A and B after the effect of the other variables are removed.
 - ▶ In our example, this means correlation between shoe size and IQ, after adjusting for age.
- ▶ The partial correlation between A and B given C is given by:

$$\rho_{AB \cdot C} \equiv \rho(A, B | C) = \frac{\rho_{AB} - \rho_{AC}\rho_{BC}}{\sqrt{1 - \rho_{AC}^2}\sqrt{1 - \rho_{BC}^2}}.$$

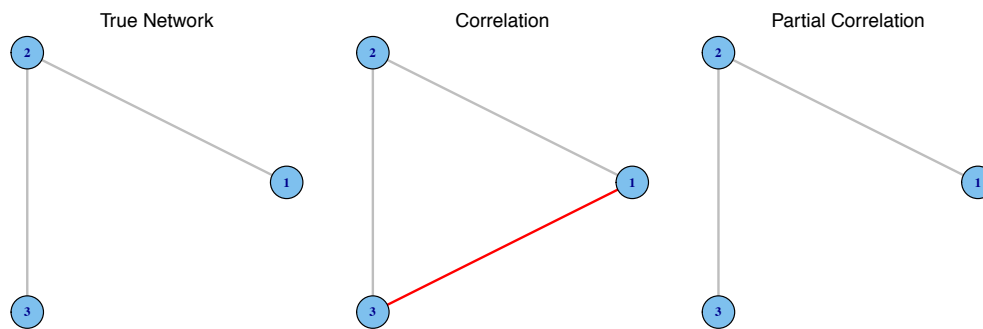
- ▶ Alternatively, regress A on C and get the residual, r_A ; do the same for B to get r_B . The partial correlation between A and B given C is $\text{Cor}(r_A, r_B)$.

Partial Correlation

- ▶ Partial correlation is symmetric \Rightarrow undirected network
- ▶ Partial correlation takes values between -1 and 1
- ▶ In partial correlation networks, we draw an edge between A and B , if the partial correlation between them is large
- ▶ Calculation of partial correlation is more involved

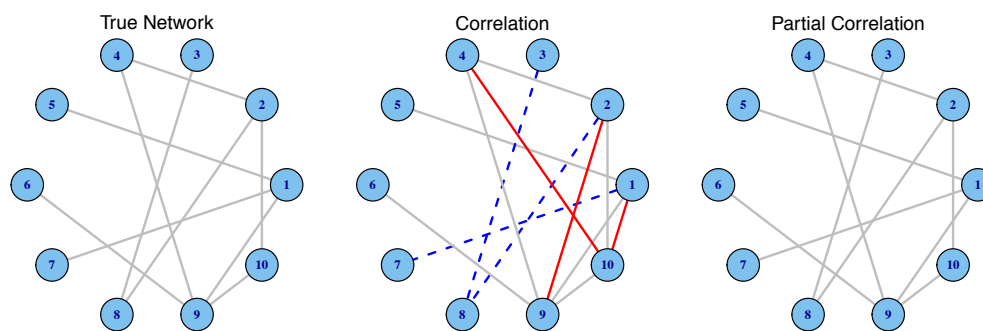
A Simple Example

$$\text{Correlation} = \begin{bmatrix} 1 & .8 & .7 \\ .8 & 1 & .8 \\ .7 & .8 & 1 \end{bmatrix} \quad \text{PartialCorr} = \begin{bmatrix} 1 & .6 & 0 \\ .6 & 1 & .6 \\ 0 & .6 & 1 \end{bmatrix}$$



A Larger Example

- ▶ A network with 10 nodes and 20 edges
- ▶ $n = 100$ observations
- ▶ Estimation using correlation & partial correlation (20 edges)

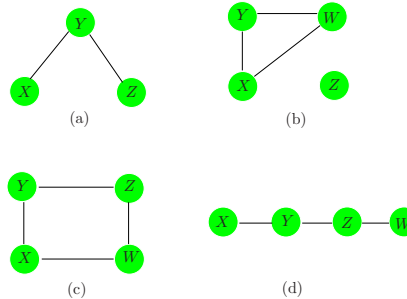


Gaussian Graphical Models (GGMs)

Partial Correlation for Gaussian Random Variables

- ▶ For Gaussian (multivariate normal) random variables, partial correlation between X_i and X_j **given all other variables** is given by **the inverse of the (standardized) covariance matrix Σ** .
 - ▶ The (i, j) entry in Σ^{-1} gives the partial correlation between X_i and X_j given all other variables $X_{\setminus i, j}$.
 - ▶ Multivariate normal: $X \sim N(0, \Sigma)$
 - ▶ $\Theta \equiv \Sigma^{-1}$ = inverse covariance/**precision**/**concentration** matrix.
 - ▶ Zeros in $\Theta \implies$ conditional independence!
 - ▶ Edges correspond to non-zeros in Θ .

Partial Correlation for Gaussian Random Variables



$$\begin{pmatrix} - & \times & 0 \\ \times & - & \times \\ 0 & \times & - \end{pmatrix} \quad \begin{pmatrix} - & \times & \times & 0 \\ \times & - & \times & 0 \\ \times & \times & - & 0 \\ 0 & 0 & 0 & - \end{pmatrix}$$

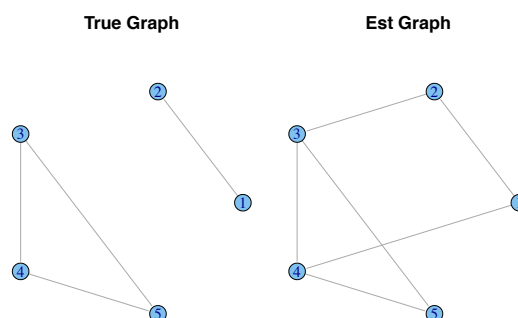
$$\begin{pmatrix} - & \times & 0 & \times \\ \times & - & \times & 0 \\ 0 & \times & - & \times \\ \times & 0 & \times & - \end{pmatrix} \quad \begin{pmatrix} - & 0 & 0 & \times \\ 0 & - & \times & 0 \\ 0 & \times & - & \times \\ \times & 0 & \times & - \end{pmatrix}$$

Estimating GGMs

From the discussion so far, to estimate the network, we can

1. Calculate the **empirical covariance matrix**: for (centered) $n \times p$ data matrix X , $S = (n - 1)^{-1} X^T X$.
2. **Get the inverse of S** . Non-zero values of S^{-1} give the edges.

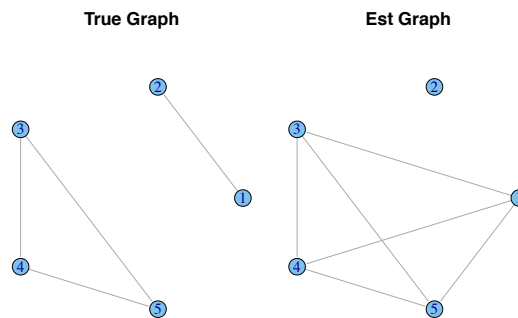
While simple, this may not work well in practice, even with large samples!



Estimating GGMs in High Dimensions

Many problems arise in high-dimensional settings, when $p \gg n$.

- ▶ First, S is not invertible if $p > n$!
- ▶ Even if $p < n$, but n is not very large, we may still get poor estimates, and many false positives/negatives.



Estimating GGMs in High Dimensions

- ▶ A number of methods have been recently proposed for estimating GGMs in high dimensions.
- ▶ The main idea in most of these methods is to use a regularization penalty, like the lasso.
- ▶ We discuss two approaches:
 - ▶ neighborhood selection
 - ▶ graphical lasso

The Lasso

- The lasso involves finding β that minimizes

$$\left\| y - \sum_{k=1}^p X_k \beta_k \right\|^2 + \lambda \sum_j |\beta_j|.$$

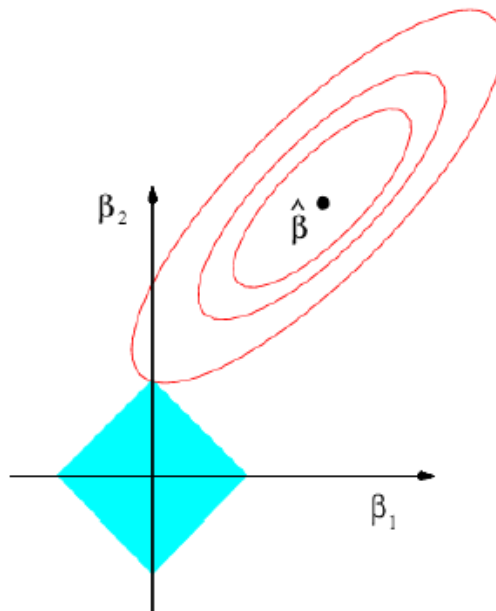
- Here λ is a **tuning parameter**
 - When $\lambda = 0$, we get least squares!
 - When λ is very large, we get $\hat{\beta} = 0$.
- Equivalently, find β that minimizes

$$\left\| y - \sum_{k=1}^p X_k \beta_k \right\|^2$$

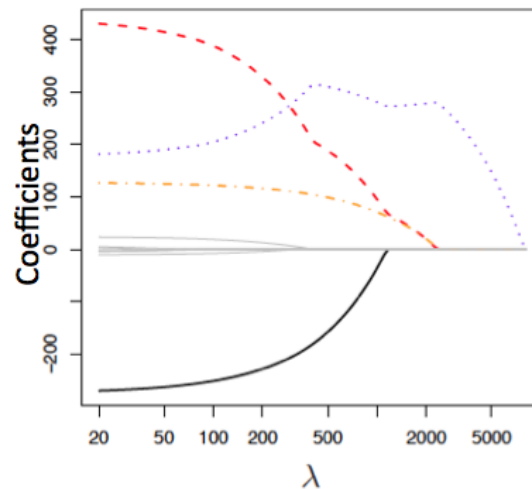
subject to the constraint that

$$\sum_{k=1}^p |\beta_k| \leq s.$$

A Geometric Interpretation



Lasso As λ Varies



Estimating GGMs in High Dimensions – Method 1

The idea behind **neighborhood selection**, is to estimate the graph by fitting a **penalized regression of each variable on all other variables**.

- Find **neighbors** of each node X_j by l_1 -penalized regression or lasso:

$$\underset{\beta^j}{\text{minimize}} \quad \|X_j - X_{\neq j}\beta^j\|_2^2 + \lambda \sum_{k \neq j} |\beta_k^j|$$

- The final estimate is found by combining all of the edges from these individual regression problems.
 - Symmetry — β_k^j not always same as β_j^k .
 - Use min or max rule.

Estimating GGMs in High Dimensions – Method 2

Estimate a sparse Θ via penalized maximum likelihood estimation (MLE).

Graphical Lasso (glasso)

$$\underset{\Theta}{\text{maximize}} \quad \log\det(\Theta) - \text{tr}(S\Theta) - \lambda \|\Theta\|_1$$

- **Blue**: Log-likelihood; $\log\det$ denotes the logarithm of the determinant of Θ and tr the trace (sum of diagonal elements) $S\Theta$.
- **Red**: Penalty term encourages zeros on the off-diagonal elements of Θ .

Comparing the Two Approaches

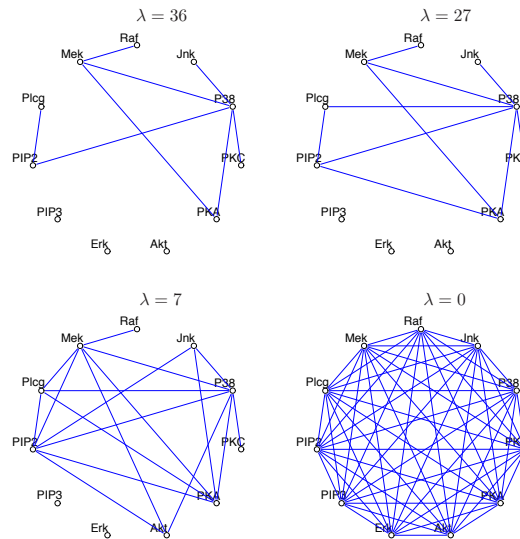
- Neighborhood selection is an **approximation for graphical lasso**:
 - Consider regression of X_j on $X_k, j \neq k$
 - Then, the regression coefficient for neighborhood selection is related to the j, k element of Θ :

$$\beta_k^j = -\frac{\Theta_{jk}}{\Theta_{jj}}$$

- Neighborhood selection is computationally more efficient, and may give better estimates, but doesn't give an estimate of Θ !

A Real Example

- **Flow cytometry** proteomics in single cells (Sachs et al, 2003).
- $p = 11$ proteins measured in $n = 7466$ cells



How to Choose λ ?

- λ modulates trade-off between **model fit** and **network sparsity**:
 - $\lambda = 0$ gives a dense network (no sparsity).
 - As λ increases, network becomes more sparse.
- A number of approaches proposed in the literature and used in practice
 1. **Cross-Validation** — tends to yield overly dense networks.
 2. **Extended BIC** — adjusted BIC for high dimensions.
 3. **Controlling the probability of falsely connecting disconnected components** at level α (Banerjee et al, 2008):

$$\lambda(\alpha) = \frac{t_{n-2}(\alpha/2p^2)}{\sqrt{n-2 + t_{n-2}(\alpha/2p^2)}},$$

($t_{n-2}(\alpha)$ is the $(100 - \alpha)\%$ quantile of t -dist with $n - 2$ d.f.)

4. **Stability selection** — Choose λ that gives the most **stable network** (R: huge package)

Other Types of Graphical Models

Nonparanormal (Gaussian Copula) Models

- ▶ Suppose $X \approx N(0, \Sigma)$, but there **exist monotone functions** $f_j, j = 1, \dots, p$ such that $[f_1(X_1), \dots, f_p(X_p)] \sim N(0, \Sigma)$
 - ▶ X has a nonparanormal distribution $X \sim NPN_p(f, \Sigma)$.
 - ▶ f and Σ are parameters of the distribution, and estimated from data.
 - ▶ For continuous distributions, the nonparanormal family is **the same as the Gaussian copula family**
- ▶ To estimate the nonparanormal network:
 - i) **transform the data:** $[f_1(X_1), \dots, f_p(X_p)]$
 - ii) **estimate the network of the transformed data** (e.g. calculate the empirical covariance matrix of the transformed data, and apply glasso or neighborhood selection)

A Related Procedure

- Liu et al (2012) and Xue & Zou (2012) proposed a closely related idea using **rank-based correlation**
 - Let r_j^i be the **rank of x_j^i** among x_j^1, \dots, x_j^n and $\bar{r}_j = (n+1)/2$ be the average rank
 - Calculate **Spearman's ρ** or **Kendall's τ**

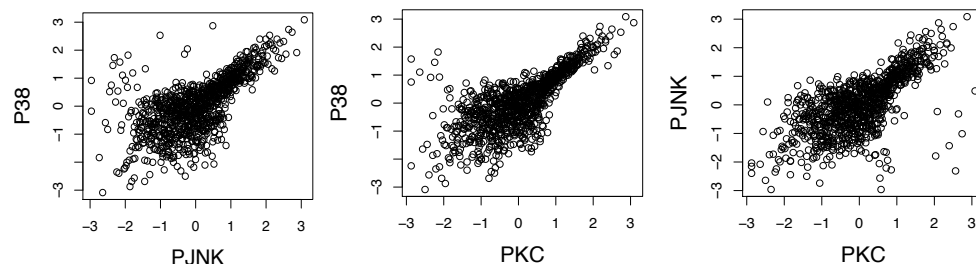
$$\hat{\rho}_{jk} = \frac{\sum_{i=1}^n (r_j^i - \bar{r}_j)(r_k^i - \bar{r}_k)}{\sqrt{\sum_{i=1}^n (r_j^i - \bar{r}_j)^2 \sum_{i=1}^n (r_k^i - \bar{r}_k)^2}}$$

$$\hat{\tau}_{jk} = \frac{2}{n(n-1)} \sum_{1 \leq i < i' \leq n} \text{sign}((x_j^i - x_j^{i'})(x_k^i - x_k^{i'}))$$

- If $X \sim NPN_p(f, \Sigma)$, then $\Sigma_{jk} = 2 \sin(\rho_{jk}\pi/6) = \sin(\tau_{jk}\pi/2)$
- Therefore, we can estimate Σ^{-1} by **plugging in rank-based correlations into graphical lasso** (R-package huge)

A Real Data Example

- Protein cytometry data for cell signaling (Sachs et al, 2005)
- Transform the data using a **Gaussian copula** (Liu et al, 2009), giving marginal normality
- Pairwise relationships still seem **non-linear**



- Shapiro-Wilk test rejects multivariate normality:
 $p < 2 \times 10^{-16}$

Graphical Models for Discrete Random Variables

- ▶ In many cases, biological data are not Gaussian: SNPs, RNAseq, etc
- ▶ Need to estimate CIG for other distributions: **binomial**, **poisson**, etc
- ▶ In this case, the estimators do not have a closed-form!
- ▶ A special case, which is computationally more tractable, is the class of **pairwise MRFs**

Pairwise Markov Random Fields

- ▶ The idea of **pairwise MRFs** is to “assume” that **only two-way interactions among variables** exist
 - ▶ The pairwise MRF associated with graph G over the random vector X is the family of probability distributions $P(X)$ that can be written as

$$P(X) \propto \exp \sum_{(j,k) \in E} \phi_{jk}(x_j, x_k)$$

- ▶ For each edge $(j, k) \in E$, ϕ_{jk} is called the **edge potential function**
- ▶ For discrete random variables, any MRF can be transformed to an MRF with pairwise interactions by introducing additional variables³

³Wainwright & Jordan (2008)

Graphical Models for Binary Random Variables

- ▶ Suppose X_1, \dots, X_p are binary random variables, corresponding to, e.g. SNPs, or DNA methylation
- ▶ A special case of discrete graphical models is the **Ising model for binary random variables**

$$P_{\theta}(x) = \frac{1}{Z(\theta)} \exp \left\{ \sum_{(j,k) \in E} \theta_{jk} x_j x_k \right\}$$

- ▶ A **pairwise MRF** for binary data, with $\phi_{jk}(x_j, x_k) = \theta_{jk} x_j x_k$
- ▶ $x^i \in \{-1, +1\}^p$
- ▶ The **partition function** $Z(\theta)$ ensures that the distribution sums to 1
- ▶ $(j, k) \in E$ iff $\theta_{jk} \neq 0$!

Graphical Models for Binary Random Variables

- ▶ We can consider a **neighborhood selection**⁴ approach with an ℓ_1 (lasso) penalty to find the neighborhood of each node $N(j) = \{k \in V : (j, k) \in E\}$
- ▶ For $j = 1, \dots, p$, need to solve (after some algebra)

$$\min_{\theta} \left\{ n^{-1} \sum_{i=1}^n \left[f(\theta; x^i) - \sum_{k \neq j} \theta_{jk} x_j^i x_k^i + \lambda \|\theta_{-j}\|_1 \right] \right\}$$

- ▶ $f(\theta; x) = \log \left\{ \exp \left(\sum_{k \neq j} \theta_{jk} x_k \right) + \exp \left(- \sum_{k \in -j} \theta_{jk} x_k \right) \right\}$
- ▶ This is equivalent to **solving p penalized logistic regression** problems, which is straightforward (R-package `glmnet`)

⁴Ravikumar et al (2010)

Other Non-Gaussian Distributions

- Assume a **pairwise graphical model**

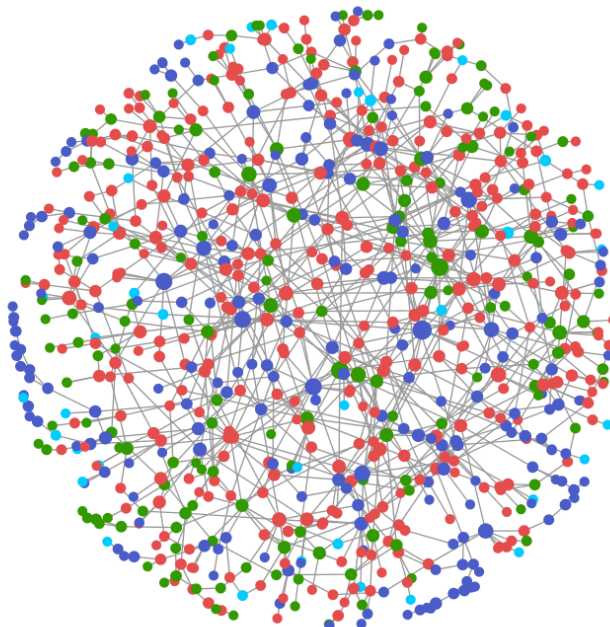
$$P(X) \propto \exp \left\{ \sum_{j \in V} \theta_j \phi_j(X_j) + \sum_{(j,k) \in E} \theta_{jk} \phi_{jk}(X_j, X_k) \right\}$$

- Then, similar to the Ising model, graphical models can be learned for other members of the **exponential family**
 - Poisson graphical models (for e.g. RNAseq), Multinomial graphical models, etc
 - All of these can be learned using a **neighborhood selection approach**, using the `glmnet` package⁵
 - We can even learn networks with multiple types of nodes (gene expression, SNPs, and CNVs)⁶

⁵Yang et al (2012)

⁶Yang et al (2014), Chen et al (2015)

Mixed Graphical Models



A General Approach for Estimation of Graphical Models

- ▶ Consider n iid observations from a p -dimensional random vector $\mathbf{x} = (X_1, \dots, X_p) \sim \mathcal{P}$
- ▶ Consider the (undirected) graph $G = (V, E)$ with vertices $V = \{1, \dots, p\}$
- ▶ Want to estimate edges $E \subset V \times V$ that satisfy $\forall j \in V, \exists N(j)$ such that:

$$p_j(X_j | \{X_k, k \neq j\}) = p_j(X_j | \{X_k : k \in N(j)\}) = p_j(X_j | \{X_k : (k, j) \in E\})$$

- ▶ $N(j)$ is the minimal set of variables on which the conditional densities depend

Estimating Conditional Independencies

Question: how to condition?

- ▶ **Approach 1:** Estimate the joint density $f(X_1, \dots, X_p)$; then get the conditionals $f_j(X_j | X_{-j})$
 - ▶ Efficient, coherent
 - ▶ Computationally challenging
 - ▶ Restrictive: how many joint distributions do you know?
 - ▶ Hard to check if assumptions hold!
- ▶ **Approach 2:** Estimate the conditionals directly $f_j(X_j | X_{-j})$
 - ▶ Computationally easy
 - ▶ Leads to easy & flexible models (regression)!
 - ▶ May not be efficient or coherent

A Semi-parametric Approach

- Consider additive non-linear relationships (additive model):

$$X_j \mid X_{-j} = \sum_{k \neq j} f_{jk}(X_k) + \varepsilon$$

- Then if $f_{jk}(X_k) = f_{kj}(X_j) = 0$, we conclude that X_j and X_k are **conditionally independent**, given the other variables
- In other words, we **assume that conditional distributions and conditional means depend on the same set of variables**
- We then use a semi-parametric approach for estimating the conditional dependencies

SpaCE JAM⁷

- Sparse Conditional Estimation with Jointly Additive Models (SpaCE JAM)

$$\underset{f_{jk} \in \mathcal{F}}{\text{minimize}} \frac{1}{2n} \sum_{j=1}^p \left\| x_j - \sum_{k \neq j} f_{jk}(x_k) \right\|_2^2 + \lambda \sum_{k > j} (\|f_{jk}(x_k)\|_2^2 + \|f_{kj}(x_j)\|_2^2)^{1/2}$$

- $f_{jk}(x_k) = \Psi_{jk} \beta_{jk}$
- Ψ_{jk} is a $n \times r$ matrix of basis functions for f_{jk}
- β_{jk} is an r -vector of coefficients
- The **standardized group lasso** penalty for functions $\|f_{jk}\|_2$
- This is a **convex** problem, and **block coordinate descent** converges to the global minimum

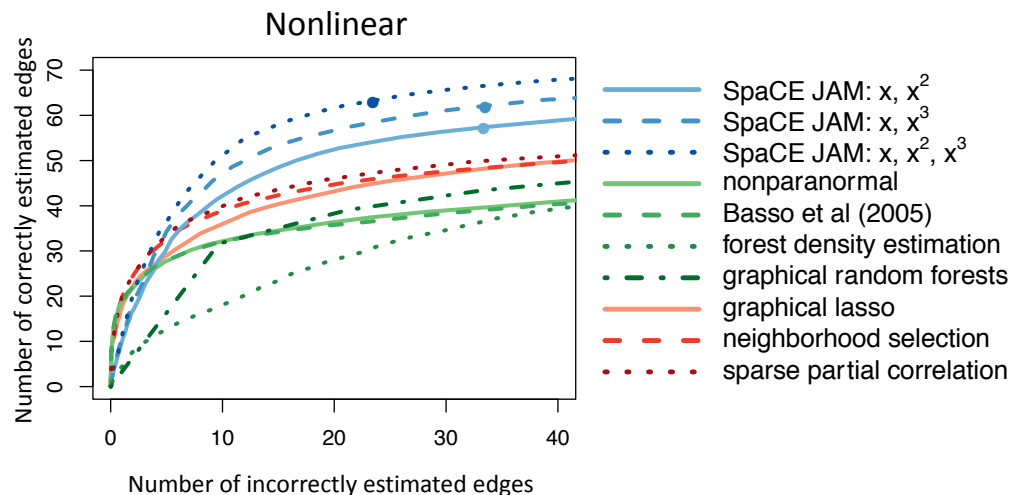
⁷Voorman et al (2014), R-package spacejam

Other Flexible Procedures

- ▶ **Forest density estimation** (Liu et al, 2011) assumes that underlying graph is a forest, and estimates the bivariate densities non-parametrically.
- ▶ **Graphical random forests** (Fellinghauer et al, 2013) uses random forests to flexibly model conditional means
 - ▶ They consider conditional dependencies through conditional mean
 - ▶ They allow for general random variables, discrete or continuous
 - ▶ Use a **random forest** to estimate $E[X_j | X_{\setminus j}]$ non-parametrically
 - ▶ Theoretical properties have not yet been justified

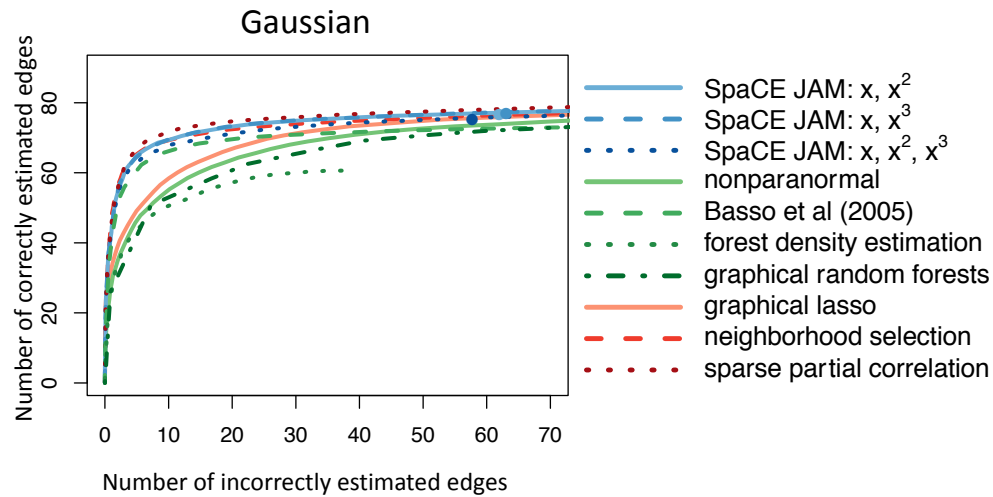
Comparison on Simulated Data

non-linear relationships ($p = 100$, $n = 50$)

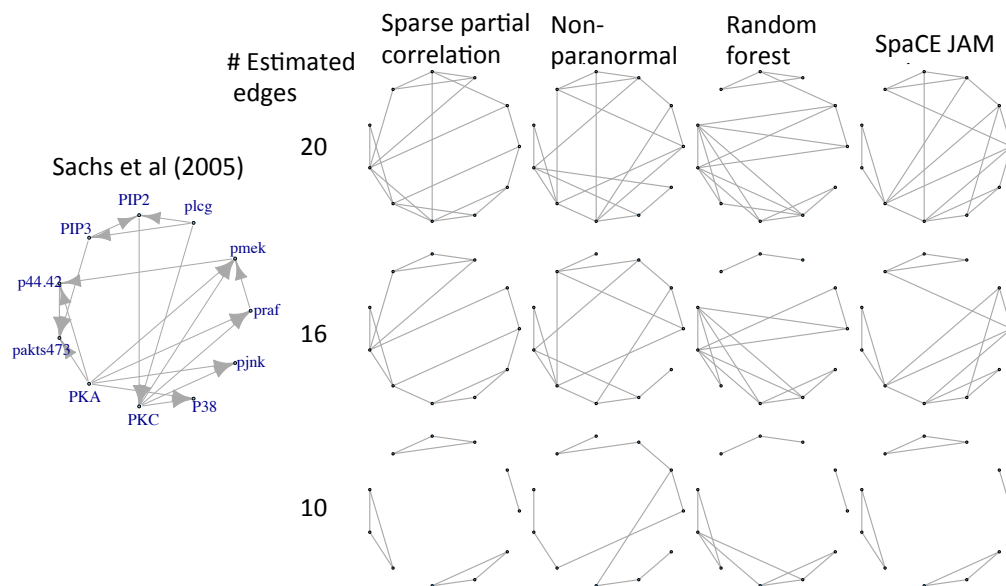


Comparison on Simulated Data

linear relationships ($p = 100$, $n = 50$)



Estimation of Cell Signaling Network



Other Extensions of GGMs

- ▶ Multiple Graphical Models
 - ▶ For groups of observations, estimate graphical models with shared structure across groups and individual structure within groups.
- ▶ Time Varying Graphical Models
 - ▶ Smoothly varying graph over time estimated via local kernel smoothers.
 - ▶ Change points in graph structure over time estimated via fusion penalties.
- ▶ Latent Variable Graphical Models
 - ▶ Assume observed features are dependent on latent variables which exhibit a low-rank effect. Estimate a sparse (graph structure) plus low-rank inverse covariance matrix.

Pathway & Network Analysis of Omics Data: Learning Directed Networks

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Summer Institute for Statistical Genetics – 2020

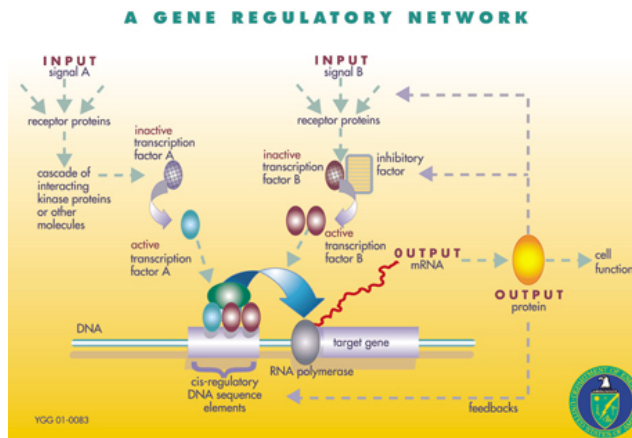
Bayesian Networks

- ▶ Bayesian networks are a special class of graphical models defined on **directed acyclic graphs**.
- ▶ Directed acyclic graphs (DAGs) are defined as graphs that:
 - i) **only have directed edges**, i.e. if $A_{ij} \neq 0$, $A_{ji} = 0$;
 - ii) there are **no cycles in the network**.
- ▶ Bayesian networks are widely used to **model causal relationships** between variables.
- ▶ Note that **correlation \neq causation!**
- ▶ Therefore, we (usually) cannot estimate Bayesian networks from (partial) correlations

Why Bayesian Networks?

Many biological networks include directed edges:

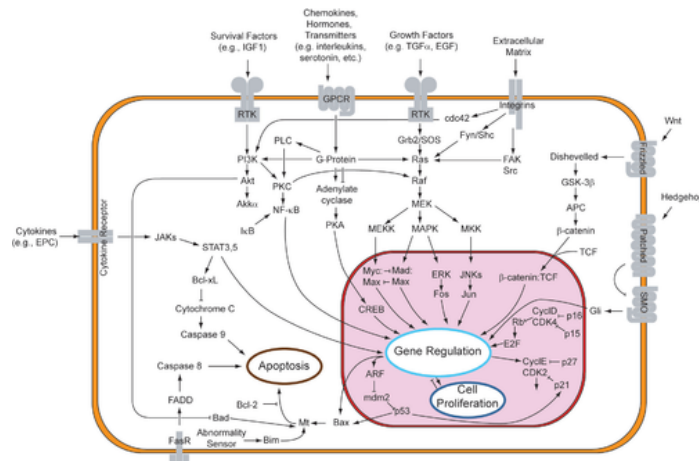
- In **gene regulatory networks**, protein products of **transcription factors** can alter the expression of **target genes**, but the target genes (usually) don't have a direct effect on the expression of transcription factors



Why Bayesian Networks?

Many biological networks include directed edges:

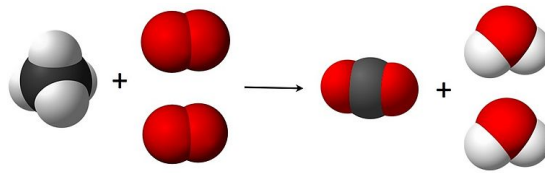
- In **cell signaling networks**, the signal from the cell's environment is transducted into the cell, and results e.g. in (global) changes in gene expression, but gene expression may not affect the environmental factors



Why Bayesian Networks?

Many biological networks include directed edges:

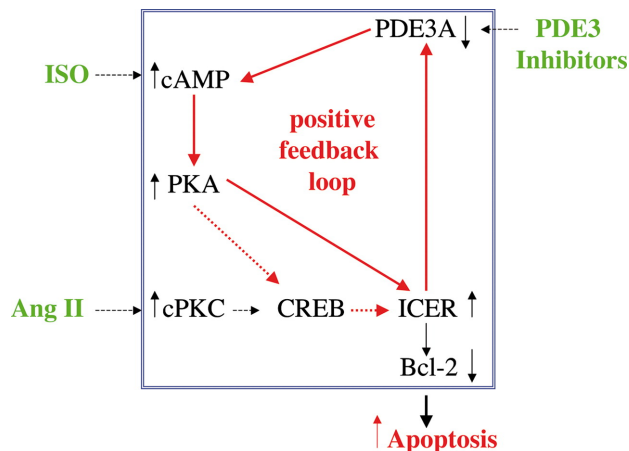
- Biochemical reactions in metabolic networks, may not be reversible, and in that case, one metabolite may affect the other, but the relationship is not reciprocated



Why Bayesian Networks?

However, biological networks may not be DAGs:

- Gene regulatory networks, signaling networks and metabolic networks, may all contain feedback loops (positive/negative)



which make estimation even more difficult!

What's the Difference?

- ▶ Bayesian networks are widely used to model **causal relationships** between variables.
- ▶ Undirected networks (e.g. GGM) provide information about **associations** among variables; while this greatly helps in the study of biological systems, in some cases, they are not enough (e.g. drug development).
- ▶ The main difference is the **direction of the edges**; however, it turns out that there are also some differences in terms of **structure/skeleton** of the network (more on this later).
- ▶ We can estimate undirected networks from **observational data**, i.e. steady-state gene expression data, but usually they are not enough for estimation of directed networks
- ▶ Finally, **estimating** directed networks is (much) **more difficult**

Why is estimation more difficult?

- ▶ Estimation of Bayesian networks requires estimating both the **skeleton** of the network (i.e. whether there is an edge between i and j) and also the **direction** of the edges.
- ▶ While estimation of skeleton is possible, **direction of edges cannot be in general learned from observational data**, no matter how many samples we have (this is referred to as *observational equivalence*). Consider this simple graph:



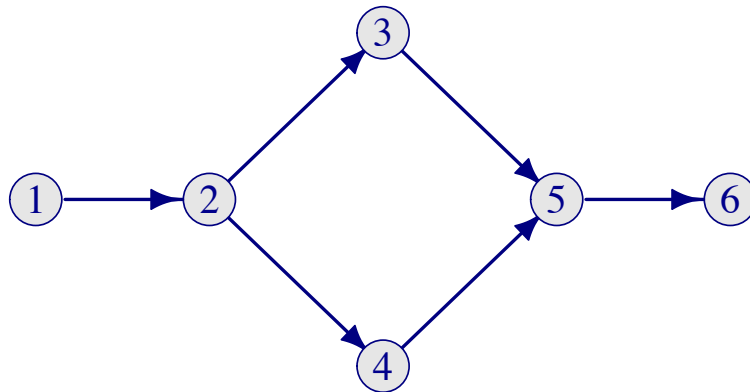
- ▶ Then, no matter what n is, we cannot distinguish between $X_1 \rightarrow X_2$ and $X_2 \rightarrow X_1$, so basically what we see is:



Directed Graphs: Some Terminology

- ▶ The **parents** of node j are $\{k : k \rightarrow j\}$, we denote this by pa_j or $\text{pa}(j)$
- ▶ The **children** of node j are $\{k : j \rightarrow k\}$
- ▶ Two vertices connected by an edge are called **adjacent**
- ▶ A **path** between two nodes i and j is a **sequence of distinct adjacent nodes**:
 - ▶ e.g. $i \leftarrow k_1 \rightarrow k_2 \rightarrow k_3 \leftarrow j$
 - ▶ In a DAG with p nodes, there cannot be a path longer than $p - 1$ (why?)
 - ▶ There can be multiple paths between two nodes
- ▶ i is an **ancestor** of j if there is a **directed path** of length ≥ 1 from i to j : $i \rightarrow \dots \rightarrow j$ (or if $i = j$)
- ▶ If i is an ancestor of j , then j is said to be a **descendant** of i

Directed Graphs: Some Terminology



- ▶ What are **parents/children** of $\{1, \dots, 5\}$?
- ▶ What are paths between 1&4, 3&4, 2&6?
- ▶ What are **ancestors** of $\{1, \dots, 5\}$?

Directed Graphs: Some Terminology

An important concept in DAGs is **colliders** (aka “inverted forks”):

- ▶ k is a **collider on a path** between i and j if it is not an end-point of the path, and the path is of the form

$$i \dots \rightarrow k \leftarrow \dots j$$

- ▶ k is a **non-collider** if it is not an end-point, and is not a collider **on a path**:

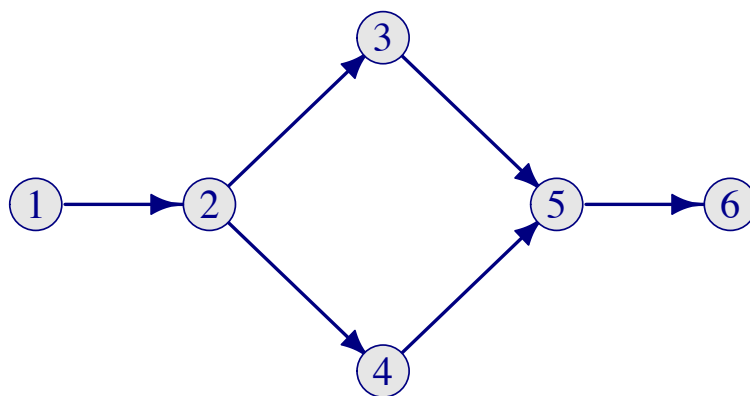
- ▶ $i \dots \leftarrow k \leftarrow \dots j$

- ▶ $i \dots \rightarrow k \rightarrow \dots j$

- ▶ $i \dots \leftarrow k \rightarrow \dots j$

- ▶ Note: **colliders and non-colliders are defined w.r.t. paths**; a collider in one path can be a non-collider in another!

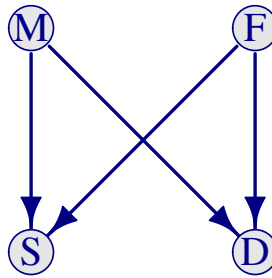
Directed Graphs: Some Terminology



- ▶ What are the **colliders** on paths between 1&4, 3&4, 2&6?
- ▶ What are the **non-colliders** on paths between 1&4, 3&4, 2&6?

Estimating Directed Graphs

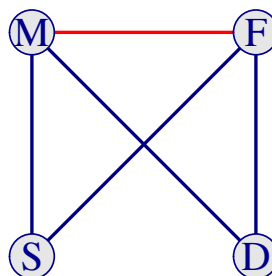
- The presence of **colliders** makes the estimation of directed graphs very challenging...



- Genetic information for **M**other, **F**ather, **D**aughter and **S**on in form of dominant/recessive genotype (A/a) for a single gene
- Then each individual can have one of three states: **AA**, **aa**, **Aa**

Estimating Directed Graphs

- **Conditioning on all other nodes**, gives additional **moral** (!!)
edges (\Rightarrow **moral graph**)



- Learning the **skeleton** of DAGs from **observational data** requires finding **right conditioning set**
 - Naively, this is done by *searching over all possible subset of other $p - 2$ nodes* — NP-hard with complexity $O(2^{p^2})$!!

Estimation of DAGs from Observational Data

Two general classes of algorithms for estimating DAGs:

- ▶ **constraint-based** methods
 - ▶ Often based on tests for CI; provide theoretical guarantees
 - ▶ PC algorithm, Grow-Shrink
- ▶ **score & search** methods
 - ▶ They assign a “score” to each estimated graph (e.g. based on likelihood, Bayes factor, AIC etc)
 - ▶ Greedy search to find the best scoring graph (Hill Climbing)
- ▶ **“hybrid” methods**
 - ▶ Usually first find the Markov blanket (e.g. the moral graph)
 - ▶ Then search in a restricted space (Max-Min Hill Climbing)

Constraint-Based Methods

- ▶ Need a conditional independence test (to test if $X \perp\!\!\!\perp Y \mid Z$)
 - ▶ For **Gaussian** data, we can use **partial correlation** (or the Fisher’s Z-transformation of it)
 - ▶ For **Binary** data, we can use **logOR**
 - ▶ In general, we can use **conditional mutual information**
- ▶ The idea is to see if there exists a set S , for each pair of nodes j, j' , such that $X_j \perp\!\!\!\perp X_{j'} \mid S$
 - ▶ S can have 0 to $p-2$ members! usually **stop at some $k \ll p$**
 - ▶ I.e., for each pair of variables (all $\binom{p}{2}$ of them), we need to look at all possible subsets of remaining variables!!
- ▶ These methods find the **DAG skeleton** (*conditional independence is symmetric*) — will talk about direction later

PC Algorithm (Spirtes et al, 1993)

- ▶ One of the first algorithms for learning structure of DAGs
- ▶ Efficient implementations that allow for learning DAG structures with p up to ~ 1000
 - ▶ R-package `pcalg` (Kalisch & Buhlmann, 2007)
- ▶ The algorithm **starts with a complete graph** (i.e. fully connected)
- ▶ Then for each pair of nodes j, j' it finds a **separating set, S** such that $X_j \perp\!\!\!\perp X_{j'} \mid S$
- ▶ If a set is found, then remove the edge, otherwise, $j - j'$

PC Algorithm (Spirtes et al, 1993)

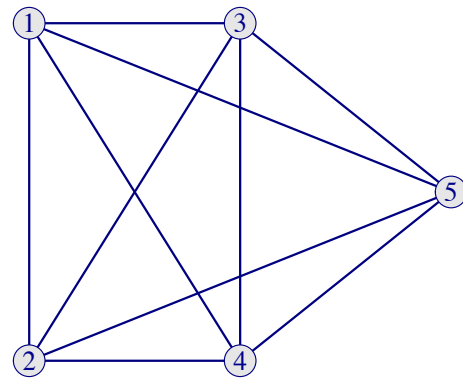
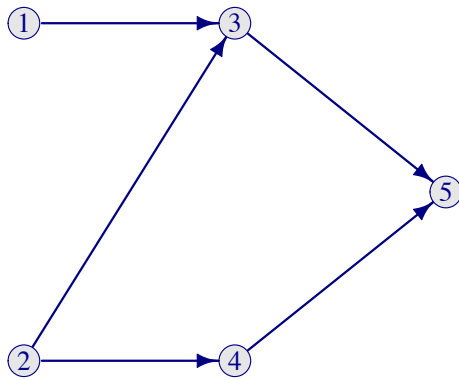
Start with a complete undirected graph, and set $i = 0$

Repeat

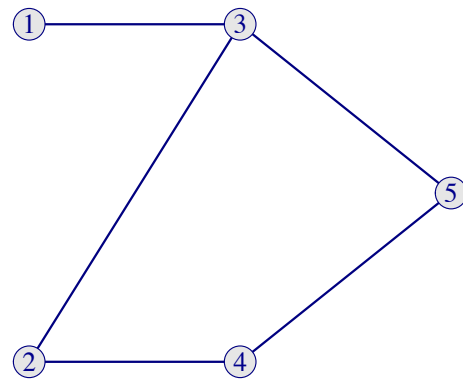
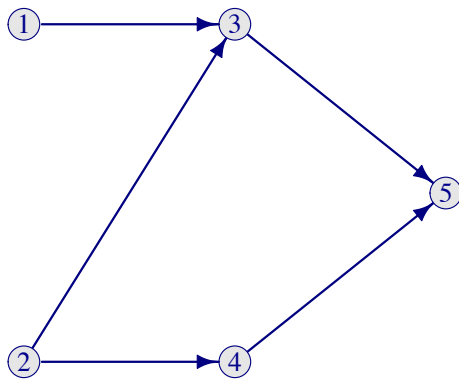
- ▶ For each $j \in V$
- ▶ For each $j' \in \text{ne}(j)$
- ▶ Determine if $\exists S \subset \text{ne}(j) \setminus \{j'\}$ with $|S| = i$
 - ▶ Test for CI: is $X_j \perp\!\!\!\perp X_{j'} \mid S$?
 - ▶ If such an S exists, then set $S_{jj'} = S$, **remove $j - j'$ edge**
- ▶ $i = i + 1$

Until $|\text{ne}(j)| < i$ for all j

Example



Example



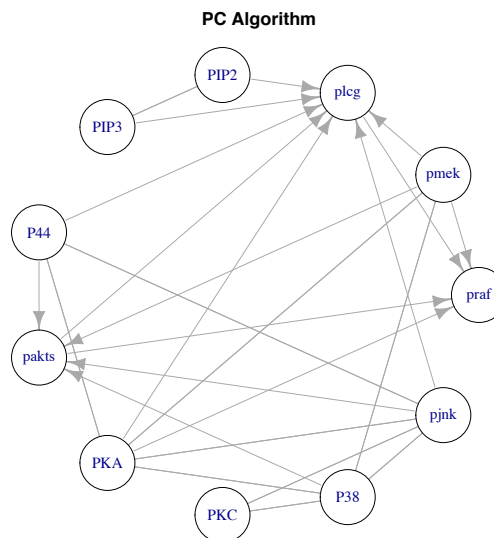
$i = 0$ $S_{1,2} = \emptyset$
 $S_{1,4} = \emptyset$
 $i = 1$ $S_{3,4} = \{2\}$
 $i = 2$ $S_{1,5} = \{3, 4\}$
 $S_{2,5} = \{3, 4\}$
 $i = 3$ **STOP** ($|ne_j| < 3 \ \forall j$)

Analysis of Protein Flow Cytometry using pcalg

```
> dat <- read.table('sachs.data')
> p <- ncol(dat)
> n <- nrow(dat)
## define independence test (partial correlations)
> indepTest <- gaussCItest
## define sufficient statistics
> suffStat <- list(C=cor(dat), n=n)
## estimate CPDAG
> pc.fit <- pc(suffStat, indepTest, p, alpha=0.1, verbose=FALSE)
> plot(pc.fit, main='PC Algorithm')
```

- Need to determine the **type of CI test** (indepTest), and **sufficient statistics** (suffStat)
- Also need to choose α (alpha), the **probability of false positive** for selecting edges.
 - Larger values of α allow more edges (not adjusted for multiple comparisons)
 - The algorithm works faster when α is small

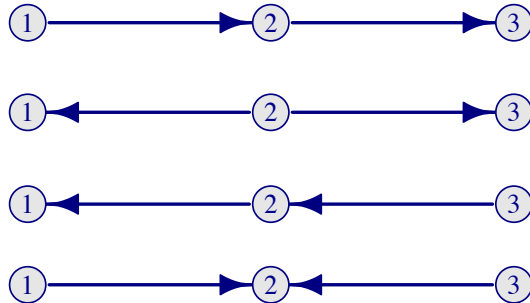
Analysis of Protein Flow Cytometry using pcalg



But wait, where did the **directions** come from? And why are only some of the edges directed?

Markov Equivalence

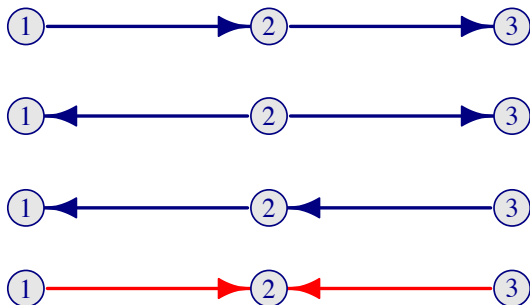
Consider the following 4 graphs



Which graphs satisfy $X_1 \perp\!\!\!\perp X_3 \mid X_2$?

Markov Equivalence

Consider the following 4 graphs



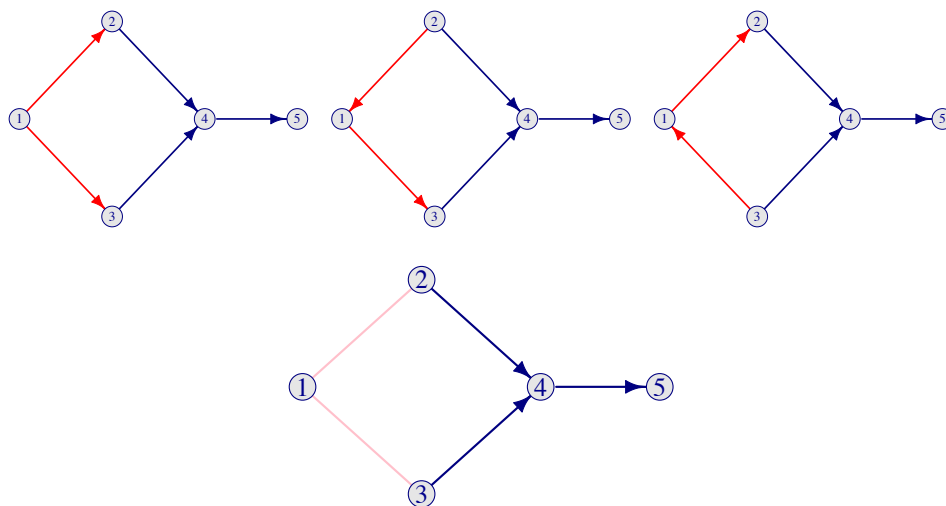
In the first 3 graphs, $X_1 \perp\!\!\!\perp X_3 \mid X_2$?

Two graphs that imply the same CI relationships via d-separation are called **Markov equivalent**

Representation of Markov Equivalence

- ▶ Markov equivalent graphs correspond to the same probability distribution and **cannot be distinguished from each other** based on observations!
- ▶ Therefore, the direction of edges that correspond to Markov equivalent graphs cannot be determined
- ▶ We show these edges using **undirected edges** in the graph
- ▶ The resulting graph is a **CPDAG** (completed partially directed acyclic graph), and is really the best we can do!

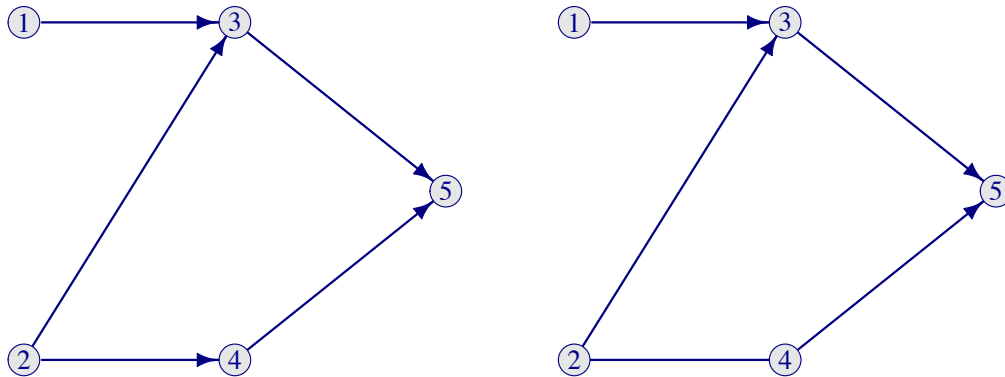
CPDAGs



Finding Partial Directions in DAGs

- ▶ Partial directions are determined from **unmarried colliders**:
 - ▶ For each unmarried collider $i - k - j$
 - ▶ If $k \notin S_{ij}$, orient $i - k - j$ as $i \rightarrow k \leftarrow j$
- ▶ In addition to the above rule,
 - ▶ Orient each **remaining unmarried collider** $i \rightarrow k - j$ as $i \rightarrow k \rightarrow j$
 - ▶ If $i \rightarrow k \rightarrow j$ and $i - j$ then orient as $i \rightarrow j$
 - ▶ If $i - m - j$ and $i \rightarrow k \leftarrow j$ are unmarried colliders and $m - k$, then orient as $m \rightarrow k$

Example



$$\begin{aligned}
 i = 0 \quad & S_{1,2} = \emptyset \\
 & S_{1,4} = \emptyset \\
 i = 1 \quad & S_{3,4} = \{2\} \\
 i = 2 \quad & S_{1,5} = \{3, 4\} \\
 & S_{2,5} = \{3, 4\}
 \end{aligned}$$

The bnlearn package

- ▶ There are a couple of R-packages for learning (CP)DAGs, including pclag, bnlearn, deal
- ▶ bnlearn implements a number of estimation methods, both constraint-based and search-based:
 - ▶ constraint-based algorithms:
 - ▶ Grow-Shrink (GS)
 - ▶ Incremental Association Markov Blanket (IAMB)
 - ▶ Fast Incremental Association (Fast-IAMB)
 - ▶ Interleaved Incremental Association (Inter-IAMB)
 - ▶ score-based algorithms:
 - ▶ Hill Climbing (HC)
 - ▶ Tabu Search (Tabu)
 - ▶ hybrid learning algorithms:
 - ▶ Max-Min Hill Climbing (MMHC)
 - ▶ General 2-Phase Restricted Maximization (RSMAX2)

Analysis of Protein Flow Cytometry using bnlearn

```
> dag1 <- gs(dat, alpha=0.01)    #GS method
> dag2 <- hc(dat2)               #Hill-Climbing search
>
> par(mfrow= c(1,2))
> plot(dag1)
> plot(dag2)
>
> compare(dag1, dag2)            #compare the two DAGs
```

- ▶ For GS need to choose α (alpha), the **false positive probability** for selecting edges
- ▶ gs (and other structure-based methods) find a PCDAG
- ▶ hc gives a directed graph (with highest score)
 - ▶ Multiple criteria for choosing the “best” graph
 - ▶ To “search” the space either a new edge is added, or a current edge is removed, or reversed (if no cycles)

Analysis of Protein Flow Cytometry using bnlearn

```
> dag1
Bayesian network learned via Constraint-based methods

model:
  [partially directed graph]
nodes:                                11
arcs:                                 26
  undirected arcs:                     3
  directed arcs:                       23
average markov blanket size:          6.00
average neighbourhood size:           4.73
average branching factor:             2.09

learning algorithm:                   Grow-Shrink
conditional independence test:         Pearson's Linear Correlation
alpha threshold:                      0.01
tests used in the learning procedure: 2029
optimized:                            TRUE
```

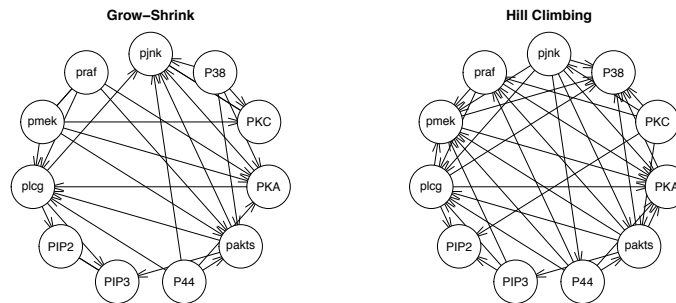
Analysis of Protein Flow Cytometry using bnlearn

```
> dag2
Bayesian network learned via Score-based methods

model:
  [PKC|pjnk|PKC] [P44|pjnk] [pakts|P44:PKC:pjnk] [praf|P44:pakts:PKC] [PIP3|pakts
  [plcg|praf:PIP3:P44:pakts:pjnk] [pmek|praf:plcg:PIP3:P44:pakts:pjnk]
  [PIP2|plcg:PIP3:PKC] [PKA|praf:pmek:plcg:P44:pakts:pjnk]
  [P38|pmek:plcg:pakts:PKA:PKC:pjnk]
nodes:                                11
arcs:                                 35
  undirected arcs:                     0
  directed arcs:                       35
average markov blanket size:          8.00
average neighbourhood size:           6.36
average branching factor:             3.18

learning algorithm:                   Hill-Climbing
score:                                Bayesian Information Criterion (Gaussian)
penalization coefficient:              4.459057
tests used in the learning procedure: 505
optimized:                            TRUE
```

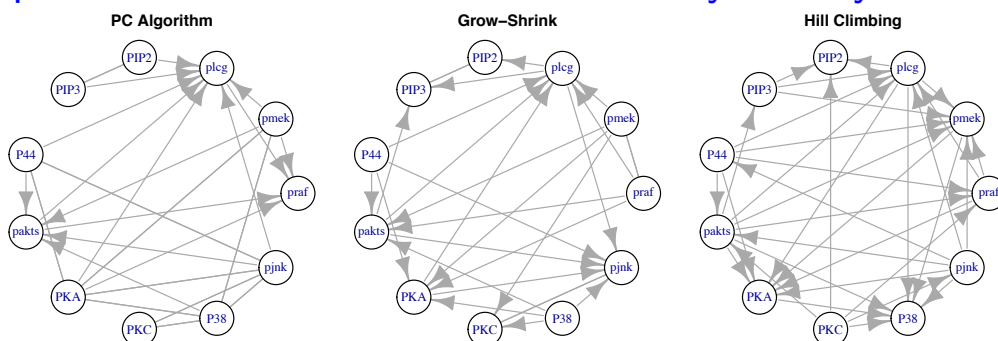

Analysis of Protein Flow Cytometry using bnlearn



The two graphs are quite different

```
> compare(dag1,dag3)
$tp
[1] 9
$fp
[1] 26
$fn
[1] 17
```

Comparison of Results for Protein Flow Cytometry Data



- The estimated graphs are quite different
- The constrained-based methods seem to have more similarities (at least in terms of structure)
- The estimate from HC has more edges; we can change e.g. the score, but cannot directly control the sparsity

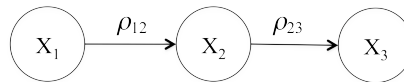
Penalized Likelihood Estimation of DAGs

- Causal relationships (and probability distributions) on DAGs can be represented using **structural equation models**

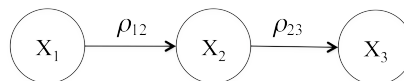
$$X_i = f_i(\text{pa}_i, \gamma_i), \quad i = 1, \dots, p$$

- And, for Gaussian random variables, we can write

$$X_i = \sum_{j \in \text{pa}_i} \rho_{ji} X_j + \gamma_i, \quad i = 1, \dots, p$$



Penalized Likelihood Estimation of DAGs



$$X_1 = \gamma_1$$

$$X_2 = \rho_{12}X_1 + \gamma_2 = \rho_{12}\gamma_1 + \gamma_2$$

$$X_3 = \rho_{23}X_2 + \gamma_3 = \rho_{23}\rho_{12}\gamma_1 + \rho_{23}\gamma_2 + \gamma_3$$

Thus $X = \Lambda\gamma$ where

$$\Lambda = \begin{pmatrix} 1 & 0 & 0 \\ \rho_{12} & 1 & 0 \\ \rho_{12}\rho_{23} & \rho_{23} & 1 \end{pmatrix}$$

Penalized Likelihood Estimation of DAGs

- It turns out that $\Lambda = (I - A)^{-1}$, where A is the weighted adjacency matrix of the DAG¹
- Thus, for Gaussian random variables, **if we know the ordering of the variables** (which is a BIG assumption!)

after some math...

we can estimate the adjacency matrix of DAGs, by minimizing the log-likelihood as a function of A :

$$\hat{A} = \arg \min_{A \in \mathcal{A}} \left\{ \text{tr}[(I - A)^T(I - A)S] \right\}$$

¹Shojaie & Michailidis (2010)

Penalized Likelihood Estimation of DAGs

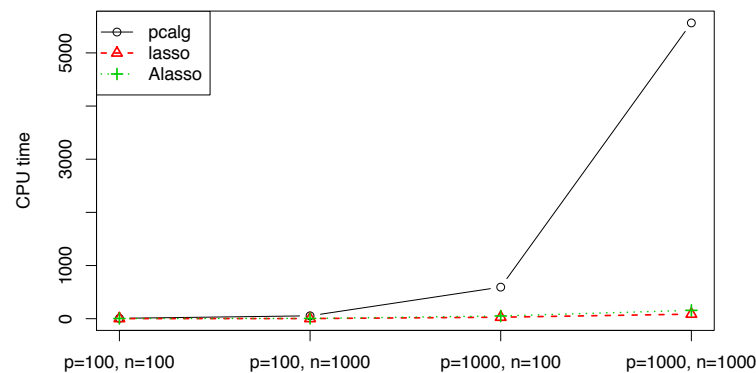
- In high dimensions, we can solve a penalized version of this problem, e.g. by adding a **lasso penalty** $\lambda \sum_{i < j} |A_{ij}|$
- It turns out that, the problem can be reformulated as $(p - 1)$ **lasso problems**, where we **regress each variable, on those appearing earlier in the ordering**:

$$\hat{A}_{k,1:k-1} = \arg \min_{\theta \in \mathbb{R}^{k-1}} \left\{ n^{-1} \|X_{1:k-1}\theta - X_{\cdot,k}\|_2^2 + \lambda \sum_{j=1}^{k-1} |\theta_j| w_j \right\}$$

- As in glasso, λ controls the sparsity; $\lambda = \frac{2}{\sqrt{n}} Z_{\alpha/(2p^2)}$ controls a false positive probability at level α

Computational Complexity

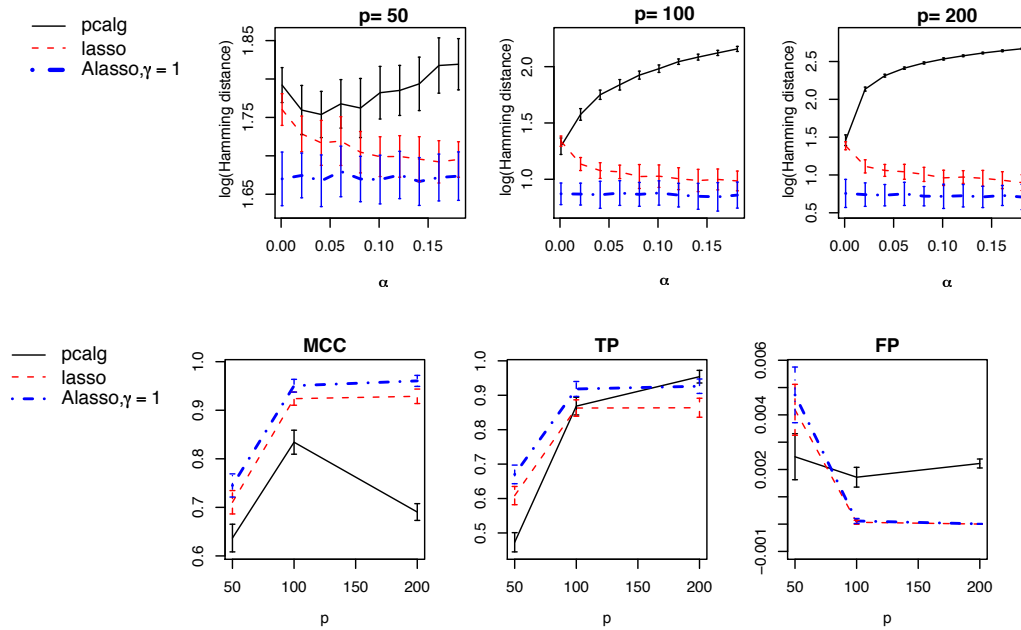
- Compared to pcalg, this method runs much faster: $\sim np^2$ operations vs $\sim p^q$ (q is the max degree)
- Can be easily implemented in R as $p - 1$ regressions using glmnet. A more general version is available in the spacejam package, which also includes estimation for non-Gaussian data



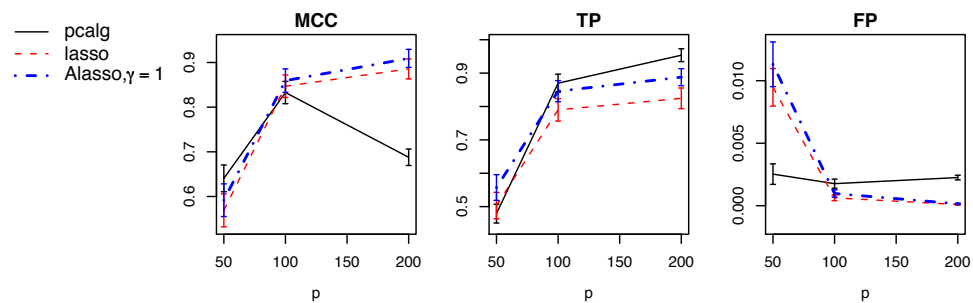
Simulations

- Settings:
 - $p = 50, 100, 200$
 - $n = 100$
 - Total number of edges in the network = n
 - 100 repetitions
- Performance Criteria
 1. Matthew's Correlation Coefficient (**MCC**): ranges between -1 (worst fit) and 1 (best fit), similar to F_1
 2. Structural Hamming Distance (**SHD**): sum of false positive and false negatives
 3. True positive and false positive rates
- Tuning parameter for both PC-Algorithm and penalized likelihood method based on false positive error α

Gaussian Observations

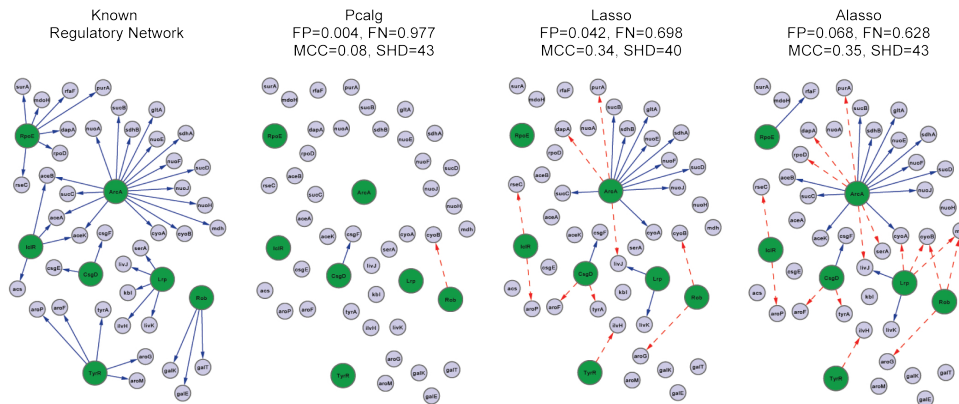


Random Ordering of Variables



Regulatory Network of E-Coli

- Regulatory network of E-coli with $p = 49$ genes (7 TFs)
- Want to identify regulatory interactions among TFs and regulated genes



DAGs for Time Series Data

Time Series Data: A setting where ordering is known

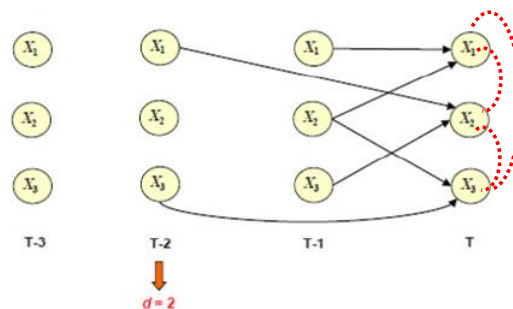
- p -dimensional, discrete time, stationary process

$$X^t = \{X_1^t, \dots, X_p^t\}$$

$$X^t = A_1 X^{t-1} + \dots + A_d X^{t-d} + \epsilon^t, \quad \epsilon^t \stackrel{i.i.d.}{\sim} N(0, \Sigma_\epsilon) \quad (1)$$

- $A_1, \dots, A_d : p \times p$ transition matrices (solid, directed edges)
- Σ_ϵ^{-1} : contemporaneous dependence (dotted, undirected edges)

DAGs for Time Series Data

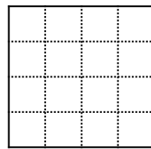



Network Granger causality (NGC)

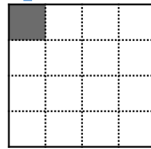
Network Granger Causality with VARs

- ▶ X_1, \dots, X_p : time series for p variables
- ▶ $X^t = (X_1^t, \dots, X_p^t)'$: realizations at time t
- ▶ **VAR** model for NGC:

$$X^T = A^1 X^{T-1} + \dots + A^d X^{T-d} + \varepsilon^T$$



 A_{11} : Autoregressive effect of X_1 on itself



 A : Autoregressive effect of X on X

NGC Estimation

Let Y be the (stacked) vector of current time points; Z be the design matrix based on previous time points; and β be

Assuming A_t are sparse, and d is known

- ▶ ℓ_1 -penalized least squares (**ℓ_1 -LS**)

$$\arg \min_{\beta \in \mathbb{R}^{dp^2}} \|Y - Z\beta\|^2 + \lambda \|\beta\|_1$$

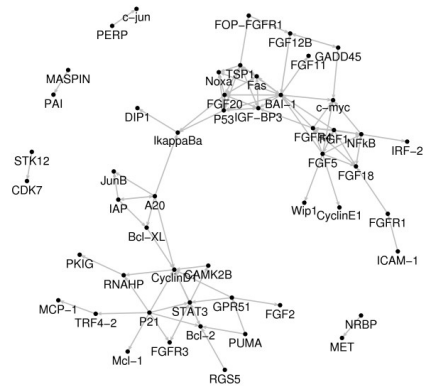
- ▶ ℓ_1 -penalized log-likelihood (**ℓ_1 -LL**) — assuming Σ_ϵ^{-1} is sparse²

$$\arg \min_{\beta \in \mathbb{R}^{dp^2}} (Y - Z\beta)' (\Sigma_\epsilon^{-1} \otimes I) (Y - Z\beta) + \lambda \|\beta\|_1$$

²Lin & Michailidis (2017)

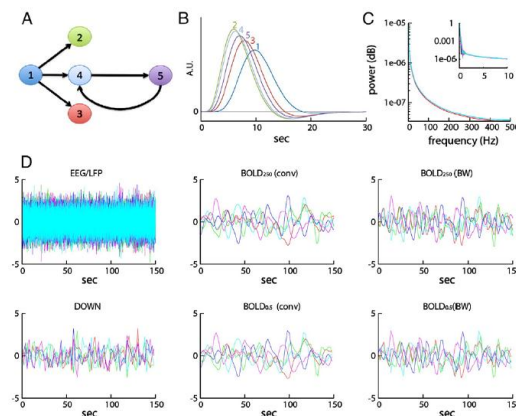
Applications — Functional Genomics

- Identifying regulatory mechanisms using transition patterns in time course expression data
- HeLa gene expression regulatory network (Fujita et al, 2007)



Applications — Neuroscience

- Connectivity among brain regions from time-course fMRI data
- Connectivity of VAR generative model (Seth et al, 2013)



Extensions

- ▶ Panel VAR Modeling (common in functional genomics and neuroscience)³
- ▶ Incorporating external information using group lasso penalties, etc⁴
- ▶ Dealing with non-stationarity (paucity of long stationary time series — T small)⁵
- ▶ Accounting for non-linearity
- ▶ ...

³S & Michailidis (2010); S, Basu & Michailidis (2012)

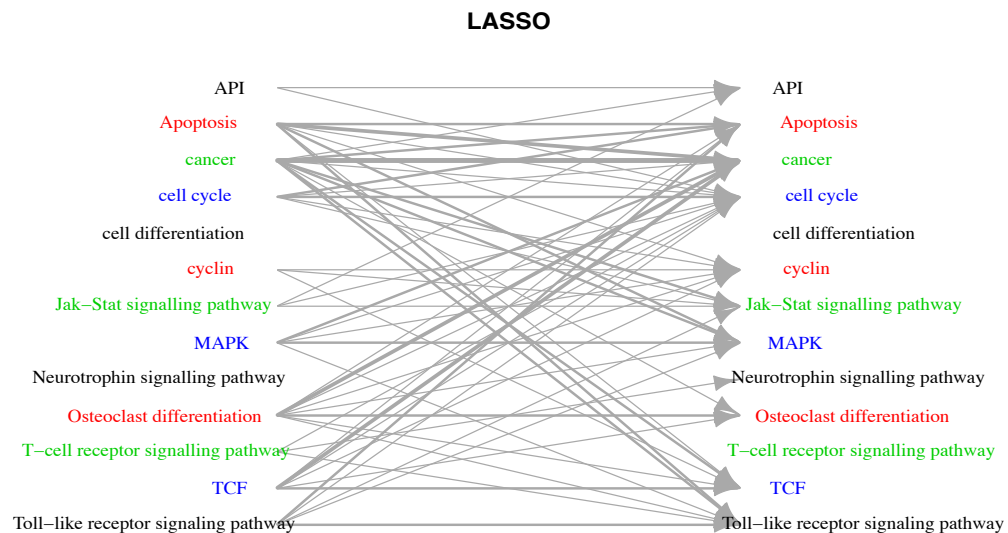
⁴Basu, S & Michailidis (2014)

⁵Safikhani & S (2020)

Example: T-cell Activation Data

- ▶ Data from Rangel et al (2004) on T-cell activation — less insight and biological knowledge regarding pathways
- ▶ $p = 58$ genes, $n = 44$ samples, and $T = 10$ time points — the first 5 time points (0, 2, 4, 6 and 8 hours) were used on a subset of 38 genes for which pathway information avail
- ▶ Goal is to estimate regulatory interactions

Estimated Network Structure



Summary

- Estimation of DAGs from observational data is both conceptually and computationally difficult
- Constraint-based & search-based algorithms — slow in high dim
- May not be able to distinguish DAGs from observational data (Markov equivalence)
- Efficient penalized likelihood methods can estimate DAGs **if the ordering is known**
- Important case is time series data, but **Granger causality \neq causality!**
- Efficient implementations in R available for most methods