Analysis and inference of gene networks from genomic data



Jean-Philippe Vert Ecole des Mines de Paris Computational Biology group Jean-Philippe.Vert@mines.org

"Complex Stochastic Systems in Biology and Medicine" workshop, Munich, Germany, October 7th, 2004.

Thanks

- Yoshihiro Yamanishi (Kyoto University)
- Computational biology at the Ecole des Mines

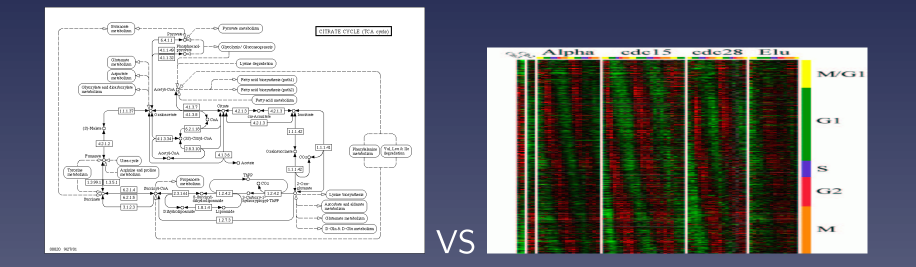


Motivations

- Many heterogeneous data about genes : sequences, expression, evolution, structures, etc...
- More and more data between genes: interactome, pathways, regulation etc...
- Goal: propose a formalism and algorithms to compare these data, and to infer gene networks from high-throughput genomic data.

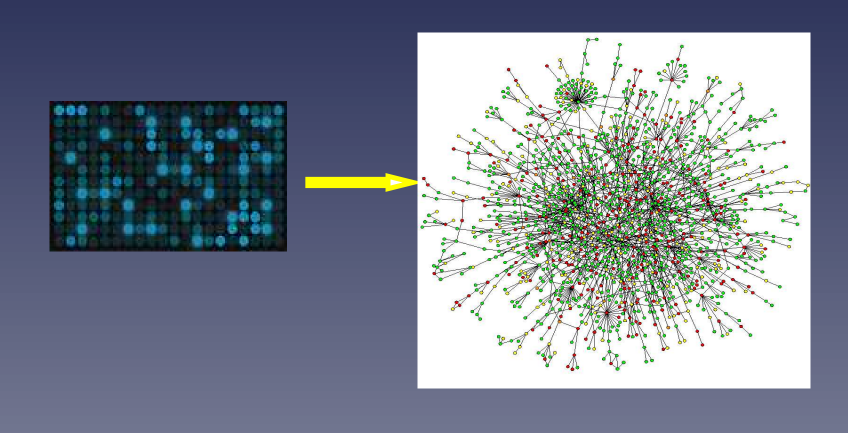
Example 1:

Comparing gene expression and pathway databases



Detect active pathways? Denoise expression data? Denoise pathway database? Find new pathways? Are there "correlations"?

Example 2: Gene network inference



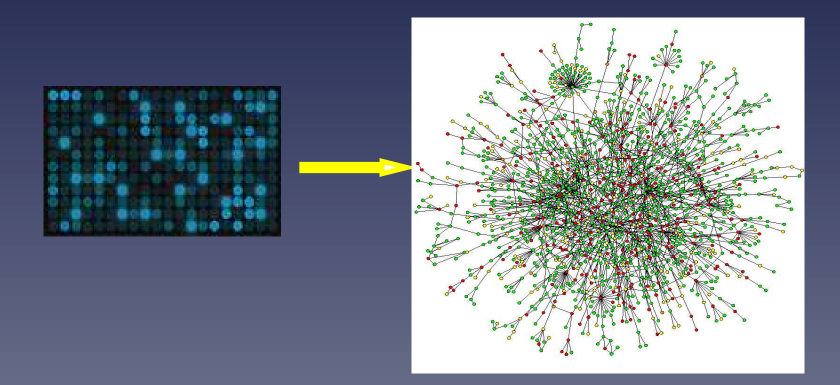
Outline

- A direct approach to network inference
- Supervised network inference
- Extraction of pathway activity
- Learning from several heterogeneous data

Part 1

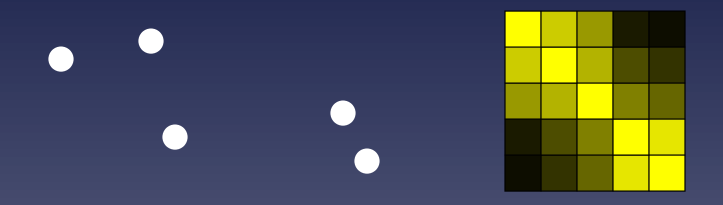
A direct approach to network inference

The problem



Related approaches

- Bayesian nets for regulatory networks (Friedman et al. 2000)
- Boolean networks (Akutsu, 2000)
- Joint graph method (Marcotte et al, 1999)



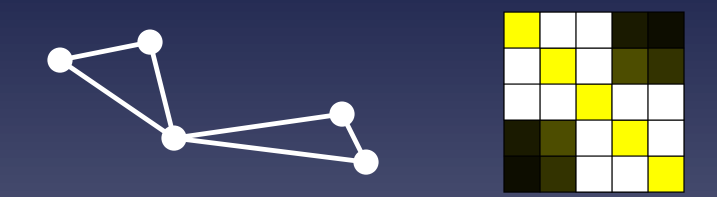


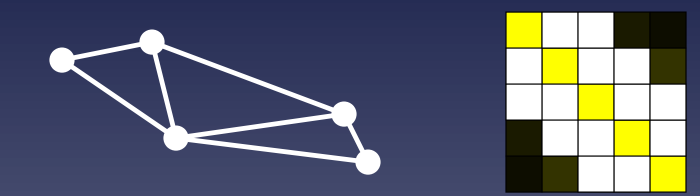


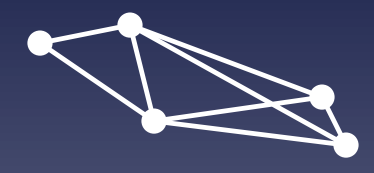


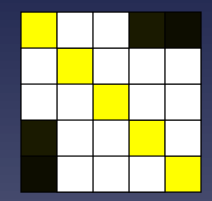


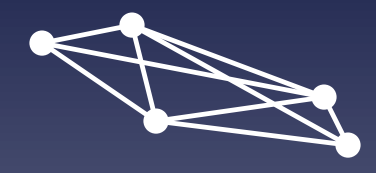


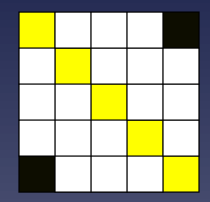


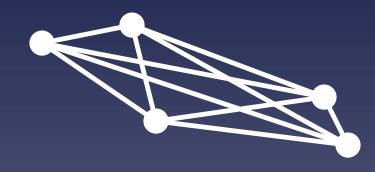


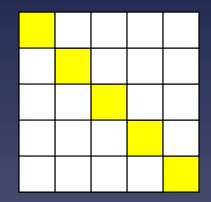


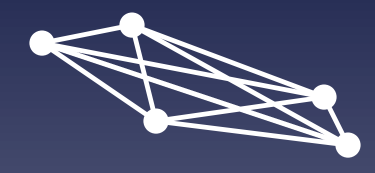


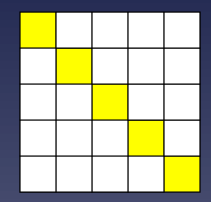


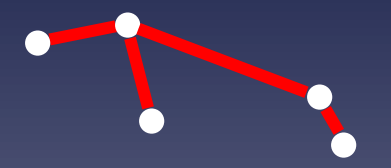


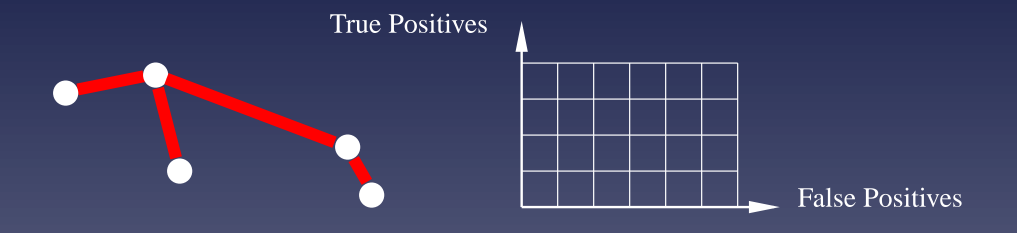


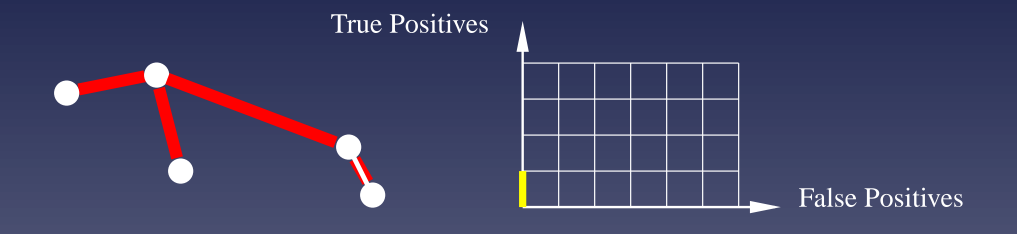


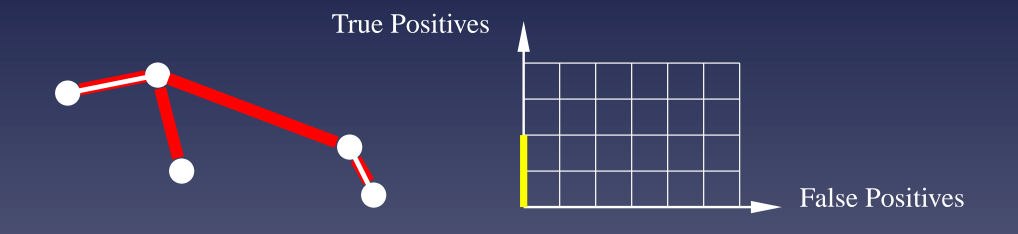


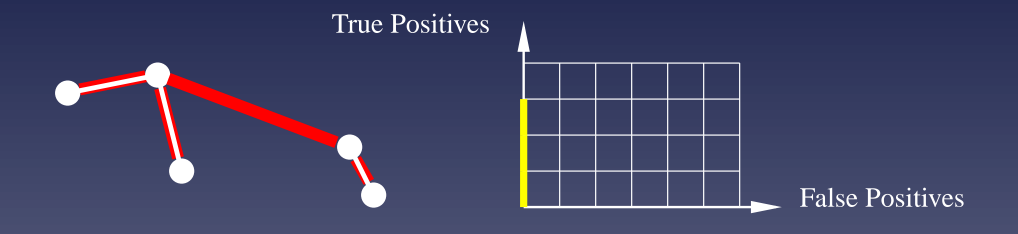


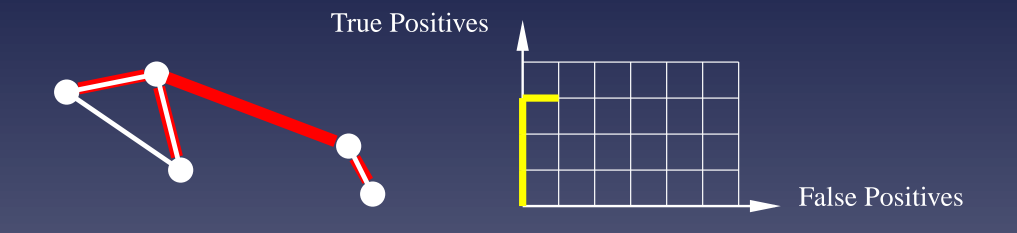


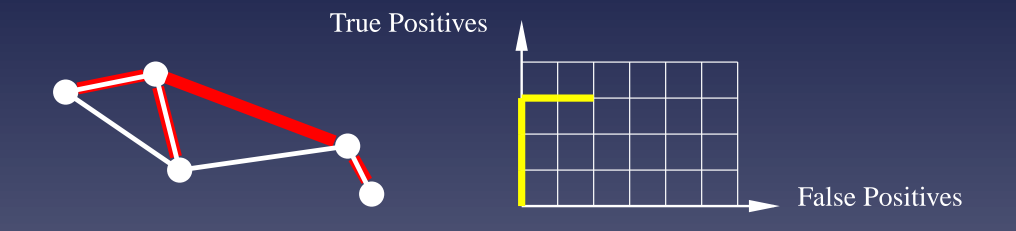


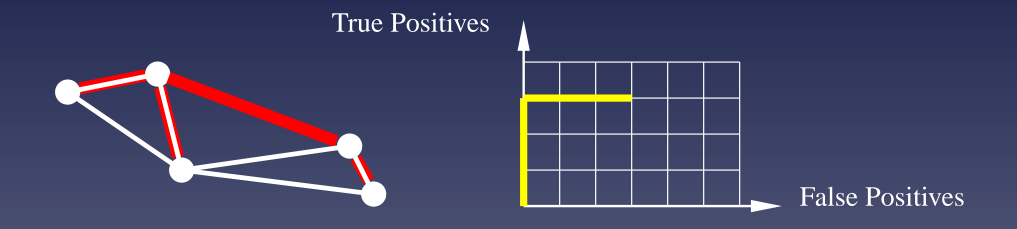


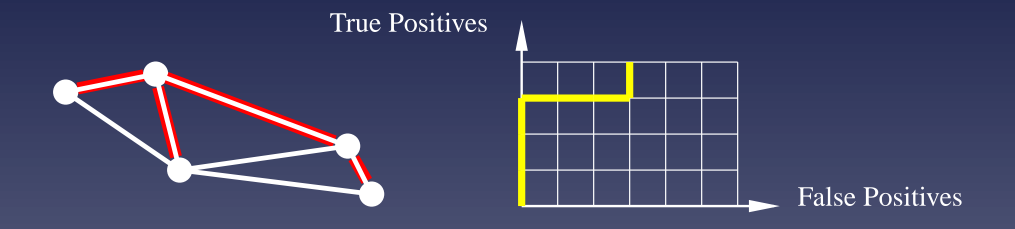


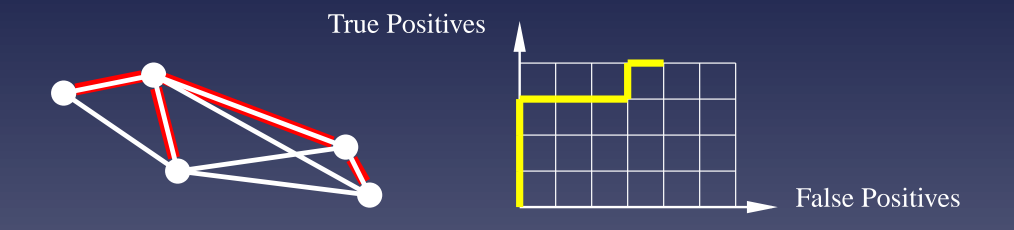


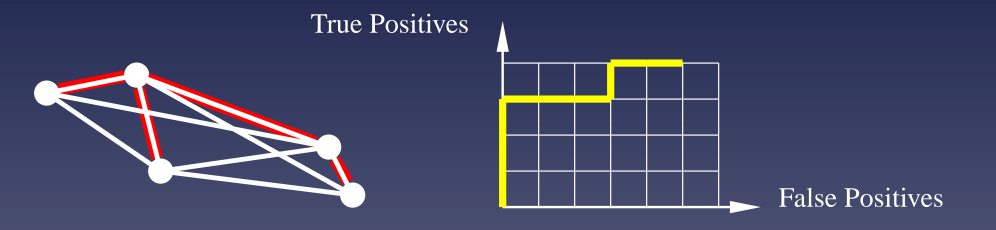


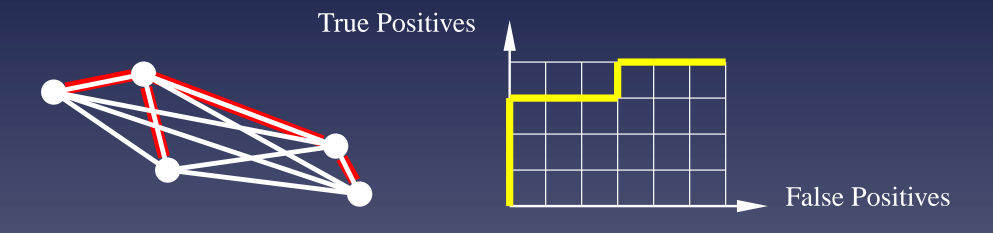






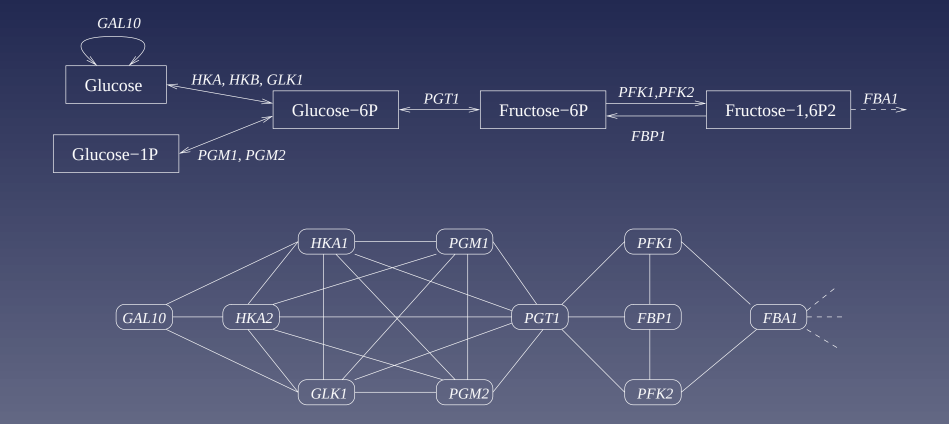






ROC = 21/24 = 87,5%

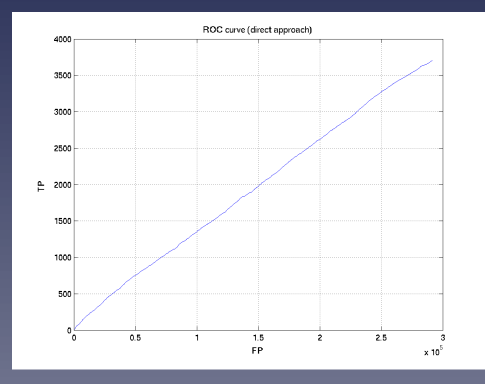
Application: the metabolic gene network



Link two genes when they can catalyze two successive reactions

Evaluation of the direct approach

The metabolic network of the yeast involves 769 genes. Each gene is represented by 157 expression measurements. (ROC=0.52)



Shortcuts of the direct approach

• What similarity measure between profiles should be use?

Shortcuts of the direct approach

- What similarity measure between profiles should be use?
- Which network are we expecting to recover?

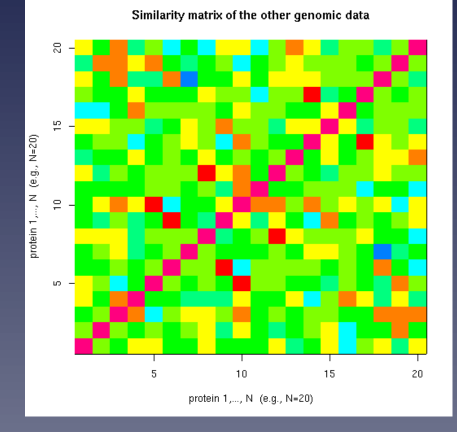
Shortcuts of the direct approach

- What similarity measure between profiles should be use?
- Which network are we expecting to recover?
- How to use prior knowledge about the network to be recovered?

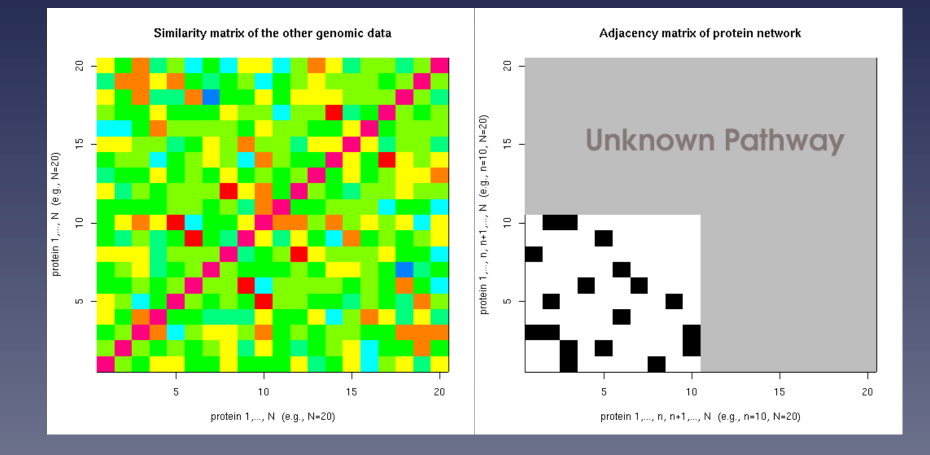
Part 2

Supervised network inference

The supervised gene inference problem



The supervised gene inference problem



The idea in a nutshell

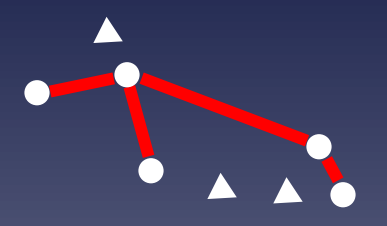
Use the known network to "learn" a more relevant measure of similarity

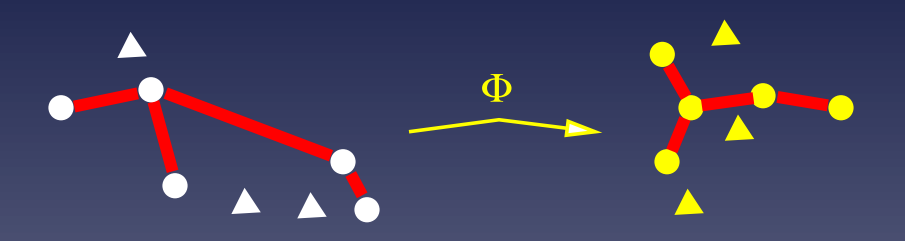
The idea in a nutshell

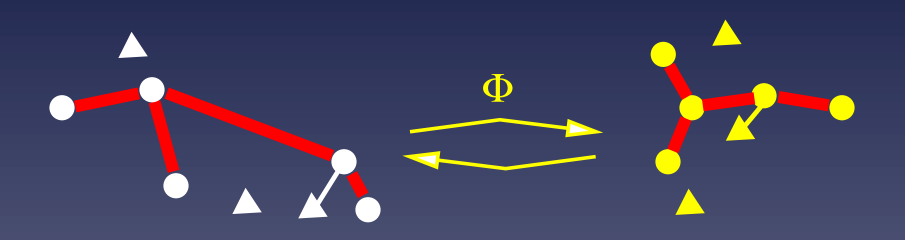
- Use the known network to "learn" a more relevant measure of similarity
- For example, map the genes expression profiles to a different space, where the natural distance better fits the known network

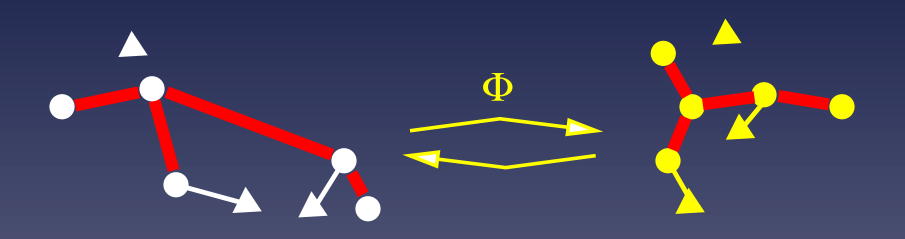
The idea in a nutshell

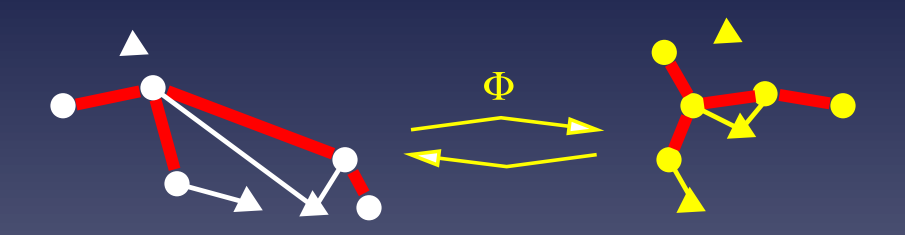
- Use the known network to "learn" a more relevant measure of similarity
- For example, map the genes expression profiles to a different space, where the natural distance better fits the known network
- Then apply the direct strategy in the second space

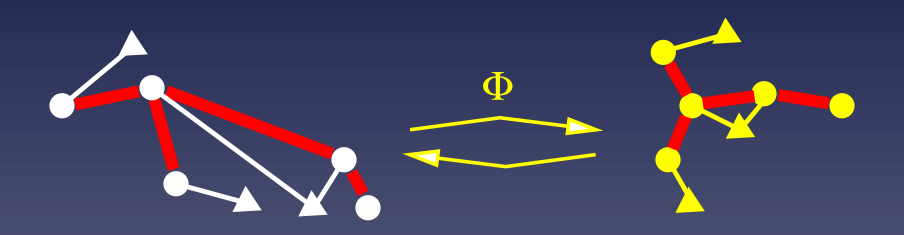


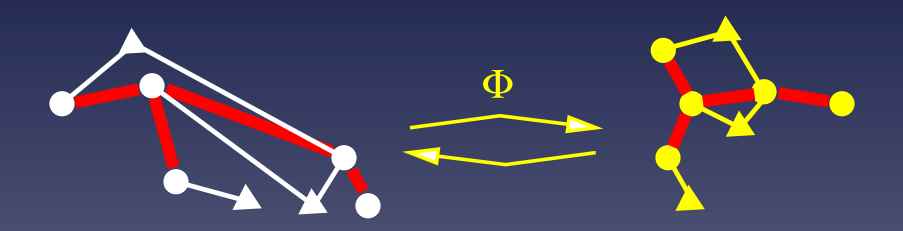












Learning the mapping Φ

• Let $x \in \mathbb{R}^p$ be an expression profile

Learning the mapping Φ

• Let $x \in \mathbb{R}^p$ be an expression profile

Let us consider linear mappings:

 $\Phi(x) = (f_1(x), \dots, f_d(x))' \in \mathbb{R}^d$

made of linear features $f_i(x) = w_i^\top x$

Learning the mapping Φ

• Let $x \in \mathbb{R}^p$ be an expression profile

• Let us consider linear mappings:

$$\Phi(x) = (f_1(x), \dots, f_d(x))' \in \mathbb{R}^d$$

made of linear features $f_i(x) = w_i^{\top} x$

 A feature f : ℝ^p → ℝ is "good" if connected genes in the known network have similar value.

"Good" features

• A "good" feature $f(x) = w^{\top}x$ should minimize:

$$R(f) = \frac{\sum_{i \sim j} \left(f(x_i) - f(x_j) \right)^2}{\sum_{i=1}^n f(x_i)^2},$$

• Regularisation: for statistical reasons, it is safer to minimize:

$$\min_{f(x)=w^{\top}x} \frac{\sum_{i\sim j} \left(f(x_i) - f(x_j)\right)^2 + \lambda ||w||^2}{\sum_{i=1}^n f(x_i)^2},$$

Influence of λ

$ightarrow \overline{\lambda ightarrow +\infty}$: PCA

★ Useful for noisy, high-dimensional data.

 Used in spectral clustering. The graph does not play any role (unsupervised)

• $\lambda \rightarrow 0$: second smallest eigenvector of the graph

- Useful to embed the graph in a Euclidean space (used in graph partitioning)
- Sensitive to noise. Mapping of points outside of the graph unstable (overfitting)

Extracting successive features

• Successive features to form Φ can be obtained by:

$$w_{i} = \operatorname*{arg\,min}_{w \perp \{w_{1}, \dots, w_{i-1}\}, \text{var}(f_{w})=1} \left\{ \sum_{i \sim j} \left(f_{w}(x_{i}) - f_{w}(x_{j}) \right)^{2} + \lambda ||w||^{2} \right\}$$

Extracting successive features

• Successive features to form Φ can be obtained by:

$$w_{i} = \operatorname*{arg\,min}_{w \perp \{w_{1}, \dots, w_{i-1}\}, \text{var}(f_{w})=1} \left\{ \sum_{i \sim j} \left(f_{w}(x_{i}) - f_{w}(x_{j}) \right)^{2} + \lambda ||w||^{2} \right\}$$

• Each features satisfies $w = \sum_i \alpha_i x_i$ (Representer theorem)

Solving the problem

• The problem can then be rewritten:

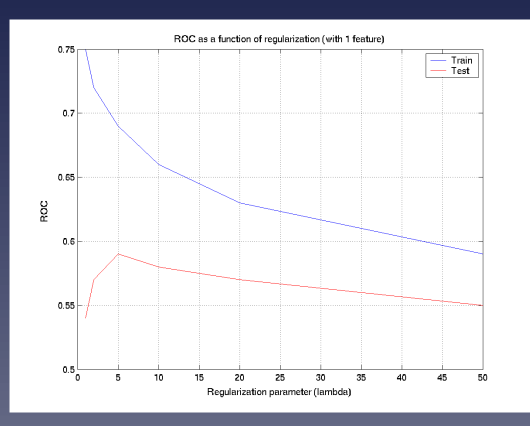
$$\alpha_{i} = \operatorname*{arg\,min}_{\alpha \in \mathbb{R}^{n}, \alpha K_{V}\alpha_{1} = \ldots = \alpha K_{V}\alpha_{i-1}} \left\{ \frac{\alpha^{\top} K_{V}LK_{V}\alpha + \lambda \alpha^{\top} K_{V}\alpha}{\alpha^{\top} K_{V}^{2}\alpha} \right\}$$

where K_V is the centered $n \times n$ matrix of inner products and L is the Laplacian of the graph

• It is equivalent to solving the generalized eigenvalue problem:

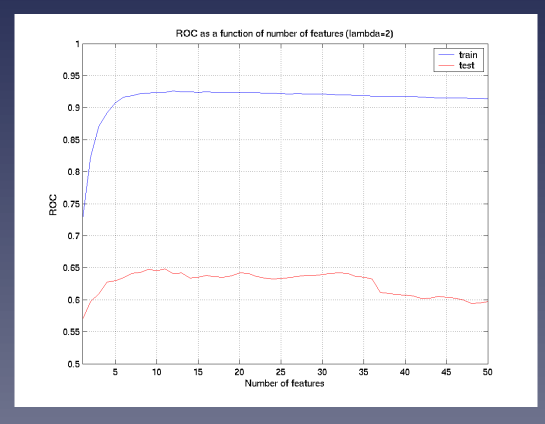
 $(LK_V + \lambda I)\alpha = \mu K_V \alpha.$

Evaluation of the supervised approach: effect of λ



Metabolic network, 10-fold cross-validation, 1 feature

Evaluation of the supervised approach: number of features ($\lambda = 2$)

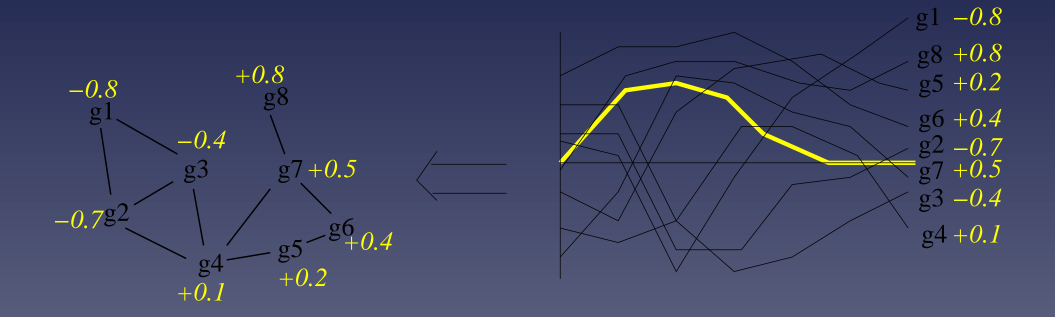


Part 3

Extraction of pathway activity

The idea

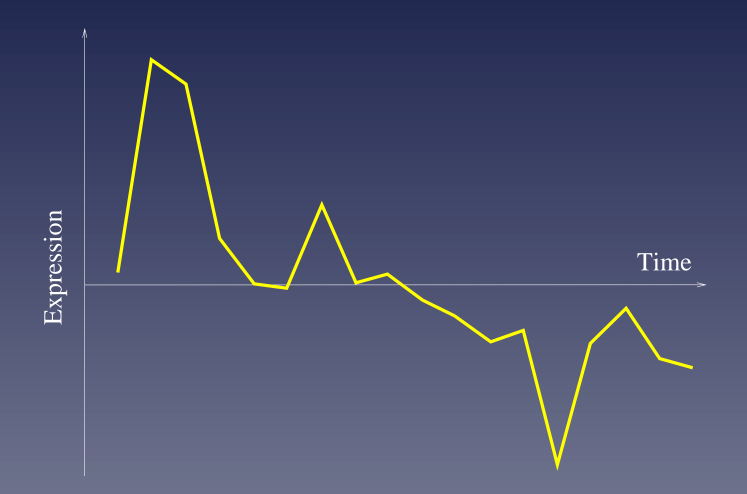
- The previous approach is a way to extract features from gene expression data: $f(x) = w^{\top}x$.
- These features are smooth on the graph: connected nodes tend to have similar values
- This is way to detect "correlations" between gene expression data and metabolic network : typical activity patterns of typical pathways



Experiment

- Gene network: two genes are linked if the catalyze successive reactions in the KEGG database (669 yeast genes)
- Expression profiles: 18 time series measures for the 6,000 genes of yeast, during two cell cycles

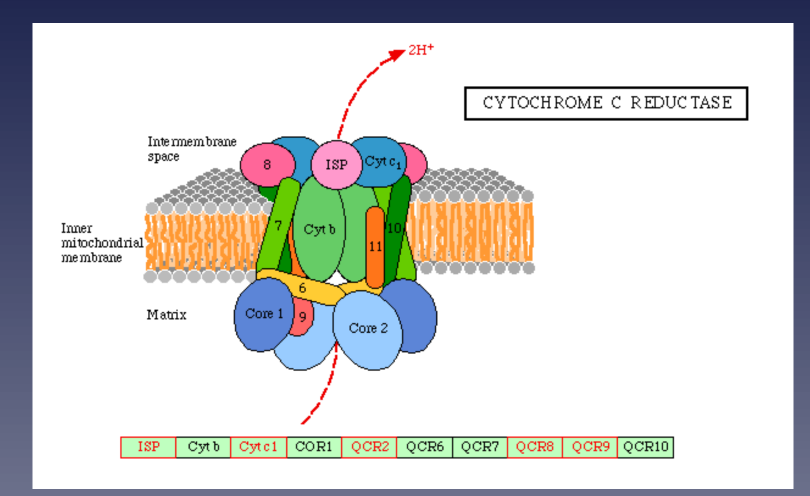
First pattern of expression



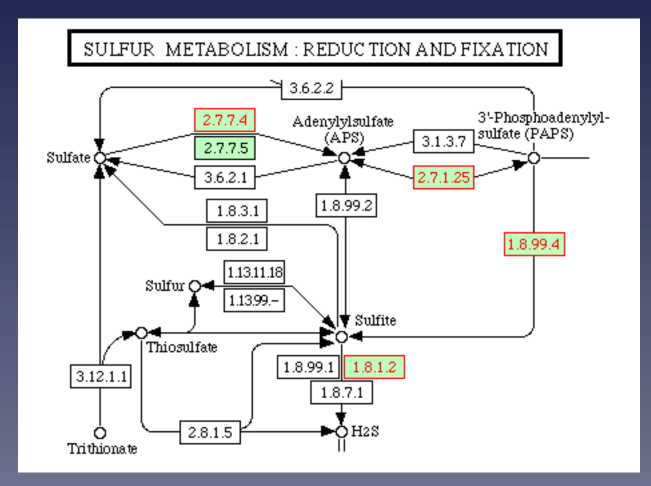
Related metabolic pathways

- 50 genes with highest $s_2 s_1$ belong to:
- Oxidative phosphorylation (10 genes)
- Citrate cycle (7)
- Purine metabolism (6)
- Glycerolipid metabolism (6)
- Sulfur metabolism (5), etc...

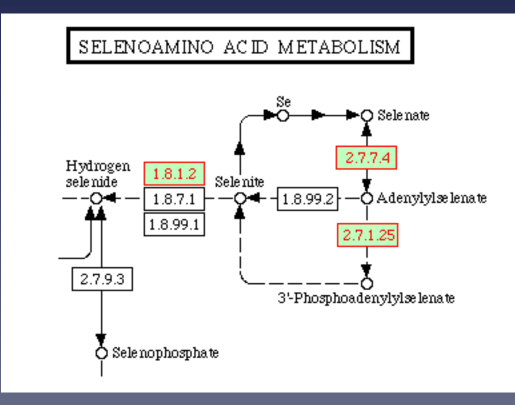
Related genes



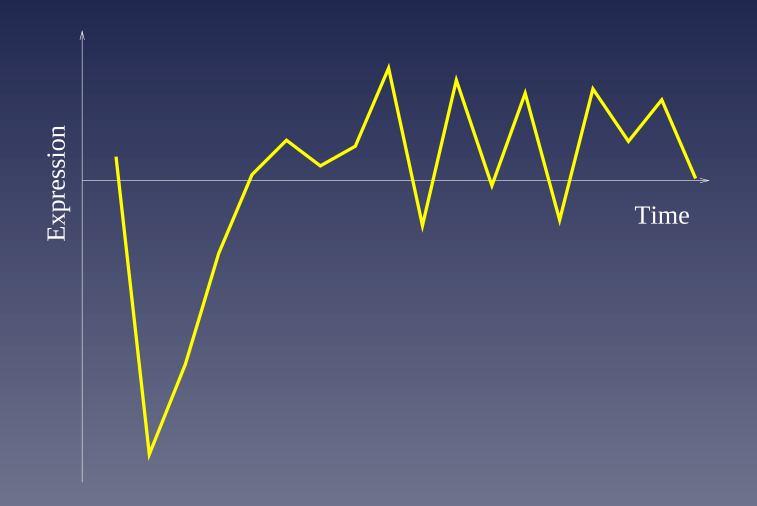
Related genes



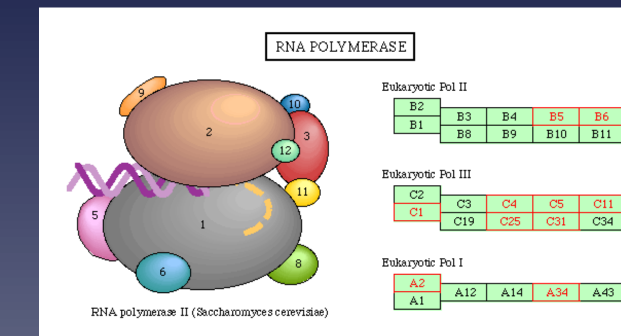
Related genes



Opposite pattern



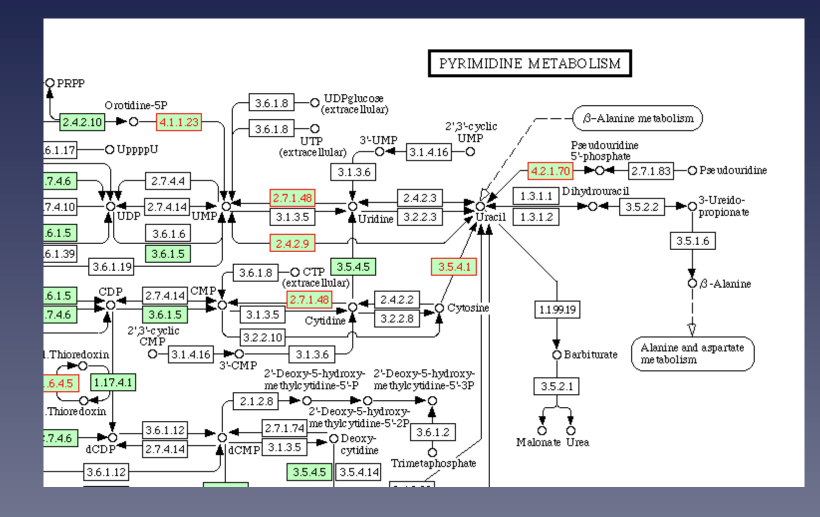
- RNA polymerase (11 genes)
- Pyrimidine metabolism (10)
- Aminoacyl-tRNA biosynthesis (7)
- Urea cycle and metabolism of amino groups (3)
- Oxidative phosphorlation (3)
- ATP synthesis(3) , etc...

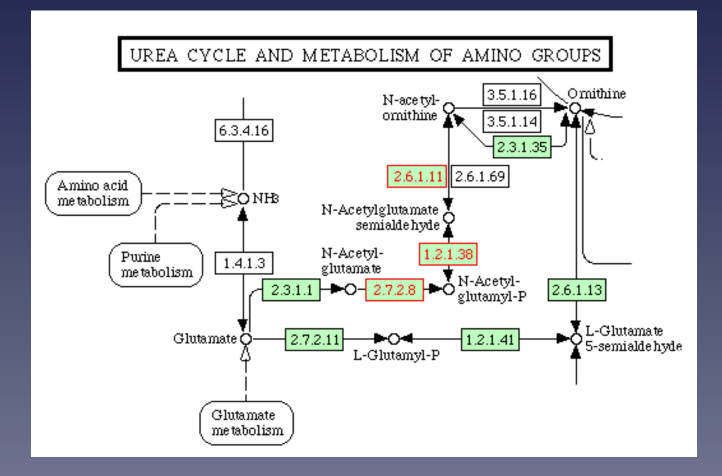


B7

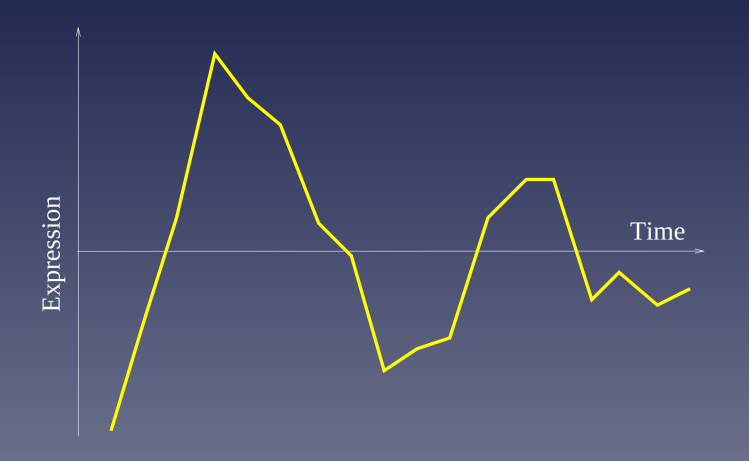
B12

A49





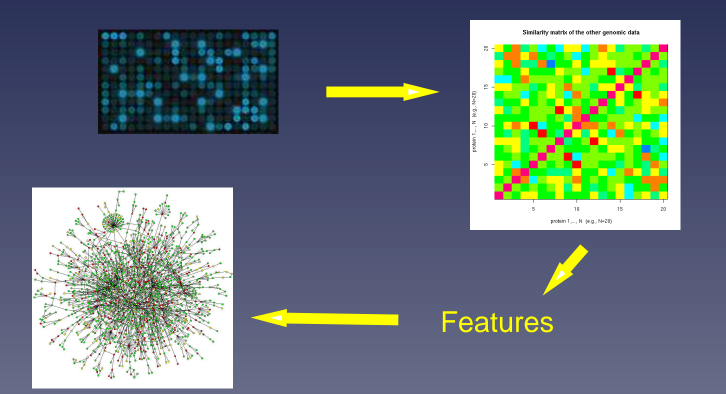
Second pattern



Part 4

Learning from several heterogeneous data

Summary of the process



The "kernel trick"

• The matrix of similarity is $K_{i,j} = x_i^{\top} x_j$

The "kernel trick"

- The matrix of similarity is $K_{i,j} = x_i^{\top} x_j$
- However, more general measures are allowed: they simply must be symetric positive definite

The "kernel trick"

- The matrix of similarity is $K_{i,j} = x_i^{\top} x_j$
- However, more general measures are allowed: they simply must be symetric positive definite
- This enables nonlinear features, as well as features from other types of data, as soon as a symetric p.d. function K(x, y) is defined

Kernels

Several kernels have been developed recently:

- for phylogenetic profiles (JPV. 2004)
- for gene sequences (Leslie et al. 2003, Saigo et al. 2004, ...)

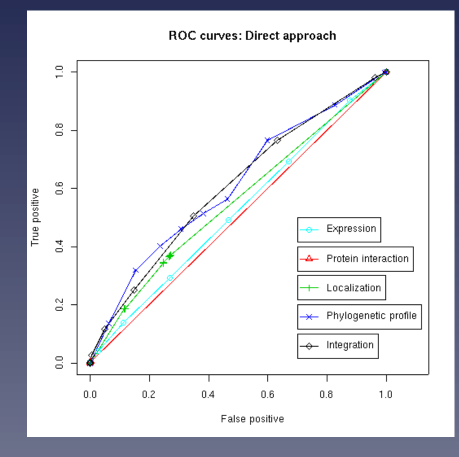
• for nodes in a network (Kondor et al. 2000)

Learning from heterogeneous data

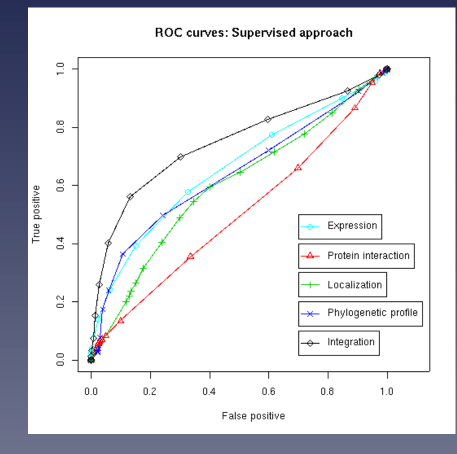
- Suppose several data are available about the genes, e.g., expression, localization, struture, predicted interaction etc...
- Each data can be represented by a positive definite similarity matrix K_1, \ldots, K_p called kernels
- Kernel can be combined by various operations, e.g., addition:

$$K = \sum_{i=1}^{p} K_i$$

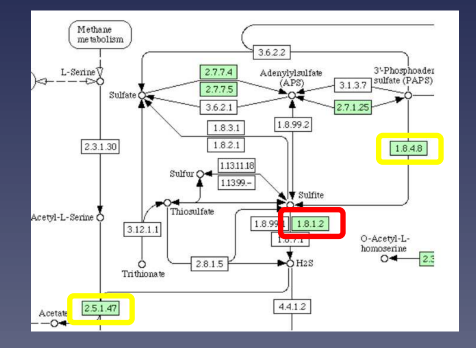
Learning from heterogeneous data (unsupervised)



Learning from heterogeneous data (supervised)



Application: missing enzyme prediction



The gene YJR137C was predicted in 09/2003 between EC : 1.8.4.8 and EC : 2.5.1.47. It was recently annotated as EC:1.8.1.2

Conclusion

Conclusion

- A new approach to feature extractions and supervised network inference, many possible variants and extensions
- Straightforward generalization to any network (e.g., interactome): the same data can be used to infer different networks
- Currently tested on characterization of tumor cells (with Institut Curie) and metabolism of P. falciparum (with Institut Pasteur).