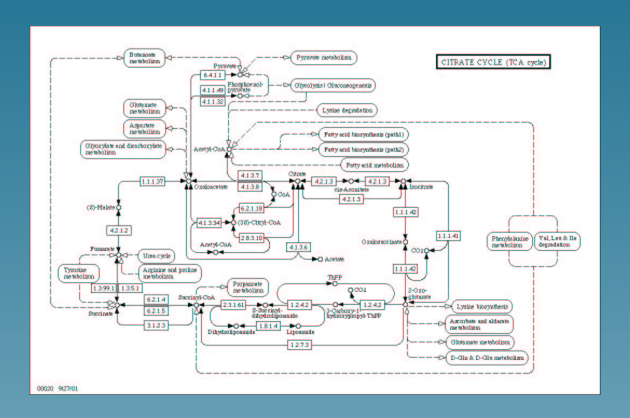
# Metabolic networks: Activity detection and Inference

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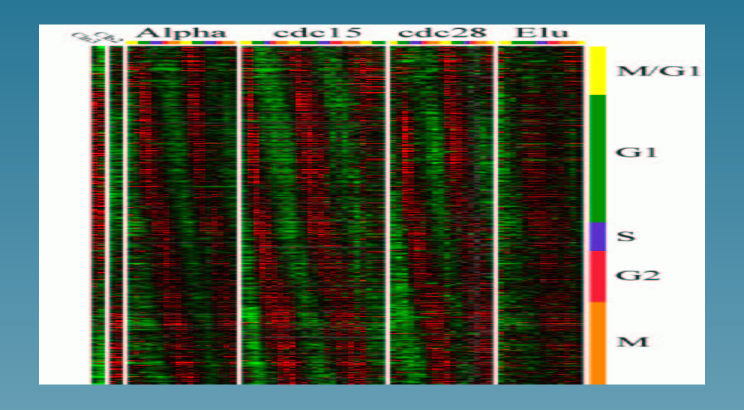
INRIA, May 5th, 2004.

#### Many metabolic pathways are known



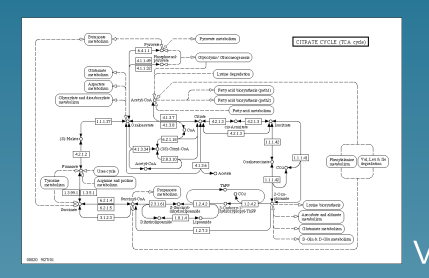
From http://www.genome.ad.jp/kegg/pathway

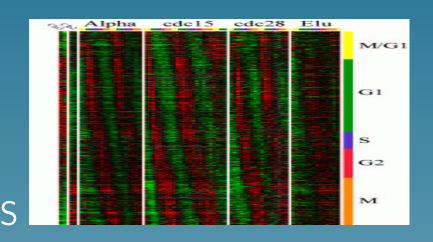
# Microarray technology monitors mRNA quantity



(From Spellman et al., 1998)

#### Comparing gene expression and pathway databases





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

#### **Overview**

- 1. Feature extractions from expression data only
- 2. Detecting correlations with the metabolic databse
- 3. Experiments
- 4. Inferring new pathways

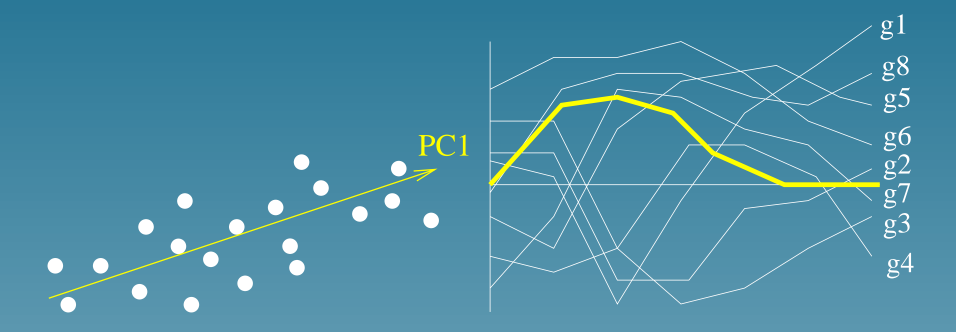
#### Part 1

# Feature extraction from expression data only

#### **Motivation**

- Pathways and biological events involve the coordinated action of several genes
- Co-regulation is an important way to coordinate the action of several genes
- Systematic variations in the set of gene expression profiles might be an indicator of an underlying biological phenomenon

## **Using microarray only**



PCA finds the directions (*profiles*) explaining the largest amount of variations among expression profiles.

#### **PCA** formulation

- Let  $f_v(i)$  be the projection of the *i*-th profile onto v.
- The amount of variation captured by  $f_v$  is:

$$h_1(v) = \sum_{i=1}^{N} f_v(i)^2$$

PCA finds an orthonormal basis by solving successively:

$$\max_{v} h_1(v)$$

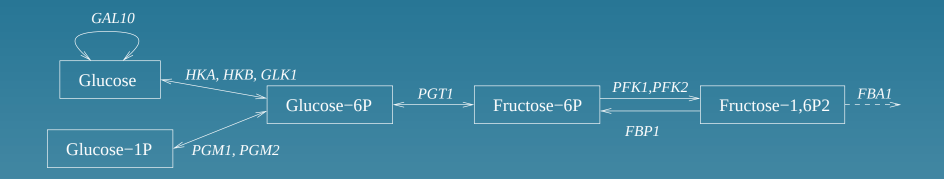
#### Part 2

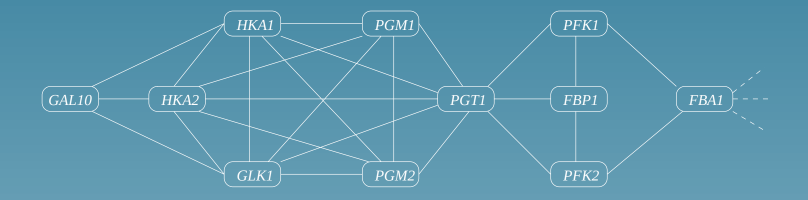
# Detecting correlations with the metabolic database

#### **Motivation**

- PCA is useful if there is a small number of strong signal
- In concrete applications, we observe a noisy superposition of many events
- Using a prior knowledge of metabolic networks can help denoising the information detected by PCA

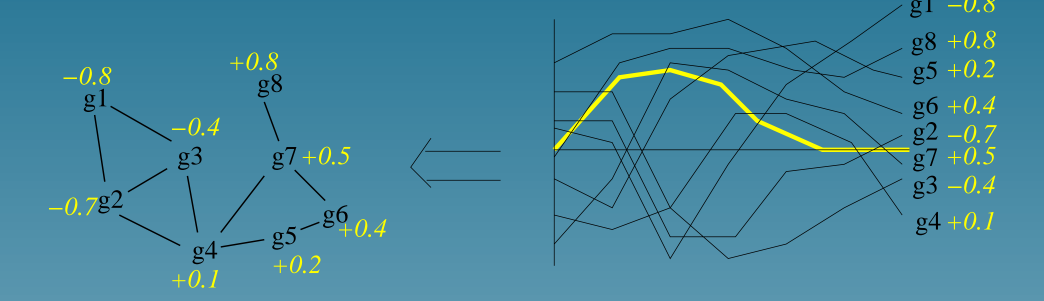
#### The metabolic gene network





Link two genes when they can catalyze two successive reactions

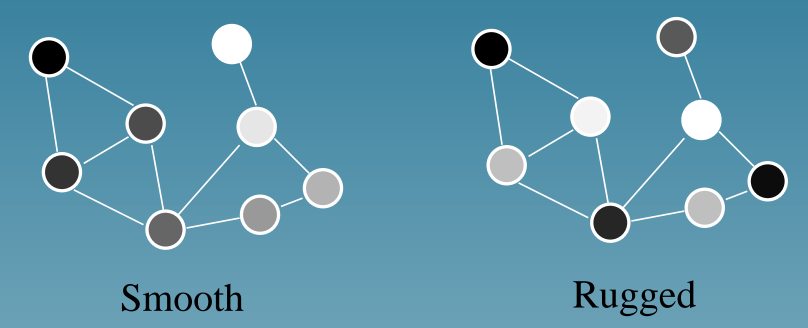
#### Mapping $f_v$ to the metabolic gene network



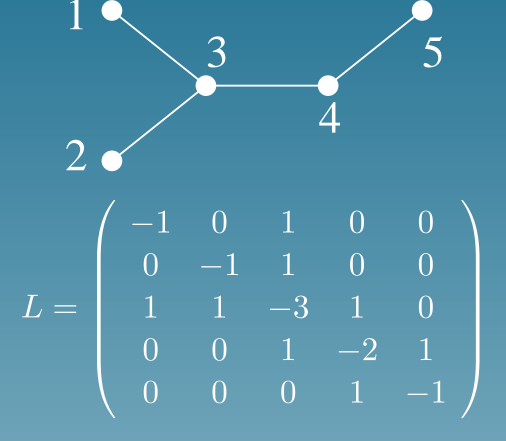
Does it look interesting or not?

#### Important hypothesis

If v is related to a metabolic activity, then  $f_v$  should vary "smoothly" on the graph



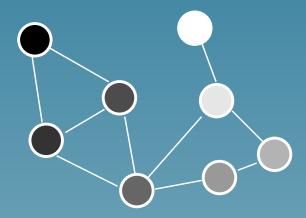
#### Graph Laplacian L = D - A



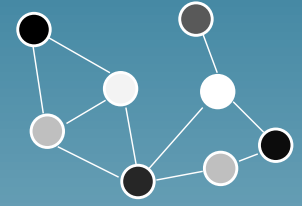
# **Smoothness quantification**

$$h_2(f) = \frac{f^{\top} f}{f^{\top} \exp(-\beta L) f}$$

is large when f is smooth



$$h(f) = 2.5$$



$$h(f) = 34.2$$

#### Where we are now...

For a candidate profile v,

- ullet  $h_1(f_v)$  is large when v captures a lot of natural variation among profiles
- $h_2(f_v)$  is large when  $f_v$  is smooth on the graph

Try to maximize both terms in the same time

#### **Problem reformulation**

Find a function  $f_v$  (and therefore a profile v) that solves:

$$\min_{v} \left\{ h_1(f_v)^{-1} + \lambda h_2(f_v)^{-1} \right\}$$

 $\lambda$  is a parameter that controls the trade-off.

## Solving the problem

ullet By the representer theorem, v can be expanded as:

$$v = \sum_{i=1}^{n} \alpha_i e(x_i).$$

#### Solving the problem (cont.)

• The problem can then be rewritten:

$$\min_{\alpha \in \mathbb{R}^n} \left\{ \alpha^{\top} K_0 K_2 K_0 \alpha + \lambda \alpha^{\top} K_0 \alpha \right\}$$

under the constraint  $\alpha^{\top} K_0^2 \alpha = 1$ , where:

- $\star K_2 = \exp(-\beta L)$  is the  $n \times n$  diffusion kernel
- $\star$   $K_0$  is the centered  $n \times n$  Gram matrix ( $[K_0]_{i,j} = e_i^{\top} e_j$ )

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- It is equivalent to solving the generalized eigenvalue problem:

$$(K_2K_0 + \lambda I)\alpha = \mu K_0\alpha.$$

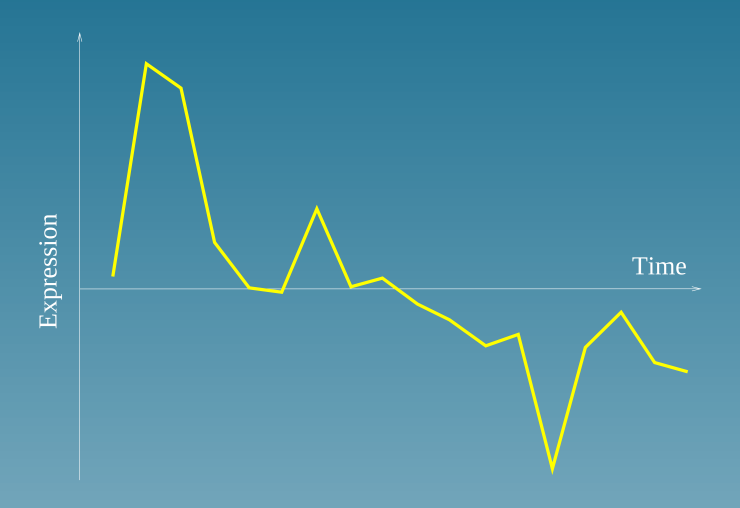
# Part 3

# Experiments

#### Data

- 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

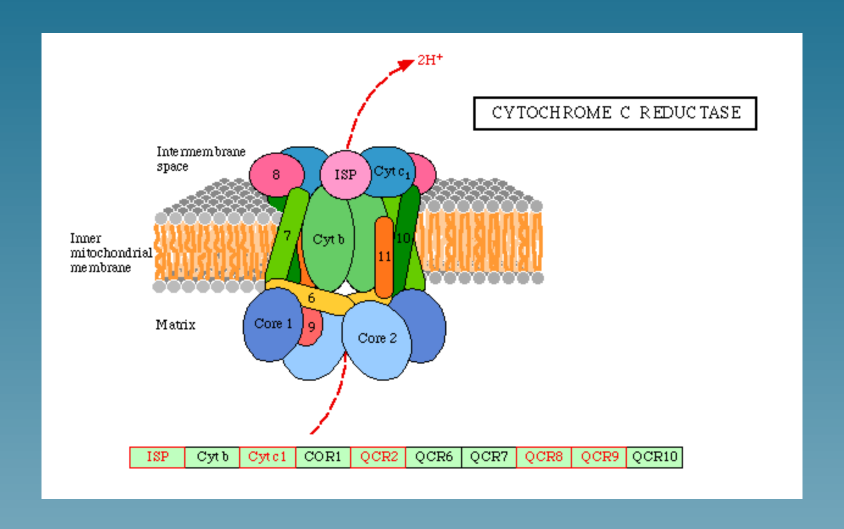
# First pattern of expression

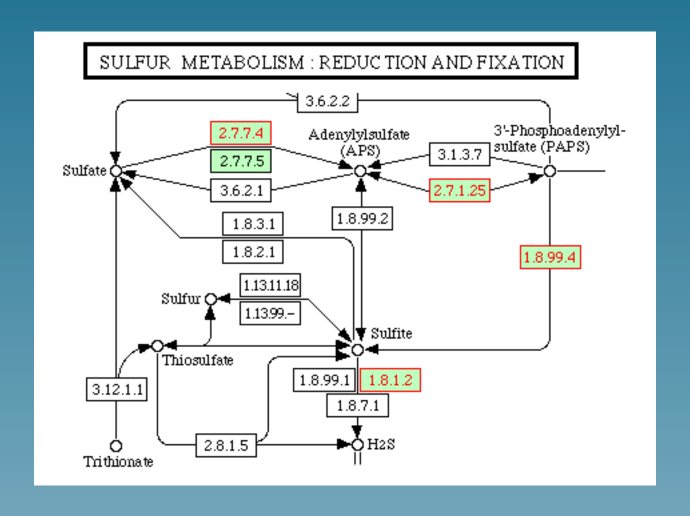


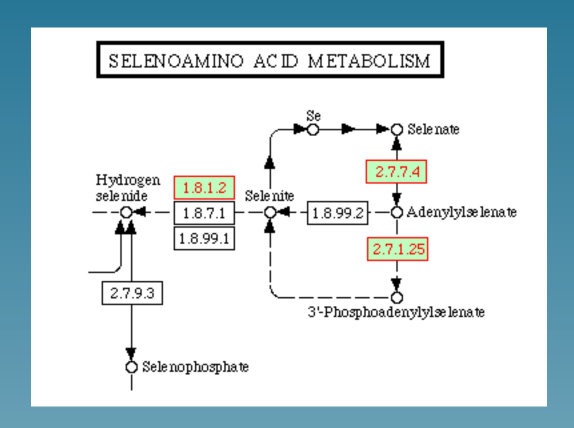
#### Related metabolic pathways

50 genes with highest  $s_2 - s_1$  belong to:

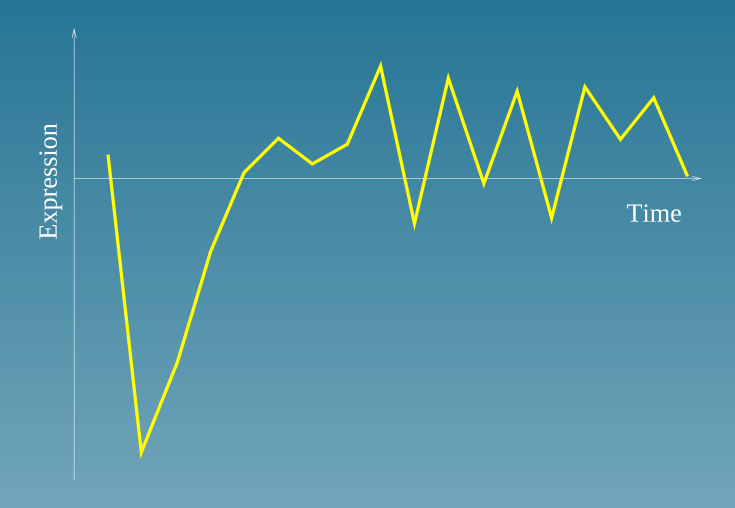
- Oxidative phosphorylation (10 genes)
- Citrate cycle (7)
- Purine metabolism (6)
- Glycerolipid metabolism (6)
- Sulfur metabolism (5), etc...



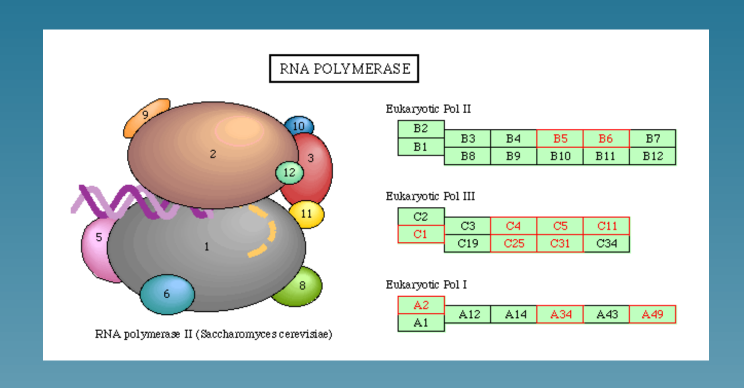


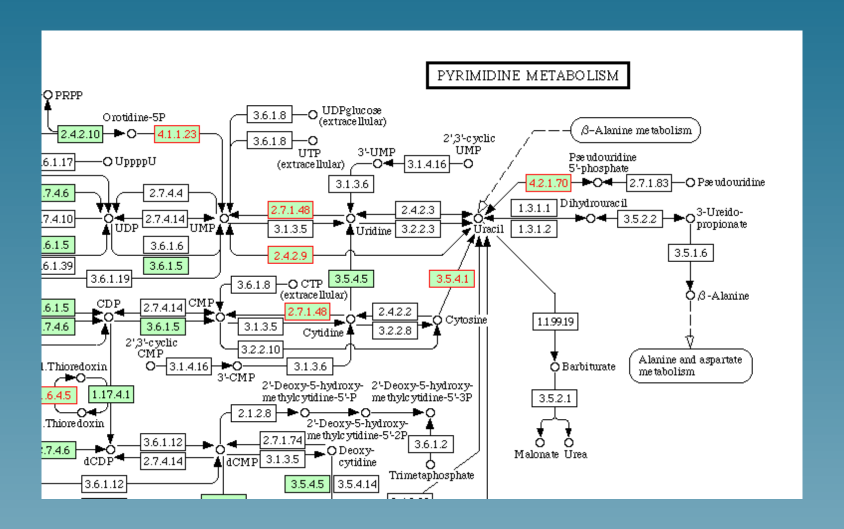


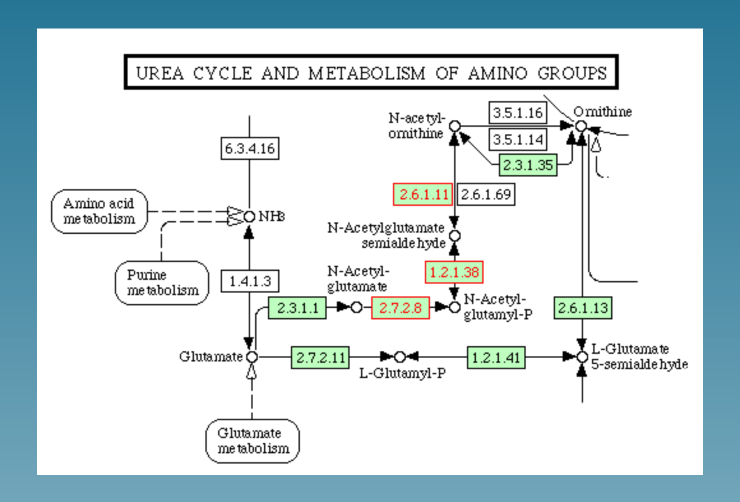
# **Opposite pattern**



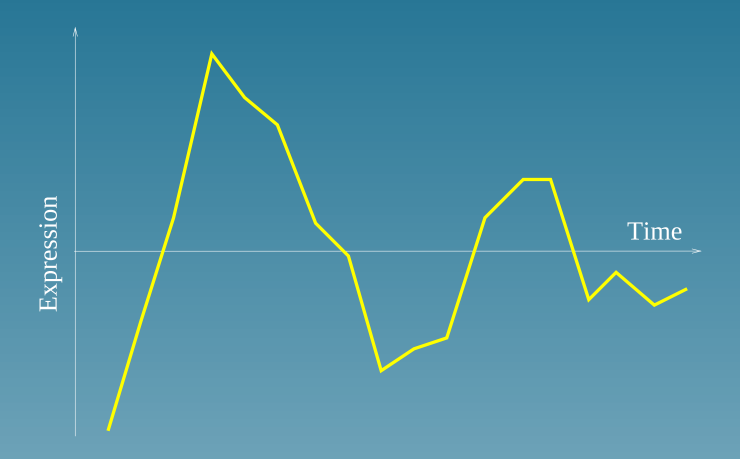
- RNA polymerase (11 genes)
- Pyrimidine metabolism (10)
- Aminoacyl-tRNA biosynthesis (7)
- Urea cycle and metabolism of amino groups (3)
- Oxidative phosphorlation (3)
- ATP synthesis(3) , etc...







# **Second pattern**



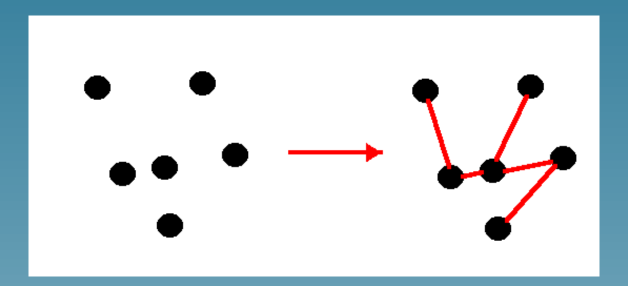
#### Part 4

# Inferring new pathways

(with Y.Yamanishi)

#### The network inference problem

Given some measurement/observation about the genes (sequences, structure, expression, ...), infer "the" gene network



#### Related approaches

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

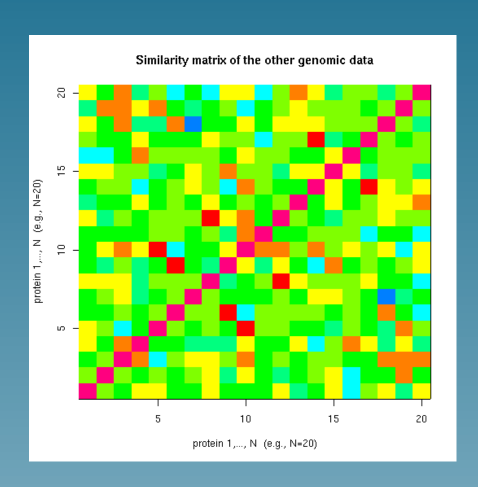
#### A direct (unsupervised) approach

• Let K(x,y) be a measure of similarity (a kernel) between genes x and y based on available measurements, e.g.,

$$K(x,y) = \exp\left(-\frac{||e(x) - e(y)||^2}{2\sigma^2}\right)$$

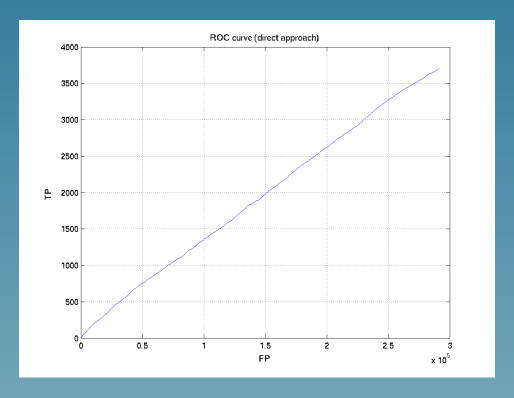
- For a set of n genes  $\{x_1, \ldots, x_n\}$ , let K be the  $n \times n$  matrix of pairwise similarity (Gram matrix)
- Direct strategy: add edges between genes by decreasing similarity.

# **Example of similarity matrix**

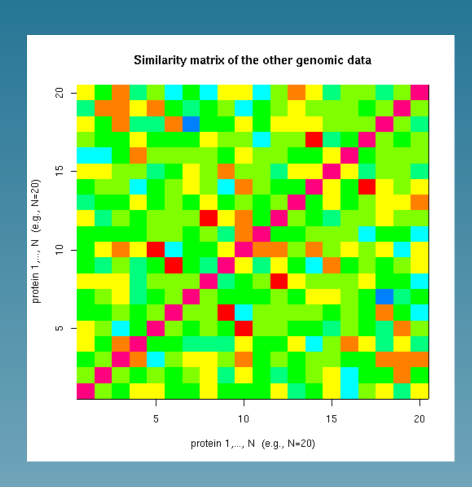


#### **Evaluation of the direct approach**

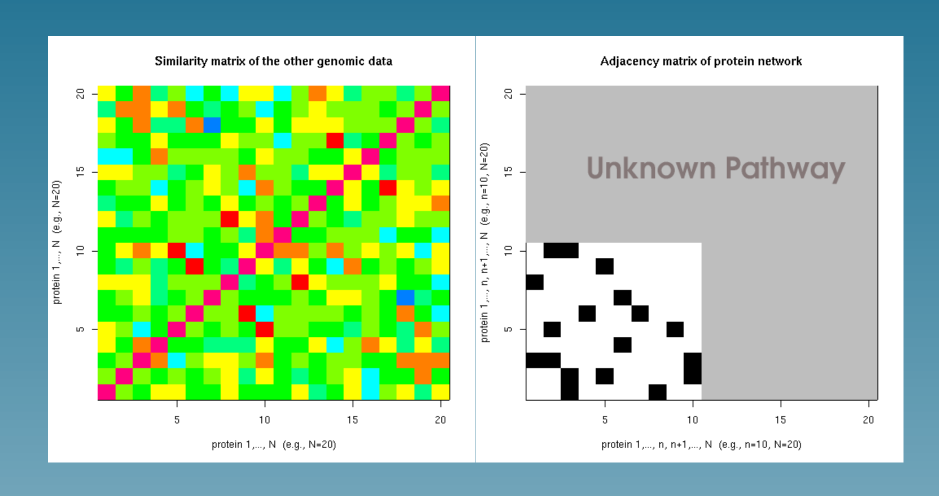
The metabolic network of the yeast involves 769 genes. Each gene is represented by 157 expression measurements. (ROC=0.52)



# The supervised gene inference problem



### The supervised gene inference problem



#### The idea in a nutshell

- Use the known network to define a more relevant measure of similarity
- For any positive definite similarity  $n \times n$  matrix, there exists a representation as n-dimensional vectors such that the matrix similarity is exactly the similarity between vectors.
- In this space, look for projections onto small-dimensional spaces that better fit the known network.

#### A two-step strategy

ullet First map any gene x onto a vector

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

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• Then apply the direct strategy to reconstruct the graph from the images  $\{\Phi(x_1), \dots, \Phi(x_n)\}$ 

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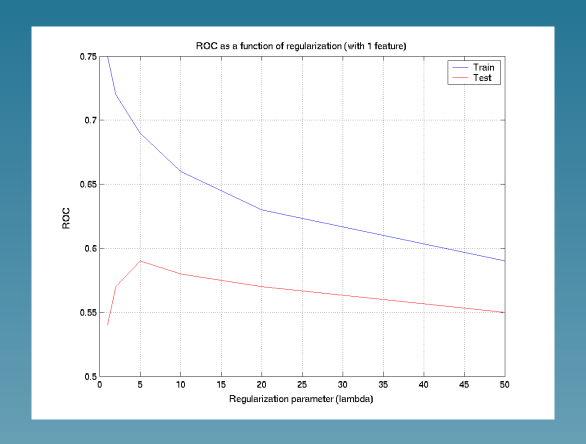
$$\Phi(x) = (f_1(x), \dots, f_d(x))' \in \mathbb{R}^d$$

- Then apply the direct strategy to reconstruct the graph from the images  $\{\Phi(x_1),\ldots,\Phi(x_n)\}$
- The functions  $f_1, \ldots, f_d$  can be learned from the knowledge of the graph on the first n genes

#### Choice of f

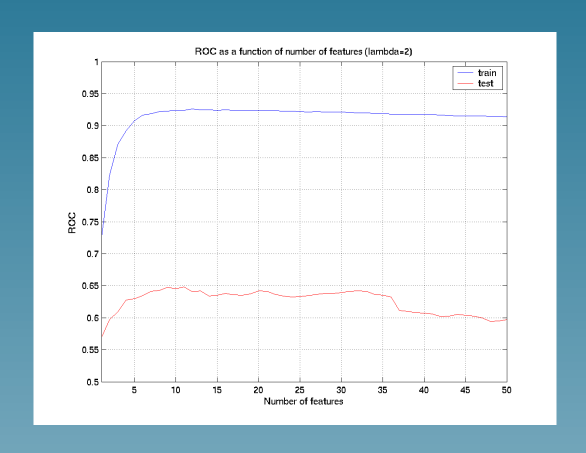
- A feature  $f: \mathcal{X} \to \mathbb{R}$  is good on the training set if connected genes have similar value.
- This is exactly what we did in the previous part!
- So use the features already extracted to map new genes onto a vector space by projection

#### Evaluation of the supervised approach: effect of $\lambda$



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

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

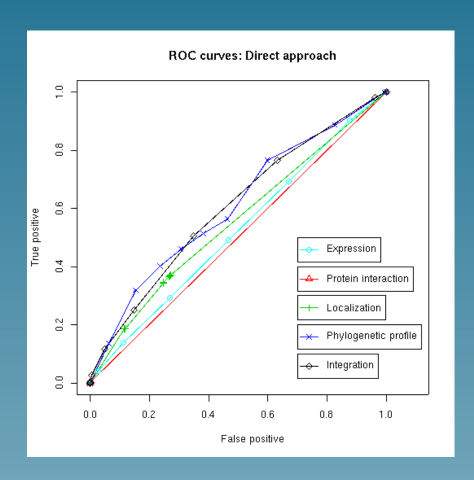


#### Learning from heterogeneous data

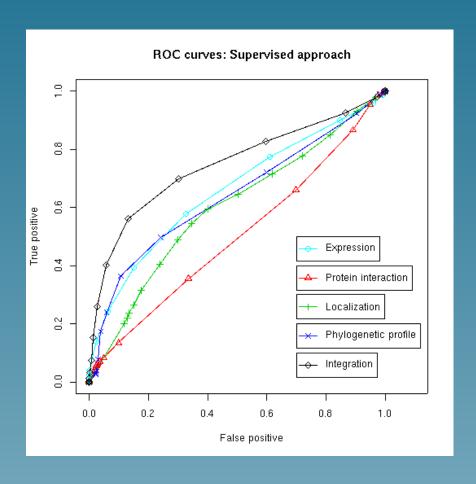
- Suppose several data are available about the genes, e.g., expression, localization, struture, predicted interaction etc...
- Each data can be represented by a positive definite similarity matrix  $K_1, \ldots, K_p$  called kernels
- Kernel can be combined by various operations, e.g., addition:

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

# Learning from heterogeneous data (unsupervised)



## Learning from heterogeneous data (supervised)



#### **Extensions**

- The diffusion kernel can be replaced by another graph kernel
- Other formulations can lead to kernel CCA (NIPS 02)

#### Open questions / Ongoing work

- What should be the number of features (problem of embedding a graph in low dimension)
- Other cost functions
- How to better integrate several similarities? (semi-definite programming?)

# 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
- Possible connections with other algorithms (SVM, kernel CCA..)