

Bioinformatics for Biologists

Functional Genomics: Microarray Data Analysis

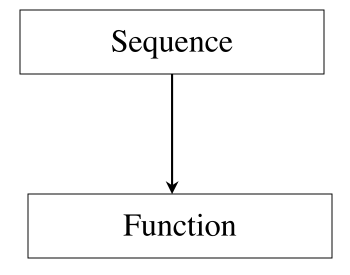
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Outline

- Introduction
- Working with microarray data
 - Normalization
 - Analysis
 - Distance metrics
 - Clustering methods

Research Trends

Genomics



- How are genes regulated?
- How do genes interact?
- What are the functional roles of different genes?
- How does expression level of a gene differ in different tissues?

Transcriptional Profiling

- Study of patterns of gene expression across many experiments that survey a wide array of cellular responses, phenotypes and conditions
- Simple analysis what's up/down regulated?
- More interesting identify patterns of expression for insight into function, etc.

Microarray Data

Collect data on *n* DNA samples (e.g. rows, genes, promotors, exons, etc.) for *p* mRNA samples of tissues or experimental conditions (eg. columns, time course, pathogen exposure, mating type, etc)

Matrix $(n \times p) = \begin{bmatrix} x_{11} & x_{12} & \dots & x_{1p} \\ x_{21} & x_{22} & \dots & x_{2p} \\ \vdots & \vdots & \vdots & \vdots \\ x_{n1} & x_{n2} & \dots & x_{np} \end{bmatrix}$

Multivariate Analysis

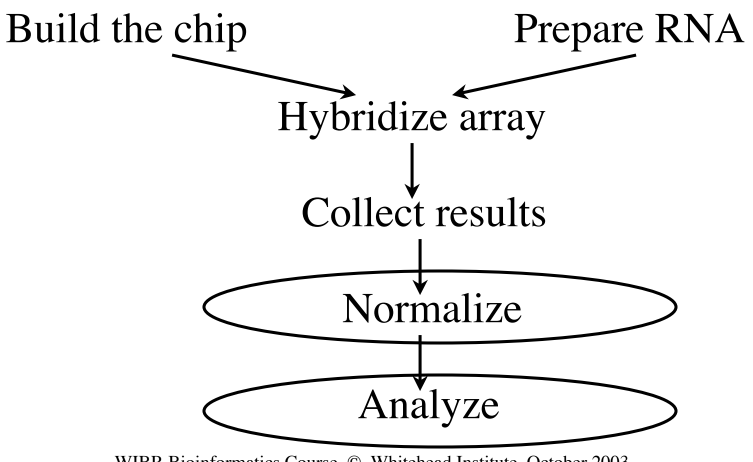
Concerned with datasets with more than one response variable for each observational or experimental unit (e.g. matrix X with *n* rows (genes) and *p* columns (tissue types))

- Hierarchical (phylogenetic trees) vs non-hierarchical (k-means)
- Divisive vs agglomerative
- Supervised vs unsupervised
 - Divide cases into groups vs discover structure of data

Multivariate Methods

- Cluster analysis discover groupings among cases of X
 - Hierarchical produces dendograms
 - K-means choose a prespecified number of clusters
 - Self Organizing Maps
- Principal component analysis (PCA)
 - Linear method, unsupervised, seeks linear combinations of the columns of X with maximal (or minimal) variance (graphical)

DNA Microarrays



Data Normalization

- Correct for systematic bias in data
 - Avoid it, recognize it, correct it, discard outliers
- First step for comparing data from one array to another

Sources of variation

wanted vs unwanted



Across experimental conditions

Chip, slide

Hybridization conditions

Imaging

Normalization Approaches

Compensate for experimental variability

- Housekeeping genes
- Spiked in controls
- Total intensity normalization
- LOWESS correction

Expression Ratios

- Let R = a query sample
- Let G = a reference sample
- Then the ratio, $T_i = R_i/G_i$
- Need to transform these to log₂
- Examples: T = 2/1 = 2; T=1/2 = .5
- Examples: $log_2(2) = 1$; $log_2(.5) = -1$

Total Intensity Normalization

<u>(Adapted from Quackenbush 2002)</u>

Assumptions: (1) start with equal amounts of RNA for the two samples; (2) arrayed elements represent random sample of genes in the organism

a.
$$N_{total} = \frac{\sum_{i=1}^{Narray} R_i}{\sum_{i=1}^{Narray} G_i}$$

c. $T_{i}' = \frac{R_{i}'}{G_{i}'} = \underbrace{\frac{1}{N_{total}} \frac{R_{i}}{G_{i}}}_{N_{total}}$

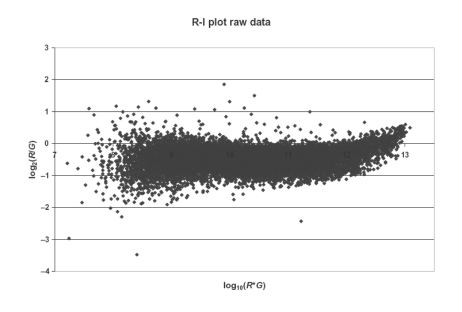
b. Rescale intensities:

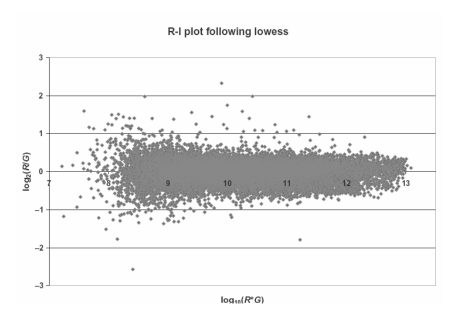
$$G'_{i} = N_{total}G_{i}$$
 and $R'_{i} = R_{i}^{d}$. $\log_{2}(T'_{i}) \neq \log_{2}(T_{i}) - \log_{2}(N_{total})$

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LOWESS - The R-I Plot

- Data exhibit an intensity-dependent structure
- Uncertainty in intensity and ratio measurements is greater at lower intensities





LOWESS - The R-I Plot

- Plot log₂(R/G) ratio as a function of log₁₀(R*G) product intensity
- Shows intensity specific artifacts in the measurements of ratios
- Correct using a local weighted linear regression

LOWESS Normalization

(From Quackenbush 2002)

If we set $x_i = \log_{10}(R_i^*G_i)$ and $y_i = \log_2(R_i/G_i)$, lowess first estimates $y(x_k)$, the dependence of the $\log_2(\text{ratio})$ on the $\log_{10}(\text{intensity})$, and then uses this function, point by point, to correct the measured $\log_2(\text{ratio})$ values so that

$$\log_2(T_i) = \log_2(T_i) - y(x_i) = \log_2(T_i) - \log_2(2^{y(x_i)}),$$

or equivalently,

$$\log_2(T_i') = \log_2\left(T_i * \frac{1}{2^{y(xi)}}\right) = \log_2\left(\frac{R_i}{G_i} * \frac{1}{2^{y(xi)}}\right).$$

As with the other normalization methods, we can make this equation equivalent to a transformation on the intensities, where

$$G'_i = G_i * 2^{y(x_i)}$$
 and $R'_i = R_i$.

After normalization

(Adapted from Quackenbush 2001)

- Data reported as an "expression ratio" or as a logarithm of the expression ratio
- Expression ratio is the normalized value of the expression level for a particular gene in the query sample divided by its normalized value for the control

Use log of expression ratio for easier comparisons

Distance Metrics

- Metric distances d_{ij} between two vectors, i and j, must obey several rules:
 - Distance must be positive definite, $dij \ge 0$
 - Distance must be symmetric, $d_{ij} = d_{ji}$, so that the distance from i to j is the same as the distance from j to i.
 - An object is zero distance from itself, $d_{ii} = 0$.
 - When considering three objects, i, j and $k, d_{ik} \le d_{ij} + d_{jk}$. This is sometimes called the 'triangle' rule.

Distance Metrics

(Adapted from Quackenbush 2001)

• The most common metric distance is Euclidean distance, which is a generalization of the familiar Pythagorean theorem. In a three-dimensional space, the Euclidean distance, d_{12} , between two points, (x_1,x_2,x_3) and (y_1,y_2,y_3) is given by:

$$d_{12} = \sqrt{(x_1 - y_1)^2 + (x_2 - y_2)^2 + (x_3 - y_3)^2}$$

• where (x_1,x_2,x_3) are the usual Cartesian coordinates (x,y,z).

More on distance

(Adapted from Quackenbush 2001)

The generalization of this to higher-dimensional expression spaces is straightforward.

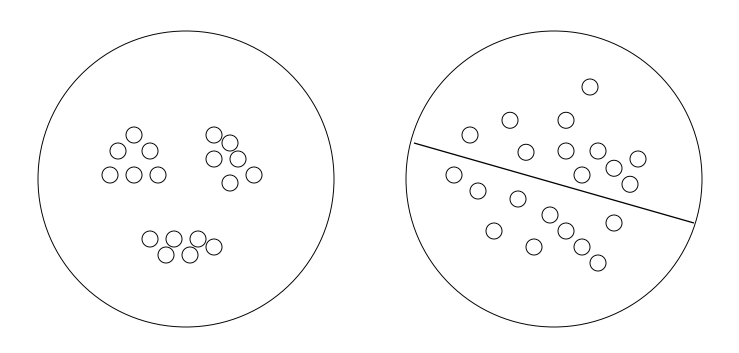
$$d = \sqrt{\sum_{i=1}^{n} (x_i - y_i)^2}$$

where x_i and y_i are the measured expression values, respectively, for genes X and Y in experiment i, and the summation runs over the n experiments under analysis.

Semi-metric distances

- Distance measures that obey the first three consistency rules, but fail to maintain the triangle rule are referred to as semi-metric.
- Pearson correlation coefficient is most commonly used semi-metric distance measure

Clustering vs Classification



Unsupervised

Supervised

Hierarchical methods

(Adapted from Dudoit and Gentleman, 2002)

- Produces a tree or dendogram
- Don't need to specify how many clusters
- The tree can be built in two distinct ways
 - bottom-up: agglomerative clustering
 - top-down: divisive clustering

Agglomerative methods

(Adapted from Dudoit and Gentleman, 2002)

- Start with *n* mRNA sample clusters
- At each step, merge two closest clusters using a measure of between-cluster dissimilarity reflecting shape of the clusters
- Between-cluster dissimilarity measures
 - Unweighted Pair Group Method with Arithmetic mean (UPGMA):
 average of pairwise dissimilarities
 - Single-link: minimum of pairwise dissimilarities
 - Complete-link: maximum of pairwise dissimilarities

Divisive methods

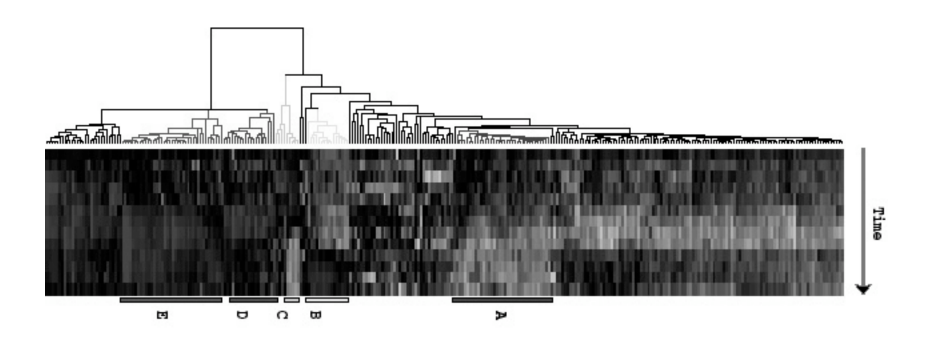
(Adapted from Dudoit and Gentleman, 2002)

- Start with only one cluster
- At each step, split clusters into parts
- Advantages: obtain main structure of the data, i.e., focus on upper levels of dendogram
- Disadvantages: computational difficulties when considering all possible divisions into two groups

Hierarchical Clustering

- Agglomerative single expression profiles are joined to form groups....forming a single tree
 - Pairwise distance matrix is calculated for all genes to be clustered
 - Distance matrix is searched for the 2 most similar genes or clusters
 - Two selected clusters are merged to produce new cluster
 - Distances calculated between this new cluster and all other clusters

Dendogram



Eisen et al 1998

K-means Clustering

- *Divisive* good if you know the number (*k*) of clusters to be represented in the data
 - Initial objects randomly assigned to one of k clusters
 - Average expression vector calculated for each cluster & compute distance between clusters
 - Objects moved between clusters and intra- and intercluster distances are measured with each move
 - Expression vectors for each cluster are recalculated
 - Shuffling proceeds until moving any more objects would make clusters more variable (> intra-cluster distances and decreasing inter-cluster dissimilarity

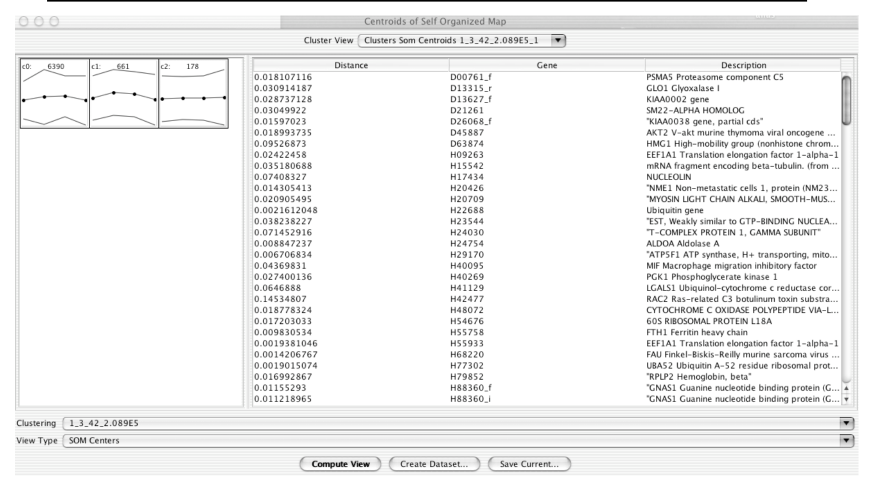
Self Organizing Maps (SOM)

- Neural-network based divisive clustering approach
 - Assigns genes to a series of partitions
 - User defines a geometric configuration for the partitions
 - Random vectors are generated for each partition
 - Vectors are first 'trained' using an iterative process until data most effectively separated

SOMs Continued

- Random vectors are constructed and assigned to each partition
- A gene is picked at random and, using a selected distance metric, the reference vector that is closest to the gene is identified
- The reference vector is then adjusted so that it is more similar to the vector of the assigned gene
- Genes are mapped to relevant partitions depending on the reference vector to which they are most similar

SOMs from GeneCluster

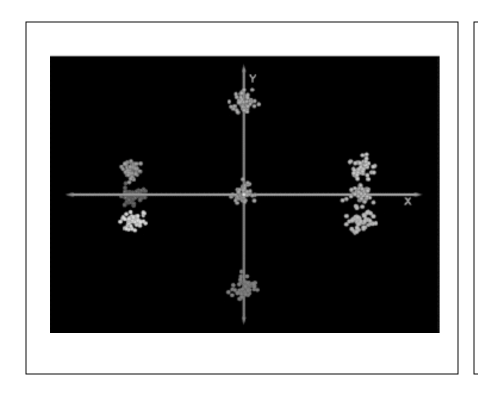


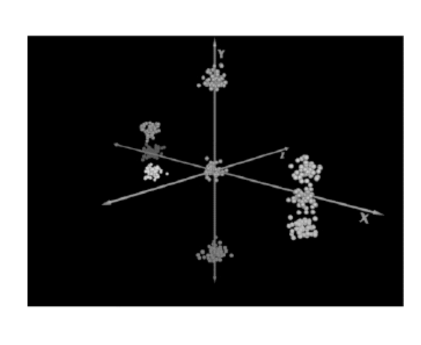
Principal Component Analysis

- Data reduction method
- AKA singular value decomposition
- Used to pick out patterns in data
- Provide projection of complex data sets onto reduced, easily visualized space
- Difficult to define precise clusters but can give you an idea of # of clusters for SOMs or k-means

Principal Component Analysis

(Quackenbush 2001)





Quackenbush 2001

"One must remember that the results of any analysis have to be evaluated in the context of other biological knowledge."

Supervised Learning

- Useful if you have some previous information about which genes are expected to cluster together
- Support Vector Machine (SVM)
- Start with training set (eg. positive and negative examples)
- SVM learns to distinguish between members and non-members of a class

Warnings

- Classification is dependent on
 - clustering method used
 - normalization of data
 - measure of similarity

Citations

- Brazma A and Vilo J. Minireview: Gene expression data analysis. *FEBS Letters* 480:17-24, 2000.
- Quackenbush J. Computational Analysis of Microarray Data. *Nature Review* | *Genetics* 2:418-427, 2001.
- Quackenbush J. Microarray data normalization and transformation. *Nature Genetics Supp.* 32:496-501, 2002.
- Dudoit S and Gentleman R. Classification in microarray experiments. Statistics and Genomics Short Course -Lecture 5, January 2002 (http://www.bioconductor.org/workshop.html)

Available Tools

- GeneCluster (WI/MIT Genome Center)
- Cluster & TreeView (Eisen)
- GeneSpring (Silicon Genetics)
- Spotfire (Spotfire)
- R Statistics Package/Bioconductor
- Matlab (modules from Churchill, JAX)

Lists of Tools

- Rockefeller University (formerly)
 - http://www.nslij-genetics.org/microarray/
- R Statistics Package Microarry Tools
 - http://www.stat.uni-muenchen.de/~strimmer/rexpress.html
- Bioconductor Project
 - http://www.bioconductor.org/
- EBI
 - http://ep.ebi.ac.uk/Links.html
 - http://ep.ebi.ac.uk/EP/