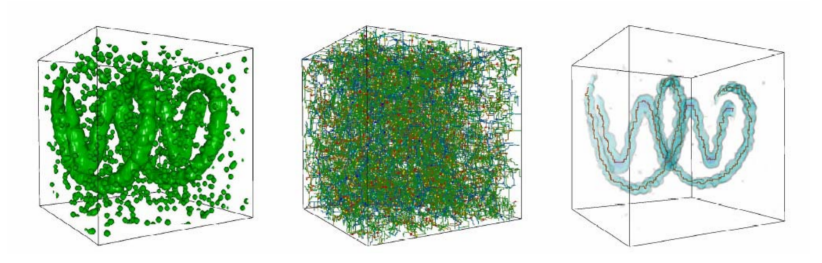
Clustering-1

Manu Madhavan

Slide Courtesy: MIT OpenCourseWare http://ocw.mit.edu

Structure in High-Dimensional Data



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Gyulassy, Atilla, et al. "Topologically Clean Distance Fields." *IEEE Transactions on Visualization and Computer Graphics* 13, no. 6 (2007): 1432-1439.

- Structure can be used to reduce dimensionality of data
- Structure can tell us something useful about the underlying phenomena
- Structure can be used to make inferences about new data

Clustering vs Classification

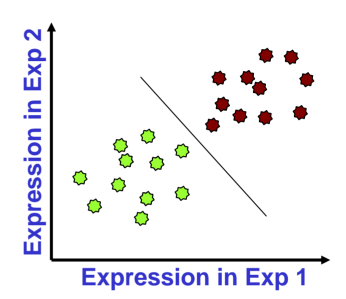
Objects characterized by one or more features

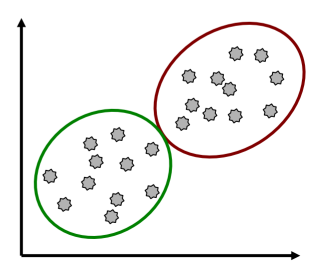
Classification

- Have <u>labels</u> for some points
- Want a "rule" that will accurately assign labels to new points
- Supervised learning

Clustering

- No labels
- Group points into clusters based on how "near" they are to one another
- Identify structure in data
- Unsupervised learning





Today

Microarray Data

K-means clustering

Expectation Maximization

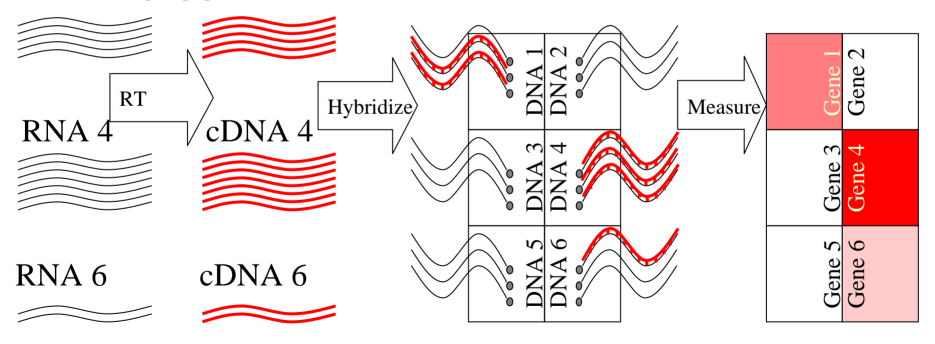
Hierarchical Clustering

Expression Microarrays

- A way to measure the levels of mRNA in every gene
- Two basic types
 - Affymetrix gene chips
 - Spotted oligonucleotides
- Both work on same principle
 - Put DNA probe on slide
 - Complementary hybridization

Expression Microarrays

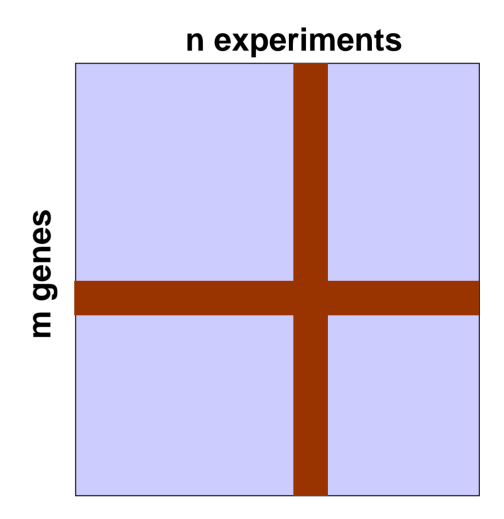
 Measure the level of mRNA messages in a cell



Expression Microarray Data Matrix

Genes are typically given as rows

 Experiment are given by columns



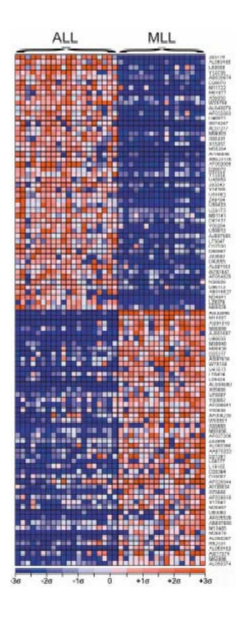
Clustering and Classification in Genomics

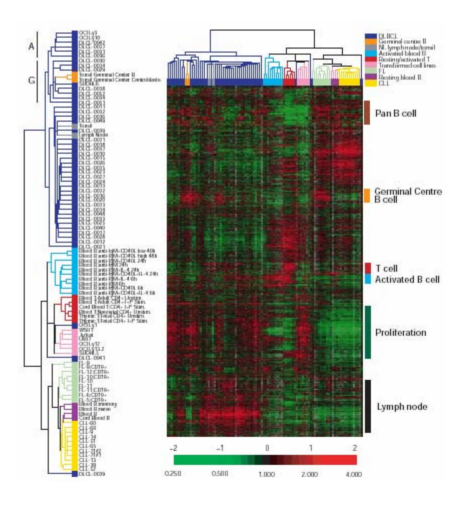
Classification

- Microarray data: classify cell state (i.e. AML vs ALL) using expression data
- Protein/gene sequences: predict function, localization, etc.

Clustering

- Microarray data: groups of genes that share similar function have similar expression patterns – identify regulons
- > Protein sequence: group related proteins to infer function
- > EST data: collapse redundant sequences

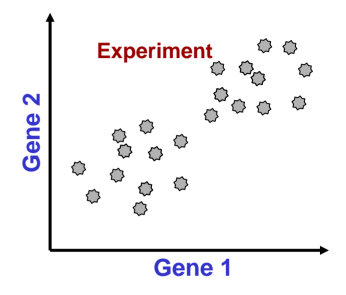




Clustering Expression Data

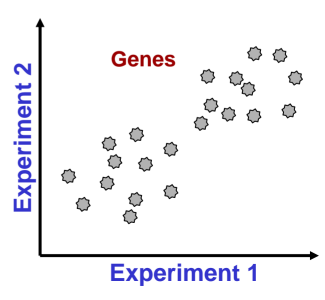
Cluster Experiments

Group by similar expression profiles



Cluster Genes

Group by similar expression in different conditions



Why Cluster Genes by Expression?

Data Exploration

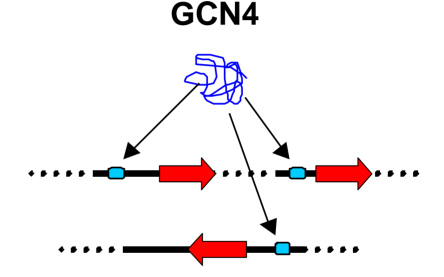
- Summarize data
- Explore without getting lost in each data point
- Enhance visualization

Co-regulated Genes

- Common expression may imply common regulation
- Predict *cis*-regulatory promoter sequences

Functional Annotation

 Similar function from similar expression



His2 His3 Unknown Amino Acids Amino Acids

Clustering Algorithms

- Partitioning
 - Divides objects into non-overlapping clusters such that each data object is in exactly one subset

- Agglomerative
 - A set of nested clusters organized as a hierarchy

K-Means Clustering

The Basic Idea

Assume a fixed number of clusters, K

Goal: create "compact" clusters

More Formally

1. Initialize K centers **u**_k

For each iteration n until convergence

 Assign each x_i the label of the nearest center, where the distance between x_i and u_k is

$$d_{i,k} = \left(\mathbf{x}_i - \mathbf{\mu}_k\right)^2$$

3. Move the position of each \mathbf{u}_k to the centroid of the points with that label

$$\mu_k(n+1) = \sum_{\mathbf{x}_i \text{ with label j}} \frac{\mathbf{X}_i}{\left|\mathbf{X}^k\right|}, \left|\mathbf{X}^k\right| = \#\mathbf{X}_i \text{ with label k}$$

Cost Criterion

We can think of K-means as trying to create clusters that minimize a cost criterion associated with the size of the cluster

$$COST(\mathbf{x}_{1},\mathbf{x}_{2},\mathbf{x}_{3},...,\mathbf{x}_{n}) = \sum_{\mathbf{\mu}_{k}} \sum_{\mathbf{x}_{i} \text{ with label k}} (\mathbf{x}_{i} - \mathbf{\mu}_{k})^{2}$$

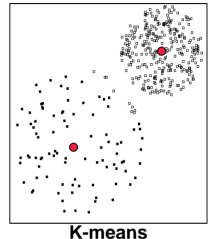
Minimizing this means minimizing each cluster term separately:

$$\sum_{\mathbf{x}: \text{with label k}} \left(\mathbf{x}_{i} - \mathbf{\mu}_{k}\right)^{2}$$

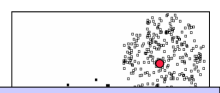
Fuzzy K-Means

- Initialize K centers **u**_k
- For each point calculate the probability of membership for each category

P(label
$$K | \mathbf{x}_i, \boldsymbol{\mu}_k$$
)



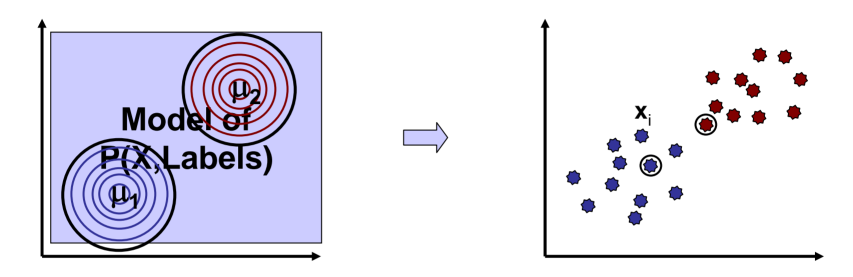
Move the position of each \mathbf{u}_k to the weighted centroid:



Of course, K-Means just special case where

P(label K |
$$\mathbf{x}_i, \boldsymbol{\mu}_k$$
) =
$$\begin{cases} 1 & \text{if } \mathbf{x}_i \text{ is closest to } \boldsymbol{\mu}_k \\ 0 & \text{otherwise} \end{cases}$$

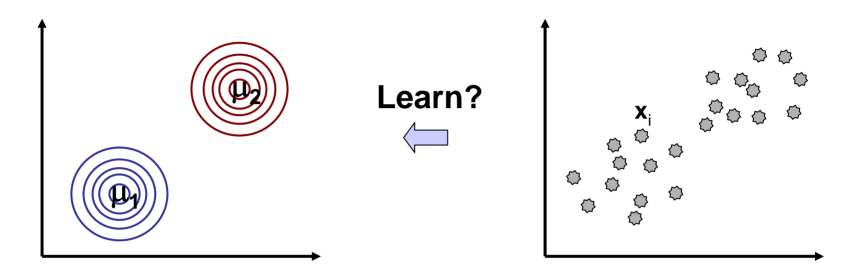
K-Means as a Generative Model



Samples drawn from two equally normal distributions with unit variance - a *Gaussian Mixture Model*

$$P(\mathbf{x}_i | \mathbf{u}_j) = \frac{1}{\sqrt{2\pi}} \exp \left\{ -\frac{\left(\mathbf{x}_i - \mathbf{u}_j\right)^2}{2} \right\}$$

Unsupervised Learning



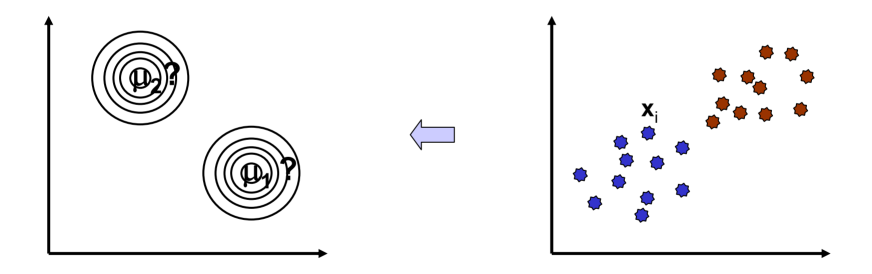
Samples drawn from two equally normal distributions with unit variance - a *Gaussian Mixture Model*

$$P(\mathbf{x}_i | \mathbf{u}_j) = \frac{1}{\sqrt{2\pi}} \exp \left\{ -\frac{\left(\mathbf{x}_i - \mathbf{u}_j\right)^2}{2} \right\}$$

If We Have Labeled Points

Need to estimate unknown gaussian centers from data

In general, how could we do this? How could we "estimate" the "best" u_{k?}



Choose u_k to maximize probability of model

If We Have Labeled Points

Need to estimate unknown gaussian centers from data

In general, how could we do this? How could we "estimate" the "best" u_{k?}

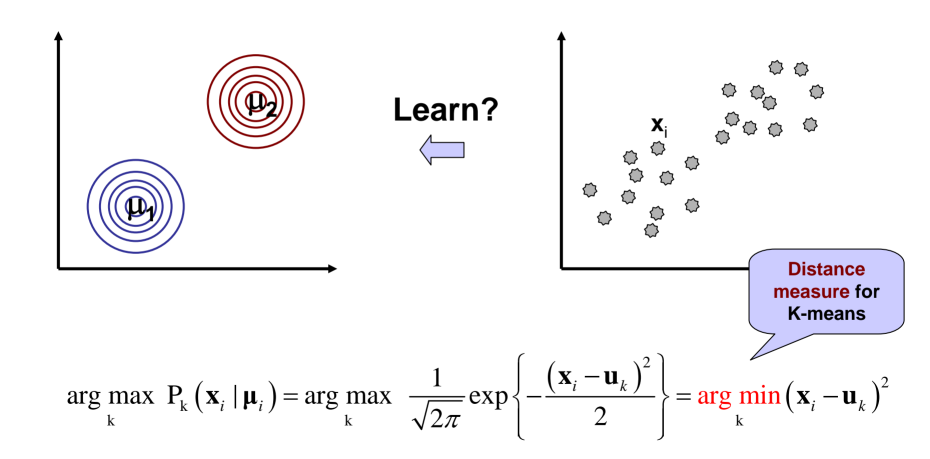
Given a set of \mathbf{x}_i , all with label k, we can find the maximum likelihood μ_k from

$$\arg \max_{\mu} \left\{ \log \prod_{i} P\left(\mathbf{x}_{i} \mid \boldsymbol{\mu}\right) \right\} = \arg \max_{\mu} \sum_{i} \left\{ -\frac{1}{2} \left(\mathbf{x}_{i} - \boldsymbol{u}\right)^{2} + \log \left(\frac{1}{\sqrt{2\pi}}\right) \right\}$$

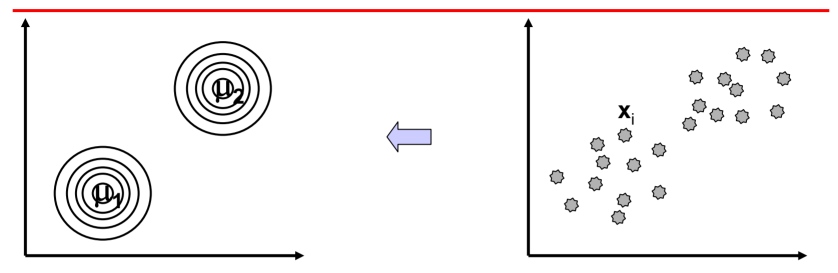
$$= \arg \min_{\mu} \sum_{i} \left(\mathbf{x}_{i} - \boldsymbol{u}\right)^{2}$$
Solution is the centroid of the \mathbf{x}_{i}

If We Know Cluster Centers

Need to estimate labels for the data



What If We Have Neither?



An idea:

- 1. Imagine we start with some u_k⁰
- 2. We *could* calculate the most likely labels for x_i^0 given these u_k^0
- 3. We *could* then use these labels to choose u_k¹
- 4. And iterate (to convergence)

Expectation Maximization (EM)

1. Initialize parameters

2. E Step Estimate probability of hidden labels, Q, given parameters and sequence

$$Q = P(label_i | x, u_i^{t-1})$$

3. M Step Choose new parameters to maximize expected likelihood of parameters given Q

$$u_k^t = \underset{u}{\operatorname{arg max}} E_Q \left[\log P(labels \mid x, u_k^{t-1}) \right]$$

4. Iterate

P(x|Model) guaranteed to increase each iteration

Expectation Maximization (EM)

Remember the basic idea!

1.Use model to estimate (distribution of) missing data 2.Use estimate to update model 3.Repeat until convergence

Model is the gaussian distributions

Missing data are the data point labels

Revisiting K-Means

Generative Model Perspective

- 1. Initialize K centers **u**_k
- 2. Assign each \mathbf{x}_i the label of the nearest center, where the distance between \mathbf{x}_i and \mathbf{u}_k is

$$\qquad \qquad \square \rangle$$

The most likely label k for a point x_i

$$d_{i,k} = \left(\mathbf{x}_i - \mathbf{\mu}_k\right)^2$$

3. Move the position of each \mathbf{u}_k to the centroid of the points with that label



Maximum likelihood parameter μ_k given most likely label

4. Iterate

Revisiting K-Means

Generative Model Perspective

- 1. Initialize K centers **u**_k
- 2. Assign each \mathbf{x}_i the label of the nearest center, where the distance between \mathbf{x}_i and \mathbf{u}_k is

$$d_{i,k} = \left(\mathbf{x}_i - \mathbf{\mu}_k\right)^2$$

- 3. Move the position of each **u**_k to the centroid of the points with that label
- 4. Iterate

- 1. Initialize parameters
- 2.E Step Estimate most likely missing label given previous parameter

- 3.M Step Choose new parameters to maximize likelihood of parameters given estimated labels
- 4. Iterate

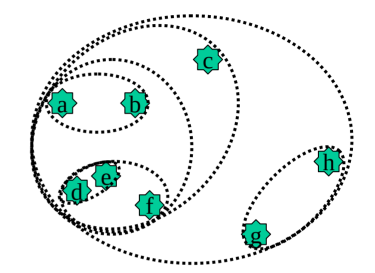
But How Many clusters?

- How do we select K?
 - We can always make clusters "more compact" by increasing K
 - e.g. What happens is if K=number of data points?
 - What is a meaningful improvement?
- Hierarchical clustering side-steps this issue

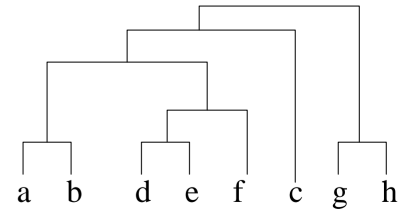
Hierarchical clustering

Most widely used algorithm for expression data

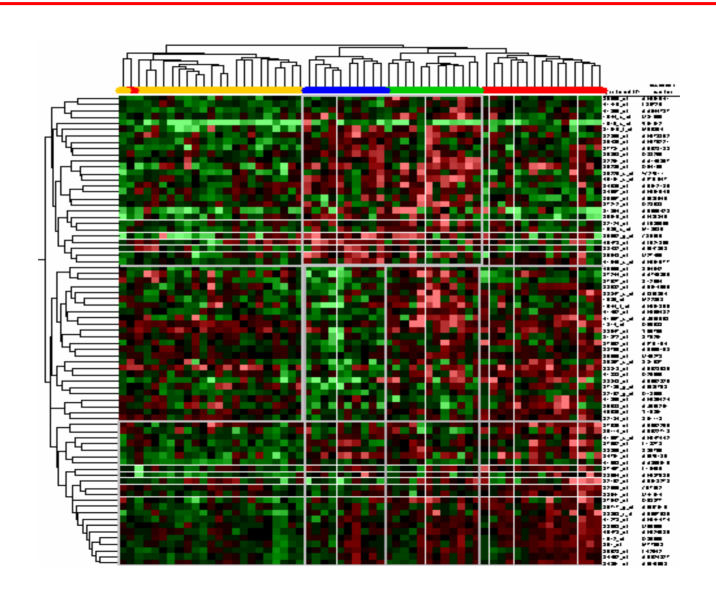
- Start with each point in a separate cluster
- At each step:
 - Choose the pair of closest clusters
 - Merge







Visualization of results

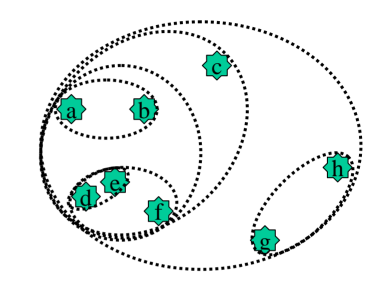


Hierarchical clustering

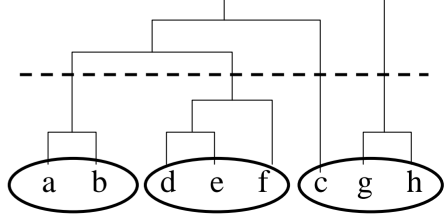
Avoid needing to select number of clusters

Produces clusters at all levels

We can always select a "cut level" to create disjoint clusters



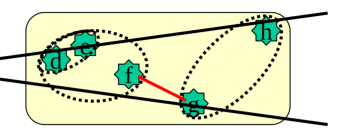
But how do we define distances between clusters?



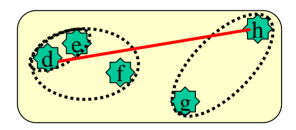
slide credits: M. Kellis

Distance between clusters

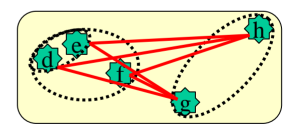
• $CD(X,Y)=min_{x \in X, y \in Y}D(x,y)$ Single-link method



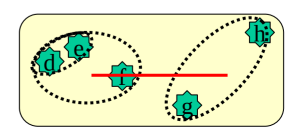
CD(X,Y)=max_{x∈X,y∈Y}D(x,y)
 Complete-link method



CD(X,Y)=avg_{x ∈X, y ∈Y} D(x,y)
 Average-link method



CD(X,Y)=D(avg(X), avg(Y))
 Centroid method



(Dis)Similarity Measures

Image removed due to copyright restrictions.

Table 1, Gene expression similarity measures. D'haeseleer, Patrik. "How Does Gene Expression Clustering Work?" *Nature Biotechnology* 23 (2005): 1499-1501.

Evaluating Cluster Performance

In general, it depends on your goals in clustering

Robustness

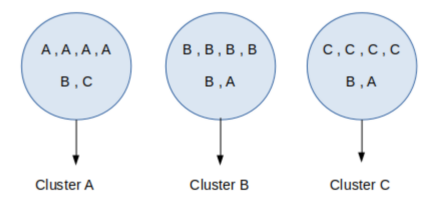
- Select random samples from data set and cluster
- Repeat
- Robust clusters show up in all clusters

Category Enrichment

- Look for categories of genes "over-represented" in particular clusters
- Also used in Motif Discovery

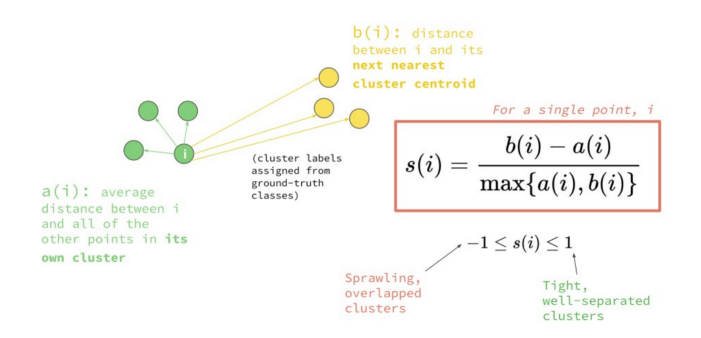
Evaluating Cluster Quality

Purity

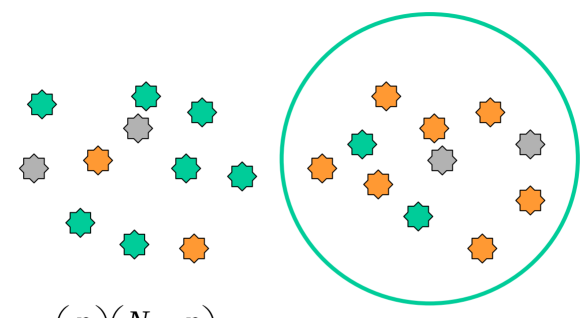


$$purity = rac{(cluster A + cluster B + cluster C)}{total} = rac{(4+5+4)}{18} = 0.722$$

Silhoutte Score



Evaluating clusters – Hypergeometric Distribution

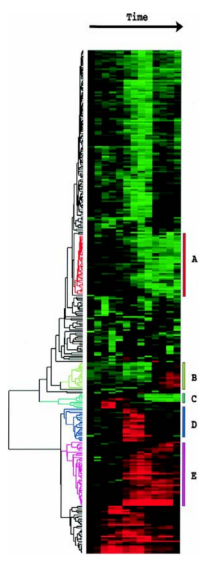


$$P(pos \ge r) = \sum_{m \ge r} \frac{\binom{p}{m} \binom{N-p}{k-m}}{\binom{N}{k}}$$
 P-value of uniformity in computed cluster

- - P-value of *single* cluster containing k elements of which at least r are +

Prob that a randomly chosen set of k experiments would result in m positive and k-m negative

Similar Genes Can Cluster



Clustered 8600 human genes using expression time course in fibroblasts

- (A) Cholesterol biosynthesis
- (B) Cell cycle
- (C) Immediate early response
- (D) Signalling and angiogenesis
- (E) Wound healing

Eisen, Michael et al. "Cluster Analysis and Display of Genome-wide Expression Patterns." *PNAS* 95, no. 25 (1998): 14863-14868. Copyright (1998) National Academy of Sciences, U.S.A.

(Eisen (1998) PNAS)

Clusters and Motif Discovery

Expression from 15 time points during yeast cell cycle

