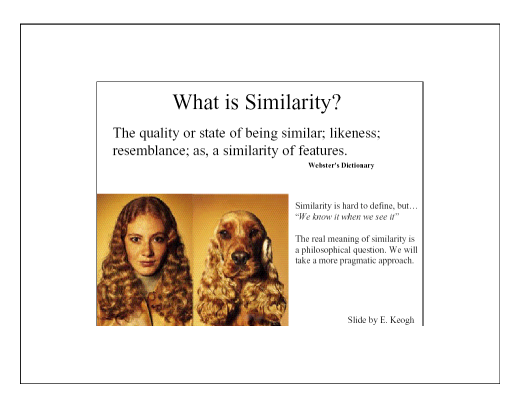


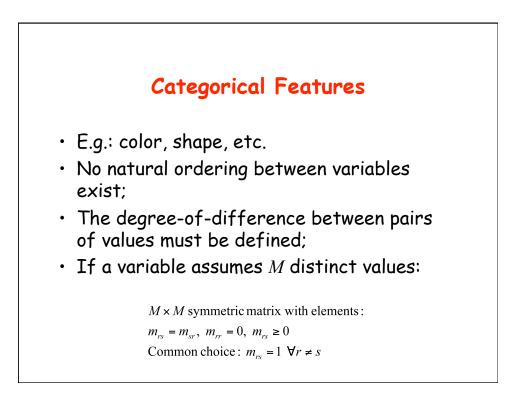
Clustering

- <u>Goal</u>: Grouping a collection of objects (data points) into subsets or "clusters", such that those within each cluster are more closely related to one other than objects assigned to different clusters.
- Fundamental to all clustering techniques is the choice of *distance or dissimilarity measure* between two objects.



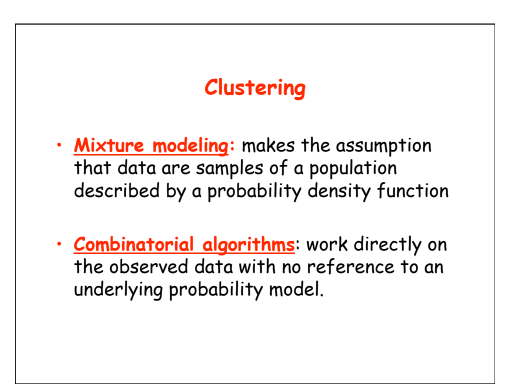
Dissimilarities based on Features

$$\begin{aligned} \mathbf{x}_{i} &= \left(x_{i1}, x_{i2}, \cdots, x_{iq}\right)^{T} \in \Re^{q}, \ i = 1, \cdots, N \\ D\left(\mathbf{x}_{i}, \mathbf{x}_{j}\right) &= \sum_{k=1}^{q} d_{k}\left(x_{ik}, x_{jk}\right) \\ d_{k}\left(x_{ik}, x_{jk}\right) &= \left(x_{ik} - x_{jk}\right)^{2} \\ \Rightarrow D\left(\mathbf{x}_{i}, \mathbf{x}_{j}\right) &= \sum_{k=1}^{q} \left(x_{ik} - x_{jk}\right)^{2} \end{aligned}$$
Squared Euclidean distance
$$D_{w}\left(\mathbf{x}_{i}, \mathbf{x}_{j}\right) &= \sum_{k=1}^{q} w_{k}\left(x_{ik} - x_{jk}\right)^{2} \end{aligned}$$
Weighted squared Euclidean distance



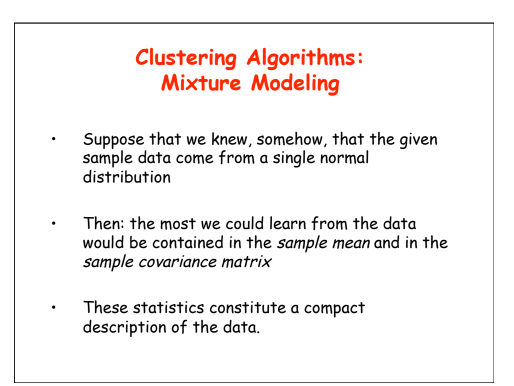
Clustering

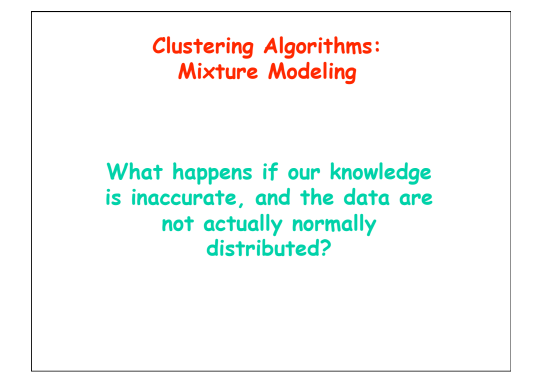
- Discovering patterns (e.g., groups) in data without any guidance (labels) sounds like an "unpromising" problem.
- The question of whether or not it is possible in principle to learn anything from unlabeled data depends upon the assumptions one is willing to accept.

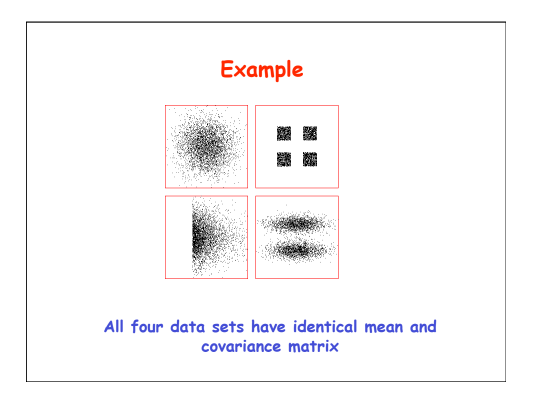


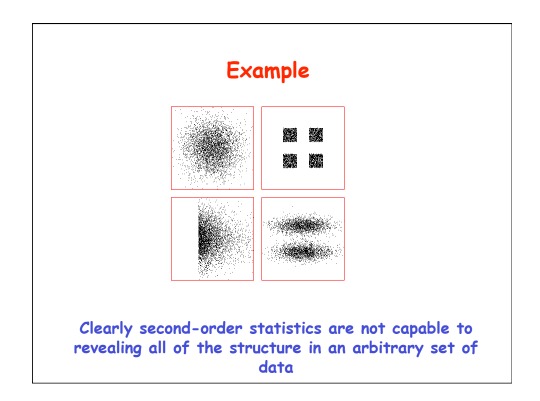
Clustering Algorithms: Mixture Modeling

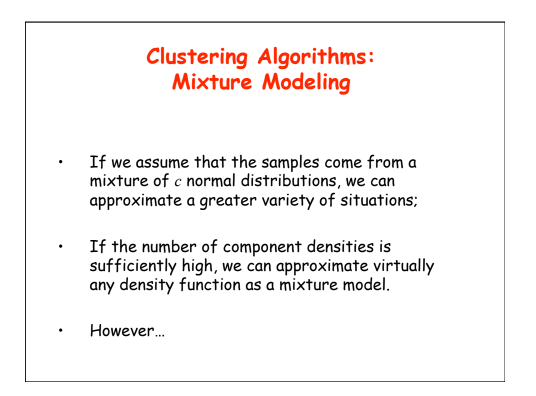
- Data is a sample from a population described by a probability density function;
- The density function is modeled as a mixture of component density functions (e.g., mixture of Gaussians). Each component density describes one of the clusters;
- The parameters of the model (e.g., means and covariance matrices for mixture of Gaussians) are estimated as to best fit the data (*maximum likelihood estimation*).





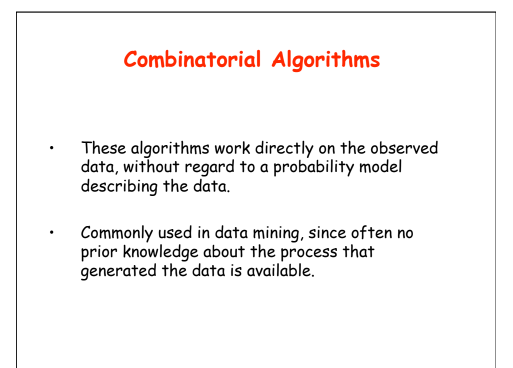






Clustering Algorithms: Mixture Modeling

- The problem of estimating the parameters of a mixture density is not trivial;
- When we have little prior knowledge about the nature of the data, the assumption of specific parametric forms may lead to poor or meaningless results.
 - There is a risk of *imposing* structure on the data instead of *finding* the structure.



Combinatorial Algorithms

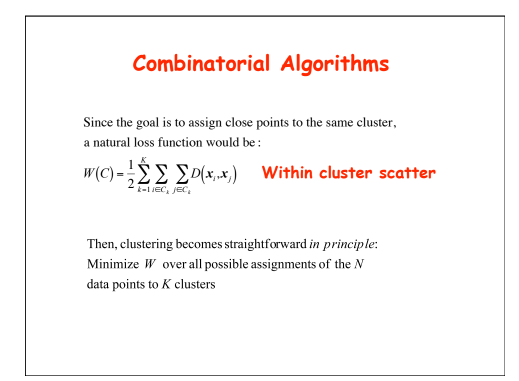
 $\mathbf{x}_i \in \mathfrak{R}^q, i = 1, \cdots, N$

Prespecified number of clusters $K, k \in \{1, \dots, K\}$

Each data point x_i is assigned to one, and only one cluster

Goal: Find a partition of the data into K clusters that achieves a required objective, defined in terms of a dissimilarity function $D(\mathbf{x}_i, \mathbf{x}_k)$

Usually, the assignment of data to clusters is done so as to **minimize** a "loss" function that measures the degree to which the clustering goal is **not** met



Combinatorial Algorithms

Unfortunately, such optimization by complete enumeration is feasible only for very small data sets.

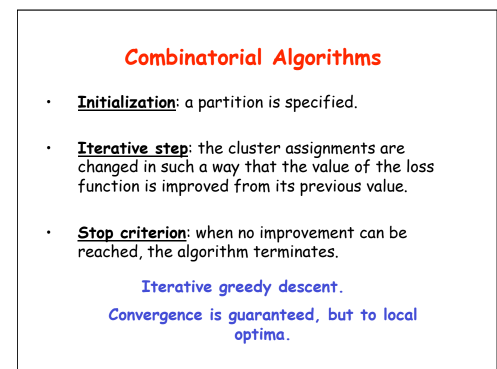
The number of distinct partitions is:

$$S(N,K) = \frac{1}{K!} \sum_{k=1}^{K} (-1)^{K-k} {\binom{K}{k}} k^{N}$$

For example:

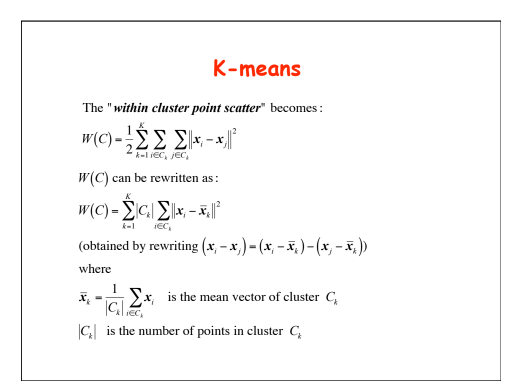
S(10,4) = 34,105 $S(19,4) \approx 10^{10}$

We need to limit the search space, and find in general a good suboptimal solution



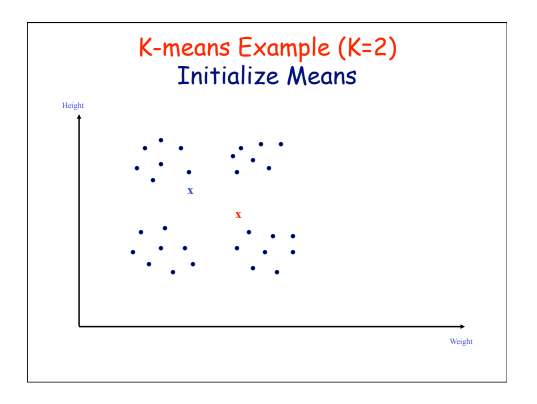
K-means

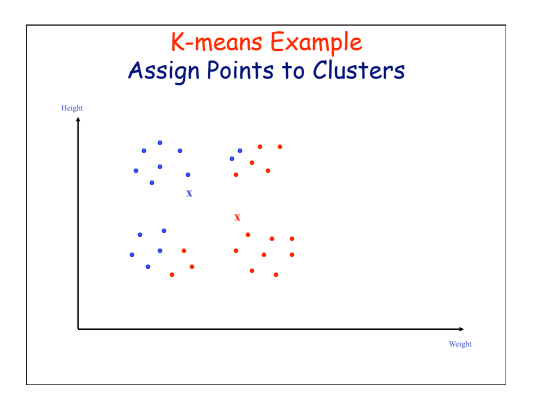
- One of the most popular iterative descent clustering methods.
- Features: quantitative type.
- Dissimilarity measure: Euclidean distance.

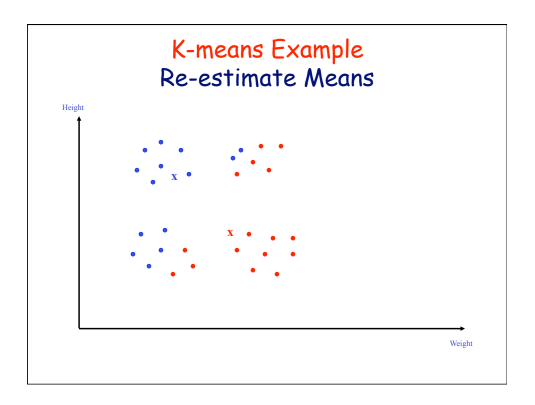


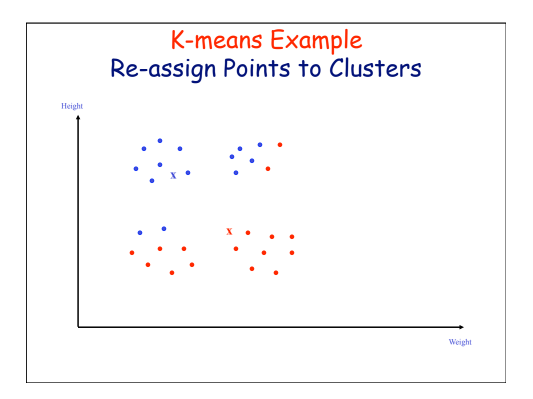
K-means The objective is: $\begin{aligned} & \min_{C} \sum_{k=1}^{K} |C_{k}| \sum_{i \in C_{k}} \|\mathbf{x}_{i} - \bar{\mathbf{x}}_{k}\|^{2} \\ & \text{We can solve this problem by noticing:} \\ & \text{for any set of data S} \\ & \overline{\mathbf{x}}_{S} = \arg_{m} \sum_{i \in S} \|\mathbf{x}_{i} - \mathbf{m}\|^{2} \\ & \text{(this is obtained by setting } \frac{\partial \sum_{i \in S} \|\mathbf{x}_{i} - \mathbf{m}\|^{2}}{\partial \mathbf{m}} = 0; \\ & \text{So we can solve the enlarged optimization problem:} \\ & \min_{C, m_{k}} \sum_{k=1}^{K} |C_{k}| \sum_{i \in C_{k}} \|\mathbf{x}_{i} - \mathbf{m}_{k}\|^{2} \end{aligned}$

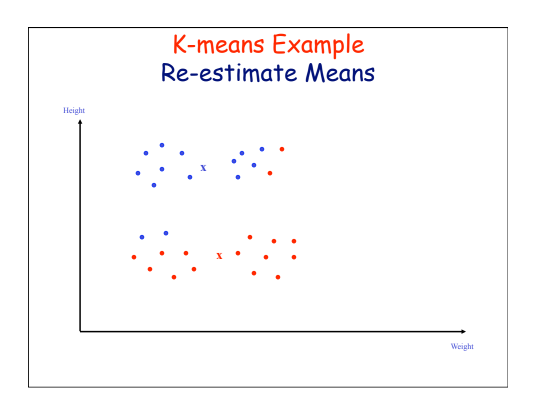
K-means: The Algorithm 1. Given a cluster assignment *C*, the total within cluster scatter $\sum_{k=1}^{K} |C_k| \sum_{i \in C_k} ||\mathbf{x}_i - \mathbf{m}_k||^2 \text{ is minimized with respect to the } \{\mathbf{m}_1, \dots, \mathbf{m}_K\}$ giving the means of the currently assigned clusters; 2. Given a current set of means $\{\mathbf{m}_1, \dots, \mathbf{m}_K\}$, $\sum_{k=1}^{K} |C_k| \sum_{i \in C_k} ||\mathbf{x}_i - \mathbf{m}_k||^2 \text{ is minimized with respect to } C$ by assigning each point to the closest current cluster mean; 3. Steps 1 and 2 are iterated until the assignments do not change.

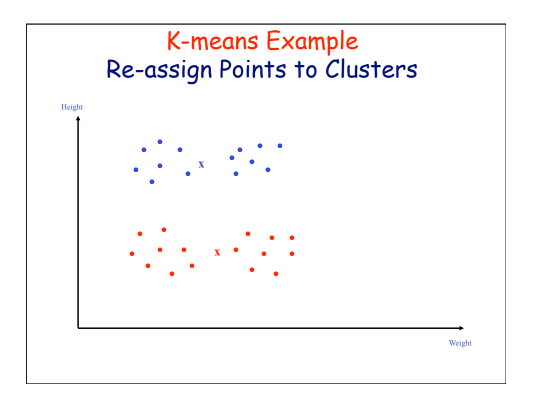


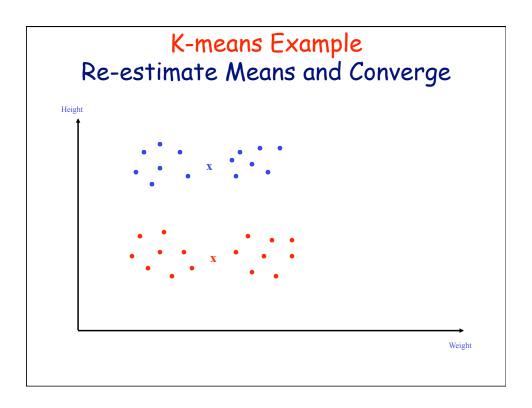


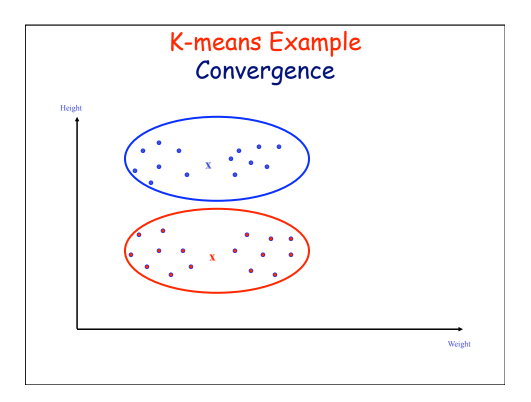










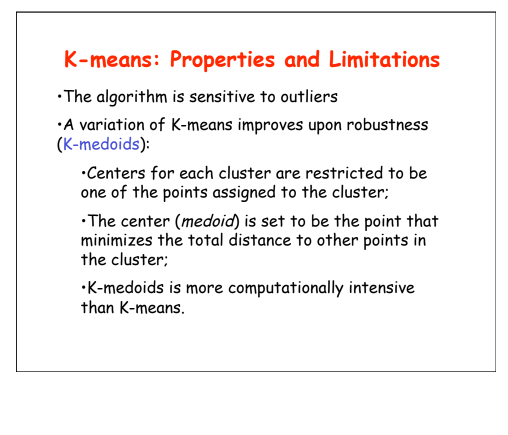


K-means: Properties and Limitations

•The algorithm converges to a local minimum

•The solution depends on the initial partition

•One should start the algorithm with many different random choices for the initial means, and choose the solution having smallest value of the objective function



K-means: Properties and Limitations

•The algorithm requires the number of clusters K;

•Often K is unknown, and must be estimated from the data:

We can test $K \in \{1, 2, \dots, K_{\max}\}$ Compute $\{W_1, W_2, \dots, W_{\max}\}$

In general: $W_1 > W_2 > \cdots > W_{max}$

 K^* = actual number of clusters in the data,

when $K < K^*$, we can expect $W_K >> W_{K+1}$

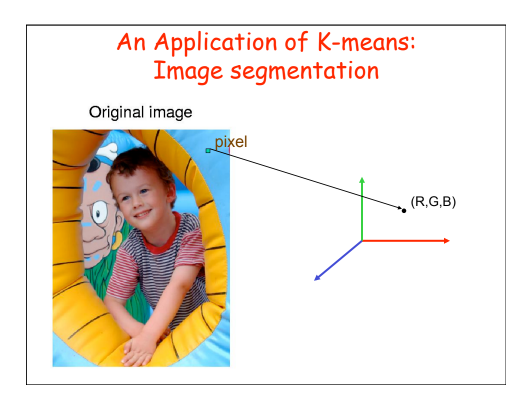
when $K > K^*$, further splits provide smaller decrease of W

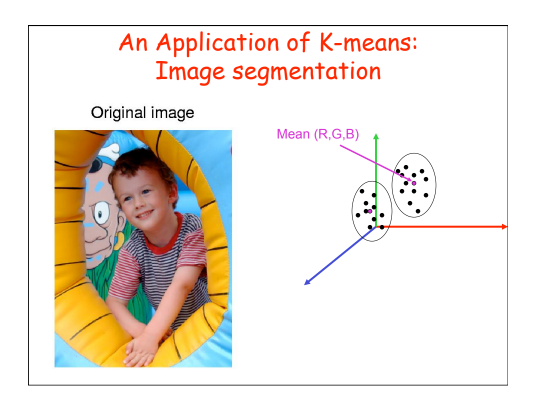
Set \hat{K}^* by identifying an "elbow shape" in the plot of W_k

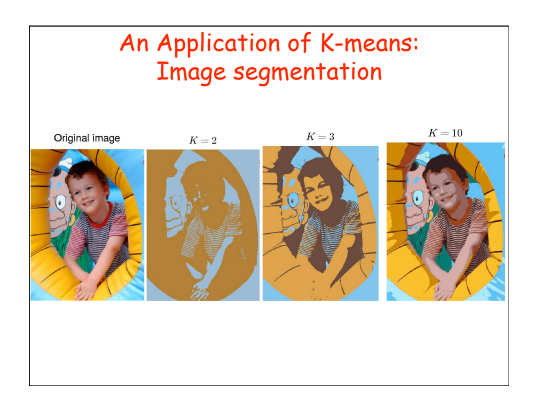
Gap Statistics:Estimating the number of clusters in a
data set via the gap statistic
Tibshirani, Walther, & Hastie, 2001Plot $\log W_{\kappa}$ Plot the curve $\log W_{\kappa}$ obtained from data uniformely distributedEstimate \hat{K}^* to be the point where the gap between the two
curves is largest

An Application of K-means: Image segmentation

- <u>Goal of segmentation</u>: partition an image into regions with homogeneous visual appearance (which could correspond to objects or parts of objects)
- <u>Image representation</u>: each pixel is represented as a three dimensional point in RGB space, where
 - R = intensity of red
 - G = intensity of green
 - B = intensity of blue







An Application of K-means: (Lossy) Data compression

- Original image has N pixels
- Each pixel \rightarrow (R,G,B) values
- Each value is stored with 8 bits of precision
- Transmitting the whole image costs <u>24N bits</u>

Compression achieved by K-means:

- Identify each pixel with the corresponding centroid
- We have K such centroids \rightarrow we need $\log_2 K$ bits per pixel
- · For each centroid we need 24 bits
- Transmitting the whole image costs $24K + N \log_2 K$ bits

<u>Original image</u> = 240x180=43,200 pixels $\rightarrow 43,200x24=1,036,800$ bits Compressed images:

K=2: 43,248 bits

K=3: 86,472

K=10: 173,040 bits