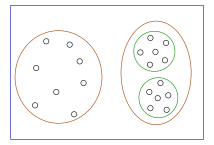
# **Clustering**

**DSE 210** 

#### Widely-used clustering methods

- 1 K-means and its many variants
- 2 EM for mixtures of Gaussians
- 3 Agglomerative hierarchical clustering

## Clustering in $\mathbb{R}^p$



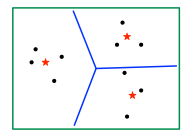
#### Two common uses of clustering:

- Vector quantization
   Find a finite set of representatives that provides good coverage of a complex, possibly infinite, high-dimensional space.
- Finding meaningful structure in data Finding salient grouping in data.

## The *k*-means optimization problem

- Input: Points  $x_1, \ldots, x_n \in \mathbb{R}^p$ ; integer k
- Output: "Centers", or representatives,  $\mu_1,\ldots,\mu_k\in\mathbb{R}^p$
- Goal: Minimize average squared distance between points and their nearest representatives:

$$cost(\mu_1, \dots, \mu_k) = \sum_{i=1}^n \min_j \|x_i - \mu_j\|^2$$

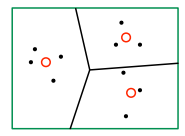


The centers carve  $\mathbb{R}^p$  up into k convex regions:  $\mu_j$ 's region consists of points for which it is the closest center.

#### Lloyd's k-means algorithm

The k-means problem is NP-hard to solve. The most popular heuristic is called the "k-means algorithm".

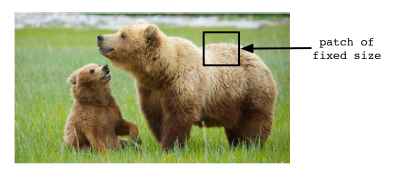
- Initialize centers  $\mu_1, \ldots, \mu_k$  in some manner.
- Repeat until convergence:
  - Assign each point to its closest center.
  - Update each  $\mu_i$  to the mean of the points assigned to it.



Each iteration reduces the cost  $\Rightarrow$  convergence to a local optimum.

# Representing images using k-means codewords

Given a collection of images, how to represent as fixed-length vectors?



- Look at all  $\ell \times \ell$  patches in all images.
- Run k-means on this entire collection to get k centers.
- Now associate any image patch with its nearest center.
- Represent an image by a histogram over  $\{1, 2, \dots, k\}$ .

Such data sets are truly enormous.

#### Initializing the *k*-means algorithm

Typical practice: choose k data points at random as the initial centers.

Another common trick: start with extra centers, then prune later.

A particularly good initializer: k-means++

- Pick a data point x at random as the first center
- Let  $C = \{x\}$  (centers chosen so far)
- Repeat until desired number of centers is attained:
  - Pick a data point x at random from the following distribution:

$$\Pr(x) \propto \operatorname{dist}(x, C)^2$$
,

where 
$$dist(x, C) = min_{z \in C} ||x - z||$$

• Add *x* to *C* 

#### Streaming and online computation

**Streaming computation**: for data sets that are too large to fit in memory.

- Make one pass (or maybe a few passes) through the data.
- On each pass:
  - See data points one at a time, in order.
  - Update models/parameters along the way.
- There is only enough space to store a tiny fraction of the data, or a perhaps short summary.

**Online computation**: an even more lightweight setup, for data that is continuously being collected.

- Initialize a model.
- Repeat forever:
  - See a new data point.
  - Update model if need be.

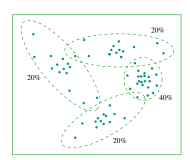
## **Example:** sequential *k*-means

- 1) Set the centers  $\mu_1, \ldots, \mu_k$  to the first k data points
- 2 Set their counts to  $n_1 = n_2 = \cdots = n_k = 1$
- 3 Repeat, possibly foreover:
  - Get next data point x
  - Let  $\mu_j$  be the center closest to x
  - Update  $\mu_i$  and  $n_i$ :

$$\mu_j = rac{n_j \mu_j + x}{n_j + 1}$$
 and  $n_j = n_j + 1$ 

#### Mixtures of Gaussians

Idea: model each cluster by a Gaussian:



Each of the k clusters is specified by:

- a Gaussian distribution  $P_j = N(\mu_j, \Sigma_j)$
- a mixing weight  $\pi_i$

Overall distribution over  $\mathbb{R}^p$ : a **mixture of Gaussians** 

$$Pr(x) = \pi_1 P_1(x) + \cdots + \pi_k P_k(x)$$

## K-means: the good and the bad

#### The good:

- · Fast and easy.
- Effective in quantization.

#### The bad:

 Geared towards data in which the clusters are spherical, and of roughly the same radius.

Is there is a similarly-simple algorithm in which clusters of more general shape are accommodated?

#### The clustering task

Given data  $x_1, \ldots, x_n \in \mathbb{R}^P$ , find the maximum-likelihood mixture of Gaussians: that is, find parameters

- $\pi_1, \ldots, \pi_k \geq 0$  summing to one
- $\mu_1, \ldots, \mu_k \in \mathbb{R}^p$
- $\Sigma_1, \ldots, \Sigma_k \in \mathbb{R}^{p \times p}$

to maximize

$$\begin{aligned} & \text{Pr}\left(\mathsf{data} \mid \pi_{1}P_{1} + \dots + \pi_{k}P_{k}\right) \\ & = \prod_{i=1}^{n} \left(\sum_{j=1}^{k} \pi_{j}P_{j}(x_{i})\right) \\ & = \prod_{i=1}^{n} \left(\sum_{j=1}^{k} \frac{\pi_{j}}{(2\pi)^{p/2}|\Sigma_{j}|^{1/2}} \exp\left(-\frac{1}{2}(x_{i} - \mu_{j})^{T}\Sigma_{j}^{-1}(x_{i} - \mu_{j})\right)\right) \end{aligned}$$

where  $P_i$  is the distribution of the *j*th cluster,  $N(\mu_i, \Sigma_i)$ .

## The EM algorithm

- **1** Initialize  $\pi_1, \ldots, \pi_k$  and  $P_1 = N(\mu_1, \Sigma_1), \ldots, P_k = N(\mu_k, \Sigma_k)$  in some manner.
- 2 Repeat until convergence:
  - Assign each point  $x_i$  fractionally between the k clusters:

$$w_{ij} = \Pr(\text{cluster } j \mid x_i) = \frac{\pi_j P_j(x_i)}{\sum_{\ell} \pi_{\ell} P_{\ell}(x_i)}$$

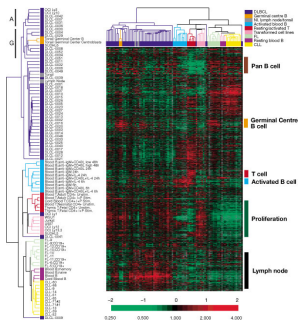
• Now update the mixing weights, means, and covariances:

$$\pi_{j} = \frac{1}{n} \sum_{i=1}^{n} w_{ij}$$

$$\mu_{j} = \frac{1}{n\pi_{j}} \sum_{i=1}^{n} w_{ij} x_{i}$$

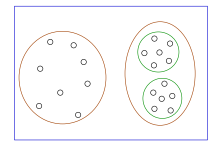
$$\Sigma_{j} = \frac{1}{n\pi_{j}} \sum_{i=1}^{n} w_{ij} (x_{i} - \mu_{j}) (x_{i} - \mu_{j})^{T}$$

# **Example:** gene expression data



#### **Hierarchical clustering**

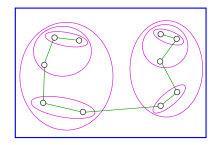
Choosing the number of clusters (k) is difficult.



Often there is no single right answer, because of multiscale structure.

Hierarchical clustering avoids these problems.

## The single linkage algorithm

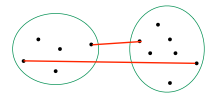


- Start with each point in its own, singleton, cluster
- Repeat until there is just one cluster:
  - Merge the two clusters with the closest pair of points
- Disregard singleton clusters

## **Linkage methods**

- Start with each point in its own, singleton, cluster
- Repeat until there is just one cluster:
  - Merge the two "closest" clusters

How to measure the distance between two clusters of points, C and C'?



• Single linkage

$$\mathsf{dist}(C,C') = \min_{x \in C, x' \in C'} \|x - x'\|$$

Complete linkage

$$\operatorname{dist}(C,C') = \max_{x \in C, x' \in C'} \|x - x'\|$$

#### Average linkage

Three commonly-used variants:

1 Average pairwise distance between points in the two clusters

$$\mathsf{dist}(C, C') = \frac{1}{|C| \cdot |C'|} \sum_{x \in C} \sum_{x' \in C'} \|x - x'\|$$

2 Distance between cluster centers

$$dist(C, C') = ||mean(C) - mean(C')||$$

**3** Ward's method: the increase in k-means cost occasioned by merging the two clusters

$$\mathsf{dist}(\mathit{C},\mathit{C}') = \frac{|\mathit{C}| \cdot |\mathit{C}'|}{|\mathit{C}| + |\mathit{C}'|} \|\mathsf{mean}(\mathit{C}) - \mathsf{mean}(\mathit{C}')\|^2$$