# Clustering

Reference book: Bishop: "Pattern Recognition and Machine Learning" Chapter 9.1

### Outline

- Unsupervised learning
- Clustering
- K-means clustering
- Discussion

#### Supervised Learning

Input: Data X and label y

Goal: Learn how to map X

to y

Examples: Classification,

regression

#### Supervised Learning

Unsupervised Learning

Input: Data X and label y

Input: Just data X, no labels

Goal: Learn how to map X to y

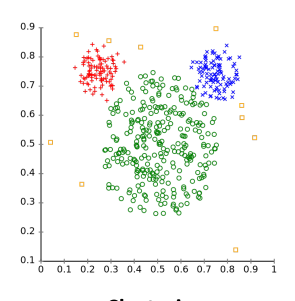
**Goal**: Learn *underlying* structure of the data

**Examples:** Classification, regression

**Examples:** Clustering, Dimensionality reduction

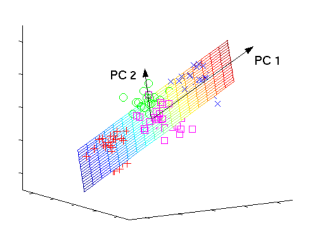
### **Unsupervised Learning**

- No labels/responses. Finding structure in data.
- Dimensionality Reduction.



Clustering

$$T: \mathbb{R}^d \to \{1, 2, \dots, K\}$$

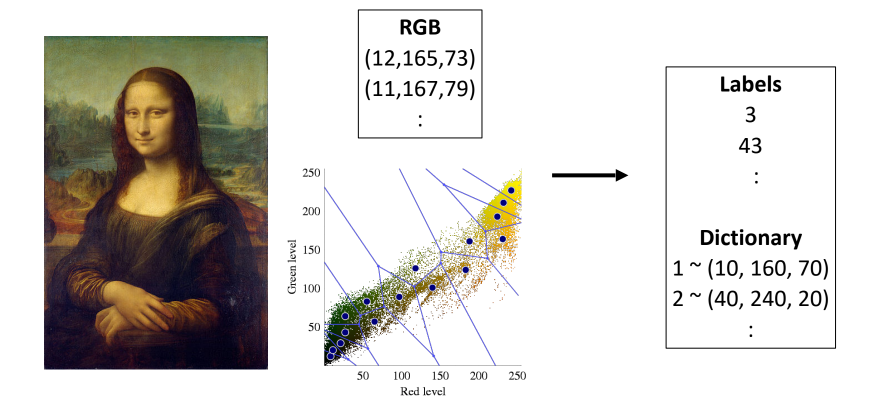


**Subspace Learning** 

$$T: \mathbb{R}^d \to \mathbb{R}^m, \ m < D$$

### Uses of Unsupervised Learning

Data compression



### Uses of Unsupervised Learning

- To improve classification/regression (semi-supervised learning).
- 1. From unlabeled data, learn a good feature  $T: \mathbb{R}^d \to \mathbb{R}^m$ .
- 2. To labeled data, apply a transformation  $T: \mathbb{R}^d \to \mathbb{R}^m$ .

$$(T(x_1), y_1), ..., (T(x_N), y_N)$$

3. Perform classification/regression on transformed low-dimensional data.

## What is clustering?

### What is Clustering?

- Unsupervised learning no information from teacher
- The process of partitioning a set of data into a set of meaningful (hopefully) sub-classes, called clusters
- Cluster:
  - collection of data points that are "similar" to one another and collectively should be treated as a group (it is not the group we learned linear algebra)
  - as a collection, are sufficiently different from other groups

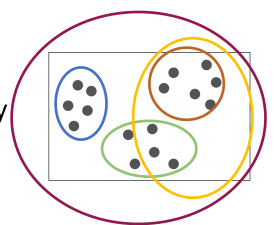
### Basic Clustering Methodology

Hierarchical Algorithms

Cluster the clusters

Agglomerative: pairs of items/clusters are successively linked to produce larger clusters

Divisive (partitioning): items are initially placed in one cluster and then divided into separate groups

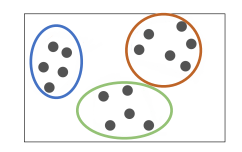


#### Flat Algorithms

Usually start with a random (partial) partitioning of points into groups

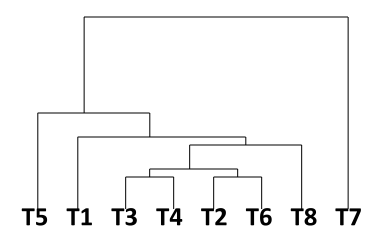
Refine Iteratively

K-Means



#### Hierarchical Agglomerative

- Based on some methods of representing hierarchy of data points
- One idea: hierarchical dendogram (connects points based on similarity)



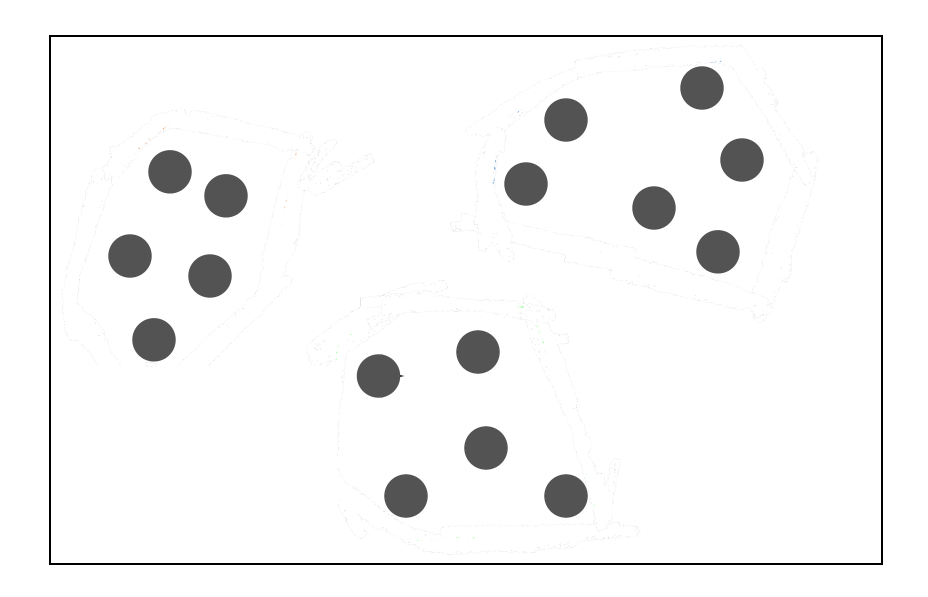
Similarity matrix of items T1 – T8
A larger number means higher similarity

#### Hierarchical Agglomerative

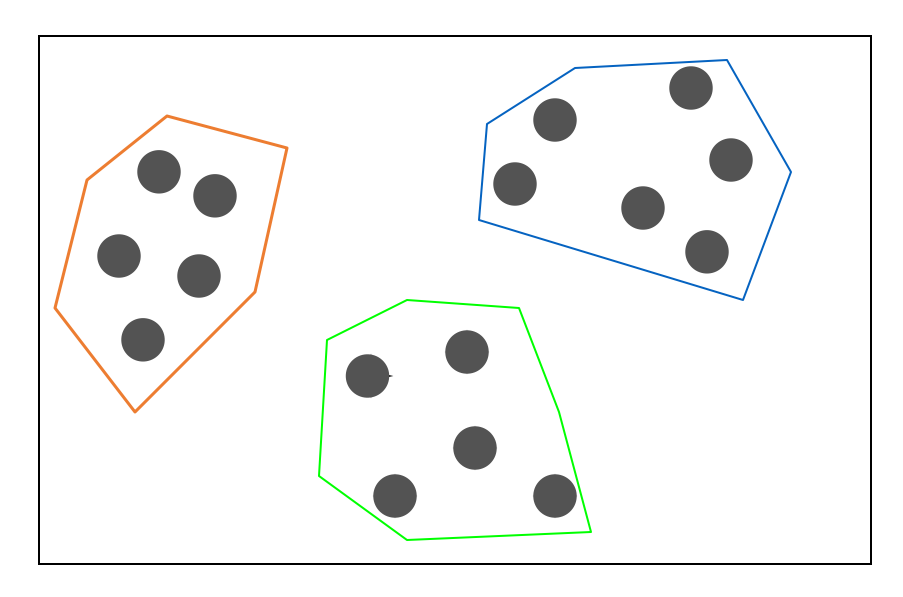
- Compute distance matrix
- Put each data point in its own cluster
- Find most similar pair of clusters
  - merge pairs of clusters (show merger in dendogram)
  - update similarity matrix
  - repeat until all data points are in one cluster

### K-Means

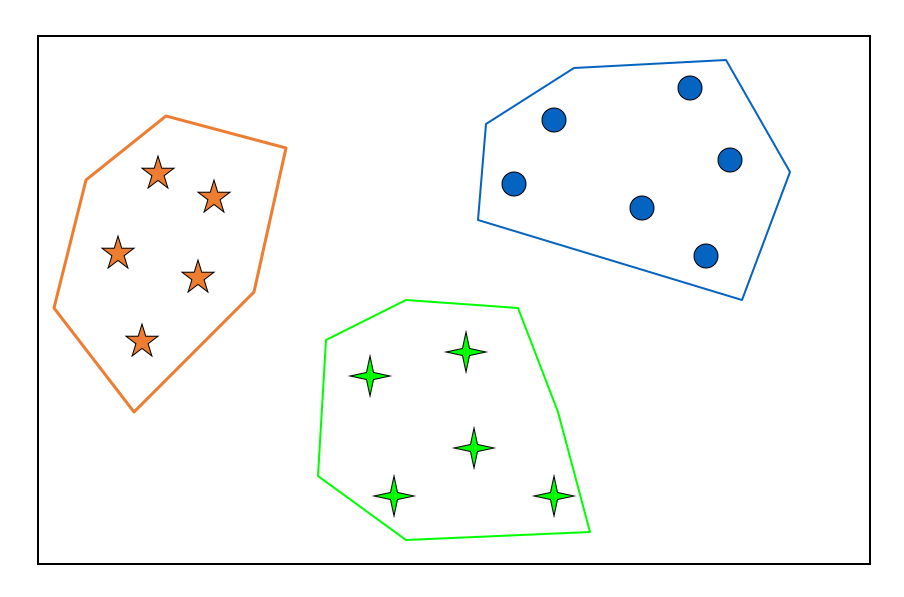
### Clusters



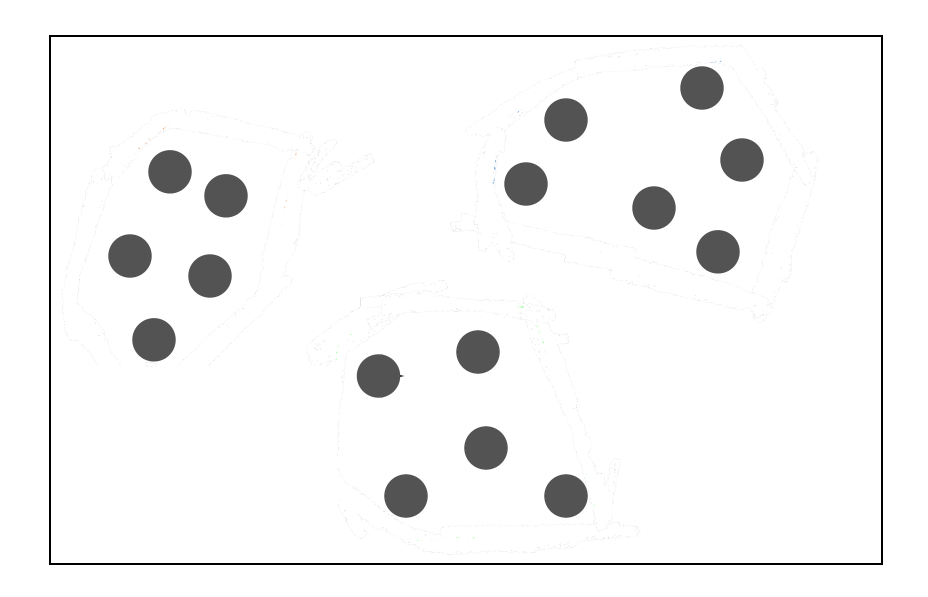
## What your brain sees



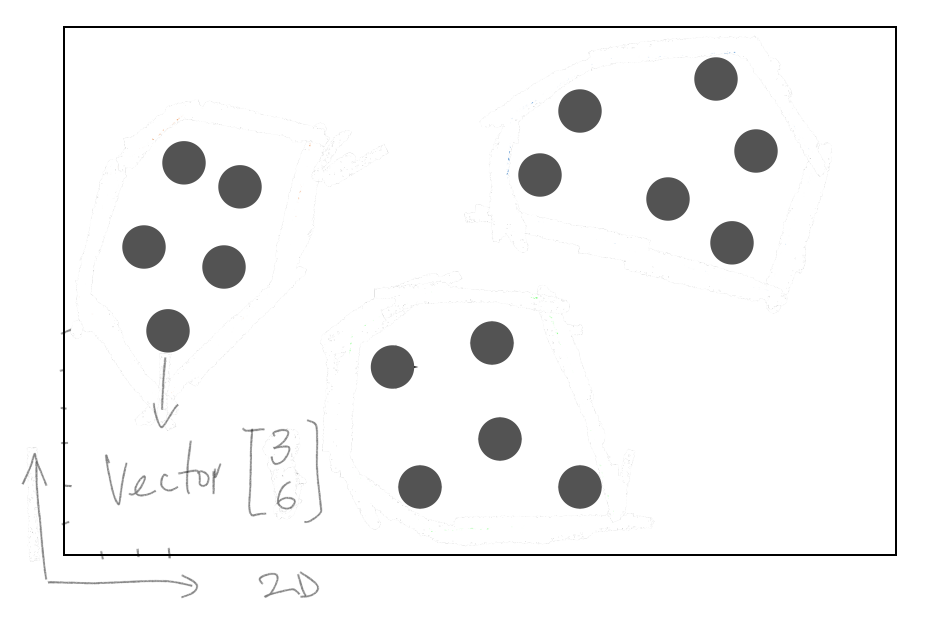
## What your brain sees



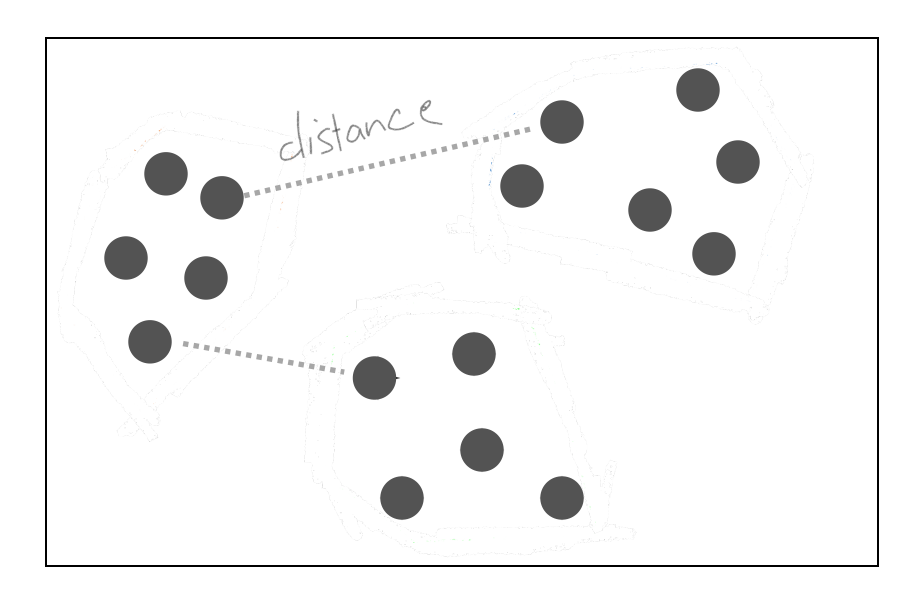
### What the machine sees



### What the machine sees



#### What the machine sees



### Formalising the Problem

Input. Training data  $S_N = \{x_1, x_2, ..., x_N\}, x_n \in \mathbb{R}^d$ . Integer K

Output. Clusters  $\mathscr{C}_1, \mathscr{C}_2, ..., \mathscr{C}_K \subset \{1, 2, ..., N\}$  such that every data point is in one and only one cluster.

Cluster representatives  $\{\mu_1, ..., \mu_K\}$ 

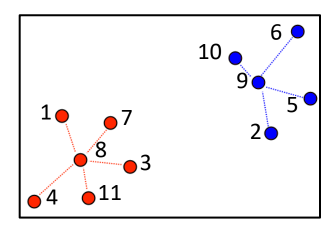
Some clusters could be empty!

### How to Specify a Cluster

By listing all its elements

$$\mathcal{C}_1 = \{1, 3, 4, 7, 8, 11\}$$

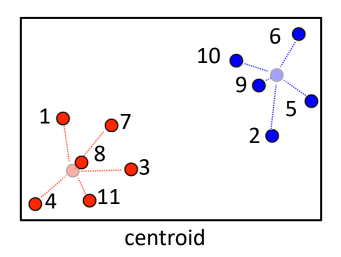
$$\mathcal{C}_2 = \{2, 5, 6, 9, 10\}$$

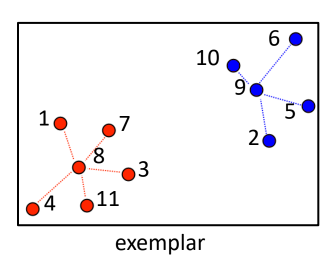


### How to Specify a Cluster

- Using a representative
  - a. A point in center of cluster (centroid)
  - b. A point in the training data (exemplar)

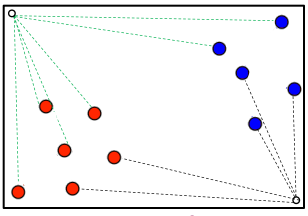
Each point  $x_i$  will be assigned the closest representative.





### **Training Loss**

Sum of squared distances to closest representative.

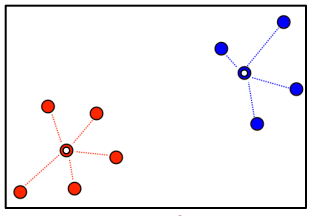


 $loss \approx 11 \times (1)^2 = 11$ 

assume length of each edge is about 1

### **Training Loss**

Sum of squared distances to closest representative (cluster center).



 $loss \approx 9 \times (0.1)^2 = 0.09$ 

assume length of each edge is about 0.1

### **Training Loss**

Optimizing both clusters and representatives.

$$\mathscr{L}(\mathbf{R}, \mathbf{M}; \mathscr{S}_n) = \sum_{n=1}^{N} \sum_{k=1}^{K} r_{nk} \|\mathbf{x}_n - \boldsymbol{\mu}_k\|^2$$

where

$$\mathcal{S}_n = \{ \boldsymbol{x}_1, \boldsymbol{x}_2, ..., \boldsymbol{x}_N \}, \ \boldsymbol{x}_i \in \mathbb{R}^d$$

 $r_{nk} \in \{0,1\}$  denotes which of the K clusters data point  $x_n$  is assigned to. If  $x_n$  is assigned to cluster k then  $r_{nk} = 1$ , and  $r_{nj} = 0$  if  $j \neq k$ .

$$\mathbf{R} = \{r_{nk}\} \in \{0,1\}^{n \times k}, \ n = 1,...,N, \ k = 1,...,K$$

$$\mathbf{M} = [\mu_1, ..., \mu_K]^{\mathrm{T}}$$

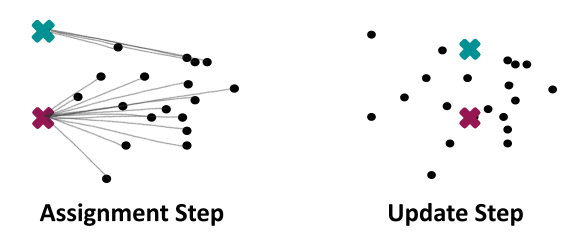
Goal: find values for R and M so as to minimize  $\mathcal{L}$ .

### **Optimization Algorithm**

#### Repeat until convergence:

- Find best clusters given centres
- Find best centres given clusters

$$\mathscr{L}(\mathbf{R}, \mathbf{M}; \mathcal{S}_n) = \sum_{n=1}^{N} \sum_{k=1}^{N} |\mathbf{r}_{nk}| |\mathbf{x}_n - \boldsymbol{\mu}_k||^2$$



#### **Optimization Algorithm**

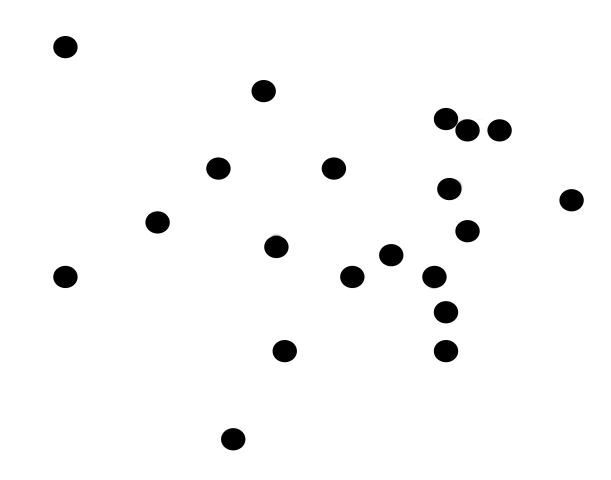
- 1. Initialize centers  $\mu_1, ..., \mu_K$  from the data.
- 2. Repeat until no further change in training loss:
  - a. For each  $n \in \{1, ..., N\}$ ,  $r_{nk} = \begin{cases} 1, & \text{if } k = \arg\min_{j} ||\mathbf{x}_{n} \boldsymbol{\mu}_{j}||^{2} \\ 0, & \text{otherwise.} \end{cases}$

We assign the  $n^{th}$  data point to the closest cluster centre.

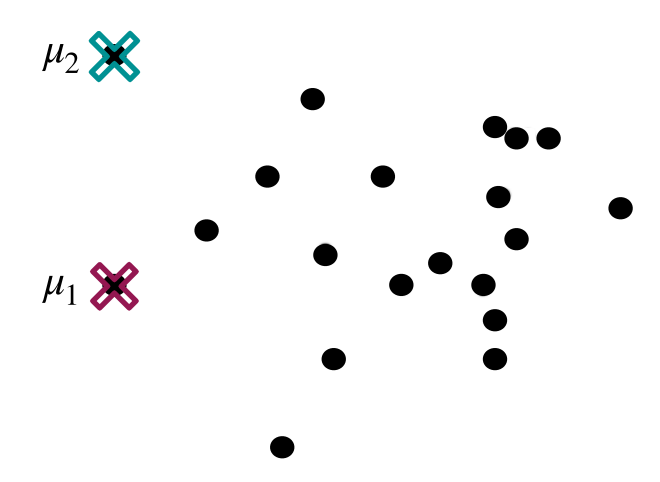
b. For each  $j \in \{1, ..., k\}$ ,  $\mu_j = \frac{\sum_n r_{nj} \mathbf{x}_n}{\sum_n r_{nj}}$ 

We recompute cluster means.

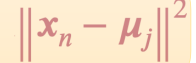
#### Goal: group data in 2 clusters

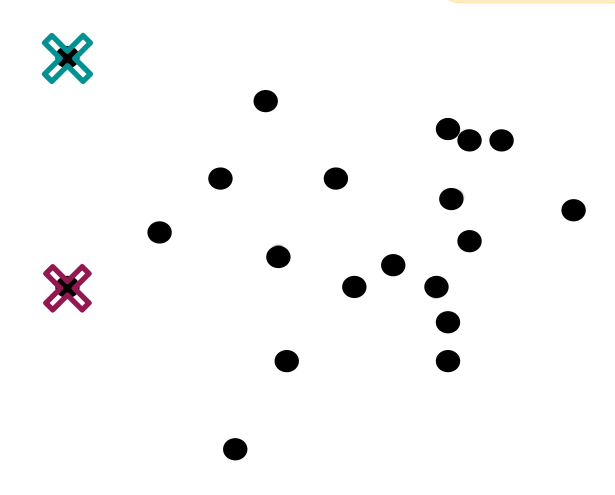


#### 1. Initialize centers from the data.

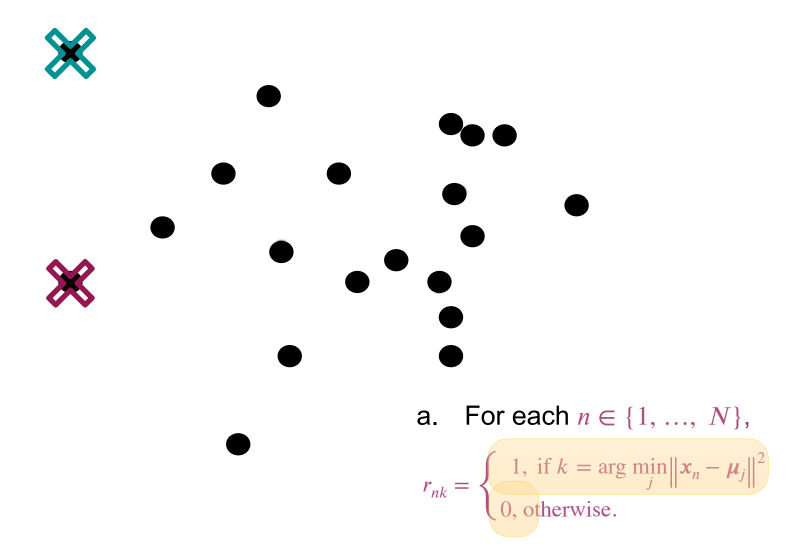


"Call centre": Each point calls to find  $\|\mathbf{x}_n - \boldsymbol{\mu}_j\|^2$ 

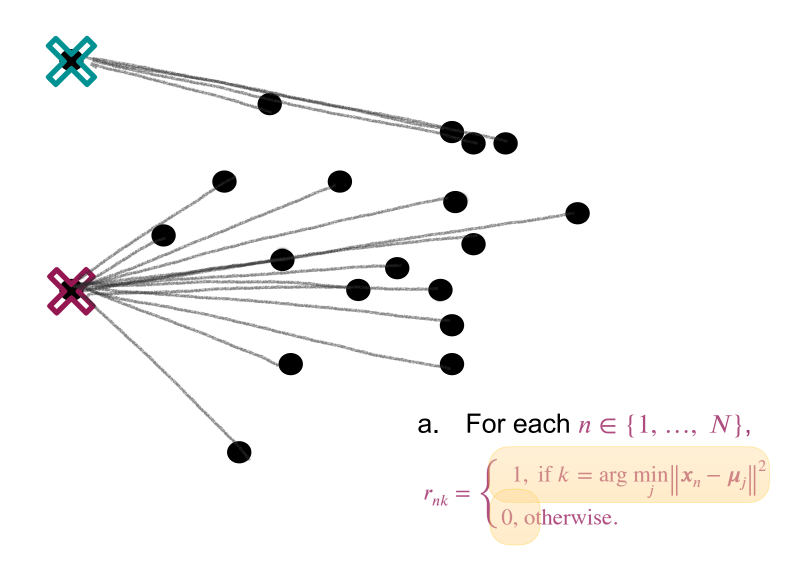




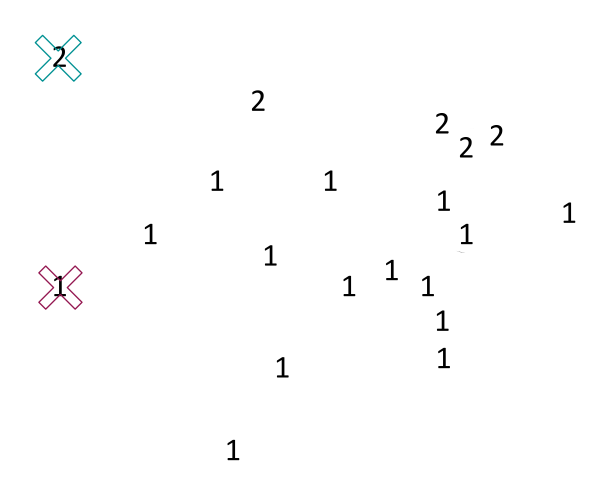
Step 1: Assign points to closest centroid.



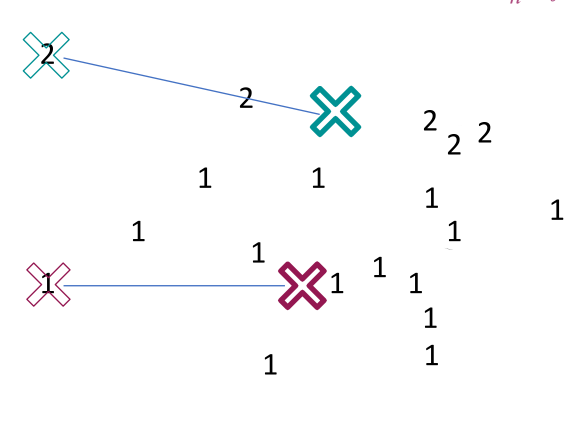
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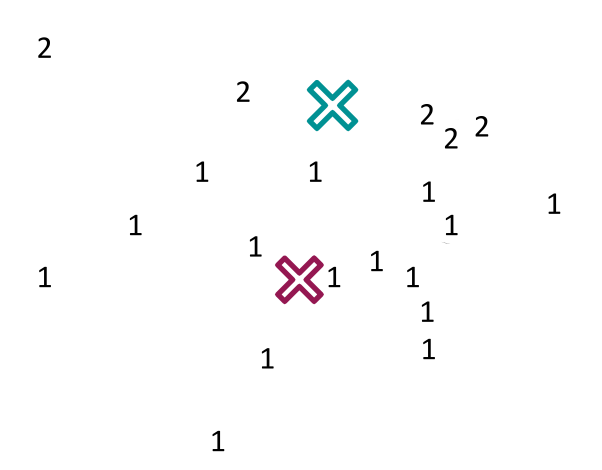
Step 1: Assign points to closest centroid.



Step 2: Compute cluster centres:  $\mu_j = \frac{\sum_n r_{nj} x_n}{\sum_n r_{nj}}$ 



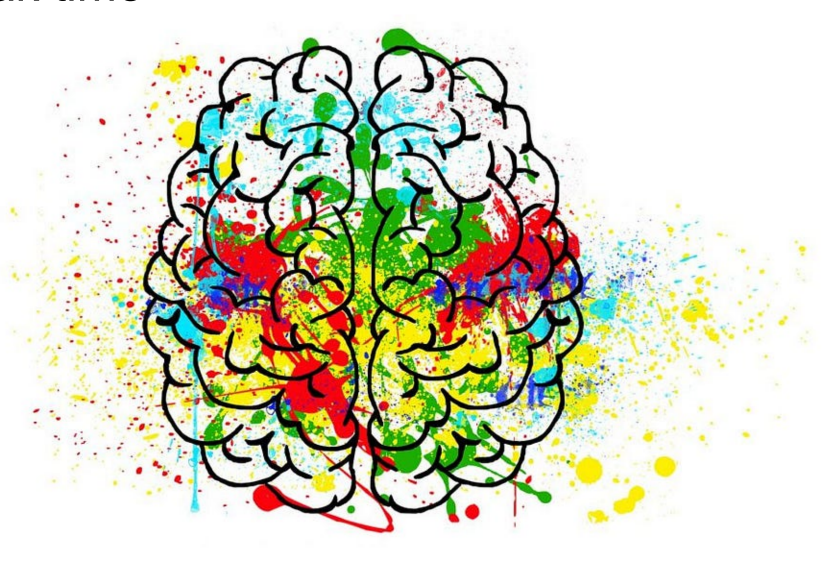
Repeat Step 1: Assign points to closest centroid.



Compute updated cluster centre.

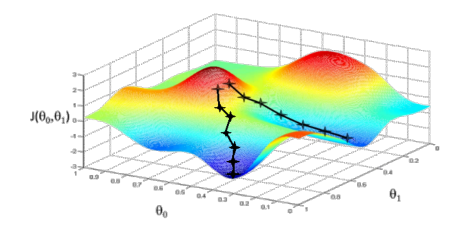
Repeat until convergence.

### Break time



## Convergence

- Training loss always decreases in each step.
- K-Means is guaranteed to converge.
- Convergence usually fast (less than 10-20 iterations).
- Converges to local minimum, not necessarily global minimum.

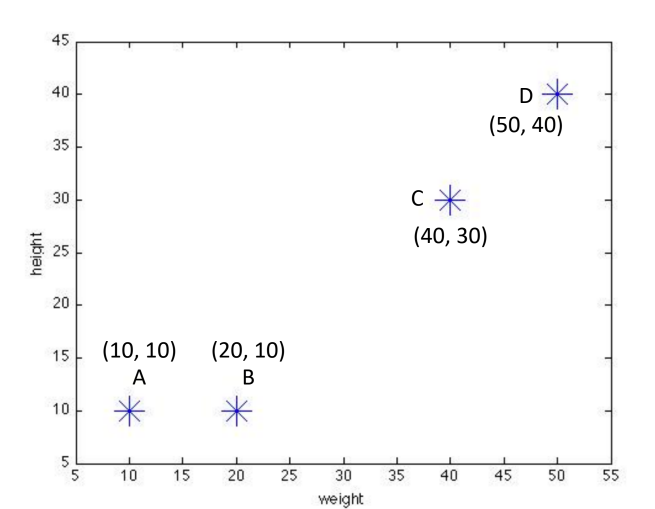


Repeat algorithm over many initial points and pick the configuration with the smallest training loss.

Let's cluster.

### Exercise – k-means clustering

- Suppose we have 4 boxes of different sizes and want to divide them into 2 classes
- Each box represents one point with two attributes (w, h):



- Initial centers: suppose we choose points A and B as the initial centers, so c1 = (10, 10) and c2 = (20, 10)
- Object centre distance: calculate the Euclidean distance between cluster centres and the objects. For example, the distance of object C from the first center is:

$$\sqrt{(40-10)^2 + (30-10)^2} = 36.06$$

We obtain the following distance matrix:

Centre 1	0	10	36.06	50
Centre 2	10	0	28.28	43.43

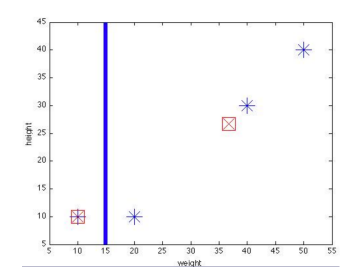
 Object clustering: We assign each object to one of the clusters based on the minimum distance from the centre:

Centre 1
Centre 2

1	0	0	0
0	1	1	1

 Determine centres: Based on the group membership, we compute the new centers

• 
$$c_1 = (10, 10), c_2 = \left(\frac{20 + 40 + 50}{3}, \frac{10 + 30 + 40}{3}\right) = (36.7, 26.7)$$



 Recompute the object-centre distances: We compute the distances of each data point from the new centres:

Centre 1	0	10	36.06	50
Centre 2	31.4	23.6	4.7	18.9

 Object clustering: We reassign the objects to the clusters based on the minimum distance from the centre:

Centre 1	1	1	0	0
Centre 2	0	0	1	1

Determine the new centres:

$$c_1 = \left(\frac{10 + 20}{2}, \frac{10 + 10}{2}\right) = (15, 10)$$
 $c_2 = \left(\frac{40 + 50}{2}, \frac{30 + 40}{2}\right) = (45, 35)$ 

Recompute the object-centres distances:

 Centre 1
 5
 5
 32
 46.1

 Centre 2
 43
 35.4
 7.1
 7.1

Object clustering:

 Centre 1
 1
 1
 0
 0

 Centre 2
 0
 0
 1
 1

 The cluster membership did not change from one iteration to another and so the k-means computation terminates.

## **Evaluating K-Means**

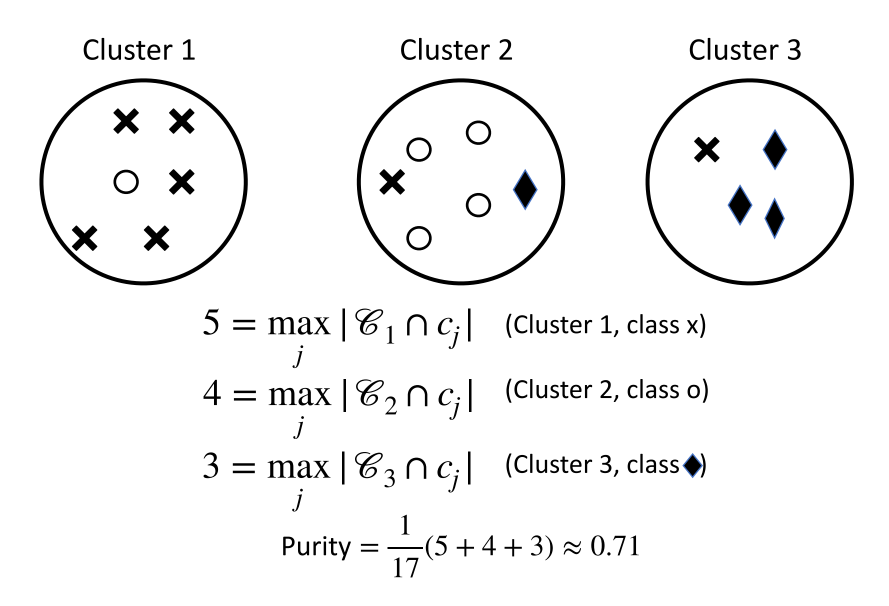
- Internal criteria: the residual sum of squares
- But doesn't evaluate the actual utility of the K-Means to an application
- Extrinsic criteria: evaluate with respect to a humandefined classification

## Extrinsic criterion: Purity

- If we have a set of truth class labels, we can use purity
- For each cluster k, find the class with the most members in the cluster
- Sum them and divide by the total number of points

Purity = 
$$\frac{1}{N} \sum_{k=1}^{K} \max_{j} |\mathscr{C}_k \cap c_j|$$

## Example

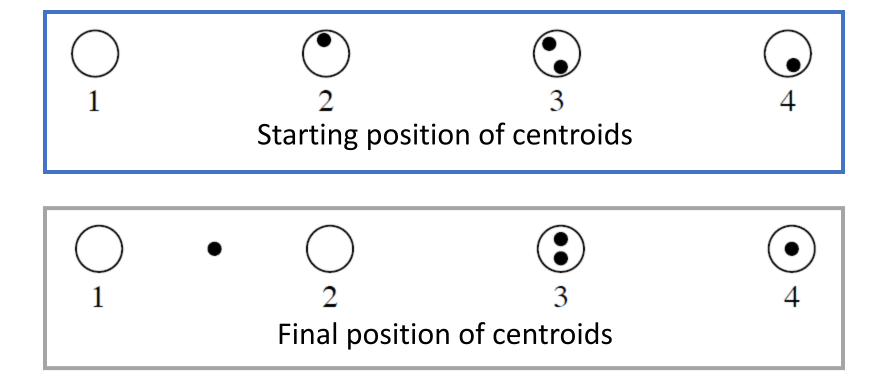


# Discussion

#### Initialization

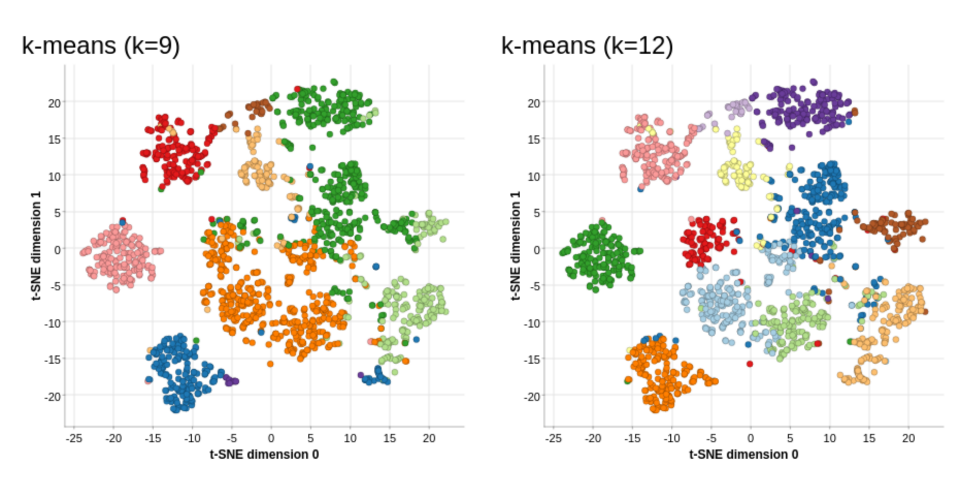
- Empty clusters
  - Pick data points to initialize clusters
- Bad local minima
  - Initialize many times and pick solution with smallest training loss
  - Pick good starting positions

### Initialization



Problem. How to choose good starting positions?Solution. Place them far apart with high probability.

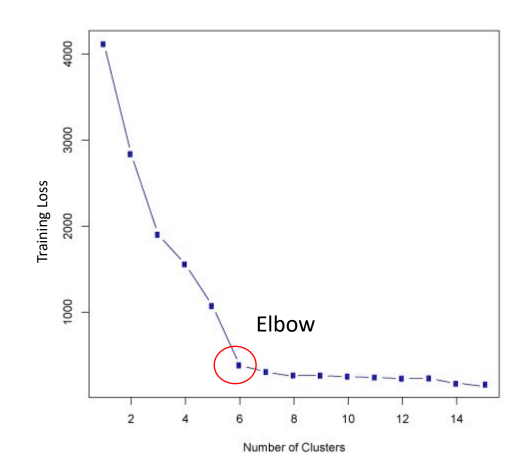
### **Number of Clusters**



### Number of Clusters

How do we choose k, the optimal number of clusters?

- Elbow method
  - Training Loss
  - Validation Loss



## Check your understanding

- Clustering gives you an idea of how the data is distributed.
- You have 1000 data vectors that are very similar to each other (e.g., Eu distance less than 0.0001). We should divide them into a few clusters.
- When you use K-Means, you usually obtain a global minimum of the loss function
- When you use K-Means, you can never achieve global minimum of the loss function.
- The number of clusters is a parameter that can be optimized by K-Means.
- In K-fold cross validation, K is a hyperparameter.
- Clustering is just a different version of classification.

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