

EE2211 Pre-Tutorial 11

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Agenda

- Recap
- Self-learning
- Tutorial 11



Recap

- Introduction of unsupervised learning
- K-means Clustering
 - The most popular clustering technique
- Fuzzy Clustering

Unsupervised Learning

Introduction

Motivation: we do not always have labeled data.

In **unsupervised learning**, the dataset is a collection of **unlabeled examples** $\{\mathbf{x}_i\}_{i=1}^M$.

Evaluation of unsupervised learning is hard:

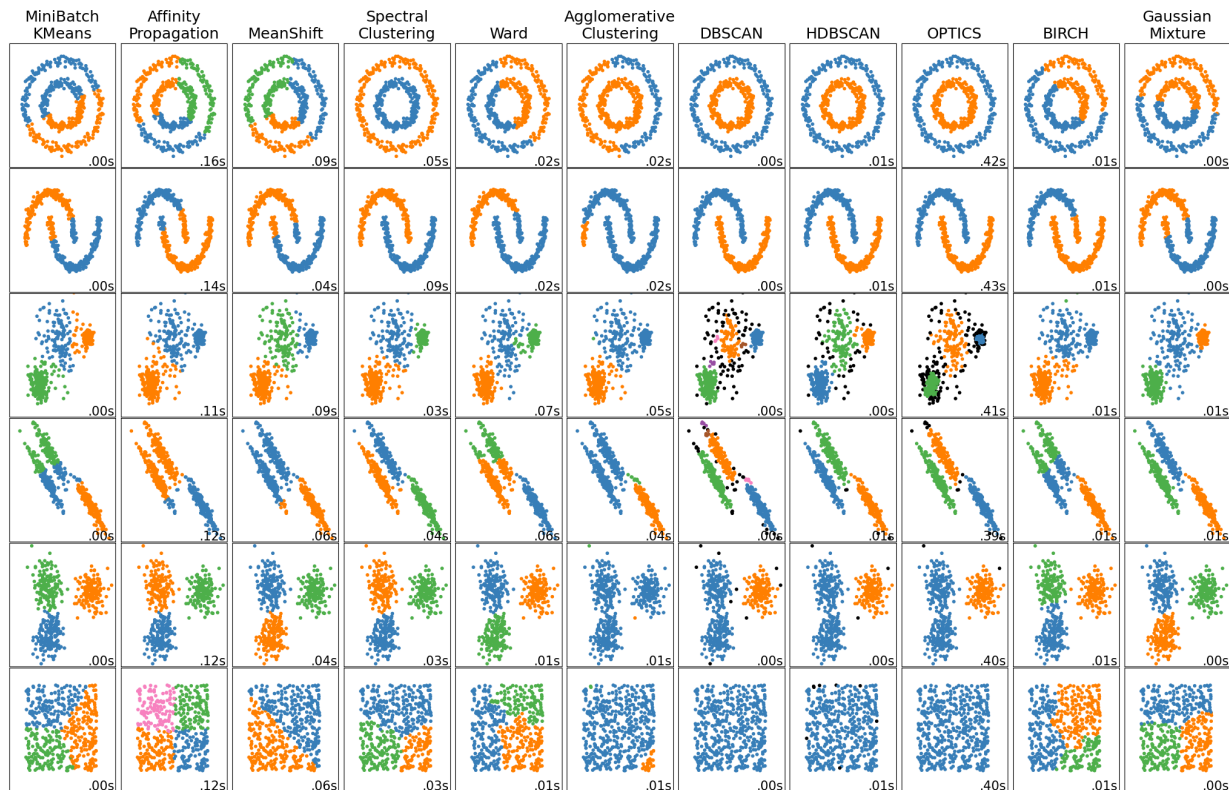
- The **absence of labels** representing the desired behavior for your model means **the absence of a solid reference point to judge the quality of your model**.

Unsupervised Learning

Main Tasks/Approaches

- **Clustering**
 - ✓ Groups a set of objects in such a way that objects in the same group (called a **cluster**) are **more similar** (in some sense) to each other than to those in other groups (clusters).
- **Density Estimation**
 - ✓ Models the probability density function (pdf) of the unknown probability distribution from which the dataset has been drawn.
- **Component Analysis**
 - ✓ Breaks down the data from the perspective of signal analysis.
- **Unsupervised Neural Networks**
 - ✓ Autoencoder

Overview of Clustering Methods





K-Means Clustering

Distance based grouping method

- Feature vectors that are closed to each other

K-means

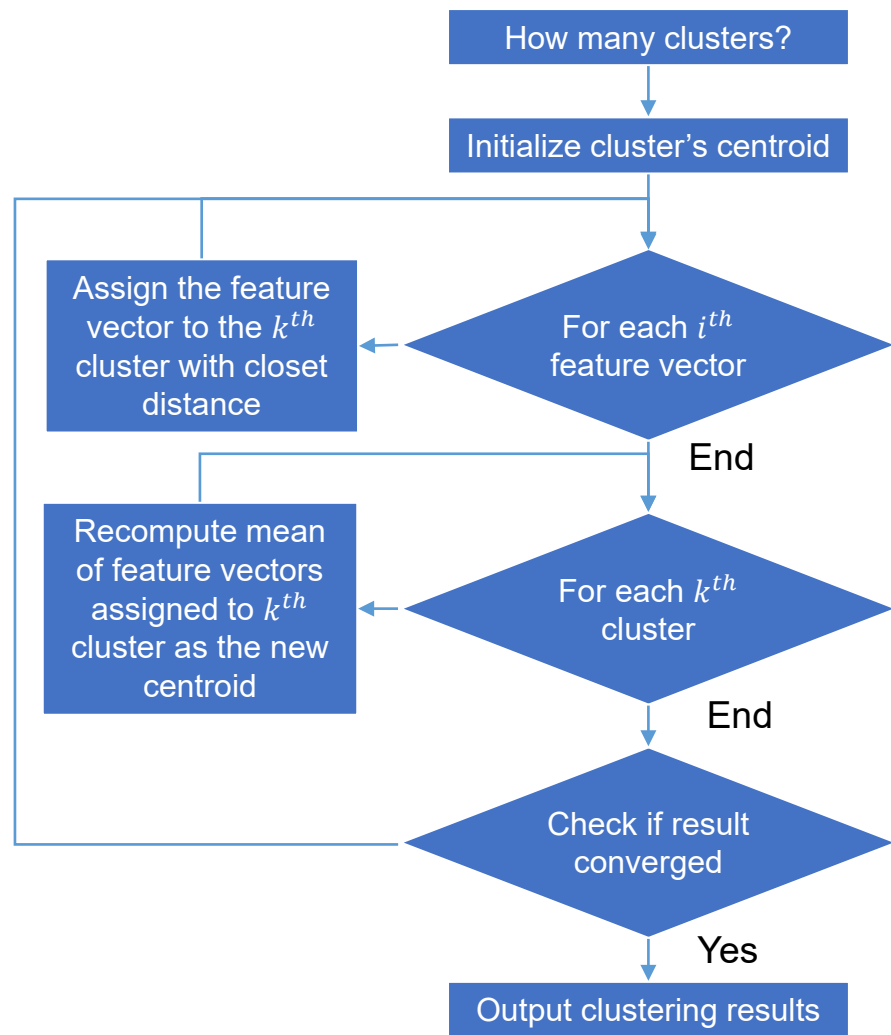
- K = number of clusters
- Means = average or centroid is the average of feature vectors

K-means Clustering

Basic/Naïve K-means Clustering

Looping between
Assignment and Centroid Update

1. First, we **choose** K — the number of clusters. Then we randomly select K feature vectors, called **centroids**, to the feature space.
2. Next, **compute the distance from each example x to each centroid c** using some metric, like the Euclidean distance. Then we **assign the closest centroid to each example** (like if we labeled each example with a centroid id as the label).
3. For each centroid, we **calculate the average feature vector** of the examples labeled with it. These average feature vectors become the **new** locations of the **centroids**.
4. We **recompute** the distance from each example to each centroid, modify the assignment and repeat the procedure until **the assignments don't change after the centroid locations are recomputed**.
5. Finally, we **conclude** the clustering with a list of assignments of centroids IDs to the examples.



K-means Clustering

Optimization Objective Function (within-cluster variance)

Minimize J

m : # of samples; i : index of samples

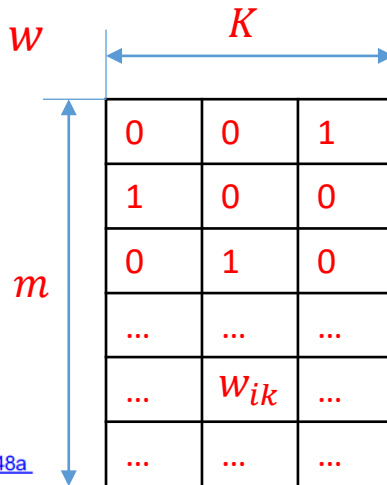
K : # of clusters; k : index of clusters

$$J = \sum_{i=1}^m \sum_{k=1}^K w_{ik} \|\mathbf{x}_i - \mathbf{c}_k\|^2 \quad (1)$$

The term w_{ik} is equal to 1 for data point \mathbf{x}_i if the data point belongs to cluster S_k , else $w_{ik} = 0$.

Note: The optimization objective function was called $\mathcal{C}(\mathbf{w})$ in Lecture 8. Here, we use J (with parameters w_{ik} and \mathbf{c}_k) so that it is differentiated from the centroids \mathbf{c}_k .

$$w_{ik} = \begin{cases} 1 & \text{if } x_i \text{ is assigned to cluster } k \\ 0 & \text{else} \end{cases}$$



	K		
w			
m	0	0	1
	1	0	0
	0	1	0

	...	w_{ik}	...

1st sample is in cluster 3

2nd sample is in cluster 1

Ref: <https://towardsdatascience.com/k-means-clustering-algorithm-applications-evaluation-methods-and-drawbacks-aa03e644b48a>
https://en.wikipedia.org/wiki/K-means_clustering

K-means Clustering

Naïve K-means Algorithm

1. **Assignment Step (fix \mathbf{c} and update w):**

Computing distances to
all centroids

$$\mathbf{x}_i \in S_k \text{ } (w_{ik} = 1) \text{ if } \|\mathbf{x}_i - \mathbf{c}_k\|^2 < \|\mathbf{x}_i - \mathbf{c}_j\|^2 \text{ (else } w_{ik} = 0), \\ i = 1, \dots, m; \quad j, k = 1, \dots, K.$$

Assign the feature vector to
the k^{th} cluster with closet
distance

2. **Update Step (fix w and update \mathbf{c}):**

$$\frac{\partial J}{\partial \mathbf{c}_k} = -2 \sum_{i=1}^m w_{ik} (\mathbf{x}_i - \mathbf{c}_k) = 0 \Rightarrow \mathbf{c}_k = \frac{\sum_{i=1}^m w_{ik} \mathbf{x}_i}{\sum_{i=1}^m w_{ik}}$$

Solving an optimization, i.e., setting derivative to 0

Recompute mean of feature vectors
assigned to k^{th} cluster as the new centroid

Note: $\|\mathbf{x} - \mathbf{c}\| = \sqrt{\sum_{d=1}^D (x_d - c_d)^2}$ is called the Euclidean distance.

where $\mathbf{x} = (x_1, x_2, \dots, x_D)$, $\mathbf{c} = (c_1, c_2, \dots, c_D)$

K-means Clustering

1. **Assignment Step (fix \mathbf{c} and update w):**

$$\mathbf{x}_i \in S_k \text{ (} w_{ik} = 1 \text{) if } \|\mathbf{x}_i - \mathbf{c}_k\|^2 < \|\mathbf{x}_i - \mathbf{c}_j\|^2 \text{ (else } w_{ik} = 0), \\ i = 1, \dots, m; \quad j, k = 1, \dots, K.$$

2. **Update Step (fix w and update \mathbf{c}):**

$$\frac{\partial J}{\partial \mathbf{c}_k} = -2 \sum_{i=1}^m w_{ik} (\mathbf{x}_i - \mathbf{c}_k) = 0 \Rightarrow \mathbf{c}_k = \frac{\sum_{i=1}^m w_{ik} \mathbf{x}_i}{\sum_{i=1}^m w_{ik}}$$

By repeating these two steps, the total loss $J = \sum_{i=1}^m \sum_{k=1}^K w_{ik} \|\mathbf{x}_i - \mathbf{c}_k\|^2$, is **guaranteed to NOT increase (i.e., remain the same or decrease)** until convergence.

Why? **At Step 2:** we compute the new mean, by solving an optimization, i.e., compute the derivative and set to zero, and solve \mathbf{c}_k . This means that, the new \mathbf{c}_k is guaranteed to give a smaller J value.

At Step 1: we only change the assignment, if the distance to the new centroid is smaller! In other words, we either remain in the old group, or change to a new group that is closer (i.e., gives a smaller J)

An Example of K-means

Consider the following unlabelled one-dimensional dataset (so that the samples are all scalar)

$$x_1 = -2, \quad x_2 = 0, \quad x_3 = x_4 = 2.$$

Consider the first initialization

$$c_1^{(1)} = -3, \quad c_2^{(1)} = 3.5$$

Then, once we run the Assignment step, we see that

$$k_1 = k_2 = 1, \quad k_3 = k_4 = 2.$$

This means that samples 1 and 2 are in group one and samples 3 and 4 are in group two. Thus,

$$c_1^{(2)} = -1, \quad c_2^{(2)} = 2.$$

The total cost function is

$$J = 1^2 + 1^2 + 0^2 + 0^2 = 2,$$

which turns out to be the optimum partitioning.

An Example of K-means

Now, instead consider the second initialization

$$c_1^{(1)} = -3, \quad c_2^{(1)} = 2.5$$

Then, once we run the Assignment step, we see that

$$k_1 = 1, \quad k_2 = k_3 = k_4 = 2.$$

Thus,

$$c_1^{(2)} = -2, \quad c_2^{(2)} = 4/3$$

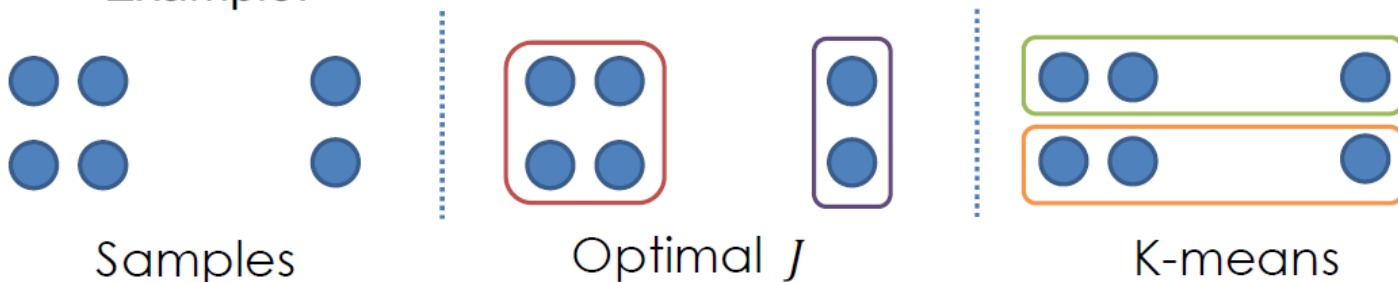
The total cost function is

$$J' = 0^2 + (4/3)^2 + 2(2 - 4/3)^2 = 24/9$$

which is suboptimal and there is no way of improving the cost anymore, i.e., we are stuck. The moral of the story is that initialization is important.

K-means Clustering

- Unfortunately, k-means is not guaranteed to find a global minimum, it finds only local minimum.
- Example:



- Finding the optimal J is NP-hard*
- In practice, k-means clustering usually performs well
- It can be very efficient, and its solution can be used as a starting point for other clustering algorithms

K-means Clustering

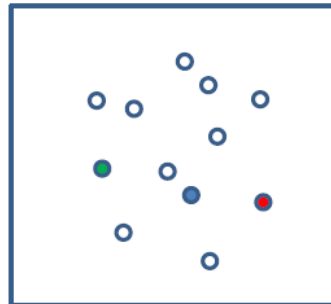
- **Initialization**

Initialization by centroid

Forgy method:

- Randomly chooses k observations from the dataset and uses these as the initial means.

$k=3$

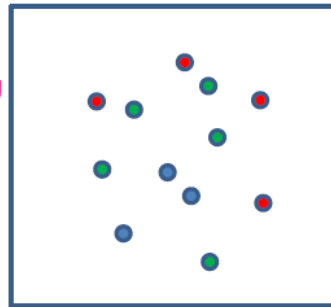


Random partition:

- First randomly assigns a cluster to each observation and then proceeds to the update step, thus computing the initial mean to be the centroid of the cluster's randomly assigned points

Initialization by grouping

$k=3$



Hard vs Soft Clustering

Hard clustering:

Each data point can belong only one cluster, e.g. K-means

- For example, an apple can be red **OR** green (hard clustering)

Soft clustering (also known as Fuzzy clustering):

Each data point can belong to more than one cluster.

- For example, an apple can be red **AND** green (fuzzy clustering)
- Here, the apple can be red to a certain degree as well as green to a certain degree.
- Instead of the apple belonging to green [green = 1] and not red [red = 0], the apple can belong to green [green = 0.3] and red [red = 0.5]. These value are normalized between 0 and 1; however, they do not represent probabilities, so the two values **do not need to add up to 1**.

Hard vs Soft Clustering

Objective Function for Fuzzy C-means

Minimize J

$$J = \sum_{i=1}^m \sum_{k=1}^c (w_{ik})^r \|\mathbf{x}_i - \mathbf{c}_k\|^2$$

$$\text{where } w_{ik} = \frac{1}{\sum_{j=1}^c \left(\frac{\|\mathbf{x}_i - \mathbf{c}_k\|}{\|\mathbf{x}_i - \mathbf{c}_j\|} \right)^{\frac{2}{r-1}}}$$

Each element, $w_{ik} \in [0,1]$, tells the degree to which element, \mathbf{x}_i , belongs to cluster \mathbf{c}_k .

The fuzzifier $r > 1$ determines the level of cluster fuzziness; usually $1.25 \leq r \leq 2$.

$$w_{11} = 0.6$$
$$w_{12} = 0.2$$

\mathbf{x}_1

\mathbf{c}_1

\mathbf{x}_2

\mathbf{c}_2

\mathbf{c}_2

\mathbf{x}_4

$$w_{41} = 0.18$$

$$w_{42} = 0.75$$

- Points **closer** to a centroid \mathbf{c}_k have **higher membership** w_{ik} in that cluster.
- Larger r leads to **softer memberships**; as r approaches 1, memberships become **sharper**, favoring the nearest cluster.



THANK YOU