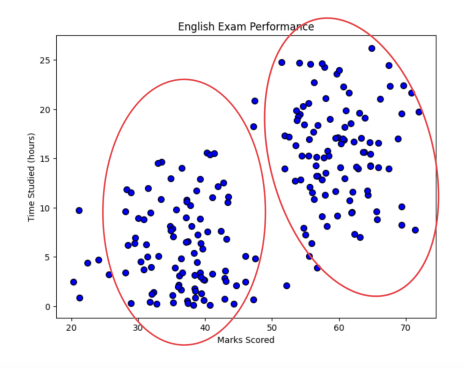
## Clustering

- The process of grouping any kind of data based on the similarity in their features, automatically, without human expertise, is called **clustering**. It is a type of unsupervised learning.
- Intuitively, clustering is dividing a population into groups such that the points in one group are similar to each other. Each group is called a **cluster**.
  - o The points in the same cluster are closer and similar to each other.
  - The points in different clusters are more distant and distinct from each other.
- So, the task in clustering is grouping the points of a similar kind based on our definition of similarity. For example,
  - Given the English exam performance of students



Each group in clustering is called a **cluster**.

- The points in the same cluster are more closer and similar to each other.
- The points in different clusters are more distant and distinct from each other.

So, the task in clustering is **grouping the points of similar kind** based on **our definition of similarity**.

• **Similarity** can be measured using different distance metrics like Euclidean distance, manhattan distance, and Hamming distance.

#### Introduction to K-Means

- The value 'K' in the K-means algorithm denotes the number of clusters.
- In k-means, data is divided into k clusters where each cluster has a centroid which is the average of all the points in the cluster.
- The centroid (C<sub>i</sub>) of the cluster (S<sub>i</sub>) can be defined as

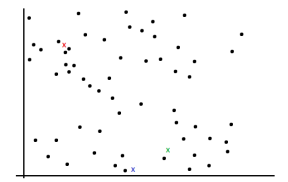
$$C_i = \frac{1}{|S_i|} \sum_{x_i \in S_i} x_j$$

where |S<sub>i</sub>| represents the number of points belonging to the i<sup>th</sup> cluster.

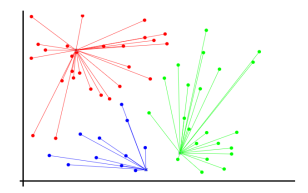
- K-Means assign only one cluster to each point.
- Steps in In K-Means:
  - Every point is assigned to the cluster centroid closest to it.
  - Update the centroid.
  - Repeat the above two steps until convergence.

### Lloyd's algorithm (K-means algorithm)

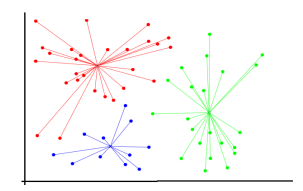
- This algorithm is used to cope with the problem of updating the centers.
- It has 4 basic steps:
  - → Initialization: Randomly initialize k centers from the dataset.



→ **Assignment**: For each point, we find the distance of existing centroids from it and assign the point to that cluster whose centroid has the minimum distance.



→ **Update** the centroids of the clusters by taking the average of points from each cluster.



→ Repeat the previous two steps until convergence (the center of new cluster centroids stops changing their positions).

#### **Mathematical Formulation**

Given the dataset D, Our task is to:

- find the k centroids (C1,C2,...Ck)
- and their corresponding clusters (S<sub>1</sub>, S<sub>2</sub>, S<sub>3</sub>)

such that each datapoint belongs to a cluster.

argmin 
$$\sum_{i=1}^{k} \sum_{x \in S_i} \|x - C_i\|^2$$

C<sub>1</sub>, C<sub>2</sub>, ... C<sub>k</sub>  $\sum_{i=1}^{k} \sum_{x \in S_i} \|x - C_i\|^2$ 

for each databaint belonging to cluster "i" cluster

ie For each cluster and for each databoint belonging cluster; we want to minimize  $||x - C_i||^2$ 

 $||x-C_i||^2$  is nothing but the squared distance between the point and the centroid  $C_i$ 

This optimization problem is very hard to solve and it's not used in real-life applications.

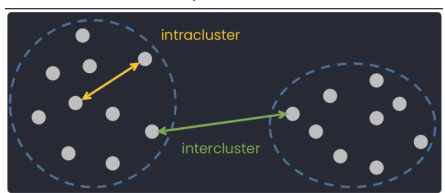
 we optimize with the approximation algorithms and find out the nearest solutions to the problems.

One such approximation algorithm is **Lloyd's algorithm**.

#### **Evaluation**

#### Distances used while clustering:

- Inter-cluster distance represents the distance between two clusters
  - Distance between average values of the clusters.
  - Distance between closest points from the clusters (min distance)
  - Distance between farthest points from the clusters (max distance)
- **Intra-cluster** distance represents the distance within a certain cluster.
  - Average distance between the points of a cluster.
  - Distance between farthest points of a cluster



 Having only one inter or intra-cluster distance won't tell us how good or bad our clusters are, therefore we need a metric to evaluate our clusters.

#### **Dunn Index**

 It is calculated as a ratio of the smallest inter-cluster distance to the largest intra-cluster distance.

i.e. 
$$D = \frac{minimum \ inter-cluster \ distance}{maximum \ intra-cluster \ distance}$$

- The objective of the Dunn index is to identify clusters that are:
  - o compact with a small variance between members of the cluster
  - o and well separated
- A higher Dunn Index means better clustering since observations in each cluster are closer together, while clusters themselves are further away from each other.
- The Dunn Index is **unbound**, so it can only be interpreted in a relative sense.

### Within-cluster sum of squares (WCSS)

Measure of the variability of the data points within each cluster.

$$wcss = \sum_{i=1}^{k} \sum_{j=1}^{m_i} (x_{ij} - c_i)^2$$

where  $\mathbf{x}_{ij}$  is the  $\mathbf{j}^{th}$  point belonging to the  $\mathbf{i}^{th}$  cluster and  $\mathbf{m}_i$  is the number of points in the  $\mathbf{i}^{th}$  cluster.

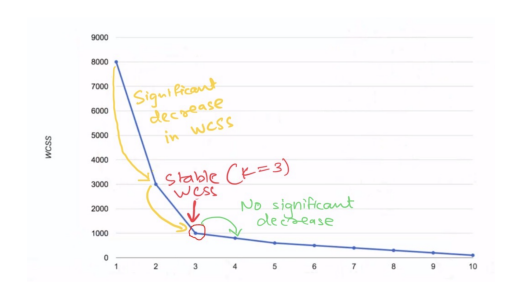
• A variation of the above formula:

$$wcss = \sum_{i=1}^k \sum_{j=1}^{m_i} d(x_{ij}, c_i)$$

where,  $d(x_{ij}, c_i)$  represents a distance metric (any of Euclidean, manhattan, etc.) that calculates the distance between the point  $x_{ij}$  and the centroid  $c_i$  of the cluster.

#### **Elbow method**

- It is a method to determine the optimal number of clusters (**k**) for k-means clustering.
- We perform the k-means clustering for a range of values of k and for each iteration, we calculate the value of the WCSS metric.
- When the value of WCSS is plotted against a range of **k** values, we get a plot that looks like an elbow.



- We can see that the WCSS value decreases as the number of clusters (k) increases.
- At some point on the graph, there is a sharp change in the slope (k = 5) after which the change in slope is very small. The k value corresponding to this point is the optimal K value or an **optimal** number of clusters.
- If we do not get a sharp change in the slope of the elbow plot while using the WCSS metric on the y-axis, we can try using the Silhouette score to get significant results or to get confidence in our decision.

#### Silhouette score

• Measure how similar an object is to its cluster (cohesion) compared to other clusters (separation).

$$S(x_i) = \frac{b-a}{max(b,a)}$$

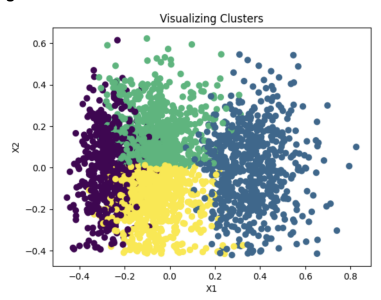
where  $\mathbf{a}$  = average distance of point  $\mathbf{x}_i$  from points in its own cluster and,  $\mathbf{b}$  = average distance of point  $\mathbf{x}_i$  from all the points of the nearest cluster.

- The range of the Silhouette score is [-1, 1].
  - → A Silhouette score near +1 indicates that the sample is far away from its neighboring cluster.
  - → A value near 0 represents overlapping clusters with samples very close to the decision boundary of the neighboring clusters.
  - → A Silhouette score of -1 indicates that the samples have been assigned to the wrong clusters.

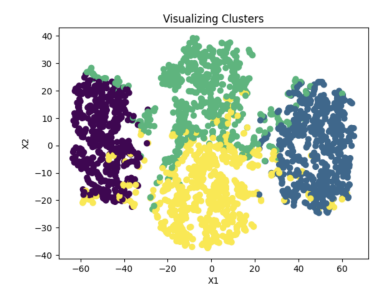
#### **Qualitative Evaluation**

It involves evaluating the clusters by visualizing them.

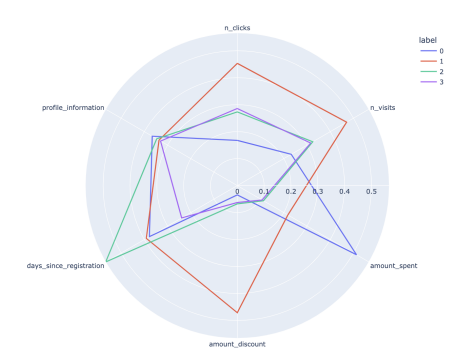
#### Visualizing using PCA:



### Visualizing using tSNE:



### Using polar plot for feature-level insights:



# **Time Complexity**

The time complexity of Kmeans means is:

O(n\*k\*d\*i)

n : number of datapoints k : number of clusters

d : number of dimension of a datapoint

i: number of iterations