## Handmade K-Means Clustering

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## **THEORY**

The aim of the K-Means Clustering algorithm is to create K groups or clusters in data using its features  $x = \{x_1, \ldots, x_P\}$ . This is an unsupervised machine learning algorithm because the aim is not to predict a target variable y, just exploring the data.

Firstly, let's see the main steps of this simple unsupervised machine learning algorithm.

- 1. Define the input data  $x_{[NxP]}$  and K or number of desired clusters.
- 2. For each cluster K initialize its *centroid*, which means the center of each cluster as a P dimensional vector.

$$C_k = [C_1^k, \dots, C_P^k]$$
 with  $1 \le k \le K$ 

- 3. For each N observation in the data  $x_{[NxP]}$ , compute the mathematical distance (*Euclidean* in this paper) to each K centroid.
- 4. For each N observation in the data  $x_{[NxP]}$ , assign it to the cluster with the nearest centroid (lower mathematical distance). Then, the result is K disjoint splits of the data x.

$$x = \bigcup_{k=1}^{K} x^k \quad with \quad x^k = \{x_1^k, \dots, x_P^k\}$$

5. For each K clusters, update its *centroid* as

$$C_k^{new} = [mean(x_1 \in x^k), \dots, mean(x_P \in x^k)]$$
 with  $1 \le k \le K$ 

6. Repeat steps 3, 4 and 5 until no relocations occur in step 4. It means when all the observations in the data are assigned to the same cluster than the actual and then any *centroid* change.

## CODE

The data used is the popular dataset *iris*. Let's create 3 groups or clusters in data using its features: Petal.Length and Petal.Width. In the paper Handmade\_KNN.pdf, the KNN algorithm tries to predict the variable Species in *iris* dataset (which has 3 categories) using the same features. But now the aim is to create 3 clusters only using the features (without Species), and let's see if the observations in each cluster correspond to the same category in Species variable.

Note: It is important to normalize (every observation minus the feature minimum value divided by the feature range) the features so that their units do not affect the mathematical distance. For example, if you compute the mathematical Euclidean distance from a person A = (1.80m, 70kg) and another B = (1.9m, 80kg) the result is

$$distance(A,B) = \sqrt{(1.8-1.9)^2 + (70-80)^2} = 10$$

But if you change the units to A = (180cm, 70000g) and B = (190cm, 80000g) the result is

$$distance(A, B) = \sqrt{(180 - 190)^2 + (70000 - 80000)^2} = 10000$$

Let's define the input data  $x_{[NxP]}$ .

```
duplicates <- duplicated(iris[,c("Petal.Length","Petal.Width")])
x <- iris[!duplicates,c("Petal.Length","Petal.Width")]
x[,1] <- (x[,1]-min(x[,1]))/(max(x[,1])-min(x[,1]))
x[,2] <- (x[,2]-min(x[,2]))/(max(x[,2])-min(x[,2]))</pre>
```

Let's create the *KMeans* algorithm function.

```
KMeans <- function(x,K,centroids){</pre>
  # Define function for mathematical distance between observations
  euclidean <- function(a,b){</pre>
    return(sqrt(sum((a-b)^2)))
  # Iteration until no relocations occur
  clusters <- rep(0,nrow(x))</pre>
  any_relocation <- TRUE
  while(any_relocation){
    any_relocation <- FALSE</pre>
    # Iterate all data observations
    distances <- matrix(NA,nrow=nrow(x),ncol=K)
    for(i in 1:nrow(x)){
      # Iterate all clusters and compute the distance from its centroid to observation i
      for(k in 1:K){
        distances[i,k] <- euclidean(x[i,],centroids[k,])</pre>
      # Assign the data observation i to the cluster with the nearest centroid (if change)
      nearest <- which.min(distances[i,])</pre>
      if(nearest!=clusters[i]){
        clusters[i] <- nearest</pre>
        any relocation <- TRUE
      }
    }
    # Update centroids
    for(k in 1:K){
      for(p in 1:ncol(x)){
        centroids[k,p] <- mean(x[clusters==k,p])</pre>
      }
    }
  }
  return(clusters)
```

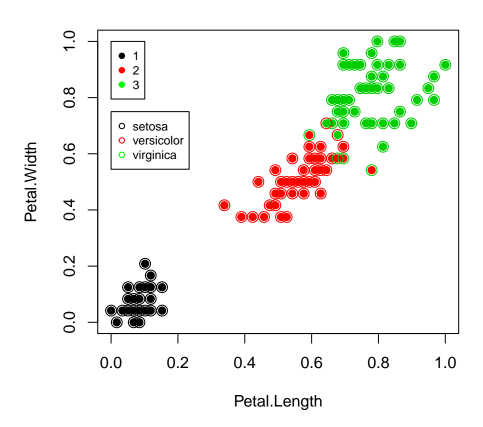
Now, let's apply the *KMeans* function with K=3. Note that we need to initialize the *centroids*, and we will use the means of *Petal.Length* and *Petal.Width* by categories in *Species* (3 equal to K).

```
K <- 3
centroids <- matrix(NA,nrow=K,ncol=ncol(x))
species <- iris[!duplicates, "Species"]
for(k in 1:K){
   for(p in 1:ncol(x)){
      centroids[k,p] <- mean(x[species==unique(species)[k],p])
   }
} clusters <- KMeans(x,K,centroids)</pre>
```

Once the clusters of each data observation are calculated, let's plot the clusters and also, the category of each observation in the variable Species (contouring by colours every observation point).

```
plot(x[,1],x[,2],col=clusters,pch=19,xlab="Petal.Length",ylab="Petal.Width",main="3-Means")
points(x[,1],x[,2],col=species,pch=1,cex=1.5)
legend(0,1,legend=unique(clusters),col=c("black","red","green"),pch=19,cex=0.75)
legend(0,0.75,legend=unique(species),col=c("black","red","green"),pch=1,cex=0.75)
```

## 3-Means



In conclusion, we can observe that the clusters created by the *K-Means* algorithm, using the features *Petal.Length* and *Petal.Width*, are quite similar to the distribution of the variable *Species*. It seems that, except some observation with different point and contour colour, the cluster 1 corresponds to the category *setosa*, the cluster 2 corresponds to the category *versicolor*, and the cluster 3 corresponds to the category *virginica*.

Note: Remember that the *KMeans* algorithm only uses the features *Petal.Length* and *Petal.Width*. Once the clusters are computed, we analyze them comparing with the variable *Species*, since it is an *unsupervised* machine learning technique that just explores data instead of predicting a target variable.