

Introduction to Machine Learning

Clustering

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Hierarchical Clustering

HIERARCHICAL CLUSTERING

Hierarchical clustering is a recursive process that builds a hierarchy of clusters. We distinguish between:

- Agglomerative (or bottom-up) clustering:
 - Start: Each observations is an individual cluster.
 - Repeat: Merge the two closest clusters.
 - Stop when there is only one cluster left.
- 2 Divisive (or top-down) clustering:
 - Start: All observations are within one cluster.
 - Repeat: Divide the cluster that results in two clusters with biggest distance.
 - Stop when each observation is an individual cluster.

HIERARCHICAL CLUSTERING

Let $X_1, ..., X_N$ be observations with P features (dimensions), where $X_i = (x_{i1}, ..., x_{iP})^{\top}$.

A data set is a (N \times P)-matrix of the form:

	feature 1			feature P
<i>X</i> ₁	<i>X</i> ₁₁			X _{1P}
÷	:	÷	÷	:
X_N	x_{N1}			x_{NP}

HIERARCHICAL CLUSTERING

For hierarchical clustering, we need a definition for

- distances $d(X_i, X_j)$ between two observations X_i and X_j :
 - manhattan distance:

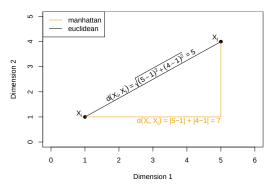
$$d(X_i, X_j) = ||X_i - X_j||_1 = \sum_{k=1}^P |x_{ik} - x_{jk}|$$

euclidean distance:

$$d(X_i, X_j) = ||X_i - X_j||_2 = \sqrt{\sum_{k=1}^{P} (x_{ik} - x_{jk})^2}$$

• distances between two clusters (called linkage).

DISTANCES BETWEEN OBSERVATIONS



- manhattan: sum up the absolute distances in each dimension. In R: dist(data, method = "manhattan")
- euclidean: remember Pythagoras theorem from school? In R: dist(data, method = "euclidean")
- gower: can be used for mixed variables (categorical and numeric). In R: gower_dist() from the gower package
- see ?dist for other distances.

GOWER DISTANCE I

- The Gower's metric calculates the distance between observations X_i and X_j for each feature separately and based on its data type (i.e., categorical or numeric).
- For a categorical feature X_k , the distance between the *i*-th and the *j*-th observation X_{ik} and X_{jk} is defined by

$$s_{ijk} = \begin{cases} 0 \text{ if } X_{ik} = X_{jk} \\ 1 \text{ if } X_{ik} \neq X_{jk}. \end{cases}$$

• For a numerical feature X_k , the distance between the *i*-th and the *j*-th observation X_{ik} and X_{jk} is defined by

$$s_{ijk} = \frac{|X_{ik} - X_{jk}|}{\max(X_k) - \min(X_k)}, \quad \text{so that } 0 \le s_{ijk} \le 1.$$

GOWER DISTANCE II

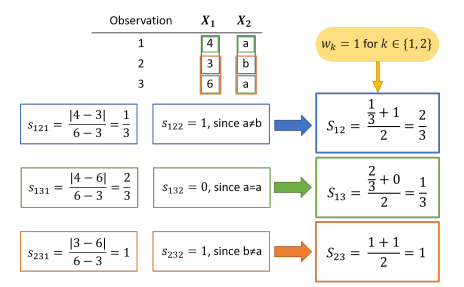
The Gower's metric S combines all individual distances of each feature by

$$S_{ij} = \frac{\sum_{k=1}^{P} w_k s_{ijk}}{\sum_{k=1}^{P} w_k},$$

where

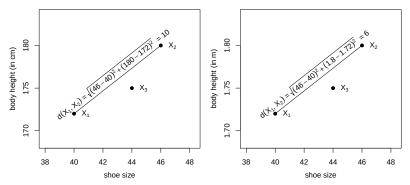
- P: number of features.
- w_k : weight for feature k (typically $w_k = 1$).
- s_{ijk} : the difference (distance) between X_{ik} and X_{jk} , i.e. the *i*-th and *j*-th observation of feature k.

GOWER DISTANCE III - EXAMPLE



DISTANCES BETWEEN OBSERVATIONS

It is often a good idea to *normalize* the data before computing distances, especially when the scale of features is different, e.g.:



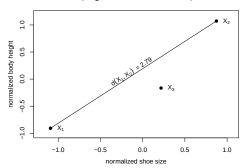
On the right plot, the distance is dominated by 'shoe size'.

DISTANCES BETWEEN OBSERVATIONS

The normalized feature $\tilde{X}_{\text{height}}$ is computed using X_{height} by <!-Normalization of the height feature means: ->

$$ilde{X}_{ ext{height}} = rac{X_{ ext{height}} - ext{mean}(X_{ ext{height}})}{ ext{sd}(X_{ ext{height}})}.$$

Distances based on normalized data are better comparable and robust in terms of linear transformations (e.g. unit conversion).



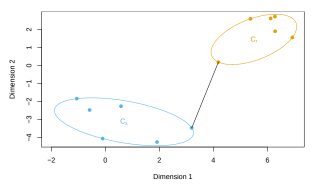
DISTANCES BETWEEN CLUSTERS (LINKAGE)

- Assume that all observations X_1, \ldots, X_N belong to K < N different clusters.
- The linkage of two clusters C_r and C_s is a "score" describing their distance.

The most popular and simplest linkages are

- Single Linkage
- Complete Linkage
- Average Linkage
- Centroid Linkage

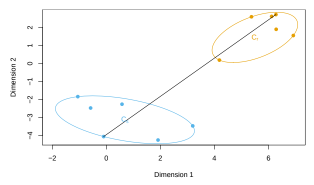
SINGLE LINKAGE



Single linkage defines the distance of the *closest point pairs* from different clusters as the distance between two clusters:

$$d_{\mathsf{single}}(C_r, C_s) = \min_{i \in C_r, j \in C_s} d(X_i, X_j)$$

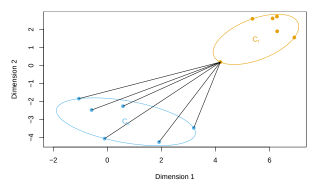
COMPLETE LINKAGE



Complete linkage defines the distance of the *furthest point pairs* of different clusters as the distance between two clusters:

$$d_{ ext{complete}}(C_r, C_s) = \max_{i \in C_r, j \in C_s} d(X_i, X_j)$$

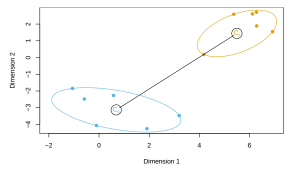
AVERAGE LINKAGE



(Note: Plot only shows distances between all green points and one red point)

In average linkage, the distance between two clusters is defined as the average distance across *all* pairs of two different clusters.

CENTROID LINKAGE



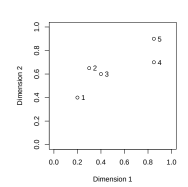
Centroid linkage defines the distance between two clusters as the distance between the two cluster centroids. The centroid of a cluster C_s with N_s points is the mean value of each dimension:

$$\bar{X}_s = \frac{1}{N_s} \sum_{i \in C_s} X_i$$

Agglomerative hierarchical clustering starts with all points forming their own cluster and iteratively merges them until all points form a single cluster containing all points.

Example:

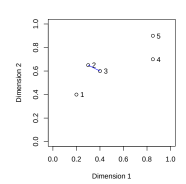
Step 1: {1}, {2}, {3}, {4}, {5}



Agglomerative hierarchical clustering starts with all points forming their own cluster and iteratively merges them until all points form a single cluster containing all points.

Example:

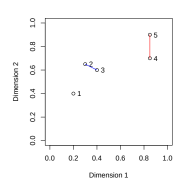
Step 1: {1}, {2}, {3}, {4}, {5} Step 2: {1}, {2,3}, {4}, {5}



Agglomerative hierarchical clustering starts with all points forming their own cluster and iteratively merges them until all points form a single cluster containing all points.

Example:

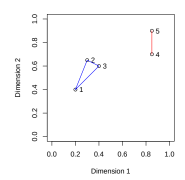
Step 1: {1}, {2}, {3}, {4}, {5} Step 2: {1}, {2,3}, {4}, {5} Step 3: {1}, {2,3}, {4,5}



Agglomerative hierarchical clustering starts with all points forming their own cluster and iteratively merges them until all points form a single cluster containing all points.

Example:

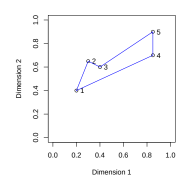
```
Step 1: {1}, {2}, {3}, {4}, {5}
Step 2: {1}, {2,3}, {4}, {5}
Step 3: {1}, {2,3}, {4,5}
Step 4: {1,2,3}, {4,5}
```



Agglomerative hierarchical clustering starts with all points forming their own cluster and iteratively merges them until all points form a single cluster containing all points.

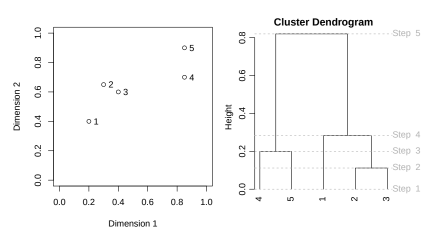
Example:

```
Step 1: {1}, {2}, {3}, {4}, {5}
Step 2: {1}, {2,3}, {4}, {5}
Step 3: {1}, {2,3}, {4,5}
Step 4: {1,2,3}, {4,5}
Step 5: {1,2,3,4,5}
```



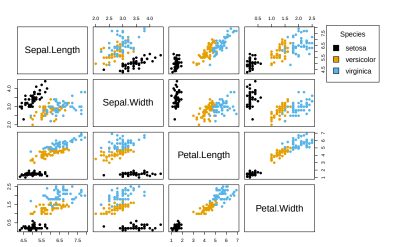
DENDROGRAM

A Dendrogram is a tree showing which clusters / observations are merged after each step. The 'height' is proportional to the distance between the two merged clusters:

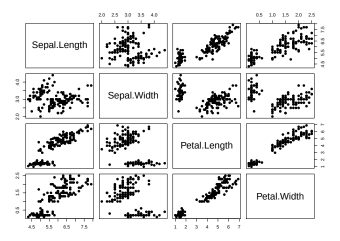


The data contains 150 leaf measurements for 3 flower species:

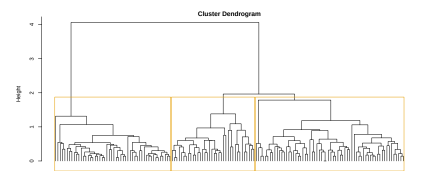
pairs(iris[1:4], col = iris\$Species)



We now "forget" the real groups specified by the 'Species' variable and try to find clusters based on the leaf measurements.

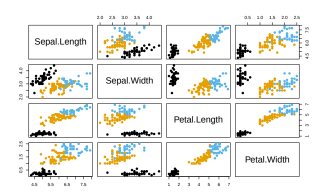


```
# compute distance matrix
d.euclid = dist(iris[1:4], method = "euclidean")
# do clustering with average linkage
cl = hclust(d.euclid, method = "average")
plot(cl, labels = FALSE, hang = -1) # plot dendrogram
rect.hclust(cl, k = 3) # highlight the k = 3 groups
```



We can extract the clustering assignments by cutting the dendrogram, e.g. using k=3 clusters:

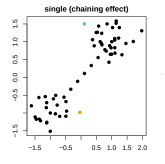
```
group = cutree(c1, k = 3) # get clusters assignments for k=3
pairs(iris[1:4], col = group) # plot clusters with different colors
```

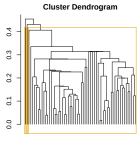


PROPERTIES: SINGLE LINKAGE

Single linkage introduces the *chaining problem*:

- Only one pair of points needs to be close to merge clusters.
- A chain of points can expand a cluster over long distances.
- Points within a cluster can be too widely spread and not dense enough.

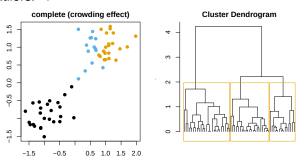




PROPERTIES: COMPLETE LINKAGE

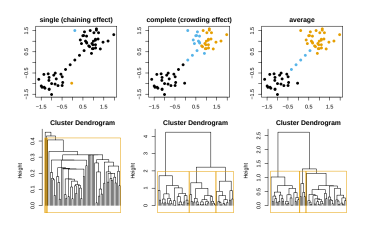
Complete linkage avoids chaining, but suffers from *crowding*:

- Merging is based on the furthest distance of point pairs from different clusters.
- Points of two different clusters can thus be closer than points within a cluster.
- Clusters are dense, but too close to each other. <!- and sensitive to outliers. ->



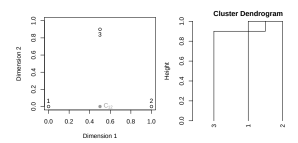
PROPERTIES: AVERAGE LINKAGE

- Average linkage is based on the average distance between clusters and tries to avoid crowding and chaining.
- Produces clusters that are quite dense and rather far apart.



PROPERTIES: CENTROID LINKAGE

- Centroid linkage defines the distance based on artificial data points (the cluster centers), which produces dendrograms with inversions, i.e., the distance between the clusters to be merged can be smaller in the next step.
- In single, complete and average linkage, the distance between the clusters to be merged increases in each step. ⇒ always produces dendrograms without inversions.



SUMMARY

- Hierarchical agglomerative clustering methods iteratively merge observations/clusters until all observations are in one single cluster.
- Results in a hierarchy of clustering assignments which can be visualized in a *dendrogram*. Each node of the dendrogram represents a cluster and its 'height' is proportional to the distance of its child nodes.
- The most common linkage functions are single, complete, average and centroid linkage. There is no perfect linkage and each linkage has its own advantages and disadvantages.

Partitioning Clustering Methods

OPTIMAL PARTITIONING CLUSTERING

Hierarchical clustering:

Stepwise merging (agglomerative methods) or dividing (divisive methods) of clusters based on distances and linkages. The number of clusters are selected by splitting the dendrogram at a specific threshold for the "height" after visual inspection.

Partitioning clustering:

Partitions the *N* observations into a predefined number of *K* clusters by optimizing a numerical criterion. The most common partitioning methods are:

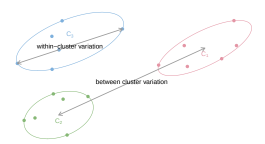
- K-means
- K-medians
- K-medoids (Partitioning Around Medoids (PAM))

K-MEANS

K-means partitions the N observations into K predefined clusters C_1, C_2, \ldots, C_K by minimizing the **compactness**, i.e. the **within-cluster variation** of all clusters using

$$\sum_{k=1}^K \sum_{i \in C_k} \|X_i - \bar{X}_k\|_2^2 o \min$$

where $\bar{X}_k = \frac{1}{N_k} \sum_{i \in C_k} X_i$ is the centroid of cluster k and N_k is the number of observations in cluster k.



K-MEANS

Idea: Consider every possible partition of N observations into K clusters and select the one with the lowest **within-cluster variation**. Problem: Requires trying all possible assignments of N observations into K clusters, which in practice is nearly impossible (Hothorn et al., 2009, p.322):

K	Number of possible partitions
3	2.375.101
4	45.232.115.901
5	10 ⁶⁸
	3

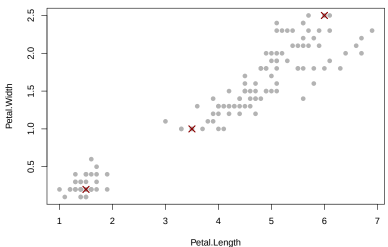
Hothorn, T., Everitt, B. S. (2009). A handbook of statistical analyses using R. Chapman and Hall/CRC.

K-MEANS

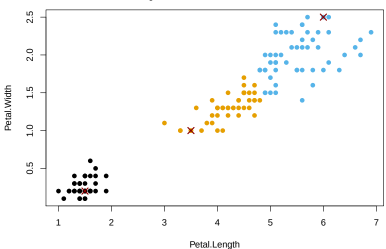
Use an approximation:

- Initialization: Choose *K* arbitrary observations to be the initial cluster centers.
- Assignment: Assign every observation to the cluster with the closest center.
- Update: Compute the new center of each cluster as the mean of its members.
- Repeat (2) and (3) until the centers do not move.

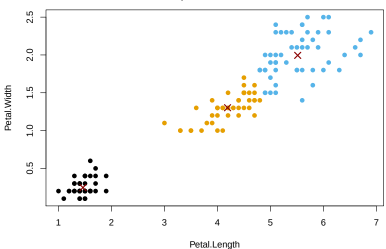
Iteration 0
Choose K arbitrary observations to be the initial cluster centers



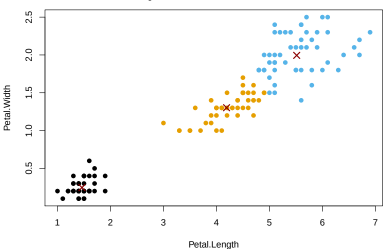
Iteration 1
Assign observations to nearest cluster center



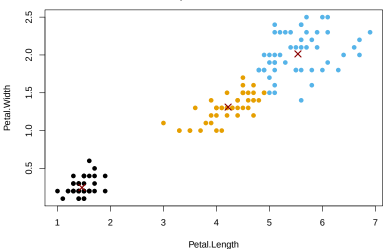




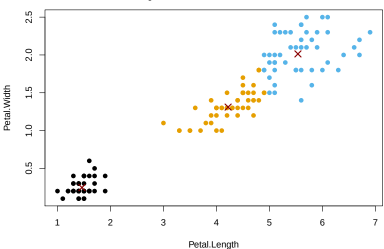
Iteration 2
Assign observations to nearest cluster center



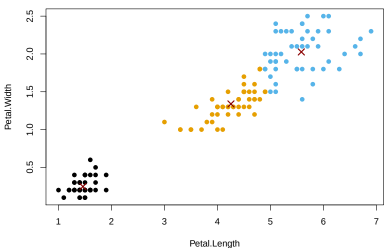




Iteration 3
Assign observations to nearest cluster center







K-MEANS IN R

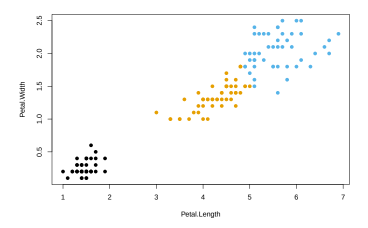
The K-means algorithm is part of the base distribution in R, given by the kmeans function (using algorithm = "Lloyd"):

```
km = kmeans(iris[,3:4], centers = 3, nstart = 100,
 iter.max = 100, algorithm = "Lloyd")
str(km)
## List of 9
## $ cluster : int [1:150] 1 1 1 1 1 1 1 1 ...
## $ centers : num [1:3, 1:2] 1.462 4.269 5.596 0.246 ...
## ..- attr(*, "dimnames")=List of 2
## ....$ : chr [1:3] "1" "2" "3"
## ....$ : chr [1:2] "Petal.Length" "Petal.Width"
## $ totss : num 551
## $ withinss : num [1:3] 2.02 13.06 16.29
## $ tot.withinss: num 31.4
## $ betweenss : num 520
## $ size : int [1:3] 50 52 48
## $ iter : int 17
## $ ifault : NULL
## - attr(*, "class")= chr "kmeans"
```

K-MEANS IN R

The final cluster assignments can be visualized by:

```
plot(iris[,3:4], pch = 19, col = km$cluster)
```

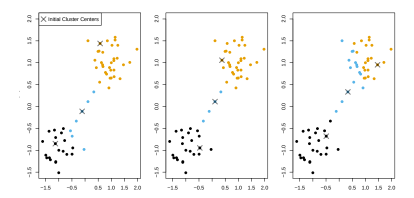


PROPERTIES OF K-MEANS

- K-means is based on computing the mean, which is sensitive to outliers and can only be computed for numerical data.
- The within-cluster variation is reduced in each iteration. In R, the maximum number of iterations is specified by iter.max.
- The final result is typically not the best result that globally minimizes the within-cluster variation.
 - → would only be possible after trying all possible partitions!
- K-means can be restarted multiple times using nstart. The clustering with the smallest within-cluster variation is then selected as the best solution.

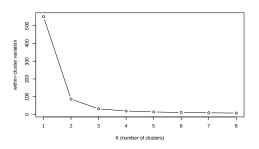
PROPERTIES OF K-MEANS

• *K*-means produces different clusters depending on the initial centers and always converges, e.g.:



CHOICE OF K

- Many methods exist for choosing the number of clusters K (there is no perfect solution).
- The easiest method is to apply K-means for different K and plot the within-cluster variation for each number of K.
- The within-cluster variation always decreases with increasing number of clusters.
- An "elbow" in the plot might indicate a useful solution.



K-MEDOIDS

- is strongly related to K-means and is realized by the Partitioning Around Medoids (PAM) algorithm.
- uses cluster medoids as representative clusters, i.e. real data points instead of artificial data points (such as the cluster centers as in K-means) are used.
- is less sensitive to outliers and more robust than K-means.
- can handle categorical features (K-means doesn't because it is based on calculating the cluster centers by taking the mean in each dimension).

THE PAM ALGORITHM

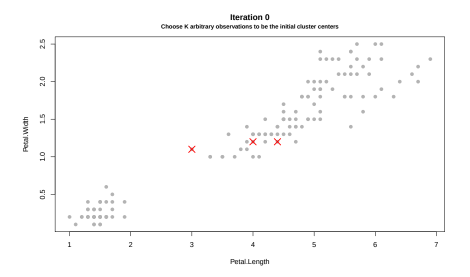
- Initialization: Randomly select K data points as the medoids.
- **2 Assignment:** Assign each data point x_i to its closest medoid m and calculate the within-cluster variation for each medoid (by summing up the distances of the current medoid m to all other data points associated to m).
- **① Update:** Swap m and x_i and recompute the within-cluster variation to see if another medoid is more appropriate. Select the medoid m with the lowest within-cluster variation.
- Repeat steps (2) and (3) until medoids do not change.

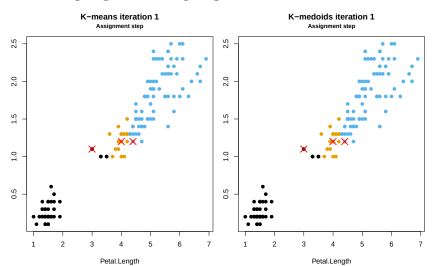
THE PAM ALGORITHM

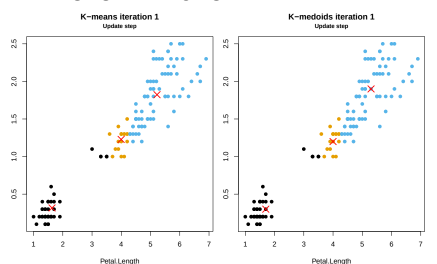
The PAM algorithm typically uses the following two metrics to compute distances:

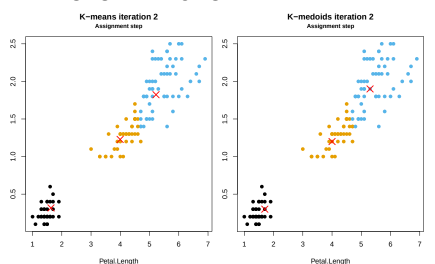
- The euclidean distance (root sum-of-squares of differences).
- The Manhattan distance (the sum of absolute distances).

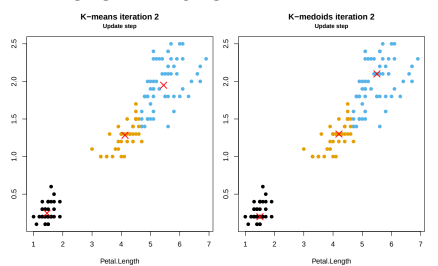
Note: The Manhattan distance should give more robust results if your data contains outliers. In all other cases, the results will be similar for both metrics.











K-MEDOIDS IN R

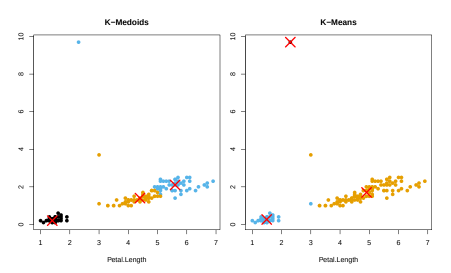
The K-medoids algorithm is implemented in the 'pam' function included in the 'cluster' R-package.

We compare *K*-means and *K*-medoids on a modified iris data with additional outliers:

```
library(cluster)
# add outliers to iris
out = data.frame(Petal.Length = c(2.3, 3), Petal.Width = c(9.7, 3.7))
iris2 = rbind(iris[, 3:4], out)

# K-medoid vs. K-means
kmedoid = pam(iris2, k = 3)
km = kmeans(iris2, centers = 3, iter.max = 100, nstart = 100)
```

K-MEDOIDS IN R



SUMMARY

- Minimizing the within-cluster variation exactly is not feasible and can be approximated by the K-means algorithm.
- K-means always converges, however, the cluster assignments strongly depend on the initial centers.
 - \rightarrow repeat it several times with different initial centers.
- A simple solution for choosing the number of clusters K is to plot the within-cluster variation for several K and look for an "elbow" which is a good guess for K.