Package: clustlearn (via r-universe)

September 9, 2024

Title Learn Clustering Techniques Through Examples and Code

Version 1.0.0

Description Clustering methods, which (if asked) can provide step-by-step explanations of the algorithms used, as described in Ezugwu et. al., (2022) [<doi:10.1016/j.engappai.2022.104743>](https://doi.org/10.1016/j.engappai.2022.104743); and datasets to test them on, which highlight the strengths and weaknesses of each technique, as presented in the clustering section of 'scikit-learn' (Pedregosa et al., 2011) <<https://jmlr.csail.mit.edu/papers/v12/pedregosa11a.html>>.

URL <https://github.com/Ediu3095/clustlearn>

BugReports <https://github.com/Ediu3095/clustlearn/issues>

License MIT + file LICENSE Encoding UTF-8 Roxygen list(markdown = TRUE) RoxygenNote 7.2.3 **Depends** R $(>= 4.3.0)$ **Imports** proxy ($> = 0.4-27$), cli ($> = 3.6.1$) **Suggests** deldir $(>= 1.0-9)$ LazyData true Repository https://ediu3095.r-universe.dev RemoteUrl https://github.com/ediu3095/clustlearn RemoteRef HEAD RemoteSha ff48cda95bd68735b4373398de3a8d41553d55ad

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agglomerative_clustering
```
Agglomerative Hierarchical Clustering

Description

Perform a hierarchical agglomerative cluster analysis on a set of observations

Usage

```
agglomerative_clustering(
  data,
 proximity = "single",
 details = FALSE,
 waiting = TRUE,
  ...
)
```
Arguments

Details

This function performs a hierarchical cluster analysis for the n objects being clustered. The definition of a set of clusters using this method follows a n step process, which repeats until a single cluster remains:

- 1. Initially, each object is assigned to its own cluster. The matrix of distances between clusters is computed.
- 2. The two clusters with closest proximity will be joined together and the proximity matrix updated. This is done according to the specified proximity. This step is repeated until a single cluster remains.

The definitions of proximity considered by this function are:

- single min $\{d(x, y) : x \in A, y \in B\}$. Defines the proximity between two clusters as the distance between the closest objects among the two clusters. It produces clusters where each object is closest to at least one other object in the same cluster. It is known as SLINK, single-link and minimum-link.
- complete $\max \{d(x, y) : x \in A, y \in B\}$. Defines the proximity between two clusters as the distance between the furthest objects among the two clusters. It is known as CLINK, completelink and maximum-link.
- average $\frac{1}{|A|\cdot|B|}\sum_{x\in A}\sum_{y\in B}d(x,y)$. Defines the proximity between two clusters as the average distance between every pair of objects, one from each cluster. It is also known as UPGMA or average-link.

Value

An [stats::hclust\(\)](#page-0-0) object which describes the tree produced by the clustering process.

Author(s)

Eduardo Ruiz Sabajanes, <eduardo.ruizs@edu.uah.es>

Examples

!! This algorithm is very slow, so we'll only test it on some datasets !!

```
### Helper function
test <- function(db, k, prox) {
 print(cl <- clustlearn::agglomerative_clustering(db, prox))
 oldpar \leq par(mfrow = c(1, 2))
 plot(db, col = cutree(cl, k), asp = 1, pch = 20)h <- rev(cl$height)[50]
 clu \leq as.hclust(cut(as.dendrogram(cl), h = h)$upper)
 ctr <- unique(cutree(cl, k)[cl$order])
 plot(clu, labels = FALSE, hang = -1, xlab = "Cluster", sub = "", main = "")
 rect.hclust(clu, k = k, border = ctr)
 par(oldpar)
}
### Example 1
test(clustlearn::db1, 2, "single")
### Example 2
# test(clustlearn::db2, 2, "sing") # same as "single"
```
 $4 \text{ d}b2$

```
### Example 3
test(clustlearn::db3, 4, "a") # same as "average"
### Example 4
test(clustlearn::db4, 6, "s") # same as "single"
### Example 5
test(clustlearn::db5, 3, "complete")
### Example 6
# test(clustlearn::db6, 3, "c") # same as "complete"
### Example 7 (with explanations, no plots)
 cl <- clustlearn::agglomerative_clustering(
 clustlearn::db5[1:6, ],
  'single',
 details = TRUE,
 waiting = FALSE
\mathcal{L}
```
db1 *Test Database 1*

Description

Test Database 1

Usage

db1

Format

db1:

A data frame with 500 rows and 2 columns. The data points form two concentric circles.

db2 *Test Database 2*

Description

Test Database 2

Usage

db2

 $d\mathbf{b}3$ 5

Format

db2:

A data frame with 500 rows and 2 columns. The data points form two moons.

db3 *Test Database 3*

Description

Test Database 3

Usage

db3

Format

db3:

A data frame with 500 rows and 2 columns.

The data points form three overlapping elliptical clusters of varying densities.

db4 *Test Database 4*

Description

Test Database 4

Usage

db4

Format

db4:

A data frame with 500 rows and 2 columns.

The data points form three diagonal parallel segments.

db5 *Test Database 5*

Description

Test Database 5

Usage

db5

Format

db5:

A data frame with 500 rows and 2 columns.

The data points form three non-overlapping circular clusters of similar density.

db6 *Test Database 6*

Description

Test Database 6

Usage

db6

Format

db6:

A data frame with 500 rows and 2 columns.

The data points are uniformly distributed on the plane.

dbscan *Density Based Spatial Clustering of Applications with Noise (DB-SCAN)*

Description

Perform DBSCAN clustering on a data matrix.

Usage

```
dbscan(data, eps, min_pts = 4, details = FALSE, waiting = TRUE, \ldots)
```
Arguments

Details

The data given by data is clustered by the DBSCAN method, which aims to partition the points into clusters such that the points in a cluster are close to each other and the points in different clusters are far away from each other. The clusters are defined as dense regions of points separated by regions of low density.

The DBSCAN method follows a 2 step process:

- 1. For each point, the neighborhood of radius eps is computed. If the neighborhood contains at least min_pts points, then the point is considered a core point. Otherwise, the point is considered an outlier.
- 2. For each core point, if the core point is not already assigned to a cluster, a new cluster is created and the core point is assigned to it. Then, the neighborhood of the core point is explored. If a point in the neighborhood is a core point, then the neighborhood of that point is also explored. This process is repeated until all points in the neighborhood have been explored. If a point in the neighborhood is not already assigned to a cluster, then it is assigned to the cluster of the core point.

Whatever points are not assigned to a cluster are considered outliers.

8 dbscan box and the set of the set

Value

A [dbscan\(\)](#page-6-1) object. It is a list with the following components:

Author(s)

Eduardo Ruiz Sabajanes, <eduardo.ruizs@edu.uah.es>

Examples

```
### Helper function
test <- function(db, eps) {
  print(cl <- clustlearn::dbscan(db, eps))
  out <- cl$cluster == 0
  plot(db[!out, ], col = cl$cluster[!out], pch = 20, asp = 1)
  points(db[out, ], col = max(cl$cluster) + 1, pch = 4, lwd = 2)}
### Example 1
test(clustlearn::db1, 0.3)
### Example 2
test(clustlearn::db2, 0.3)
### Example 3
test(clustlearn::db3, 0.25)
### Example 4
test(clustlearn::db4, 0.2)
### Example 5
test(clustlearn::db5, 0.3)
### Example 6
test(clustlearn::db6, 0.3)
### Example 7 (with explanations, no plots)
  cl <- clustlearn::dbscan(
  clustlearn::db5[1:20, ],
  0.3,
  details = TRUE,
  waiting = FALSE
\lambda
```
Description

Perform a hierarchical Divisive cluster analysis on a set of observations

Usage

```
divisive_clustering(data, details = FALSE, waiting = TRUE, ...)
```
Arguments

Details

This function performs a hierarchical cluster analysis for the n objects being clustered. The definition of a set of clusters using this method follows a n step process, which repeats until n clusters remain:

- 1. Initially, each object is assigned to the same cluster. The sum of squares of the distances between objects and their cluster center is computed.
- 2. The cluster with the highest sum of squares is split into two using the k-means algorithm. This step is repeated until n clusters remain.

Value

An [stats::hclust\(\)](#page-0-0) object which describes the tree produced by the clustering process.

Author(s)

Eduardo Ruiz Sabajanes, <eduardo.ruizs@edu.uah.es>

Examples

```
### !! This algorithm is very slow, so we'll only test it on some datasets !!
### Helper function
test \leq function(db, k) {
 print(cl <- clustlearn::divisive_clustering(db, max_iterations = 5))
 par(mfrow = c(1, 2))plot(db, col = cutree(cl, k), asp = 1, pch = 20)h <- rev(cl$height)[50]
 clu \leq as.hclust(cut(as.dendrogram(cl), h = h)$upper)
 ctr <- unique(cutree(cl, k)[cl$order])
 plot(clu, labels = FALSE, hang = -1, xlab = "Cluster", sub = "", main = "")
 rect.hclust(clu, k = k, border = ctr)}
### Example 1
# test(clustlearn::db1, 2)
### Example 2
# test(clustlearn::db2, 2)
### Example 3
# test(clustlearn::db3, 3)
### Example 4
# test(clustlearn::db4, 3)
### Example 5
test(clustlearn::db5, 3)
### Example 6
test(clustlearn::db6, 3)
### Example 7 (with explanations, no plots)
 cl <- clustlearn::divisive_clustering(
 clustlearn::db5[1:6, ],
 details = TRUE,
 waiting = FALSE
)
```
gaussian_mixture *Gaussian mixture model*

Description

Perform Gaussian mixture model clustering on a data matrix.

Usage

```
gaussian_mixture(data, k, max_iter = 10, details = FALSE, waiting = TRUE, \ldots)
```


gaussian_mixture 11

Arguments

Details

The data given by data is clustered by the model-based algorithm that assumes every cluster follows a normal distribution, thus the name "Gaussian Mixture".

The normal distributions are parameterized by their mean vector, covariance matrix and mixing proportion. Initially, the mean vector is set to the cluster centers obtained by performing a kmeans clustering on the data, the covariance matrix is set to the covariance matrix of the data points belonging to each cluster and the mixing proportion is set to the proportion of data points belonging to each cluster. The algorithm then optimizes the gaussian models by means of the Expectation Maximization (EM) algorithm.

The EM algorithm is an iterative algorithm that alternates between two steps:

- Expectation Compute how much is each observation expected to belong to each component of the GMM.
- Maximization Recompute the GMM according to the expectations from the E-step in order to maximize them.

The algorithm stops when the changes in the expectations are sufficiently small or when a maximum number of iterations is reached.

Value

A [gaussian_mixture\(\)](#page-9-1) object. It is a list with the following components:

Author(s)

Eduardo Ruiz Sabajanes, <eduardo.ruizs@edu.uah.es>

Examples

```
### !! This algorithm is very slow, so we'll only test it on some datasets !!
### Helper functions
dmnorm \leq function(x, mu, sigma) {
  k \leq ncol(sigma)
  x \leq -as.matrix(x)diff \leq t(t(x) - mu)
 num <- exp(-1 / 2 * diag(diff %*% solve(sigma) %*% t(diff)))
  den <- sqrt(((2 * pi)^k) * det(sign)num / den
}
test <- function(db, k) {
  print(cl <- clustlearn::gaussian_mixture(db, k, 100))
  x \leq -\text{seq}(\text{min}(db[, 1]), \text{max}(db[, 1]), \text{length.out} = 100)y <- seq(min(db[, 2]), max(db[, 2]), length.out = 100)
  plot(db, col = cl$cluster, asp = 1, pch = 20)for (i in seq_len(k)) {
   m <- cl$mu[i, ]
    s \leq cl$sigma[i, , ]
    f <- function(x, y) cl$lambda[i] * dmnorm(cbind(x, y), m, s)
    z \le outer(x, y, f)
    contour(x, y, z, col = i, add = TRUE)}
}
### Example 1
test(clustlearn::db1, 2)
### Example 2
# test(clustlearn::db2, 2)
### Example 3
test(clustlearn::db3, 3)
### Example 4
test(clustlearn::db4, 3)
### Example 5
test(clustlearn::db5, 3)
### Example 6
# test(clustlearn::db6, 3)
### Example 7 (with explanations, no plots)
cl <- clustlearn::gaussian_mixture(
  clustlearn::db5[1:20, ],
```
kmeans and the state of the

```
3,
  details = TRUE,
  waiting = FALSE
\mathcal{L}
```
kmeans *K-Means Clustering*

Description

Perform K-Means clustering on a data matrix.

Usage

```
kmeans(
  data,
  centers,
 max_iterations = 10,
  initialization = "kmeans++",
  details = FALSE,
  waiting = TRUE,
  ...
)
```
Arguments

Details

The data given by data is clustered by the k -means method, which aims to partition the points into k groups such that the sum of squares from points to the assigned cluster centers is minimized. At the minimum, all cluster centers are at the mean of their Voronoi sets (the set of data points which are nearest to the cluster center).

The k -means method follows a 2 to n step process:

- 1. The first step can be subdivided into 3 steps:
	- (a) Selection of the number k of clusters, into which the data is going to be grouped and of which the centers will be the representatives. This is determined through the use of the centers parameter.
	- (b) Computation of the distance from each data point to each center.
	- (c) Assignment of each observation to a cluster. The observation is assigned to the cluster represented by the nearest center.
- 2. The next steps are just like the first but for the first sub-step:
	- (a) Computation of the new centers. The center of each cluster is computed as the mean of the observations assigned to said cluster.

The algorithm stops once the centers in step $n + 1$ are the same as the ones in step n. However, this convergence does not always take place. For this reason, the algorithm also stops once a maximum number of iterations max_iterations is reached.

The initialization methods provided by this function are:

random A set of centers observations is chosen at random from the data as the initial centers.

kmeans++ The centers observations are chosen using the **kmeans++** algorithm. This algorithm chooses the first center at random and then chooses the next center from the remaining observations with probability proportional to the square distance to the closest center. This process is repeated until centers centers are chosen.

Value

A [stats::kmeans\(\)](#page-0-0) object.

Author(s)

Eduardo Ruiz Sabajanes, <eduardo.ruizs@edu.uah.es>

Examples

```
### Voronoi tesselation
voronoi <- suppressMessages(suppressWarnings(require(deldir)))
\text{cols} \leq c(
  "#00000019",
 "#DF536B19",
  "#61D04F19",
  "#2297E619",
  "#28E2E519",
  "#CD0BBC19",
  "#F5C71019",
  "#9E9E9E19"
)
### Helper function
test <- function(db, k) {
 print(cl <- clustlearn::kmeans(db, k, 100))
 plot(db, col = c1$cluster, asp = 1, pch = 20)
```
kmeans and the state of the

```
points(cl$centers, col = seq_length(k), pch = 13, cex = 2, lwd = 2)if (voronoi) {
   x \leftarrow c(min(db[, 1]), max(db[, 1]))dx <- c(x[1] - x[2], x[2] - x[1])
    y <- c(min(db[, 2]), max(db[, 2]))
    dy \leftarrow c(y[1] - y[2], y[2] - y[1])tesselation <- deldir(
      cl$centers[, 1],
      cl$centers[, 2],
      rw = c(x + dx, y + dy)\lambdatiles <- tile.list(tesselation)
    plot(
      tiles,
      asp = 1,add = TRUE,showpoints = FALSE,
      border = "#00000000",
      fillcol = cols
    \lambda}
}
### Example 1
test(clustlearn::db1, 2)
### Example 2
test(clustlearn::db2, 2)
### Example 3
test(clustlearn::db3, 3)
### Example 4
test(clustlearn::db4, 3)
### Example 5
test(clustlearn::db5, 3)
### Example 6
test(clustlearn::db6, 3)
### Example 7 (with explanations, no plots)
cl <- clustlearn::kmeans(
  clustlearn::db5[1:20, ],
  3,
  details = TRUE,
  waiting = FALSE
\lambda
```
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