Package 'Evacluster'

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Description Contains a set of clustering methods and evaluation metrics to select the best number of the clusters based on clustering stability. Two references describe the methodology: Fahimeh Nezhadmoghadam, and Jose Tamez-Pena (2021) <doi:10.1016 j.compbiomed.2021.104753="">, and Fahimeh Nezhadmoghadam, et al.(2021)<doi:10.2174 1567205018666210831145825="">.</doi:10.2174></doi:10.1016>
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Description

Contains a set of clustering methods and evaluation metrics to select the best number of the clusters based on clustering stability.

Details

Package: Evacluster
Type: Package
Version: 0.1.0
Date: 2022-03-25
License: LGPL (>= 2)

Purpose: The design of clustering models and evaluation metrics for finding the cluster's number via computing clustering stability. The best number of clusters is selected via consensus clustering and clustering stability.

Author(s)

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References

Nezhadmoghadam, Fahimeh, and Jose Tamez-Pena. "Risk profiles for negative and positive COVID-19 hospitalized patients.(2021) *Computers in biology and medicine* 136: 104753.

Fahimeh Nezhadmoghadam, et al., Robust Discovery of Mild Cognitive impairment subtypes and their Risk of Alzheimer's Disease conversion using unsupervised machine learning and Gaussian Mixture Modeling (2021), *Current Alzheimer Research*, 18 (7), 595-606.

Examples

Not run:

```
### Evacluster Package Examples ####
 library(datasets)
 data(iris)
 # Split data to training set and testing set
 rndSamples <- sample(nrow(iris),100)</pre>
 trainData <- iris[rndSamples,]</pre>
 testData <- iris[-rndSamples,]</pre>
 ## Expectation Maximization Clustering
# Perform Expectation Maximization Clustering on training set with 3 clusters
clsut <- EMCluster(trainData[,1:4],3)</pre>
# Predict the labels of the cluster for new data based on cluster labels of the training set
pre <- predict(clsut,testData[,1:4])</pre>
 ## Fuzzy C-means Clustering
 # Perform Fuzzy C-means Clustering on training set with 3 clusters
clsut <- FuzzyCluster(trainData[,1:4],3)</pre>
 # Predict the labels of the new data
pre <- predict(clsut,testData[,1:4])</pre>
 ## hierarchical clustering
 # Perform hierarchical clustering on training set with 3 clusters
clsut <- hierarchicalCluster(trainData[,1:4],distmethod="euclidean",clusters=3)</pre>
 # Predict the labels of the new data
pre <- predict(clsut,testData[,1:4])</pre>
 ## K-means Clustering
 # Perform K-means Clustering on training set with 3 clusters
clsut <- kmeansCluster(trainData[,1:4],3)</pre>
 # Predict the labels of the new data
pre <- predict(clsut,testData[,1:4])</pre>
 ## Partitioning Around Medoids (PAM) Clustering
# Perform pam Clustering on training set with 3 clusters
clsut <- pamCluster(trainData[,1:4],3)</pre>
# Predict the labels of the new data
pre <- predict(clsut,testData[,1:4])</pre>
## Non-negative matrix factorization (NMF)
\mbox{\# Perform nmf Clustering on training set with 3 clusters}
clsut <- nmfCluster(trainData[,1:4],rank=3)</pre>
```

```
# Predict the labels of the new data
 pre <- predict(clsut,testData[,1:4])</pre>
 ## t-Distributed Stochastic Neighbor Embedding (t-SNE)
 library(mlbench)
 data(Sonar)
 rndSamples <- sample(nrow(Sonar),150)</pre>
 trainData <- Sonar[rndSamples,]</pre>
 testData <- Sonar[-rndSamples,]</pre>
 # Perform tSNE dimensionality reduction method on training data
 tsne_trainData <- tsneReductor(trainData[,1:60],dim = 3,perplexity = 10,max_iter = 1000)
 # performs an embedding of new data using an existing embedding
 tsne_testData <- predict(tsne_trainData,k=3,testData[,1:60])</pre>
 ## clustering stability function
 # Compute the stability of clustering to select the best number of clusters.
 library(mlbench)
 data(Sonar)
 Sonar$Class <- as.numeric(Sonar$Class)</pre>
 Sonar$Class[Sonar$Class == 1] <- 0</pre>
 Sonar$Class[Sonar$Class == 2] <- 1</pre>
 # Compute the stability of clustering using kmeans clustering, UMAP as
 dimensionality reduction method, and feature selection technique
ClustStab <- clusterStability(data=Sonar, clustermethod=kmeansCluster, dimenreducmethod="UMAP",
                             n_components = 3,featureselection="yes", outcome="Class",
                        fs.pvalue = 0.05,randomTests = 100,trainFraction = 0.7,center=3)
 # Get the labels of the subjects that share the same connectivity
 clusterLabels <- getConsensusCluster(ClustStab,who="training",thr=seq(0.80,0.30,-0.1))</pre>
    # Compute the stability of clustering using PAM clustering, tSNE as
   dimensionality reduction method, and feature selection technique
ClustStab <- clusterStability(data=Sonar, clustermethod=pamCluster, dimenreducmethod="tSNE",
                             n_components = 3, perplexity=10,max_iter=100,k_neighbor=2,
                            featureselection="yes", outcome="Class",fs.pvalue = 0.05,
                              randomTests = 100,trainFraction = 0.7,k=3)
  # Get the labels of the subjects that share the same connectivity
 clusterLabels <- getConsensusCluster(ClustStab,who="training",thr=seq(0.80,0.30,-0.1))</pre>
```

```
# Compute the stability of clustering using hierarchical clustering,
   PCA as dimensionality reduction method, and without applying feature selection
 ClustStab <- clusterStability(data=Sonar, clustermethod=hierarchicalCluster,</pre>
                          dimenreducmethod="PCA", n_components = 3,featureselection="no",
                            randomTests = 100,trainFraction = 0.7,distmethod="euclidean",
# Get the labels of the subjects that share the same connectivity
 clusterLabels <- getConsensusCluster(ClustStab,who="training",thr=seq(0.80,0.30,-0.1))</pre>
  # Show the clustering stability resuldts
 mycolors <- c("red", "green", "blue", "yellow")</pre>
  ordermatrix <- ClustStab$dataConcensus</pre>
 heatmapsubsample <- sample(nrow(ordermatrix),70)</pre>
  orderindex <- 10*clusterLabels + ClustStab$trainJaccardpoint
 orderindex <- orderindex[heatmapsubsample]</pre>
 orderindex <- order(orderindex)</pre>
 ordermatrix <- ordermatrix[heatmapsubsample, heatmapsubsample]</pre>
 ordermatrix <- ordermatrix[orderindex,orderindex]</pre>
  rowcolors <- mycolors[1+clusterLabels[heatmapsubsample]]</pre>
  rowcolors <- rowcolors[orderindex]</pre>
 hplot <- gplots::heatmap.2(as.matrix(ordermatrix),Rowv=FALSE,Colv=FALSE,</pre>
                  RowSideColors = rowcolors,ColSideColors = rowcolors,dendrogram = "none",
                            trace="none",main="Cluster Co-Association \n (k=3)")
  # Compare the PAC values of clustering stability with different numbers of clusters
{\tt ClustStab2} \gets {\tt clusterStability(data=Sonar, clustermethod=kmeansCluster, dimenreducmethod="UMAP", and {\tt clustStab2})} \\
                               n_components = 3,featureselection="yes", outcome="Class",
                         fs.pvalue = 0.05,randomTests = 100,trainFraction = 0.7,center=2)
ClustStab3 <- clusterStability(data=Sonar, clustermethod=kmeansCluster, dimenreducmethod="UMAP",
                                n_components = 3,featureselection="yes", outcome="Class",
                         fs.pvalue = 0.05,randomTests = 100,trainFraction = 0.7,center=3)
ClustStab4 <- clusterStability(data=Sonar, clustermethod=kmeansCluster, dimenreducmethod="UMAP",
                                n_components = 3,featureselection="yes", outcome="Class",
                         fs.pvalue = 0.05,randomTests = 100,trainFraction = 0.7,center=4)
  color_range<- c(black="#FDFC74", orange="#76FF7A", skyblue="#B2EC5D")</pre>
```

6 clusterStability

Description

This function computes the stability of clustering that helps to select the best number of clusters. Feature selection and dimensionality reduction methods can be used before clustering the data.

Usage

```
clusterStability(
  data = NULL,
  clustermethod = NULL,
  dimenreducmethod = NULL,
  n_components = 3,
  perplexity = 25,
  max_iter = 1000,
  k_neighbor = 3,
  featureselection = NULL,
  outcome = NULL,
  fs.pvalue = 0.05,
  randomTests = 20,
  trainFraction = 0.5,
  pac.thr = 0.1,
  ...
)
```

Arguments

data	A Data set								
clustermethod	The clustering method. This can be one of "Mclust", "pamCluster", "kmeansCluster", "hierarchicalCluster", and "FuzzyCluster".								
dimenreducmethod									
	The dimensionality reduction method. This must be one of "UMAP", "tSNE", and "PCA".								
n_components	The dimension of the space that data embed into. It can be set to any integer value in the range of 2 to 100.								
perplexity	The Perplexity parameter that determines the optimal number of neighbors in tSNE method.(it is only used in the tSNE reduction method)								

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max_iter The maximum number of iterations for performing tSNE reduction method.

k_neighbor The k_neighbor is used for computing the means of #neighbors with min dis-

tance (#Neighbor=sqrt(#Samples/k) for performing an embedding of new data

using an existing embedding in the tSNE method.

featureselection

This parameter determines whether feature selection is applied before clustering

data or not. if used, it should be "yes", otherwisw "no".

outcome The outcome feature is used for feature selection.

fs.pvalue The threshold pvalue used for feature selection process. The default value is

0.05

randomTests The number of iterations of the clustering process for computing the cluster

stability.

trainFraction This parameter determines the ratio of training data. The default value is 0.5.

pac.thr The pac.thr is the thresold to use for computing the proportion of ambiguous

clustering (PAC) score. It is as the fraction of sample pairs with consensus

indices falling in the interval. The default value is 0.1.

... Additional arguments passed to clusterStability().

Value

A list with the following elements:

- randIndex A vector of the Rand Index that computes a similarity measure between two clusterings.
- jaccIndex A vector of jaccard Index that measures how frequently pairs of items are joined together in two clustering data sets.
- randomSamples A vector with indexes of selected samples for training in each iteration.
- clusterLabels A vector with clusters' labels in all iterations. jaccardpoint
- jaccardpoint The corresponding Jaccard index for each data point of testing set
- averageNumberofClusters The mean Number of Clusters.
- testConsesus A vector of consensus clustering results of testing set.
- trainRandIndex A vector of the Rand Index for training set.
- trainJaccIndex A vector of the jaccard Index for training set.
- trainJaccardpoint The corresponding Jaccard index for each data point of training set.
- PAC The proportion of ambiguous clustering (PAC) score.
- dataConcensus A vector of consensus clustering results of training set.

```
library("mlbench")
data(Sonar)
Sonar$Class <- as.numeric(Sonar$Class)</pre>
```

8 EMCluster

EMCluster

Expectation Maximization Clustering

Description

This function perform EM algorithm for model-based clustering of finite mixture multivariate Gaussian distribution. The general purpose of clustering is to detect clusters of data and to assign the data to the clusters.

Usage

```
EMCluster(data = NULL, ...)
```

Arguments

data A Data set
... k: The number of Clusters

Value

A list of cluster labels and a returned object from init.EM

```
library(datasets)
data(iris)
rndSamples <- sample(nrow(iris),100)</pre>
```

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```
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

clsut <- EMCluster(trainData[,1:4],3)</pre>
```

FuzzyCluster

Fuzzy C-means Clustering Algorithm

Description

This function works by assigning membership to each data point corresponding to each cluster center based on the distance between the cluster center and the data point. A data object is the member of all clusters with varying degrees of fuzzy membership between 0 and 1.

Usage

```
FuzzyCluster(data = NULL, ...)
```

Arguments

```
data A Data set
... k: The number of Clusters
```

Value

A list of cluster labels and a R object of class "fcm ppclust"

```
library(datasets)
data(iris)

rndSamples <- sample(nrow(iris),100)
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

cls <- FuzzyCluster(trainData[,1:4],3)</pre>
```

10 getConsensusCluster

getConsensusCluster Consensus Clustering Results

Description

This function gets the labels of the subjects that share the same connectivity.

Usage

```
getConsensusCluster(object, who = "training", thr = seq(0.8, 0.3, -0.1))
```

Arguments

object A object of "clusterStability" function result

who This value shows the consensus clustering result of training and testing sets. If

who="training" for training set, otherwise other sets.

thr This is the seq function with three arguments that are: initial value, final value,

and increment (or decrement for a declining sequence). This produces ascending

or descending sequences.

Value

A list of samples' labels with same connectivity.

hierarchicalCluster 11

hierarchicalCluster hierarchical clustering

Description

This function seeks to build a hierarchy of clusters

Usage

```
hierarchicalCluster(data = NULL, distmethod = NULL, clusters = NULL, ...)
```

Arguments

clusters The number of Clusters

... Additional parameters passed to helust function

Value

A list of cluster labels

Examples

```
library(datasets)
data(iris)

rndSamples <- sample(nrow(iris),100)
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

cls <- hierarchicalCluster(trainData[,1:4],distmethod="euclidean",clusters=3)</pre>
```

kmeansCluster K-means Clustering

Description

This function classifies unlabeled data by grouping them by features, rather than pre-defined categories. It splits the data into K different clusters and describes the location of the center of each cluster. Then, a new data point can be assigned a cluster (class) based on the closed center of mass.

```
kmeansCluster(data = NULL, ...)
```

nmfCluster

Arguments

data A Data set
... center: The number of centers

Value

A list of cluster labels and a R object of class "kmeans"

Examples

```
library(datasets)
data(iris)

rndSamples <- sample(nrow(iris),100)
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

cls <- kmeansCluster(trainData[,1:4],3)</pre>
```

nmfCluster

Non-negative matrix factorization (NMF)

Description

This function factorizes samples matrix into (usually) two matrices W the cluster centroids and H the cluster membership,

Usage

```
nmfCluster(data = NULL, rank = NULL)
```

Arguments

data A Data set

rank Specification of the factorization rank

Value

A list of cluster labels, a R object of class "nmf" and the centers of the clusters

```
library(datasets)
data(iris)

rndSamples <- sample(nrow(iris),100)
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

cls <- nmfCluster(trainData[,1:4],rank=3)</pre>
```

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pamCluster

Partitioning Around Medoids (PAM) Clustering

Description

This function partitions (clustering) of the data into k clusters "around medoids". In contrast to the k-means algorithm, this clustering methods chooses actual data points as centers

Usage

```
pamCluster(data = NULL, ...)
```

Arguments

```
data A Data set
... k: The number of clusters
```

Value

A list of cluster labels and a R object of class "pam cluster"

Examples

```
library(datasets)
data(iris)

rndSamples <- sample(nrow(iris),100)
trainData <- iris[rndSamples,]
testData <- iris[-rndSamples,]

cls <- pamCluster(trainData[,1:4],3)</pre>
```

predict.EMCluster

EMCluster prediction function

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set.

```
## S3 method for class 'EMCluster'
predict(object, ...)
```

Arguments

object A returned object of EMCluster
... New sample set

Value

A list of cluster labels

```
predict.FuzzyCluster FuzzyCluster prediction function
```

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set.

Usage

```
## S3 method for class 'FuzzyCluster'
predict(object, ...)
```

Arguments

object A returned object of FuzzyCluster function

... New samples set

Value

A list of cluster labels

```
predict.hierarchicalCluster
```

hierarchicalCluster prediction function

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set.

```
## S3 method for class 'hierarchicalCluster'
predict(object, ...)
```

predict.kmeansCluster 15

Arguments

object A returned object of hierarchicalCluster function

... New samples set

Value

A list of cluster labels

```
{\tt predict.kmeansCluster}\ \textit{kmeansCluster prediction function}
```

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set.

Usage

```
## S3 method for class 'kmeansCluster'
predict(object, ...)
```

Arguments

object A returned object of kmeansCluster function
... New samples set

Value

A list of cluster labels

```
predict.nmfCluster \ nmfCluster \ prediction \ function
```

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set

```
## S3 method for class 'nmfCluster'
predict(object, ...)
```

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Arguments

object A returned object of nmfCluster

... New samples set

Value

A list of cluster labels

predict.pamCluster

pamCluster prediction function

Description

This function predicts the labels of the cluster for new data based on cluster labels of the training set.

Usage

```
## S3 method for class 'pamCluster'
predict(object, ...)
```

Arguments

object A returned object of pamCluster function

... New samples set

Value

A list of cluster labels

```
{\tt predict.tsneReductor} \quad \textit{tsneReductor prediction function}
```

Description

This function performs an embedding of new data using an existing embedding.

```
## S3 method for class 'tsneReductor'
predict(object, k = NULL, ...)
```

tsneReductor 17

Arguments

object A returned object of tsneReductor function

k The number is used for computing the means of #neighbors with min distance

(#Neighbor=sqrt(#Samples/k).

... New samples set

Value

tsneY:An embedding of new data

Examples

```
library("mlbench")
data(Sonar)

rndSamples <- sample(nrow(Sonar),150)
trainData <- Sonar[rndSamples,]
testData <- Sonar[-rndSamples,]

tsne_trainData <- tsneReductor(trainData[,1:60],dim = 3,perplexity = 10,max_iter = 1000)
tsne_testData <- predict(tsne_trainData,k=3,testData[,1:60])</pre>
```

tsneReductor

t-Distributed Stochastic Neighbor Embedding (t-SNE)

Description

This method is an unsupervised, non-linear technique used for data exploration and visualizing high-dimensional data. This function constructs a low-dimensional embedding of high-dimensional data, distances, or similarities.

Usage

```
tsneReductor(data = NULL, dim = 2, perplexity = 30, max_iter = 500)
```

Arguments

data Data matrix (each row is an observation, each column is a variable)

dim Integer number; Output dimensional (default=2)

perplexity numeric; Perplexity parameter (should not be bigger than 3 * perplexity < nrow(X)

- 1, default=30)

max_iter Integer; Number of iterations (default: 500)

Value

tsneY: A Matrix containing the new representations for the observation with selected dimensions by user

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```
library("mlbench")
data(Sonar)

rndSamples <- sample(nrow(Sonar),150)
trainData <- Sonar[rndSamples,]
testData <- Sonar[-rndSamples,]

tsne_trainData <- tsneReductor(trainData[,1:60],dim = 3,perplexity = 10,max_iter = 1000)</pre>
```

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