# Package 'CLARA.seq'

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Type Package

Title CLARA Clustering
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Description  Clustering algorithm (CLARA) to handle big datasets and index to evaluate the quality of the partition. This package works with the help of TraMineR's package for distance calulations.
Imports TraMineR, cluster, dplyr, doParallel, parallel, foreach
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R topics documented:
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clarans\_clust

CLARANS clustering

#### **Description**

With the help of TraMineR package, CLARANS clustering provide a clustering of big dataset. The main objective is to cluster state sequences with the "LCS" distance calculation method to find the best partition in N clusters.

WARNING: this function is less efficient than cLARA.

#### Usage

```
clarans_clust(
  data,
  nb_cluster,
  distargs = list(method = "LCS"),
  maxneighbours,
  numlocal,
  plot = FALSE,
  cores = detectCores() - 1
)
```

#### **Arguments**

data The dataset to use. In case of sequences, use seqdef (from TraMineR package)

to create such an object.

nb\_cluster The number of medoids

distargs List with method parameters to apply. (See the function seqdist in TraMineR

package)

maxneighbours Number of neighbours to explore to find a better clustering

numlocal Number of initialisation of the starting medoids
plot Boolean variable to plot the research convergence

cores Number of cores to use for parallelism

## Value

An object of class clarans\_seq

```
#creating sequences
library(TraMineR)
data(mvad)
mvad.labels <- c("employment", "further education", "higher education","joblessness", "school", "training")
mvad.scode <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, 17:86, states = mvad.scode,abels = mvad.labels, xtstep = 6)

#CLARANS Clustering
my_cluster <- clarans_clust(mvad.seq,nb_cluster = 4, maxneighbours = 20, numlocal = 4, plot = TRUE)</pre>
```

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clara\_clust

CLARA clustering

#### **Description**

With the help of TraMineR package, CLARA clustering provide a clustering of big dataset. The main objective is to cluster state sequences with the "LCS" distance calculation method to find the best partition in N clusters.

#### Usage

```
clara_clust(
  data,
  nb_sample = 100,
  size_sample = 40 + 2 * nb_cluster,
  nb_cluster = 4,
  distargs = list(method = "LCS"),
  plot = FALSE,
  find_best_method = "Distance",
  with.diss = TRUE,
  cores = detectCores() - 1
)
```

## **Arguments**

The dataset to use. In case of sequences, use seqdef (from TraMineR package) data to create such an object. nb\_sample The number of subsets to test. size\_sample The size of each subset nb\_cluster The number of medoids List with method parameters to apply. (See the function segdist in TraMineR distargs package) plot Boolean variable to plot the result of clustering find\_best\_method Method to select the best subset. "Distance" is for the mean distance and "DB" is for Davies-Bouldin value. with.diss Boolean if the distance matrix should be returned cores Number of cores to use for parallelism

#### Value

An object of class clara\_seq

```
#creating sequences
library(TraMineR)
data(mvad)
mvad.labels <- c("employment", "further education", "higher education", "joblessness", "school", "training")</pre>
```

davies\_bouldin

```
mvad.scode <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, 17:86, states = mvad.scode,abels = mvad.labels, xtstep = 6)

#CLARA Clustering
my_cluster <- clara_clust(mvad.seq,nb_cluster = 4, nb_sample = 10, size_sample = 20, with.diss = TRUE)

#CLARA Clustering with Davies-Bouldin Method
my_cluster <- clara_clust(mvad.seq,nb_cluster = 4, nb_sample = 10, size_sample = 20, with.diss = TRUE, find_bes</pre>
```

davies\_bouldin

Davies Bouldin Index

## Description

Implementation of Davies-Bouldin index to evaluate the quality of CLARA Clustering.

## Usage

```
davies_bouldin(
  seq_obj,
  distargs = list(method = "LCS"),
  diss = TRUE,
  plot = TRUE,
  cores = detectCores() - 1
)
```

## **Arguments**

seq_obj	The object generated with CLARA Clustering
distargs	List with method parameters to apply. (See the function seqdist in TraMineR package)
diss	Boolean to express if the parameter diss from CLARA.seq clustering has been returned (Matrix size must be k columns and n rows - see refseq function from TraMineR package)
plot	Boolean variable to plot the research convergence
cores	Number of cores to use for parallelism

## Value

The value of the index

```
my_index <- davies_bouldin(my_cluster)</pre>
```

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# Description

With the help of TraMineR package, FUZZY-CLARA clustering provide a clustering of big dataset. The main objective is to cluster state sequences with the "LCS" distance calculation method to find the best partition in N clusters.

This function is a mix between CLARA and the ROBUST FUZZY C-MEDOIDS.

WARNING: This function is not finished yet!

## Usage

```
fuzzy_clust(
  data,
  nb_sample = 100,
  size_sample = 40 + 2 * nb_cluster,
  nb_cluster = 4,
  distargs = list(method = "LCS"),
  fuzzyfier = 2,
  p = 5,
  threshold = 10,
  max_iter = 10,
  noise = 0.5,
  plot = FALSE,
  cores = detectCores() - 1
)
```

#### **Arguments**

data	The dataset to use. In case of sequences, use seqdef (from TraMineR package) to create such an object.
nb_sample	The number of subsets to test.
size_sample	The size of each subset
nb_cluster	The number of medoids
distargs	List with method parameters to apply. (See the function seqdist in TraMineR package)
fuzzyfier	Value of the fuzzifier (default is 2, which is the traditionnal value)
р	Number of candidate to test to be a better medoid
threshold	Variable to exclude outliers, whose values are greater than threshold
max_iter	Number of maximal iteration to do to find the set of medoids
noise	Small value to avoid divisions by 0 error
plot	Boolean variable to plot the research convergence
cores	Number of cores to use for parallelism

## Value

An object of class clarafuzzy\_seq

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#### **Examples**

```
#creating sequences
library(TraMineR)
data(mvad)
mvad.labels <- c("employment", "further education", "higher education","joblessness", "school", "training")
mvad.scode <- c("EM", "FE", "HE", "JL", "SC", "TR")
mvad.seq <- seqdef(mvad, 17:86, states = mvad.scode,abels = mvad.labels, xtstep = 6)

#CLARA-FUzZY Clustering
my_cluster <- fuzzy_clust(mvad.seq,nb_sample = 14, size_sample = 50, plot = TRUE, threshold = 7, max_iter = 10,</pre>
```

plot.clarafuzzy\_seq CLARA-FUZZY Plot Method

#### **Description**

Create graph from CLARA-FUZZY object to show the evolution of the index value

## Usage

```
## S3 method for class 'clarafuzzy_seq'
plot(obj)
```

## **Arguments**

obj

An object from the class clarafuzzy\_seq (create by the function fuzzy\_clust())

### Value

A plot of the index evolution

## **Examples**

```
plot(my_cluster)
```

plot.clarans\_seq

CLARANS Plot Method

#### **Description**

Create graph from CLARANS object to show the evolution of the mean distance

# Usage

```
## S3 method for class 'clarans_seq'
plot(obj)
```

#### **Arguments**

obj

An object from the class clarans\_seq (create by the function clarans\_clust())

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## Value

A plot of the mean distance's evolution

## **Examples**

```
plot(my_cluster)
```

plot.clara\_seq

CLARA Plot Method

# Description

Create graph from CLARA object to show the evolution of the mean distance

## Usage

```
## S3 method for class 'clara_seq'
plot(obj)
```

# Arguments

obj

An object from the class clara\_seq (create by the function clara\_clust())

## Value

A plot of the mean distance's evolution

```
plot(my_cluster)
```

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