# Package 'fdakmapp'

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Title Functional Data Analysis: K-mean and K-medoid with Alignemnt

Version 2.1

| Type Package   |
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| Description  The fdakmapp package provides the kmap function that jointly performs clustering and alignment of a functional dataset (multidimensional or unidimensional). The centers can be computed by mean and medoid center methods. Many options are available as parallal version. |
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aneurisk65

Data for Examples.

#### Description

A dataset containing 65 multidimensional curves from aneurisk project. The curves represent the first derivatives of the 65 ICA centerlines.

#### Usage

aneurisk65

#### **Format**

A list with abscissas x and values y:

x matrix 65 x 1380

y array 65 x1380 x 3

fdakmapp

Functional Data Analysis Plus: K-Mean/Medoid Alignment:

#### **Description**

Fdakmapp is a package that allows to jointly perform clustering and alignment of a functional dataset (multidimensional or unidimensional functions).

#### References

- Sangalli, L.M., Secchi, P., Vantini, S., Vitelli, V., 2010. "K-mean alignment for curve clustering". Computational Statistics and Data Analysis, 54, 1219-1233.
- Sangalli, L.M., Secchi, P., Vantini, S., 2014. "Analysis of AneuRisk65 data: K-mean Alignment". Electronic Journal of Statistics, Special Section on "Statistics of Time Warpings and Phase Variations", Vol. 8, No. 2, 1891-1904.

#### See Also

kmap.

#### **Examples**

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kmap

K-mean algorithm for clustering and alignment of functional data

#### **Description**

kmap jointly performs clustering and alignment of a functional dataset (multidimensional or unidimensional functions). To run kmap function with different numbers of clusters and/or different alignment methods see the input options.

#### Usage

```
res<-kmap( x, y, y1, n_clust,warping_method, center_method,similarity_method,
optim_method, seeds, span, delta, d_max, s_max, n_out, toll, fence,
iter_max,show_iter, check_total_similarity)</pre>
```

#### **Arguments**

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|   |   |  |

numeric matrix [n.func X grid.size] or vector [grid.size]: the abscissa values where each function is evaluated. n.func: number of functions in the dataset. grid.size: maximal number of abscissa values where each function is evaluated. The abscissa points may be unevenly spaced and they may differ from function to function. x can also be a vector of length grid.size. In this case, x will be used as abscissa grid for all functions. Furthermore if the grid's size differs from one function to another the matrix must be completed with NA values. The parameter x must be provided.

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numeric matrix  $[n.func \ X \ grid.size]$  or array  $[n.func \ X \ grid.size \ X \ d]$ : evaluations of the set of original functions on the abscissa grid x. n.func: number of functions in the dataset. grid.size: maximal number of abscissa values where each function is evaluated. d: (only if the sample is multidimensional) number of function components, i.e. each function is a d-dimensional curve. The parameter y must be provided.

seeds

numeric vector [n.clust] indexes of the functions to be used as initial centers. In the case where the values of seeds are not provided, they are randomly chosen among the *n.func* original functions. If seeds=NULL all the centers are randomly chosen. Default value of seeds is NULL.

n\_clust

scalar: required number of clusters. Default value is 1. Note that if n.clust=1 kma performs only alignment without clustering.

warping\_method

character: type of alignment required. The implemented options are: "affine", "dialation", "shift" and "noalign". If warping.method='noalign' kma performs only clustering (without alignment). If warping.method='affine' kma performs alignment (and possibly clustering) of functions using linear affine transformation as warping functions, i.e., x.final = dilation\*x + shift. If warping.method='shift' kma allows only shift, i.e., x.final = x + shift. If warping.method='dilation' kma allows only dilation, i.e., x.final = dilation\*x. Default value is 'affine'.

center\_method

character: type of clustering method to be used. Possible choices are: 'mean', 'medoid' and 'pseudomedoid'. Default value is 'mean'.

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similarity\_method

character: required similarity measure. Possible choices are: 'pearson','12'. Default value is 'pearson'.

raunt value is pearson

 $\hbox{\tt optim\_method} \qquad \hbox{\tt character: optimization method chosen to solve the minimization problems at}$ 

each iteration. Possible choices are: 'bobyqa'. Default method is 'bobyqa'

warping\_opt numeric vector. The parameters depend on the warping\_method chosen. If

 $warping\_method = `affine' \ warping\_opt <- \ c( \ max\_dilation \ , \ max\_shift).$ 

If warping\_method <- 'dilation' warping\_opt <- c(max\_dilation).

If warping\_method <- 'shift' warping\_opt <- c(max\_shift). If warping\_method <- 'noalign' warping\_opt <- as.numeric().

Default value is warping\_opt<-c(0.15,0.15).

center\_opt numeric vector. The parameters depend on the center\_method chosen. If cen-

ter\_method = 'mean' center\_opt <- c(span, delta).

If center\_method = 'medoid' center\_opt <- as.numeric().

If center\_method = 'pseudomedoid' center\_opt <- as.numeric().

Default value is center opt<-c(0.01,0.1).

out\_opt numeric vector. The parameters to set are (n\_out, tollerance, max\_iteration).

 $n\_out$  is the size of the grid where the centers will be computed. tollerance is a stop condition parameter. max\_iterationa is a stop condition parameter. The

defaut value is out\_opt <- c(100, 0.001, 100).

fence boolean: if fence=TRUE a control is activated at the end of each iteration. The

aim of the control is to avoid warping outliers with respect to their computed distributions. If fence=TRUE the running time can increase considerably. Default

value of fence is FALSE.

check\_total\_similarity

boolean: if check.total.similarity=TRUE at each iteration the algorithm checks if the total similarity is improving and stops if it's not true. In this case the results

obtained in the penultimate iteration are returned. Defaultvalue is TRUE.

show\_iter boolean: if show.iter=TRUE kmap shows the current iteration of the algorithm.

Default value is TRUE.

comp\_original\_center

boolean: if comp\_original\_center=TRUE the initial center with relative dissimilarities is computed otherwise this step is skipped.It can by computationally

expensive. Default value is FALSE.

par\_opt numeric vector: Parallel options. The parameters to set are (num\_threads, paral-

lel\_version) parallel\_version available are 0 and 1 : 0 is a trivial parallelization in which each thread compute the center of a cluster; 1 is a more efficient parallelization in which all the threads compute the centers sequentially (available

only with center\_method = 'medoid').

#### Value

The function output is a list containing the following elements:

x as input.

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y as input.

seeds vector with the indeces used in the algorithm.

warping.method as input.

similarity.method

as input.

center.method as input.

iterations scalar: total number of iterations performed by kma function.

n.clust as input.

x.center.orig numeric vector *n\_out*: abscissa of the center computed if *comp\_original\_center*=TRUE.

y.center.orig numeric vector  $n\_out$  or matrix  $n\_out \times n\_dim$ : value of the center computed if

comp\_original\_center=TRUE.

similarity.origin

numeric vector n\_obs dissimilarity, similarity or distance of the original center

respect the obserbations computed if *comp\_original\_center*=TRUE.

x.final matrix [n.func X grid.size]: aligned abscissas.

n.clust.final scalar: final number of clusters. Note that n.clust.final may differ from initial

number of clusters (i.e., from n.clust) if some clusters are found to be empty.

x.centers.final

matrix [n.clust.final X grid.size]: abscissas of the final function centers.

y.centers.final

matrix [n.clust.final X n.out] or array [n.clust.final X n.out x n\_dim], contain

the evaluations of the final functions centers.

templates\_vec list iteration : each element of the list contain centers of that iteration.

x\_out\_vec list iteration : each element of the list contain the abscissa of the centers of that

iteration.

labels vector n\_obs: cluster assignments.

similarity.final

vector [n\_obs]: similarities, dissimilarities or distance between each function and

the center of the cluster the function is assigned to.

parameters.list

list [iterations]: warping parameters at each iteration.

parameters matrix [n\_par X n\_obs]: warping parameters applied to the original abscissas x

to obtain the aligned abscissas x.final.

timer vector: time of execution by step.

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#### Description

Show results of the clustering with alignment of functional data

#### Usage

```
kma_show_results <-function (Result, bp_sim,wr_fun)</pre>
```

#### Arguments

Result output of kmap.

bp\_sim boolean: if TRUE dissimilarity, similarity or distance boxplot are plotted. De-

fault value is FALSE.

wr\_fun boolean: if TRUE the warping functions applied to x are plotted. Default value

is FALSE.

simulated30 Data for Examples.

#### Description

A dataset containing 30 simulated unidimensional curves.

#### Usage

simulated30

#### **Format**

A list with abscissas x and values y:

x matrix 30 x 200

y array 30 x 200 x 1

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