Package 'clustEff'

June 8, 2020

Type Package

Title Clusters of Effects Curves in Quantile Regression Models	
Version 0.2.0	
Description Clustering method to cluster both effects curves, through quantile regression coefficient modeling, and curves in functional data analysis. Sottile G. and Adelfio G. (2019) <doi:10.1007 s00180-018-0817-8="">.</doi:10.1007>	
Depends qrcm, cluster, fda, ggpubr, ggplot2	
License GPL-2	
Encoding UTF-8	
LazyData true	
RoxygenNote 7.1.0	
NeedsCompilation no	
Author Gianluca Sottile [aut, cre], Giada Adelfio [aut]	
Maintainer Gianluca Sottile <gianluca.sottile@unipa.it></gianluca.sottile@unipa.it>	
Repository CRAN	
Date/Publication 2020-06-08 15:00:15 UTC	
R topics documented:	
fpcac	2 3 8 10 11 13 15 17 18 19
Index	21

2 clustEff-package

clustEff-package

Clusters of effects curves

Description

This package implements a general algorithm to cluster coefficient functions (i.e. clusters of effects) obtained from a quantile regression (qrcm; Frumento and Bottai, 2016). This algorithm is also used for clustering curves observed in time, as in functional data analysis. The objectives of this algorithm vary with the scenario in which it is used, i.e. in the case of a cluster of effects, in a univariate case the objective may be to reduce its dimensionality or in the multivariate case to group similar effects on a covariate. In the case of a functional data analysis the main objective is to cluster waves or any other function of time or space. Sottile G. and Adelfio G. (2019) https://doi.org/10.1007/s00180-018-0817-8.

Details

Package: clustEff
Type: Package
Version: 0.2.0
Date: 2020-05-31
License: GPL-2

The function clustEff allows to specify the type of the curves to apply the proposed clustering algorithm. The function extract.object extracts the matrices, in case of multivariate response, through the quantile regression coefficient modeling, useful to run the main algorithm. The auxiliary functions summary.clustEff and plot.clustEff can be used to extract information from the main algorithm. In the new version of the package you can also find a PCA-based clustering approach called Functional Principal Components Analysis Clustering (FPCAC). Main function of this algorithm is fpcac, and some auxiliary functions are summary.fpcac and plot.fpcac.

Author(s)

Gianluca Sottile

Maintainer: Gianluca Sottile <gianluca.sottile@unipa.it>

References

Sottile, G., Adelfio, G. *Clusters of effects curves in quantile regression models*. Comput Stat 34, 551–569 (2019). https://doi.org/10.1007/s00180-018-0817-8

Sottile, G and Adelfio, G (2017). Clustering of effects through quantile regression. Proceedings 32nd International Workshop of Statistical Modeling, Groningen (NL), vol.2 127-130, https://iwsm2017.webhosting.rug.nl/IV

Frumento, P., and Bottai, M. (2015). *Parametric modeling of quantile regression coefficient functions*. Biometrics, doi: 10.1111/biom.12410.

Adelfio, G., Chiodi, M., D'Alessandro, A. and Luzio, D. (2011) FPCA algorithm for waveform clustering. Journal of Communication and Computer, 8(6), 494-502.

Examples

clustEff

Cluster Effects Algorithm

Description

This function implements the algorithm to cluster curves of effects obtained from a quantile regression (qrcm; Frumento and Bottai, 2015) in which the coefficients are described by flexible parametric functions of the order of the quantile. This algorithm can be also used for clustering of curves observed in time, as in functional data analysis.

Usage

```
clustEff(Beta, Beta.lower = NULL, Beta.upper = NULL,
    k = c(2, min(5, (ncol(Beta)-1))), ask = FALSE, diss.mat, alpha = .5,
    step = c("both", "shape", "distance"),
    cut.method = c("mindist", "length", "conf.int"),
    method = "ward.D2", approx.spline = FALSE, nbasis = 50,
    conf.level = 0.9, stand = FALSE, plot = TRUE, trace = TRUE)
```

Arguments

Beta

A matrix n x q. q represents the number of curves to cluster and n is either the length of percentiles used in the quantile regression or the length of the time vector.

Beta.lower

A matrix n x q. q represents the number of lower interval of the curves to cluster and n the length of percentiles used in quantile regression. Used only if cluster.effects=TRUE.

Beta.upper	A matrix $n \times q$. q represents the number of upper interval of the curves to cluster and n the length of percentiles used in quantile regression. Used only if cluster.effects=TRUE.
k	It represents the number of clusters to look for. If it is two-length vector (k.min - k.max) an optimization is performed, if it is a unique value it is fixed.
ask	If TRUE, after plotting the dendrogram, the user make is own choice about how many cluster to use.
diss.mat	a dissimilarity matrix, obtained by using distshape function.
alpha	It is the alpha-percentile used for computing the dissimilarity matrix. The default value is alpha=.5.
step	The steps used in computing the dissimilarity matrix. Default is "both"=("shape" and "distance")
cut.method	The method used in optimization step to look for the optimal number of clusters. Default is "mindist", however if Beta.lower and Beta.upper are available the suggested method is "conf.int".
method	The agglomeration method to be used.
approx.spline	If TRUE, Beta is approximated by a smooth spline.
nbasis	An integer variable specifying the number of basis functions. Only when approx.spline=TRUE
conf.level	the confidence level required.
stand	If TRUE, the argument Beta is standardized.
plot	If TRUE, dendrogram, boxplot and clusters are plotted.
trace	If TRUE, some informations are printed.

Details

Quantile regression models conditional quantiles of a response variabile, given a set of covariates. Assume that each coefficient can be expressed as a parametric function of p in the form:

$$\beta(p|\theta) = \theta_0 + \theta_1 b_1(p) + \theta_2 b_2(p) + \dots$$

where $b_1(p), b_2(p, ...)$ are known functions of p.

Value

An object of class "clustEff", a list containing the following items:

call the matched call.

p The percentiles used in quantile regression coefficient modeling or the time oth-

erwise.

X The curves matrix.
clusters The vector of clusters.

X. mean The mean curves matrix of dimension $n \times k$.

X.mean.dist The within cluster distance from the mean curve.

X. lower The lower bound matrix.

X.mean.lower The mean lower bound of dimension n x k.

X. upper The upper bound matrix.

X.mean.upper The mean upper bound of dimension $n \times k$.

Signif.interval

The matrix of dimension n x k containing the intervals in which each mean lower

and upper bounds don't include the zero.

k The number of selected clusters.

diss.matrix The dissimilarity matrix.

X.mean.diss The within cluster dissimilarity.

oggSilhouette An object of class "silhouette".

oggHclust An object of class "hclust".

distance A vector of goodness measures used to select the best number of clusters.

step The selected step.

method The used agglomeration method.

cut.method The used method to select the best number of clusters.

alpha The selected alpha-percentile.

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.it>

References

Sottile, G., Adelfio, G. Clusters of effects curves in quantile regression models. Comput Stat 34, 551–569 (2019). https://doi.org/10.1007/s00180-018-0817-8

Sottile, G and Adelfio, G (2017). *Clustering of effects through quantile regression*. Proceedings 32nd International Workshop of Statistical Modeling, Groningen (NL), vol.2 127-130, https://iwsm2017.webhosting.rug.nl/IV

Frumento, P., and Bottai, M. (2015). *Parametric modeling of quantile regression coefficient functions*. Biometrics, doi: 10.1111/biom.12410.

See Also

summary.clustEff, plot.clustEff, for summary and plotting. extract.object to extract useful objects for the clustering algorithm through a quantile regression coefficient modeling in a multivariate case.

Examples

```
# CURVES EFFECTS CLUSTERING
```

```
set.seed(1234)
n <- 300
q <- 2
```

```
k <- 5
x1 <- runif(n, 0, 5)
x2 <- runif(n, 0, 5)
X \leftarrow cbind(x1, x2)
rownames(X) <- 1:n
colnames(X) <- paste0("X", 1:q)</pre>
theta1 <- matrix(c(1, 1, 0, 0, 0, .5, 0, .5, 1, 2, .5, 0, 2, 1, .5),
                 ncol=k, byrow=TRUE)
theta2 <- matrix(c(1, 1, 0, 0, 0, -.3, 0, .5, 1, .5, -1.5, 0, -1, -.5, 1),
                 ncol=k, byrow=TRUE)
theta3 <- matrix(c(1, 1, 0, 0, 0, .3, 0, -.5, -1, 2, -.5, 0, 1, -.5, -1),
                 ncol=k, byrow=TRUE)
rownames(theta3) <- rownames(theta2) <- rownames(theta1) <-</pre>
    c("(intercept)", paste("X", 1:q, sep=""))
colnames(theta3) <- colnames(theta2) <- colnames(theta1) <-</pre>
    c("(intercept)", "qnorm(p)", "p", "p^2", "p^3")
Theta <- list(theta1, theta2, theta3)
B <- function(p, k){matrix(cbind(1, qnorm(p), p, p^2, p^3), nrow=k, byrow=TRUE)}
Q \leftarrow function(p, theta, B, k, X)\{rowSums(X * t(theta %*% B(p, k)))\}
Y <- matrix(NA, nrow(X), 15)
for(i in 1:15){
  if(i <= 5) Y[, i] \leftarrow Q(runif(n), Theta[[1]], B, k, cbind(1, X))
  if(i \le 10 \& i > 5) Y[, i] < Q(runif(n), Theta[[2]], B, k, cbind(1, X))
  if(i \le 15 \& i > 10) Y[, i] <- Q(runif(n), Theta[[3]], B, k, cbind(1, X))
XX <- extract.object(Y, X, intercept=TRUE, formula.p= ~ I(p) + I(p^2) + I(p^3))
obj <- clustEff(XX$X$X1, Beta.lower=XX$X1$X1, Beta.upper=XX$Xr$X1, cut.method = "conf.int")
summary(obj)
plot(obj, xvar="clusters", col = 1:3)
plot(obj, xvar="dendrogram")
plot(obj, xvar="boxplot")
obj2 <- clustEff(XX$X$X2, Beta.lower=XX$X1$X2, Beta.upper=XX$Xr$X2, cut.method = "conf.int")
summary(obj2)
plot(obj2, xvar="clusters", col=1:3)
plot(obj2, xvar="dendrogram")
plot(obj2, xvar="boxplot")
## Not run:
set.seed(1234)
n <- 300
q <- 15
```

```
k <- 5
X <- matrix(rnorm(n*q), n, q); X <- scale(X)</pre>
rownames(X) <- 1:n
colnames(X) <- paste0("X", 1:q)</pre>
.5, 0, .5, 1, 1,
                  .5, 0, 1, 2, .5,
                   .5, 0, 1, 1, .5,
                  .5, 0, .5, 1, 1,
                   .5, 0, .5, 1, .5,
                 -1.5, 0, -.5, 1, 1,
                  -1, 0, .5, -1, -1,
                 -.5, 0, -.5, -1, .5,
                  -1, 0, .5, -1, -.5,
                -1.5, 0, -.5, -1, -.5,
                  2, 0, 1, 1.5, 2,
                  2, 0, .5, 1.5, 2,
                  2.5, 0, 1, 1, 2,
                  1.5, 0, 1.5, 1, 2,
                  3, 0, 2, 1, .5),
                 ncol=k, byrow=TRUE)
rownames(Theta) <- c("(intercept)", paste("X", 1:q, sep=""))</pre>
colnames(Theta) <- c("(intercept)", "qnorm(p)", "p", "p^2", "p^3")
B <- function(p, k){matrix(cbind(1, qnorm(p), p, p^2, p^3), nrow=k, byrow=TRUE)}
Q \leftarrow function(p, theta, B, k, X)\{rowSums(X * t(theta %*% B(p, k)))\}
s \leftarrow matrix(1, q+1, k)
s[2:(q+1), 2] \leftarrow 0
s[1, 3:k] <- 0
Y <- Q(runif(n), Theta, B, k, cbind(1, X))
XX <- extract.object(Y, X, intercept = TRUE, formula.p= \sim I(p) + I(p^2) + I(p^3))
obj3 <- clustEff(XX$X, Beta.lower=XX$X1, Beta.upper=XX$Xr, cut.method = "conf.int")
summary(obj3)
# changing the alpha-percentile clusters are correctly identified
obj4 <- clustEff(XX$X, Beta.lower=XX$X1, Beta.upper=XX$Xr, cut.method = "conf.int",
                 alpha = 0.25)
summary(obj4)
# CURVES CLUSTERING IN FUNCTIONAL DATA ANALYSIS
set.seed(1234)
n <- 300
x < -1:n/n
Y <- matrix(0, n, 30)
sigma2 < -4*pmax(x-.2, 0) - 8*pmax(x-.5, 0) + 4*pmax(x-.8, 0)
```

8 distshape

```
mu <- sin(3*pi*x)
for(i in 1:10) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))
mu <- cos(3*pi*x)
for(i in 11:23) Y[,i] <- mu + rnorm(length(x), 0, pmax(sigma2,0))
mu <- sin(3*pi*x)*cos(pi*x)
for(i in 24:28) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))
mu <- 0 #sin(1/3*pi*x)*cos(2*pi*x)
for(i in 29:30) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))
obj5 <- clustEff(Y)
summary(obj5)
plot(obj5, xvar="clusters", col=1:4)
plot(obj5, xvar="dendrogram")
plot(obj5, xvar="boxplot")
## End(Not run)</pre>
```

distshape

Dissimilarity matrix

Description

This function implements the dissimilarity matrix based on shape and distance of curves.

Usage

```
distshape(Beta, alpha=.5, step=c("both", "shape", "distance"), trace=TRUE)
```

Arguments

Beta	A matrix n x q. q represents the number of curves to cluster and n is either the length of percentiles used in the quantile regression or the length of the time vector.
alpha	It is the alpha-percentile used for computing the dissimilarity matrix. If not fixed, the algorithm choose alpha=.25 (cluster.effects=TRUE) or alpha=.5 (cluster.effects=FALSE).
step	The steps used in computing the dissimilarity matrix. Default is "both"=("shape" and "distance")
trace	If TRUE, some informations are printed.

Value

The dissimilarity matrix of class "dist".

distshape 9

Author(s)

Gianluca Sottile < gianluca.sottile@unipa.it>

References

Sottile, G., Adelfio, G. Clusters of effects curves in quantile regression models. Comput Stat 34, 551–569 (2019). https://doi.org/10.1007/s00180-018-0817-8

Sottile, G and Adelfio, G (2017). *Clustering of effects through quantile regression*. Proceedings 32nd International Workshop of Statistical Modeling, Groningen (NL), vol.2 127-130, https://iwsm2017.webhosting.rug.nl/IV

Frumento, P., and Bottai, M. (2015). *Parametric modeling of quantile regression coefficient functions*. Biometrics, doi: 10.1111/biom.12410.

See Also

clustEff, summary.clustEff, plot.clustEff, for summary and plotting. extract.object to extract useful objects for the clustering algorithm through a quantile regression coefficient modeling in a multivariate case.

Examples

```
set.seed(1234)
n <- 300
x <- 1:n/n

Y <- matrix(0, n, 30)

sigma2 <- 4*pmax(x-.2, 0) - 8*pmax(x-.5, 0) + 4*pmax(x-.8, 0)

mu <- sin(3*pi*x)
for(i in 1:10) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

mu <- cos(3*pi*x)
for(i in 11:23) Y[,i] <- mu + rnorm(length(x), 0, pmax(sigma2,0))

mu <- sin(3*pi*x)*cos(pi*x)
for(i in 24:28) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

mu <- 0 #sin(1/3*pi*x)*cos(2*pi*x)
for(i in 29:30) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

diss <- distshape(Y)
diss</pre>
```

10 extract.object

extract.object extract.object fits a multivariate quantile regression and extra objects for the cluster effects algorithm.
--

Description

extract.object fits a multivariate quantile regression and extracts objects for the cluster effects algorithm.

Usage

```
extract.object(Y, X, intercept=TRUE, formula.p=~slp(p, 3), s, object, p, which)
```

Arguments

Υ	A multivariate response matrix of dimension n x q1, or a vector of length n.
Χ	The covariates matrix of dimension n x q2.
intercept	If TRUE, the intercept is included in the model.
formula.p	a one-sided formula of the form $\sim b1(p,) + b2(p,) +$
S	An optional 0/1 matrix that allows to exclude some model coefficients (see 'Examples').
object	An object of class "iqr". If missing, Y and X have to be supplied.
р	The percentiles used in quantile regression coefficient modeling. If missing a default sequence is choosen.
which	If fixed, only the selected covariates are extraced from the model. If missing all the covariates are extracted.

Details

A list of objects useful to run the cluster effect algorithm is created.

Value

р	The percentiles used in the quantile regression.
X	A list containing as many matrices as covariates, where for each matrix the number of columns corresponds to the number of the responses. Each column of a matrix corresponds to one curve effect. In the case of a univariate model it is a unique matrix.
X1	A list as X. Each column of a matrix corresponds to the lower interval of the curve effect. In the case of a univariate model it is a unique matrix.
Xr	A list as X. Each column of a matrix corresponds to the upper interval of the curve effect. In the case of a univariate model it is a unique matrix.

fpcac 11

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.it>

See Also

clustEff, for clustering algorithm; summary.clustEff and plot.clustEff, for summarizing and plotting clustEff objects.

Examples

```
# using simulated data
# see the documentation for 'clustEff'
```

fpcac

Functional Principal Components Analysis Clustering

Description

This function implements the algorithm FPCAC for curves clustering as a variant of a k-means algorithm based on the principal component rotation of data

Usage

Arguments

disp

Χ	Matrix of 'curves' of dimension n x q.
•	·
K	the number of clusters.
fd	If not NULL it overrides X and must be an object of class fd.
nbasis	an integer variable specifying the number of basis functions. The default value is 5.
norder	an integer specifying the order of b-splines, which is one higher than their degree. The default value is 3.
nharmonics	the number of harmonics or principal components to use. The default value is 3.
alpha	trimming size, that is the given proportion of observations to be discarded.
niter	the number or random restarting (larger values provide more accurate solutions.
Ksteps	the number of k-mean steps (not too many ksteps are needed).
conf.level	the confidence level required.
seed	the seed used for reproducibility.

if TRUE, it is used to print some information across the algorithm.

12 fpcac

Details

FPCAC is a functional PCA-based clustering approach that provides a variation of the algorithm for curves clustering proposed by Garcia-Escudero and Gordaliza (2005).

The starting point of the proposed FPCAC is to find a linear approximation of each curve by a finite \$p\$ dimensional vector of coefficients defined by the FPCA scores.

The number of starting clusters k is obtained on the basis of the scores volume, such that we assign events to the clusters defined by events that have a distance less than a fixed threshold (e.g. 90-th percentile) in the space of PCA scores. Once k is obtained we use a modified version of the trimmed k-means algorithm, that considers the matrix of FPCA scores instead of the coefficients of a linear fitting to B-spline bases.

The trimmed k-means clustering algorithm looks for the k centers $C_1, ..., C_k$ that are solution of the minimization problem:

$$O_k(\alpha) = \min_{Y} \min_{C_1, \dots, C_k} \frac{1}{[n(1-\alpha)]} \sum_{X_i \in Y} \inf_{1 \le j \le k} ||X_i - C_j||^2$$

We think that the proposed approach has the advantage of an immediate use of PCA for functional data avoiding some objective choices related to spline fitting as in RCC. Simulations and applications suggest also the well behavior of the FPCAC algorithm, both in terms of stable and easily interpretable results.

Value

An object of class "fpcac", a list containing the following items:

call the matched call.

obj. function The percentiles used in the quantile regression coefficient modeling or objective

function $0_k(\alpha)$.

centers The curves matrix.
radius The vector of clusters.

clusters The mean curves matrix of dimension n x k.

Xorig The atrix of 'curves' of dimension n x q.

fd The object obtained by the call of FPCA of class 'fd'

X The matrix of 'curves' transformed through FPCA of dimension p x nharmonics.

X. mean The mean curves matrix of dimension n x k.

diss.matrix The Euclidean distance matrix of the transformed curves.

oggSilhouette An object of class 'silhouette'.

Author(s)

Gianluca Sottile < gianluca.sottile@unipa.it>

opt.fpcac 13

References

Adelfio, G., Chiodi, M., D'Alessandro, A. and Luzio, D. (2011) FPCA algorithm for waveform clustering. Journal of Communication and Computer, 8(6), 494-502.

Adelfio, G., Chiodi, M., D'Alessandro, A., Luzio, D., D'Anna, G., Mangano, G. (2012) Simultaneous seismic wave clustering and registration. Computers & Geosciences 44, 60-69.

Garcia-Escudero, L. A. and Gordaliza, A. (2005). A proposal for robust curve clustering, Journal of classification, 22, 185-201.

See Also

```
opt.fpcac.
```

Examples

```
set.seed(1234)
n <- 300
x <- 1:n/n

Y <- matrix(0, n, 30)

sigma2 <- 4*pmax(x-.2, 0) - 8*pmax(x-.5, 0) + 4*pmax(x-.8, 0)

mu <- sin(3*pi*x)
for(i in 1:10) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

mu <- cos(3*pi*x)
for(i in 11:23) Y[,i] <- mu + rnorm(length(x), 0, pmax(sigma2,0))

mu <- sin(3*pi*x)*cos(pi*x)
for(i in 24:28) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

mu <- 0 #sin(1/3*pi*x)*cos(2*pi*x)
for(i in 29:30) Y[, i] <- mu + rnorm(length(x), 0, pmax(sigma2, 0))

obj <- fpcac(Y, K = 4, disp = FALSE)
obj</pre>
```

opt.fpcac

Optimal cluster selection in Functional Principal Components Analysis Clustering

Description

This function provides the optimal selection of clusters for the algorithm FPCAC, as a variant of a k-means algorithm based on the principal component rotation of data

14 opt.fpcac

Usage

Arguments

Χ Matrix of 'curves' of dimension n x q. the number of cluster used in the optimization step to select the optimal one. k.max the method used to select the optimal number of clusters, "silhouette" or "wss" method (whithin sum of squares. fd If not NULL it overrides X and must be an object of class fd. nbasis an integer variable specifying the number of basis functions. The default value norder an integer specifying the order of b-splines, which is one higher than their degree. The default value is 3. nharmonics the number of harmonics or principal components to use. The default value is 3. trimming size, that is the given proportion of observations to be discarded. alpha niter the number or random restarting (larger values provide more accurate solutions. the number of k-mean steps (not too many ksteps are needed). Ksteps the seed used for reproducibility. seed the dissimilarity matrix used to compute measures "silhouette" or "wss". diss

Details

trace

Silhouette is a method for validate the consistency within clusters, providing a measure of how similar an object is to its own cluster compared to other clusters. The silhouette score S belongs to the interval [-1,1]. S close to one means that the data is appropriately clustered. If S is close to negative one, datum should be clustered in its neighbouring cluster. S near zero means that the datum is on the border of two natural clusters.

if TRUE, it is used to print some information across the algorithm.

The wss is obtained as the classical sum of the squared deviations from each observation and the cluster centroid, providing a measure of the variability of the observations within each cluster. Clusters with higher values exhibit greater variability of the observations within the cluster.

Value

a list containing the following items:

obj.function the sequence of objective functions.

clusters the matrix in which each columns identify clusters for each fixed K.

K the sequence of K used.

K. opt the optimal number of clusters

plot a ggplot object to plot the curve of silhouette or whithin sum of squares.

plot.clustEff 15

Author(s)

Gianluca Sottile < gianluca.sottile@unipa.it>

References

Peter J. Rousseeuw (1987). Silhouettes: a Graphical Aid to the Interpretation and Validation of Cluster Analysis. Computational and Applied Mathematics. 20, 53-65

K. V. Mardia, J. T. Kent and J. M. Bibby (1979). Multivariate Analysis. Academic Press.

See Also

fpcac.

Examples

```
set.seed(1234)
n <- 300
x <- 1:n/n
Y <- matrix(0, n, 30)
sigma2 < -4*pmax(x-.2, 0) - 8*pmax(x-.5, 0) + 4*pmax(x-.8, 0)
mu <- sin(3*pi*x)
for(i in 1:10) Y[, i] \leftarrow mu + rnorm(length(x), 0, pmax(sigma2, 0))
mu <- cos(3*pi*x)
for(i in 11:23) Y[,i] \leftarrow mu + rnorm(length(x), 0, pmax(sigma2,0))
mu <- sin(3*pi*x)*cos(pi*x)</pre>
for(i in 24:28) Y[, i] \leftarrow mu + rnorm(length(x), 0, pmax(sigma2, 0))
mu <- 0 \#sin(1/3*pi*x)*cos(2*pi*x)
for(i in 29:30) Y[, i] \leftarrow mu + rnorm(length(x), 0, pmax(sigma2, 0))
num.clust <- opt.fpcac(Y)</pre>
obj2 <- fpcac(Y, K = num.clust$K.opt, disp = FALSE)</pre>
obj2
```

plot.clustEff

Plot Clustering Effects

Description

Produces a dendrogram, a cluster plot and a boxplot of average distance cluster of an object of class "clustEff".

plot.clustEff

Usage

Arguments

x	An object of class "clustEdd", typically the result of a call to clustEff.	
xvar	Clusters: plot of the k clusters; Dendrogram: plot of the tree after computing the dissimilarity measure and applying a hierarchical clustering algorithm; Boxplot: plot the average distance within clusters; Numclust: plot the curve to minimize to select the best number of clusters;	
which	If missing all curves effect are plotted.	
polygon	If TRUE confidence intervals are represented by shaded areas via polygon. Otherwise, dashed lines are used. If NULL no confidence intervals are represented	
dissimilarity	If TRUE dissimilarity measure within each cluster is used to do boxplot representation.	
par	If TRUE the screen is automaticcaly splitted.	
	additional graphical parameters, that can include xlim, ylim, xlab, ylab,	

Details

Different plot for the clustering algorithm.

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.ot>

col, lwd, lty. See par.

See Also

clustEff for cluster algorithm; extract.object for extracting information through a quantile
regression coefficient modeling in a multivariate case; summary.clustEff for clustering summary.

Examples

```
# using simulated data
# see the documentation for 'clustEff'
```

plot.fpcac 17

pl	ot		fp	ca	c
P -	·	•	י ר	Cu	·

Plot Functional Principal Component Analysis Clustering

Description

Produces a cluster plot of an object of class "fpcac".

Usage

```
## S3 method for class 'fpcac'
plot(x, which, polygon=TRUE, ...)
```

Arguments

x	An object of class "clustEdd", typically the result of a call to fpcac.	
which	If missing all curves effect are plotted.	
polygon	If TRUE confidence intervals are represented by shaded areas via polygon. Otherwise, dashed lines are used. If NULL no confidence intervals are represented	
	additional graphical parameters, that can include xlim, ylim, xlab, ylab, col, lwd, lty. See par.	

Details

Different plot for the clustering algorithm.

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.ot>

See Also

```
fpcac, summary.fpcac, opt.fpcac.
```

Examples

```
# using simulated data
# see the documentation for 'fpcac'
```

18 summary.clustEff

summary.clustEff	Summary clustEff algorithm
Julillar y.Clubceri	Summary Custer at Somme

Description

Summary of an object of class "clustEff".

Usage

```
## S3 method for class 'clustEff'
summary(object, ...)
```

Arguments

object An object of class "clustEff", the result of a call to clustEff.
... for future methods.

Details

A summary of the clustering algorithm is printed.

Value

The following items is returned:

k The number of selected clusters.n The number of observations.p The number of curves.

step The selected step for computing the dissimilarity matrix.

alpha The alpha-percentile used for computing the dissimilarity matrix.

method The selected method to compute the hierarchical cluster analysis.

cut.method The selected method to choose the best number of clusters.

tabClust The table of clusters.

avClust The average distance within clusters.

avSilhouette Silhouette widths for clusters.

avDiss The average dissimilarity measure within clusters.

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.it>

See Also

clustEff, for cluster algorithmextract.object for extracting information through a quantile regression coefficient modeling in a multivariate case and plotting objects of class "clustEff".

summary.fpcac 19

Examples

```
# using simulated data
# see the documentation for 'clustEff'
```

summary.fpcac

Summary FPCAC algorithm

Description

Summary of an object of class "fpcac".

Usage

```
## S3 method for class 'fpcac'
summary(object, ...)
```

Arguments

object An object of class "fpcac", the result of a call to fpcac.
... for future methods.

Details

A summary of the clustering algorithm is printed.

Value

The following items is returned:

k The number of selected clusters.

n The number of curves.

p The number of harmonics used. trimmed The number of trimmed curves.

tabClust The table of clusters.

avClust The average distance within clusters.

Author(s)

Gianluca Sottile <gianluca.sottile@unipa.it>

See Also

```
fpcac, opt.fpcac
```

20 summary.fpcac

Examples

```
# using simulated data
```

see the documentation for 'fpcac'

Index

```
*Topic clustering algorithm
    clustEff, 3
    distshape, 8
    fpcac, 11
    opt.fpcac, 13
*Topic methods
    plot.clustEff, 15
    plot.fpcac, 17
*Topic models
    clustEff, 3
    distshape, 8
    fpcac, 11
    opt.fpcac, 13
*Topic package
    clustEff-package, 2
clustEff, 2, 3, 9, 11, 16, 18
clustEff-package, 2
distshape, 8
extract.object, 2, 5, 9, 10, 16, 18
fpcac, 2, 11, 15, 17, 19
opt.fpcac, 13, 13, 17, 19
par, 16, 17
plot.clustEff, 2, 5, 9, 11, 15
plot.fpcac, 2, 17
summary.clustEff, 2, 5, 9, 11, 16, 18
summary.fpcac, 2, 17, 19
```