

Package ‘amap’

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Title Another Multidimensional Analysis Package

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Suggests Biobase

Description Tools for Clustering and Principal Component Analysis
(With robust methods, and parallelized functions).

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Author Antoine Lucas [aut, cre]

Maintainer Antoine Lucas <antoinelucas@gmail.com>

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acp

*Principal component analysis***Description**

Principal component analysis

Usage

```
acp(x,center=TRUE,reduce=TRUE,wI=rep(1,nrow(x)),wV=rep(1,ncol(x)))
pca(x,center=TRUE,reduce=TRUE,wI=rep(1,nrow(x)),wV=rep(1,ncol(x)))
## S3 method for class 'acp'
print(x, ...)
```

Arguments

x	Matrix / data frame
center	a logical value indicating whether we center data
reduce	a logical value indicating whether we "reduce" data i.e. divide each column by standard deviation
wI, wV	weighth vector for individuals / variables
...	arguments to be passed to or from other methods.

Details

This function offer a variant of [princomp](#) and [prcomp](#) functions, with a slightly different graphic representation (see [plot.acp](#)).

Value

An object of class **acp** The object is a list with components:

sdev	the standard deviations of the principal components.
loadings	the matrix of variable loadings (i.e., a matrix whose columns contain the eigen-vectors). This is of class "loadings": see loadings for its print method.
scores	if scores = TRUE, the scores of the supplied data on the principal components.
eig	Eigen values

Author(s)

Antoine Lucas

See Also

[plot.acp](#), [acpgen](#), [princomp](#)

Examples

```
data(lubisch)
lubisch <- lubisch[,-c(1,8)]
p <- acp(lubisch)
plot(p)
```

acpgen

*Generalised principal component analysis***Description**

Generalised principal component analysis

Usage

```
acpgen(x, h1, h2, center=TRUE, reduce=TRUE, kernel="gaussien")
K(u, kernel="gaussien")
W(x, h, D=NULL, kernel="gaussien")
```

Arguments

x	Matrix or data frame
h	Scalar: bandwidth of the Kernel
h1	Scalar: bandwidth of the Kernel for W
h2	Scalar: bandwidth of the Kernel for U
kernel	The kernel used. This must be one of "gaussien", "quartic", "triweight", "epanechikov", "cosinus" or "uniform"
center	A logical value indicating whether we center data
reduce	A logical value indicating whether we "reduce" data i.e. divide each column by standard deviation
D	A product scalar matrix / une matrice de produit scalaire
u	Vector

Details

acpgen compute generalised pca. i.e. spectral analysis of $U_n.W_n^{-1}$, and project X_i with W_n^{-1} on the principal vector sub-spaces.

X_i a column vector of p variables of individu i (input data)

W compute estimation of noise in the variance.

$$W_n = \frac{\sum_{i=1}^{n-1} \sum_{j=i+1}^n K(\|X_i - X_j\|_{V_n^{-1}}/h)(X_i - X_j)(X_i - X_j)'}{\sum_{i=1}^{n-1} \sum_{j=i+1}^n K(\|X_i - X_j\|_{V_n^{-1}}/h)}$$

with V_n variance estimation;

U compute robust variance. $U_n^{-1} = S_n^{-1} - 1/hV_n^{-1}$

$$S_n = \frac{\sum_{i=1}^n K(\|X_i\|_{V_n^{-1}}/h)(X_i - \mu_n)(X_i - \mu_n)'}{\sum_{i=1}^n K(\|X_i\|_{V_n^{-1}}/h)}$$

with μ_n estimator of the mean.

K compute kernel, i.e.

gaussien:

$$\frac{1}{\sqrt{2\pi}} e^{-u^2/2}$$

quartic:

$$\frac{15}{16}(1-u^2)^2 I_{|u|\leq 1}$$

triweight:

$$\frac{35}{32}(1-u^2)^3 I_{|u|\leq 1}$$

epanechikov:

$$\frac{3}{4}(1-u^2) I_{|u|\leq 1}$$

cosinus:

$$\frac{\pi}{4} \cos\left(\frac{\pi}{2}u\right) I_{|u|\leq 1}$$

Value

An object of class **acp** The object is a list with components:

sdev	the standard deviations of the principal components.
loadings	the matrix of variable loadings (i.e., a matrix whose columns contain the eigenvectors). This is of class "loadings": see loadings for its print method.
scores	if scores = TRUE, the scores of the supplied data on the principal components.
eig	Eigen values

Author(s)

Antoine Lucas

References

H. Caussinus, M. Fekri, S. Hakam and A. Ruiz-Gazen, *A monitoring display of multivariate outliers* Computational Statistics & Data Analysis, Volume 44, Issues 1-2, 28 October 2003, Pages 237-252

Caussinus, H and Ruiz-Gazen, A. (1993): *Projection Pursuit and Generalized Principal Component Analyses*, in *New Directions in Statistical Data Analysis and Robustness* (eds. Morgenthaler et al.), pp. 35-46. Birkhäuser Verlag Basel.

Caussinus, H. and Ruiz-Gazen, A. (1995). *Metrics for Finding Typical Structures by Means of Principal Component Analysis*. In *Data Science and its Applications* (eds Y. Escoufier and C. Hayashi), pp. 177-192. Tokyo: Academic Press.

Antoine Lucas and Sylvain Jasson, *Using amap and ctc Packages for Huge Clustering*, R News, 2006, vol 6, issue 5 pages 58-60.

See Also

[acp](#) [acprob](#) [princomp](#)

Examples

```
data(lubisch)
lubisch <- lubisch[,-c(1,8)]
p <- acpgen(lubisch,h1=1,h2=1/sqrt(2))
plot(p,main='ACP robuste des individus')

# See difference with acp

p <- princomp(lubisch)
class(p)<- "acp"
```

acprob

Robust principal component analysis

Description

Robust principal component analysis

Usage

```
acprob(x,h,center=TRUE,reduce=TRUE,kernel="gaussien")
```

Arguments

x	Matrix / data frame
h	Scalar: bandwidth of the Kernel
kernel	The kernel used. This must be one of "gaussien", "quartic", "triweight", "epanechikov", "cosinus" or "uniform"
center	A logical value indicating whether we center data
reduce	A logical value indicating whether we "reduce" data i.e. divide each column by standard deviation

Details

acpgen compute robust pca. i.e. spectral analysis of a robust variance instead of usual variance. Robust variance: see [varrob](#)

Value

An object of class **acp** The object is a list with components:

sdev	the standard deviations of the principal components.
loadings	the matrix of variable loadings (i.e., a matrix whose columns contain the eigenvectors). This is of class "loadings": see loadings for its print method.
scores	if scores = TRUE, the scores of the supplied data on the principal components.
eig	Eigen values

Author(s)

Antoine Lucas

References

H. Caussinus, M. Fekri, S. Hakam and A. Ruiz-Gazen, *A monitoring display of multivariate outliers* Computational Statistics & Data Analysis, Volume 44, Issues 1-2, 28 October 2003, Pages 237-252

Caussinus, H and Ruiz-Gazen, A. (1993): *Projection Pursuit and Generalized Principal Component Analyses*, in *New Directions in Statistical Data Analysis and Robustness* (eds. Morgenthaler et al.), pp. 35-46. Birkhäuser Verlag Basel.

Caussinus, H. and Ruiz-Gazen, A. (1995). *Metrics for Finding Typical Structures by Means of Principal Component Analysis*. In *Data Science and its Applications* (eds Y. Escoufier and C. Hayashi), pp. 177-192. Tokyo: Academic Press.

Antoine Lucas and Sylvain Jasson, *Using amap and ctc Packages for Huge Clustering*, R News, 2006, vol 6, issue 5 pages 58-60.

See Also

[princomp](#) [acpgen](#)

afc

Correspondance factorial analysis.

Description

Compute an acp on a contingency table tacking into account weight of rows and columns

Usage

afc(x)

Arguments

x A contingency table, or a result of function burt or matlogic

Author(s)

Antoine Lucas

Examples

```
## Not run:
color <- as.factor(c('blue','red','red','blue','red'))
size <- as.factor(c('large','large','small','medium','large'))
x <- data.frame(color,size)

afc.1 <- afc(burt(x))
afc.2 <- afc(matlogic(x))

plotAll(afc.1)
plotAll(afc.2)

## End(Not run)
```

burt

Compute burt table from a factor dataframe.

Description

matlogic returns for all variables a matrix of logical values for each levels. burt is defined as `t(matlogic).matlogic`

Usage

```
burt(x)
matlogic(x)
```

Arguments

x A dataframe that contents only factors

Author(s)

Antoine Lucas

Examples

```
color <- as.factor(c('blue','red','red','blue','red'))
size <- as.factor(c('large','large','small','medium','large'))
x <- data.frame(color,size)

matlogic(x)
## color.blue color.red size.large size.medium size.small
##1          1          0          1          0          0
##2          0          1          1          0          0
```

```
##3      0      1      0      0      1
##4      1      0      0      1      0
##5      0      1      1      0      0

burt(x)
##           color.blue color.red size.large size.medium size.small
## color.blue           2         0         1         1         0
## color.red            0         3         2         0         1
## size.large           1         2         3         0         0
## size.medium          1         0         0         1         0
## size.small           0         1         0         0         1
```

diss *Compute a dissimilarity matrix*

Description

Compute a dissimilarity matrix from a data set (containing only factors).

Usage

```
diss(x, w=rep(1,ncol(x)) )
```

Arguments

x A matrix or data frame containing only factors.
w A vector of weight, by default each variable has got same weight

Details

Case of N individuals described by P categorical variables: each element (i,j) of the signed similarities array is computed by summation over the P variables of the contributions of each variable, multiplied by the weight of the variable. The contribution of a given categorical variable is +1 if the individual i and j are in the same class, and is -1 if they are not.

Value

A dissimilarity matrix.

Author(s)

Antoine Lucas

See Also

[Dist](#), [pop](#)

Examples

```

data <-
matrix(c(1,1,1,1,1
,1,2,1,2,1
,2,3,2,3,2
,2,4,3,3,2
,1,2,4,2,1
,2,3,2,3,1),ncol=5,byrow=TRUE)

diss(data)

## With weights
diss(data,w=c(1,1,2,2,3))

```

Dist

Distance Matrix Computation

Description

This function computes and returns the distance matrix computed by using the specified distance measure to compute the distances between the rows of a data matrix.

Usage

```
Dist(x, method = "euclidean", nbproc = 2, diag = FALSE, upper = FALSE)
```

Arguments

x	numeric matrix or (data frame) or an object of class "exprSet". Distances between the rows of x will be computed.
method	the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary", "pearson", "abspearson", "correlation", "abscorrelation", "spearman" or "kendall". Any unambiguous substring can be given.
nbproc	integer, Number of subprocess for parallelization
diag	logical value indicating whether the diagonal of the distance matrix should be printed by <code>print.dist</code> .
upper	logical value indicating whether the upper triangle of the distance matrix should be printed by <code>print.dist</code> .

Details

Available distance measures are (written for two vectors x and y):

euclidean: Usual square distance between the two vectors (2 norm).

maximum: Maximum distance between two components of x and y (supremum norm)

manhattan: Absolute distance between the two vectors (1 norm).

canberra: $\sum_i |x_i - y_i| / |x_i + y_i|$. Terms with zero numerator and denominator are omitted from the sum and treated as if the values were missing.

binary: (aka *asymmetric binary*): The vectors are regarded as binary bits, so non-zero elements are 'on' and zero elements are 'off'. The distance is the *proportion* of bits in which only one is on amongst those in which at least one is on.

pearson: Also named "not centered Pearson" $1 - \frac{\sum_i x_i y_i}{\sqrt{\sum_i x_i^2 \sum_i y_i^2}}$.

abspearson: Absolute Pearson $1 - \left| \frac{\sum_i x_i y_i}{\sqrt{\sum_i x_i^2 \sum_i y_i^2}} \right|$.

correlation: Also named "Centered Pearson" $1 - \text{corr}(x, y)$.

abs correlation: Absolute correlation $1 - |\text{corr}(x, y)|$ with

$$\text{corr}(x, y) = \frac{\sum_i x_i y_i - \frac{1}{n} \sum_i x_i \sum_i y_i}{\sqrt{(\sum_i x_i^2 - \frac{1}{n} (\sum_i x_i)^2) (\sum_i y_i^2 - \frac{1}{n} (\sum_i y_i)^2)}}.$$

spearman: Compute a distance based on rank. $\sum (d_i^2)$ where d_i is the difference in rank between x_i and y_i .

```
Dist(x, method="spearman")[i, j] =
```

```
cor.test(x[i, ], x[j, ], method="spearman")$statistic
```

kendall: Compute a distance based on rank. $\sum_{i,j} K_{i,j}(x, y)$ with $K_{i,j}(x, y)$ is 0 if x_i, x_j in same order as y_i, y_j , 1 if not.

Missing values are allowed, and are excluded from all computations involving the rows within which they occur. If some columns are excluded in calculating a Euclidean, Manhattan or Canberra distance, the sum is scaled up proportionally to the number of columns used. If all pairs are excluded when calculating a particular distance, the value is NA.

The functions `as.matrix.dist()` and `as.dist()` can be used for conversion between objects of class "dist" and conventional distance matrices and vice versa.

Value

An object of class "dist".

The lower triangle of the distance matrix stored by columns in a vector, say `do`. If n is the number of observations, i.e., $n <- \text{attr}(do, "Size")$, then for $i < j \leq n$, the dissimilarity between (row) i and j is `do[n*(i-1) - i*(i-1)/2 + j-i]`. The length of the vector is $n * (n - 1) / 2$, i.e., of order n^2 .

The object has the following attributes (besides "class" equal to "dist"):

Size	integer, the number of observations in the dataset.
Labels	optionally, contains the labels, if any, of the observations of the dataset.
Diag, Upper	logicals corresponding to the arguments <code>diag</code> and <code>upper</code> above, specifying how the object should be printed.
call	optionally, the <code>call</code> used to create the object.
methods	optionally, the distance method used; resulting form <code>dist()</code> , the <code>(match.arg())</code> ed method argument.

Note

Multi-thread (parallelisation) is disable on Windows.

References

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

Wikipedia https://en.wikipedia.org/wiki/Kendall_tau_distance

See Also

[daisy](#) in the 'cluster' package with more possibilities in the case of *mixed* (contiuous / categorical) variables. [dist](#) [hcluster](#).

Examples

```
x <- matrix(rnorm(100), nrow=5)
Dist(x)
Dist(x, diag = TRUE)
Dist(x, upper = TRUE)

## compute dist with 8 threads
Dist(x,nbproc=8)

Dist(x,method="abscorrelation")
Dist(x,method="kendall")
```

hcluster

Hierarchical Clustering

Description

Hierarchical cluster analysis.

Usage

```
hcluster(x, method = "euclidean", diag = FALSE, upper = FALSE,
         link = "complete", members = NULL, nbproc = 2,
         doubleprecision = TRUE)
```

Arguments

x A numeric matrix of data, or an object that can be coerced to such a matrix (such as a numeric vector or a data frame with all numeric columns). Or an object of class "exprSet".

method	the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary", "pearson", "abspearson", "correlation", "abscorrelation", "spearman" or "kendall". Any unambiguous substring can be given.
diag	logical value indicating whether the diagonal of the distance matrix should be printed by <code>print.dist</code> .
upper	logical value indicating whether the upper triangle of the distance matrix should be printed by <code>print.dist</code> .
link	the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid", "centroid2", "ward.d2".
members	NULL or a vector with length size of <code>d</code> .
nbproc	integer, number of subprocess for parallelization [Linux & Mac only]
doubleprecision	True: use of double precision for distance matrix computation; False: use simple precision

Details

This function is a mix of function `hclust` and function `dist`. `hcluster(x, method = "euclidean", link = "complete") = hclust(dist(x, method = "euclidean"), method = "complete")` It use twice less memory, as it doesn't store distance matrix.

For more details, see documentation of `hclust` and `Dist`.

Value

An object of class **hclust** which describes the tree produced by the clustering process. The object is a list with components:

merge	an $n - 1$ by 2 matrix. Row i of merge describes the merging of clusters at step i of the clustering. If an element j in the row is negative, then observation $-j$ was merged at this stage. If j is positive then the merge was with the cluster formed at the (earlier) stage j of the algorithm. Thus negative entries in merge indicate agglomerations of singletons, and positive entries indicate agglomerations of non-singletons.
height	a set of $n - 1$ non-decreasing real values. The clustering <i>height</i> : that is, the value of the criterion associated with the clustering method for the particular agglomeration.
order	a vector giving the permutation of the original observations suitable for plotting, in the sense that a cluster plot using this ordering and matrix merge will not have crossings of the branches.
labels	labels for each of the objects being clustered.
call	the call which produced the result.
method	the cluster method that has been used.
dist.method	the distance that has been used to create <code>d</code> (only returned if the distance object has a "method" attribute).

There is a `print` and a `plot` method for `hclust` objects. The `plclust()` function is basically the same as the `plot` method, `plot.hclust`, primarily for back compatibility with S-plus. Its extra arguments are not yet implemented.

Note

Multi-thread (parallelisation) is disabled on Windows.

Author(s)

The `hcluster` function is based on C code adapted from Cran Fortran routine by Antoine Lucas.

References

Antoine Lucas and Sylvain Jasson, *Using amap and ctc Packages for Huge Clustering*, R News, 2006, vol 6, issue 5 pages 58-60.

See Also

[Dist](#), [hclust](#), [kmeans](#).

Examples

```
data(USArrests)
hc <- hcluster(USArrests,link = "ave")
plot(hc)
plot(hc, hang = -1)

## Do the same with centroid clustering and squared Euclidean distance,
## cut the tree into ten clusters and reconstruct the upper part of the
## tree from the cluster centers.
hc <- hclust(dist(USArrests)^2, "cen")
memb <- cutree(hc, k = 10)
cent <- NULL
for(k in 1:10){
  cent <- rbind(cent, colMeans(USArrests[memb == k, , drop = FALSE]))
}
hc1 <- hclust(dist(cent)^2, method = "cen", members = table(memb))
opar <- par(mfrow = c(1, 2))
plot(hc, labels = FALSE, hang = -1, main = "Original Tree")
plot(hc1, labels = FALSE, hang = -1, main = "Re-start from 10 clusters")
par(opar)

## other combinaison are possible

hc <- hcluster(USArrests,method = "euc",link = "ward", nbproc= 1,
doubleprecision = TRUE)
hc <- hcluster(USArrests,method = "max",link = "single", nbproc= 2,
doubleprecision = TRUE)
hc <- hcluster(USArrests,method = "man",link = "complete", nbproc= 1,
doubleprecision = TRUE)
```

```

hc <- hcluster(USArrests,method = "can",link = "average", nbproc= 2,
doubleprecision = TRUE)
hc <- hcluster(USArrests,method = "bin",link = "mcquitty", nbproc= 1,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "pea",link = "median", nbproc= 2,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "abspea",link = "median", nbproc= 2,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "cor",link = "centroid", nbproc= 1,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "absacor",link = "centroid", nbproc= 1,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "spe",link = "complete", nbproc= 2,
doubleprecision = FALSE)
hc <- hcluster(USArrests,method = "ken",link = "complete", nbproc= 2,
doubleprecision = FALSE)

```

Kmeans

K-Means Clustering

Description

Perform k-means clustering on a data matrix.

Usage

```

Kmeans(x, centers, iter.max = 10, nstart = 1,
method = "euclidean")

```

Arguments

x	A numeric matrix of data, or an object that can be coerced to such a matrix (such as a numeric vector or a data frame with all numeric columns). Or an object of class "exprSet".
centers	Either the number of clusters or a set of initial cluster centers. If the first, a random set of rows in x are chosen as the initial centers.
iter.max	The maximum number of iterations allowed.
nstart	If centers is a number, how many random sets should be chosen?
method	the distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary", "pearson", "abspearson", "absacorrelation", "correlation", "spearman" or "kendall". Any unambiguous substring can be given.

Details

The data given by `x` is clustered by the k-means algorithm. When this terminates, all cluster centres are at the mean of their Voronoi sets (the set of data points which are nearest to the cluster centre).

The algorithm of Lloyd–Forgy is used; `method="euclidean"` should return same result as with function [kmeans](#).

Value

A list with components:

<code>cluster</code>	A vector of integers indicating the cluster to which each point is allocated.
<code>centers</code>	A matrix of cluster centres.
<code>withinss</code>	The within-cluster sum of square distances for each cluster.
<code>size</code>	The number of points in each cluster.

Note

An objective: to allow NA values.

See Also

[hcluster](#), [kmeans](#).

Examples

```
## a 2-dimensional example
x <- rbind(matrix(rnorm(100, sd = 0.3), ncol = 2),
           matrix(rnorm(100, mean = 1, sd = 0.3), ncol = 2))
colnames(x) <- c("x", "y")
(cl <- Kmeans(x, 2))
plot(x, col = cl$cluster)
points(cl$centers, col = 1:2, pch = 8, cex=2)

## random starts do help here with too many clusters
(cl <- Kmeans(x, 5, nstart = 25))
plot(x, col = cl$cluster)
points(cl$centers, col = 1:5, pch = 8)

Kmeans(x, 5, nstart = 25, method="abscorrelation")
```

lubisch	<i>Dataset Lubischew</i>
---------	--------------------------

Description

Lubischew data (1962): 74 insects, 6 morphologic size. 3 supposed classes

Usage

```
data(lubisch)
```

plot	<i>Graphics for Principal component Analysis</i>
------	--

Description

Graphics for Principal component Analysis

Usage

```
## S3 method for class 'acp'
plot(x,i=1,j=2,text=TRUE,label='Composants',col='darkblue',
main='Individuals PCA',variables=TRUE,individual.label=NULL,...)
## S3 method for class 'acp'
biplot(x,i=1,j=2,label='Composants',col='darkblue',length=0.1,
main='Variables PCA',circle=TRUE,...)
plot2(x,pourcent=FALSE,eigen=TRUE,label='Comp.',col='lightgrey',
main='Scree Graph',ylab='Eigen Values')
plotAll(x)
```

Arguments

x	Result of acp or princomp
i	X axis
j	Y axis
text	a logical value indicating whether we use text or points for plot
pourcent	a logical value indicating whether we use pourcentage of values
eigen	a logical value indicating whether we use eigen values or standard deviation
label	label for X and Y axis
individual.label	labels naming individuals
col	Color of plot
main	Title of graphic

ylab	Y label
length	length of arrows
variables, circle	a logical value indicating whether we display circle or variables
...	cex, pch, and other options; see points.

Value

Graphics:
 plot.acp PCA for lines (individuals)
 plot.acp PCA for columns (variables)
 plot2 Eigen values diagram (Scree Graph)
 plotAll Plot both 3 graphs

Author(s)

Antoine Lucas

See Also

[acpgen](#), [acprob](#), [princomp](#)

Examples

```
data(lubisch)
lubisch <- lubisch[,-c(1,8)]
p <- acp(lubisch)
plotAll(p)
```

pop *Optimal Partition (classification).*

Description

Classification: Computing an Optimal Partition from Weighted Categorical Variables or from an Array of Signed Similarities.

Usage

```
pop(x, fmbvr=TRUE, triabs=TRUE, allsol=TRUE)
```

Arguments

x	A dissimilarity matrix
fmbvr	Logical, TRUE: look for the exact solution
triabs	Logical, TRUE: try to init with absolute values
allsol	Logical, TRUE all solutions, FALSE only one solution

Author(s)

Michel Petitjean, <http://petitjeanmichel.free.fr/itoweb.petitjean.class.html>

R port by Antoine Lucas,

References

Theory is explained at <http://petitjeanmichel.free.fr/itoweb.petitjean.class.html>

Marcotorchino F. *Agrégation des similarités en classification automatique*. Thèse de Doctorat d'Etat en Mathématiques, Université Paris VI, 25 June 1981.

Petitjean M. *Agrégation des similarités: une solution oubliée*. RAIRO Oper. Res. 2002,36[1],101-108.

Examples

```
## pop from a data matrix
data <-
matrix(c(1,1,1,1,1
        ,1,2,1,2,1
        ,2,3,2,3,2
        ,2,4,3,3,2
        ,1,2,4,2,1
        ,2,3,2,3,1),ncol=5,byrow=TRUE)
```

```
pop(diss(data))
```

```
## pop from a dissimilarity matrix
```

```
d <- 2 * matrix(c(9, 8, 5, 7, 7, 2
, 8, 9, 2, 5, 1, 7
, 5, 2, 9, 8, 7, 1
, 7, 5, 8, 9, 3, 2
, 7, 1, 7, 3, 9, 6
, 2, 7, 1, 2, 6, 9),ncol=6,byrow=TRUE) - 9
```

```
pop(d)
```

```
## Not run:
```

```
d <- 2 * matrix(c(57, 15, 11, 32, 1, 34, 4, 6, 17, 7
, 15, 57, 27, 35, 27, 27, 20, 24, 30, 15
, 11, 27, 57, 25, 25, 20, 34, 25, 17, 15
, 32, 35, 25, 57, 22, 44, 13, 22, 30, 11
, 1, 27, 25, 22, 57, 21, 28, 43, 20, 13
, 34, 27, 20, 44, 21, 57, 18, 27, 21, 8
, 4, 20, 34, 13, 28, 18, 57, 31, 28, 13
, 6, 24, 25, 22, 43, 27, 31, 57, 30, 15
, 17, 30, 17, 30, 20, 21, 28, 30, 57, 12
```

```
, 7, 15, 15, 11, 13, 8, 13, 15, 12, 57),ncol=10,byrow=TRUE) - 57
pop(d)

## End(Not run)
```

 VarRob

Robust variance

Description

Compute a robust variance

Usage

```
varrob(x,h,D=NULL,kernel="gaussien")
```

Arguments

x	Matrix / data frame
h	Scalar: bandwidth of the Kernel
kernel	The kernel used. This must be one of "gaussien", "quartic", "triweight", "epanechikov", "cosinus" or "uniform"
D	A product scalar matrix / une matrice de produit scalaire

Details

U compute robust variance. $U_n^{-1} = S_n^{-1} - 1/hV_n^{-1}$

$$S_n = \frac{\sum_{i=1}^n K(\|X_i\|_{V_n^{-1}}/h)(X_i - \mu_n)(X_i - \mu_n)'}{\sum_{i=1}^n K(\|X_i\|_{V_n^{-1}}/h)}$$

with μ_n estimator of the mean.

K compute a kernel.

Value

A matrix

Author(s)

Antoine Lucas

References

H. Caussinus, S. Hakam, A. Ruiz-Gazen Projections revelatrices controlees: groupements et structures diverses. 2002, to appear in Rev. Statist. Appli.

See Also

[acp princomp](#)

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