Package 'gclus'

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Title Clustering Graphics

Description Orders panels in scatterplot matrices and parallel coordinate displays by some merit index. Package contains various indices of merit, ordering functions, and enhanced versions of pairs and parcoord which color panels according to their merit level.

Depends R (\geq 2.10), cluster

License GPL (>= 2)

Suggests knitr, rmarkdown

VignetteBuilder knitr

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ac

Clustering coefficients from package cluster.

Description

Computes clustering coefficients from cluster, where x and y give the object coordinates.

Usage

ac(x, y, ...) sil(x, y, groups, ...)

Arguments

х	is a numeric vector.
У	is a numeric vector.
groups	is a vector of group memberships, used by sil only.
	are passed to agnes in ac and to dist in sil.

Details

ac - Computes clustering coefficient from agnes{cluster}.

sil - Computes the silhouette coefficient from from package cluster.

Value

The clustering coefficient is returned.

Author(s)

Catherine B. Hurley

References

Kaufman, L. and Rousseeuw, P.J. (1990). Finding Groups in Data: An Introduction to Cluster Analysis . Wiley, New York.

bank

See Also

agnes, silhouette, dist

Examples

```
x <- runif(20)
y <- runif(20)
g <- rep(c("a","b"),10)
ac(x,y)
sil(x,y,g)</pre>
```

bank

Swiss bank notes data

Description

Data from "Multivariate Statistics A practical approach", by Bernhard Flury and Hans Riedwyl, Chapman and Hall, 1988, Tables 1.1 and 1.2 pp. 5-8. Six measurements made on 100 genuine Swiss banknotes and 100 counterfeit ones.

Usage

data(bank)

Format

This data frame contains the following columns:

Status: 0 = genuine, 1 = counterfeit

Length: Length of bill, mm

Left: Width of left edge, mm

Right: Width of right edge, mm

Bottom: Bottom margin width, mm

Top: Top margin width, mm

Diagonal: Length of image diagonal, mm

Source

Flury, B. and Riedwyl, H. (1988), *Multivariate Statistics A Practical Approach*, London: Chapman and Hall.

Description

This dataset contains 21 body dimension measurements as well as age, weight, height, and gender on 507 individuals. The 247 men and 260 women were primarily individuals in their twenties and thirties, with a scattering of older men and women, all exercising several hours a week.

Measurements were initially taken by Grete Heinz and Louis J. Peterson - at San Jose State University and at the U.S. Naval Postgraduate School in Monterey, California. Later, measurements were taken at dozens of California health and fitness clubs by technicians under the supervision of one of these authors.

Usage

data(body)

Format

This data frame contains the following columns:

Biacrom: Biacromial diameter (cm)

Biiliac: Biiliac diameter, or "pelvic breadth" (cm)

Bitro: Bitrochanteric diameter (cm)

ChestDp: Chest depth between spine and sternum at nipple level, mid-expiration (cm)

ChestD: Chest diameter at nipple level, mid-expiration (cm)

ElbowD: Elbow diameter, sum of two elbows (cm)

WristD: Wrist diameter, sum of two wrists (cm)

KneeD: Knee diameter, sum of two knees (cm)

AnkleD: Ankle diameter, sum of two ankles (cm)

ShoulderG: Shoulder girth over deltoid muscles (cm)

ChestG: Chest girth, nipple line in males and just above breast tissue in females, mid-expiration (cm)

WaistG: Waist girth, narrowest part of torso below the rib cage, average of contracted and relaxed position (cm)

AbdG: Navel (or "Abdominal") girth at umbilicus and iliac crest, iliac crest as a landmark (cm)

HipG: Hip girth at level of bitrochanteric diameter (cm)

ThighG: Thigh girth below gluteal fold, average of right and left girths (cm)

BicepG: Bicep girth, flexed, average of right and left girths (cm)

ForearmG: Forearm girth, extended, palm up, average of right and left girths (cm)

KneeG: Knee girth over patella, slightly flexed position, average of right and left girths (cm)

body

colpairs

CalfG: Calf maximum girth, average of right and left girths (cm)
AnkleG: Ankle minimum girth, average of right and left girths (cm)
WristG: Wrist minimum girth, average of right and left girths (cm)
Age: in years
Weight: in kg
Height: in cm
Gender: 1 - male, 0 - female

Source

Heinz, G., Peterson, L.J., Johnson, R.W. and Kerk, C.J. (2003), "Exploring Relationships in Body Dimensions", *Journal of Statistics Education*, 11.

References

The data file is taken from http://jse.amstat.org/datasets/body.dat.txt This information file is based on http://jse.amstat.org/datasets/body.txt

colpairs

Applies a function to all pairs of columns

Description

Given an nxp matrix m and a function f, returns the pxp matrix got by applying f to all pairs of columns of m.

Usage

colpairs(m, f, diag = 0, na.omit = FALSE, ...)

Arguments

m	a matrix
f	a function of two vectors, which returns a single result.
diag	if supplied, this value is placed on the diagonal of the result.
na.omit	If TRUE, rows with missing values are omitted for each pair of columns.
	argments are passed to f.

Value

a matrix matrix got by applying f to all pairs of columns of m.

Author(s)

Catherine B. Hurley

cpairs

See Also

gave, partition.crit, order.single,order.endlink

Examples

```
data(state)
state.m <- colpairs(state.x77,
function(x,y) cor.test(x,y,"two.sided","kendall")$estimate, diag=1)
state.col <- dmat.color(state.m)
# This is equivalent to state.m <- cor(state.x77,method="kendall")</pre>
```

```
layout(matrix(1:2,nrow=1,ncol=2))
cparcoord(state.x77, panel.color= state.col)
# Get rid of the panels with lots of line crossings (yellow) by reorderings
cparcoord(state.x77, order.endlink(state.m), state.col)
layout(matrix(1,1))
```

```
# m is a homogeneity measure of each pairwise variable plot
m <- -colpairs(scale(state.x77), gave)</pre>
```

```
o<- order.single(m)
pcols = dmat.color(m)
# Color panels by level of m and reorder variables so that
# pairs with high m are near the diagonal.
cpairs(state.x77,order=o, panel.colors=pcols)</pre>
```

```
# In this case panels showing either of Area or Population
# exhibit the most clumpiness because these variables
# are skewed.
```

cpairs

Enhanced scatterplot matrix

Description

This function draws a scatterplot matrix of data. Variables may be reordered and panels colored in the display.

Usage

```
cpairs(data, order = NULL, panel.colors = NULL, border.color = "grey70",
show.points = TRUE, ...)
```

cpairs

Arguments

data	a numeric matrix
order	the order of variables. Default is the order in data.
panel.colors	a matrix of panel colors. If supplied, dimensions should match those of the pairs plot. Diagonal entries are ignored.
border.color	used for panel border.
show.points	If FALSE, no points are drawn.
•••	graphical parameters passed to pairs.default.

Author(s)

Catherine B. Hurley

References

Hurley, Catherine B. "Clustering Visualisations of Multidimensional Data", to appear in JCGS.

See Also

pairs, cparcoord, dmat.color,colpairs, order.single.

Examples

```
data(USJudgeRatings)
judge.cor <- cor(USJudgeRatings)</pre>
judge.color <- dmat.color(judge.cor)</pre>
# Colors variables by their correlation.
cpairs(USJudgeRatings,panel.colors=judge.color,pch=".",gap=.5)
judge.o <- order.single(judge.cor)</pre>
# Reorder variables so that those with highest correlation
# are close to the diagonal.
cpairs(USJudgeRatings,judge.o,judge.color,pch=".",gap=.5)
# Specify your own color scheme
judge.color <- dmat.color(judge.cor, breaks=c(-1,0,.5,.9,1), colors =</pre>
cm.colors(4))
data(bank)
# m is a homogeneity measure of each pairwise variable plot
m <- -colpairs(scale(bank[,-1]), partition.crit,gfun=gave,groups=bank[,1])</pre>
# Color panels by level of m and reorder variables so that
# pairs with high m are near the diagonal. Panels shown
# in pink have the highest amount of group homogeneity, as measured by
# gave.
cpairs(bank[,-1],order=order.single(m), panel.colors=dmat.color(m),
gap=.3,col=c("purple","black")[bank[,"Status"]+1],
pch=c(5,3)[bank[,"Status"]+1])
```

cparcoord

Description

This function draws a parallel coordinate plot of data. Variables may be reordered and panels colored in the display. It is a modified version of parcoord {MASS}.

Usage

```
cparcoord(data, order = NULL, panel.colors = NULL, col = 1, lty = 1,
horizontal = FALSE, mar = NULL, ...)
```

Arguments

data	a numeric matrix
order	the order of variables. Default is the order in data.
panel.colors	either a vector or a matrix of panel colors. If a vector is supplied, the ith color is used for the ith panel. If a matrix, dimensions should match those of the variables. Diagonal entries are ignored.
col	a vector of colours, recycled as necessary for each observation.
lty	a vector of line types, recycled as necessary for each observation.
horizontal	If TRUE, orientation is horizontal.
mar	margin parameters, passed to par.
	graphics parameters which are passed to matplot.

Details

If panel.colors is a matrix and order is supplied, panel.colors is reordered.

Author(s)

Catherine B. Hurley

References

Hurley, Catherine B. "Clustering Visualisations of Multidimensional Data", Journal of Computational and Graphical Statistics, vol. 13, (4), pp 788-806, 2004.

See Also

cpairs, parcoord, dmat.color, colpairs, order.endlink.

diameter

Examples

```
data(state)
state.m <- colpairs(state.x77,
function(x,y) cor.test(x,y,"two.sided","kendall")$estimate, diag=1)
# OR, Works only in R1.8, state.m <-cor(state.x77,method="kendall")
state.col <- dmat.color(state.m)
cparcoord(state.x77, panel.color= state.col)
# Get rid of the panels with lots of line crossings (yellow) by reordering:
cparcoord(state.x77, order.endlink(state.m), state.col)
# To get rid of the panels with lots of long line segments:
# use a different panel merit measure- pclen:
mins <- apply(state.x77,2,min)
ranges <- apply(state.x77,2,max) - mins
state.m <- -colpairs(scale(state.x77,mins,ranges), pclen)
cparcoord(state.x77, order.endlink(state.m), dmat.color(state.m))</pre>
```

diameter

Cluster heterogeneity of 2-d data

Description

Computes measures of cluster heterogeneity of 2-d data, where x and y give the object coordinates.

Usage

```
diameter(x, y, ...)
star(x, y, ...)
km2(x,y)
gtot(x,y, ...)
gave(x,y, ...)
```

Arguments

х	is a numeric vector.
У	is a numeric vector.
	are passed to dist.

Details

diameter computes the cluster diameter- the maximum distance between objects.

star computes the cluster star distance- the smallest total distance from one object to another.

km2 computes the kmeans distance.

gtot computes the sum of all inter-object distances.

gave computes the per-object average of all inter-object distances.

Value

The cluster measure is returned.

Author(s)

Catherine B. Hurley

References

See Gordon, A. D. (1999). "Classification". Second Edition. London: Chapman and Hall / CRC

See Also

colpairs, cpairs, order.single

Examples

```
x <- runif(20)
y <- runif(20)
diameter(x,y)</pre>
```

dmat.color

Colors a symmetric matrix

Description

Accepts a dissimilarity matrix or dist m, and returns a matrix of colors. Values in m are cut into categories using breaks (ranked distances if byrank is TRUE) and categories are assigned the values in colors.

Usage

```
dmat.color(m, colors = default.dmat.color, byrank = NULL, breaks = length(colors))
```

Arguments

m	a dissimilarity matrix or the result of dist
colors	a vector of colors. The default is default.dmat.color.
byrank	boolean, default TRUE is unless breaks has length > 1.
breaks	the number of break points.

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dmat.color

Details

breaks are passed to the functioncut. If byrank is TRUE, values in m are ranked before they are categorized. If byrank is TRUE and breaks is an integer, then there are breaks equal-sized categories.

Value

Returns a matrix of colors. The matrix is symmetric, with NAs on the diagonal.

Author(s)

Catherine B. Hurley

See Also

cut, cpairs, cparcoord

Examples

```
data(longley)
longley.cor <- cor(longley)</pre>
# A matrix with equal (or nearly equal) number of entries of each color.
longley.color <- dmat.color(longley.cor)</pre>
# Plot the colors
plotcolors(longley.color,dlabels=rownames(longley.color))
# Try different color schemes
# A matrix where each color represents an equal-length interval.
longley.color <- dmat.color(longley.cor, byrank=FALSE)</pre>
# Specify colors and breaks
longley.color <- dmat.color(longley.cor, breaks=c(-1,0,.5,.8,1),</pre>
cm.colors(4))
# Could also reorder variables prior to plotting:
longley.o <- order.single(longley.cor)</pre>
longley.color <- longley.color[longley.o,longley.o]</pre>
# The colors can be used in a scatterplot matrix or parallel
# coordinate display:
cpairs(longley, panel.color= longley.color)
```

cparcoord(longley, panel.color= longley.color)

order.clusters

Description

Reorders objects so that similar (or high-merit) object pairs are adjacent. The clusters argument specifies (possibly ordered) groups, and objects within a group are kept together.

Usage

Arguments

merit	is either a symmetric matrix of merit or similarity score, or a dist.
clusters	specifies a partial grouping. It should either be a list whose ith element contains the indices of the objects in the ith cluster, or a vector of integers whose ith element gives the cluster membership of the ith object. Either representation may be used to specify grouping, the first is preferrable to specify adjacencies.
within.order	is a function used to order the objects within each cluster.
between.order	is a function used to order the clusters.
	arguments are passed to within.order.

Details

within.order may be NULL, in which case objects within a cluster are assumed to be in order. Otherwise, within.order should be one of the ordering functions order.single,order.endlink or order.hclust.

between.order may be NULL, in which case cluster order is preserved. Otherwise, betweem.order should be one of the ordering functions that uses a partial ordering, order.single or order.endlink.

Value

A permutation of the objects represented by merit is returned.

Author(s)

Catherine B. Hurley

See Also

order.single,order.endlink,order.hclust.

order.clusters

Examples

```
data(state)
state.d <- dist(state.x77)</pre>
# Order the states, keeping states in a division together.
state.o <- order.clusters(-state.d, as.numeric(state.division))</pre>
cmat <- dmat.color(as.matrix(state.d), rev(cm.colors(5)))</pre>
op <- par(mar=c(1,6,1,1))
rlabels <- state.name[state.o]</pre>
plotcolors(cmat[state.o,state.o], rlabels=rlabels)
par(op)
# Alternatively, use kmeans to place the states into 6 clusters
state.km <- kmeans(state.d,6)$cluster</pre>
# An ordering obtained from the kmeans clustering...
state.o <- unlist(memship2clus(state.km))</pre>
layout(matrix(1:2,nrow=1,ncol=2),widths=c(0.1,1))
op <- par(mar=c(1,1,1,.2))
state.colors <- cbind(state.km,state.km)</pre>
plotcolors(state.colors[state.o,])
par(mar=c(1,6,1,1))
rlabels <- state.name[state.o]</pre>
plotcolors(cmat[state.o,state.o], rlabels=rlabels)
par(op)
layout(matrix(1,1))
# In the ordering above, the ordering of clusters and the
# ordering of objects within the clusters is arbitrary.
# order.clusters gives an improved order but preserves the kmeans clusters.
state.o <- order.clusters(-state.d, state.km)</pre>
# and replot
layout(matrix(1:2,nrow=1,ncol=2),widths=c(0.1,1))
op <- par(mar=c(1,1,1,.2))
state.colors <- cbind(state.km,state.km)</pre>
plotcolors(state.colors[state.o,])
par(mar=c(1,6,1,1))
rlabels <- state.name[state.o]</pre>
plotcolors(cmat[state.o,state.o], rlabels=rlabels)
```

```
par(op)
layout(matrix(1,1))
```

order.single

Orders objects using hierarchical clustering

Description

Reorders objects so that similar (or high-merit) object pairs are adjacent. A permutation vector is returned.

Usage

```
order.single(merit,clusters=NULL)
order.endlink(merit,clusters=NULL)
order.hclust(merit, reorder=TRUE,...)
```

Arguments

merit	is either a symmetric matrix of merit or similarity score, or a dist.
clusters	if non-null, specifies a partial ordering. It should be a list whose ith element contains the indices the objects in the ith ordered cluster.
reorder	if TRUE, reorders the default ordering from hclust.
	arguments are passed to hclust.

Details

order.single performs a variation on single-link cluster analysis, devised by Gruvaeus and Wainer (1972). When two ordered clusters are merged, the new cluster is formed by placing the most similar endpoints of the joining clusters adjacent to each other. When applied to variables, the resulting order is useful for scatterplot matrices.

order.endlink is another variation on single-link cluster analysis, where the similarity between two ordered clusters is defined as the minimum distance between their endpoints. When two ordered clusters are merged, the new cluster is formed by placing the most similar endpoints of the joining clusters adjacent to each other. When applied to variables, the resulting order is useful for parallel coordinate displays.

order.hclust returns the order of objects from hclust if reorder is FALSE. Otherwise, it reorders the objects using hclust.reorder so that when two ordered clusters are merged, the new cluster is formed by placing the most similar endpoints of the joining clusters adjacent to each other. order.hclust(m,method="single") is equivalent to order.single when clusters is NULL. The default method of hclust is "complete", see hclust for other possibilities.

Value

A permutation of the objects represented by merit is returned.

ozone

Author(s)

Catherine B. Hurley

References

Hurley, Catherine B. "Clustering Visualisations of Multidimensional Data", Journal of Computational and Graphical Statistics, vol. 13, (4), pp 788-806, 2004.

Gruvaeus, G. and Wainer, H. (1972), "Two Additions to Hierarchical Cluster Analysis", British Journal of Mathematical and Statistical Psychology, 25, 200-206.

See Also

cpairs, cparcoord, plotcolors, reorder.hclust, order.clusters, hclust.

Examples

```
data(state)
state.cor <- cor(state.x77)
order.single(state.cor)
order.endlink(state.cor)
order.hclust(state.cor,method="average")</pre>
```

Use for plotting...

cpairs(state.x77, panel.colors=dmat.color(state.cor), order.single(state.cor),pch=".",gap=.4)
cparcoord(state.x77, order.endlink(state.cor),panel.colors=dmat.color(state.cor))

Order the states instead of the variables...

```
state.d <- dist(state.x77)
state.o <- order.single(-state.d)</pre>
```

```
op <- par(mar=c(1,6,1,1))
cmat <- dmat.color(as.matrix(state.d), rev(cm.colors(5)))
plotcolors(cmat[state.o,state.o], rlabels=state.name[state.o])
par(op)</pre>
```

ozone

Ozone data from Breiman and Friedman, 1985

Description

This is the Ozone data discussed in Breiman and Friedman (JASA, 1985, p. 580). These data are for 330 days in 1976. All measurements are in the area of Upland, CA, east of Los Angeles.

Usage

data(ozone)

Format

This data frame contains the following columns:

Ozone: Ozone conc., ppm, at Sandbug AFB. Temp: Temperature F. (max?). InvHt: Inversion base height, feet Pres: Daggett pressure gradient (mm Hg) Vis: Visibility (miles) Hgt: Vandenburg 500 millibar height (m) Hum: Humidity, percent InvTmp: Inversion base temperature, degrees F. Wind: Wind speed, mph

Source

Breiman, L and Friedman, J. (1985), "Estimating Optimal Transformations for Multiple Regression and Correlation", *Journal of the American Statistical Association*, 80, 580-598.

partition.crit	<i>Combines the results of appplying an index to each group of observa-</i> <i>tions</i>

Description

Applies the function gfun to each group of x and y values and combines the results using the function cfun

Usage

```
partition.crit(x, y, groups, gfun = gave, cfun = sum, ...)
```

Arguments

х	is a numeric vector.
У	is a numeric vector.
groups	is a vector of group memberships.
gfun	is applied to the x and y data in each group.
cfun	combines the values returned by gfun.
	arguements are passed to gfun.

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partition.crit

Details

The function gfun is applied to each group of x and y values. The function cfun is applied to the vector or matrix of gfun results.

Value

The result of applying cfun.

Author(s)

Catherine B. Hurley

References

See Gordon, A. D. (1999). Classification. Second Edition. London: Chapman and Hall / CRC

See Also

gave, colpairs, order.single

Examples

```
x <- runif(20)</pre>
y <- runif(20)
g <- rep(c("a","b"),10)
partition.crit(x,y,g)
data(bank)
# m is a homogeneity measure of each pairwise variable plot
m <- -colpairs(scale(bank[,-1]), partition.crit,gfun=gave,groups=bank[,1])</pre>
# Color panels by level of m and reorder variables so that
# pairs with high m are near the diagonal. Panels shown
# in pink have the highest amount of group homogeneity, as measured by
# gave.
cpairs(bank[,-1],order=order.single(m), panel.colors=dmat.color(m),
gap=.3,col=c("purple","black")[bank[,"Status"]+1],
pch=c(5,3)[bank[,"Status"]+1])
# Try a different measure
m <- -colpairs(scale(bank[,-1]), partition.crit,gfun=diameter,groups=bank[,1])</pre>
cpairs(bank[,-1],order=order.single(m), panel.colors=dmat.color(m),
gap=.3,col=c("purple","black")[bank[,"Status"]+1],
pch=c(5,3)[bank[,"Status"]+1])
```

Result is the same, in this case.

pclen

Description

Computes measures of profile smoothness of 2-d data, where x and y give the object coordinates.

Usage

pclen(x, y)
pcglen(x, y)

Arguments

х	is a numeric vector.
У	is a numeric vector.

Details

pclen computes the total line length in a parallel coordinate plot of x and y.

pcglen computes the average (per object) line length in a parallel coordinate plot where all pairs of objects are connected.

Usually, the data is standardized prior to using these functions.

Value

The panel measure is returned.

Author(s)

Catherine B. Hurley

References

Hurley, Catherine B. "Clustering Visualisations of Multidimensional Data", Journal of Computational and Graphical Statistics, vol. 13, (4), pp 788-806, 2004.

See Also

cparcoord, colpairs, order.endlink.

plotcolors

Examples

```
x <- runif(20)
y <- runif(20)
pclen(x,y)</pre>
```

```
data(state)
mins <- apply(state.x77,2,min)
ranges <- apply(state.x77,2,max) - mins
state.m <- -colpairs(scale(state.x77,mins,ranges), pclen)
state.col <- dmat.color(state.m)
cparcoord(state.x77, panel.color= state.col)
# Get rid of the panels with long line segments (yellow) by reordering:
cparcoord(state.x77, order.endlink(state.m), state.col)</pre>
```

plotcolors

Plots a matrix of colors

Description

plotcolors plots a matrix of colors as an image or as points.

imageinfo is a utility that given a matrix of colors, returns a structure useful for the image function.

Usage

```
plotcolors(cmat, na.color = "white", dlabels = NULL, rlabels = FALSE, clabels = FALSE,
ptype = "image", border.color = "grey70", pch = 15, cex = 3, label.cex = 0.6, ...)
```

imageinfo(cmat)

Arguments

cmat	a matrix of numbers, nas are allowed.
na.color	used for NAs in cmat.
dlabels	vector of labels for the diagonals.
rlabels	vector of labels for the rows.
clabels	vector of labels for the columns.
ptype	should be "image" or "points"
border.color	color of border drawn around the plot.
pch	point type used when ptype="points".
cex	point cex used when ptype="points".
label.cex	cex parameter used for labels.
	graphical parameters

Value

imageinfo returns a list with components:

х	a vector of x coordinates.
У	a vector of y coordinates.
z	a matrix containing values to be plotted.
col	the colors to be used.

Author(s)

Catherine B. Hurley

See Also

plot, image

Examples

plotcolors(matrix(1:20,nrow=4,ncol=5))

plotcolors(matrix(1:20,nrow=4,ncol=5),ptype="points",cex=6)

plotcolors(matrix(1:20,nrow=4,ncol=5),rlabels = c("a","b","c","d"))

```
data(longley)
longley.cor <- cor(longley)
# A matrix with equal (or nearly equal) number of entries of each color.
longley.color <- dmat.color(longley.cor)</pre>
```

plotcolors(longley.color, dlabels=rownames(longley.color))

```
# Could also reorder variables prior to plotting:
longley.o <- order.single(longley.cor)
longley.color <- longley.color[longley.o,longley.o]</pre>
```

```
op <- par(mar=c(1,6,6,1))
plotcolors(longley.color,rlabels=rownames(longley.color),clabels=rownames(longley.color))
par(op)</pre>
```

reorder.hclust

Reorders object order of hclust, keeping objects within a cluster contiguous to each other.

Description

Reorders objects so that nearby object pairs are adjacent.

reorder.hclust

Usage

S3 method for class 'hclust'
reorder(x,dis,...)

Arguments

x	is the result of hclust.
dis	is a distance matrix or dist.
	additional arguments.

Details

In hierarchical cluster displays, a decision is needed at each merge to specify which subtree should go on the left and which on the right. This algorithm uses the order suggested by Gruvaeus and Wainer (1972). At a merge of clusters A and B, the new cluster is one of (A,B), (A',B), (A,B'), (A',B'), where A' denotes A in reverse order. The new cluster is chosen to minimize the distance between the object in A placed adjacent to an object from B.

Value

A permutation of the objects represented by dis is returned.

Author(s)

Catherine B. Hurley

References

Hurley, Catherine B. "Clustering Visualisations of Multidimensional Data", Journal of Computational and Graphical Statistics, vol. 13, (4), pp 788-806, 2004.

Gruvaeus, G. and Wainer, H. (1972), "Two Additions to Hierarchical Cluster Analysis", British Journal of Mathematical and Statistical Psychology, 25, 200-206.

See Also

hclust, order.hclust.

Examples

```
data(eurodist)
dis <- as.dist(eurodist)
hc <- hclust(dis, "ave")
```

```
layout(matrix(1:2,nrow=2,ncol=1))
op <- par(mar=c(1,1,1,1))
plot(hc)
hc1 <- reorder.hclust(hc, dis)
plot(hc1)</pre>
```

vec2distm

```
par(op)
layout(matrix(1,1))
# Both dedrograms correspond to the same tree structure,
# but the second one shows that
# Paris is closer to Cherbourg than Munich, and
# Rome is closer to Gibralter than to Barcelona.
# We can also compare both orderings with an
# image plot of the colors.
# The second ordering seems to place nearby cities
# closer to each other.
layout(matrix(1:2,nrow=2,ncol=1))
op <- par(mar=c(1,6,1,1))
cmat <- dmat.color(eurodist, rev(cm.colors(5)))</pre>
plotcolors(cmat[hc$order,hc$order], rlabels=labels(eurodist)[hc$order])
plotcolors(cmat[hc1$order,hc1$order], rlabels=labels(eurodist)[hc1$order])
layout(matrix(1,1))
par(op)
```

vec2distm

Various utility functions

Description

vec2distm converts a vector to a distance matrix.

vec2dist converts a vector to a dist structure.

lower2upper.tri.inds is the same as lower.to.upper.tri.inds from package cluster. It computes an index vector for extracting or reordering a lower triangular matrix that is stored as a contiguous vectors.

diag.off returns a vector of off-diagonal elements of a matrix. off specifies the distance above the main (0) diagonal.

clus2memship converts a list whose ith element contains the indices of objects in the ith cluster into a vector whose ith element gives the cluster number of the ith object.

memship2clus converts a vector whose ith element gives the cluster number of the ith object into a list whose ith element contains the indices of objects in the ith cluster.

Usage

```
vec2distm(vec)
vec2dist(vec)
```

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wine

```
lower2upper.tri.inds(n)
diag.off(m,off=1)
clus2memship(clusters)
memship2clus(memship)
```

Arguments

vec	is a vector.
n	is an integer > 1.
m	is a matrix.
clusters	is a list whose ith element contains the indices of the objects belonging to the ith cluster.
off	is an integer specifying the distance above the main (0) diagonal.
memship	is a vector whose ith element gives the cluster number of the ith object.

Author(s)

Catherine B. Hurley

See Also

dist, diag.

Examples

```
vec <- 1:15
vec2distm(vec)
vec2dist(vec)
diag.off(vec2distm(vec))
lower2upper.tri.inds(5)
clus2memship(list(c(1,3,5),c(2,6),4))
memship2clus(c(1,3,4,2,1,4,2,3,2,3))
```

wine

Wine recognition data

Description

Data from the machine learning repository. A chemical analysis of 178 Italian wines from three different cultivars yielded 13 measurements. This dataset is often used to test and compare the performance of various classification algorithms.

Usage

data(wine)

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Format

This data frame contains the following columns:

Class: There are 3 classes Alcohol: Alcohol Malic: Malic acid Ash: Malic acid Ash: Ash Alcalinity: Alcalinity of ash Magnesium: Magnesium Phenols: Total phenols Flavanoids: Flavanoids Nonflavanoid: Nonflavanoid phenols Proanthocyanins: Proanthocyanins Intensity: Color intensity Hue: Hue OD280: OD280/OD315 of diluted wines Proline: Proline

Source

Forina, M. et al, PARVUS - An Extendible Package for Data Exploration, Classification and Correlation. Institute of Pharmaceutical and Food Analysis and Technologies, Via Brigata Salerno, 16147 Genoa, Italy.

References

Blake, C.L. and Merz, C.J. (1998), UCI Repository of machine learning databases, \https://archive.ics.uci.edu/datasets. Irvine, CA: University of California, Department of Information and Computer Science.

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