Package ‘made4’

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Title     Multivariate analysis of microarray data using ADE4
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Suggests  affy

Description Multivariate data analysis and graphical display of microarray data. Functions include between group analysis and co-inertia analysis. It contains functions that require ADE4.


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URL      http://www.hsph.harvard.edu/aedin-culhane/
bioCViews Clustering, Classification, MultipleComparison

NeedsCompilation no

R topics documented:

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array2ade4

Converts microarray input data into a data frame suitable for analysis in ADE4.

Description

Converts input data into a data frame suitable for analysis in ADE4. This function is called by bga and other made4 function.

Usage

array2ade4(dataset, pos = FALSE, trans = FALSE)

Arguments

dataset A matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
pos Logical indicating whether to add an integer to dataset, to generate positive data.frame. Required for dudi.coa or dudi.nsc.
trans Logical indicating whether dataset should be transposed. Default is FALSE.

Details

bga and other functions in made4 call this function and it is generally not necessary to call array2ade4 this directly.
array2ade4 calls getdata, and will accept a matrix, data.frame, ExpressionSet or marrayRaw-class format. It will also transpose data or add a integer to generate a positive data matrix.

If the input data contains missing values (NA), these must first be removed or imputed (see the R libraries impute() or pamr()).

Value

Returns a data.frame suitable for analysis by ade4 or made4 functions.

Author(s)

Aedin Culhane
**bet.coinertia**

See Also

gedata.as in the Bioconductor package convert.

Examples

```r
library(affy)
data(geneData)
class(geneData)
dim(geneData)
dim(array2ade4(geneData))
class(array2ade4(geneData))

bet.coinertia
```

**Description**

Between class coinertia analysis. **cia** of 2 datasets where covariance between groups or classes of cases, rather than individual cases are maximised.

Usage

```r
bet.coinertia(df1, df2, fac1, fac2, cia.nf = 2, type = "nsc", ...)
```

Arguments

- **df1**: First dataset. A **matrix**, **data.frame**, **ExpressionSet** or **marrayRaw-class**. If the input is gene expression data in a **matrix** or **data.frame**. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
- **df2**: Second dataset. A **matrix**, **data.frame**, **ExpressionSet** or **marrayRaw-class**. If the input is gene expression data in a **matrix** or **data.frame**. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
- **fac1**: A factor or vector which describes the classes in df1.
- **fac2**: A factor or vector which describes the classes in df2.
- **cia.nf**: Integer indicating the number of coinertia analysis axes to be saved. Default value is 2.
- **type**: A character string. accepted options are type="nsc" or type="pca".
- **...**: further arguments passed to or from other methods.

Value

A list of class **bet.cia** of length 5

- **coin**: An object of class 'coinertia', sub-class **dudi**. See cointertia
- **coa1, pca1**: An object of class 'nsc' or 'pca', with sub-class 'dudi'. See dudi, dudi.pca or dudi.nsc
- **coa2, pca2**: An object of class 'nsc' or 'pca', with sub-class 'dudi'. See dudi, dudi.pca or dudi.nsc
- **bet1**: An object of class 'bga', with sub-class 'dudi'. See dudi, bga or bca
- **bet2**: An object of class 'bga', with sub-class 'dudi'. See dudi, bga or bca.
between.graph

Note
This is very computational intensive. The authors of ade4 are currently re-writing the code for coinertia analysis, so that it should substantially improve the computational requirements (May 2004).

Author(s)
Aedin Culhane

References
Culhane AC, et al., 2003 Cross platform comparison and visualisation of gene expression data using co-inertia analysis. BMC Bioinformatics. 4:59

See Also
See Also as coinertia, cia.

Examples
### NEED TO DO
if (require(ade4, quiet = TRUE)) {}

between.graph Object of the class bga resulting from a bga analysis.

Arguments

- `x` Numeric. The column number of principal component ($ls$ and $li$) to be used. Default is 1. This is the first component of the analysis.
- `ax` Vector of colours. By default colours are obtained using `getcol`.
- `hor` Logical, indicating whether the graph should be plotted horizontally or vertically. The default is a horizontal plot.
- `scaled` Logical, indicating whether the coordinates in the graph should be scaled to fit optimally in plot. Default is TRUE.
- `centnames` A vector of variables labels. Default is NULL, if NULL the row names of the centroid $li$ coordinates will be used.
- `varnames` A vector of variables labels. Default is NULL, if NULL the row names of the variable $ls$ coordinates will be used.
- `...` further arguments passed to or from other methods.
Details

This will produce a figure similar to Figure 1 in the paper by Culhane et al., 2002.

between.graph requires both samples and centroid co-ordinates (\$ls, \$li) which are passed to it via an object of class bga. If cases are to be coloured by class, it also requires a \$fac factor which is also passed to it via an object of class bga.

To plot a 1D graph from other multivariate analysis such as PCA (\texttt{dudi.pca}), COA (\texttt{dudi.coa}), or \texttt{coinertia} analysis. Please use \texttt{graph1D}.

Author(s)

Aedin Culhane

References


See Also

\texttt{graph1D}

Examples

data(khan)
if (require(ade4, quiet = TRUE)) {
    khan.bga<-bga(khan$train, khan$train.classes)
}
between.graph(khan.bga)
between.graph(khan.bga, ax=2, lwd=3, cex=0.5, col=c("green","blue", "red", "yellow"))
between.graph(khan.bga, ax=2, hor=FALSE, col=c("green","blue", "red", "yellow"))

---

**bga**

*Between group analysis*

Description

Discrimination of samples using between group analysis as described by Culhane et al., 2002.

Usage

bga(dataset, classvec, type = "coa", ...)

## S3 method for class 'bga'
plot(x, axis1=1, axis2=2, arraycol=NULL, genecol="gray25", nlab=10, genelabels= NULL, ...)

---
**Arguments**

- **dataset**: Training dataset. A `matrix`, `data.frame`, `ExpressionSet` or `marrayRaw-class`. If the input is gene expression data in a `matrix` or `data.frame`. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.

- **classvec**: A factor or vector which describes the classes in the training dataset.

- **type**: Character, "coa", "pca" or "nsc" indicating which data transformation is required. The default value is type="coa".

- **x**: An object of class `bga`. The output from `bga` or `bga.suppl`. It contains the projection coordinates from `bga`, the `$ls`, `$co` or `$li` coordinates to be plotted.

- **arraycol, genecol**: Character, colour of points on plot. If arraycol is NULL, arraycol will obtain a set of contrasting colours using `getcol`, for each classes of cases (microarray samples) on the array (case) plot. genecol is the colour of the points for each variable (genes) on gene plot.

- **nlab**: Numeric. An integer indicating the number of variables (genes) at the end of axes to be labelled, on the gene plot.

- **axis1, axis2**: Integer, the column number for the x-axis. The default is 1.

- **genelabels**: A vector of variables labels, if genelabels=NULL the row.names of input matrix `dataset` will be used.

- **...**: further arguments passed to or from other methods.

**Details**

`bga` performs a between group analysis on the input dataset. This function calls `bca`. The input format of the dataset is verified using `array2ade4`.

Between group analysis is a supervised method for sample discrimination and class prediction. BGA is carried out by ordinating groups (sets of grouped microarray samples), that is, groups of samples are projected into a reduced dimensional space. This is most easily done using PCA or COA, of the group means. The choice of PCA, COA is defined by the parameter type.

The user must define microarray sample groupings in advance. These groupings are defined using the input `classvec`, which is a factor or vector.

**Cross-validation and testing of bga results:**

`bga` results should be validated using one leave out jack-knife cross-validation using `bga.jackknife` and by projecting a blind test datasets onto the `bga` axes using `suppl`. `bga` and `suppl` are combined in `bga.suppl` which requires input of both a training and test dataset. It is important to ensure that the selection of cases for a training and test set are not biased, and generally many cross-validations should be performed. The function `randomiser` can be used to randomise the selection of training and test samples.

**Plotting and visualising bga results:**

1D plots, show one axis only: 1D graphs can be plotted using `between.graph` and `graph1D`. `between.graph` is used for plotting the cases, and required both the co-ordinates of the cases (`$ls`) and their centroids (`$li`). It accepts an object `bga`. `graph1D` can be used to plot either cases (microarrays) or variables (genes) and only requires a vector of coordinates.

2D plots: Use `plot.bga` to plot results from `bga`. `plot.bga` calls the functions `plotarrays` to draw an xy plot of cases (`$ls`), `plotgenes`, is used to draw an x-y plot of the variables (genes). `plotgenes`, is used to draw an x-y plot of the variables (genes).
3D plots: 3D graphs can be generated using do3D and html3D. html3D produces a web page in which a 3D plot can be interactively rotated, zoomed, and in which classes or groups of cases can be easily highlighted.

**Analysis of the distribution of variance among axes:**

It is important to know which cases (microarray samples) are discriminated by the axes. The number of axes or principal components from a bga will equal the number of classes - 1, that is length(levels(classvec))-1.

The distribution of variance among axes is described in the eigenvalues (\$eig) of the bga analysis. These can be visualised using a scree plot, using scatterutil.eigen as it done in plot.bga.

It is also useful to visualise the principal components from a using a bga or principal components analysis dudi.pca, or correspondence analysis dudi.coa using a heatmap. In MADE4 the function heatplot will plot a heatmap with nicer default colours.

**Extracting list of top variables (genes):**

Use topgenes to get list of variables or cases at the ends of axes. It will return a list of the top n variables (by default n=5) at the positive, negative or both ends of an axes. sumstats can be used to return the angle (slope) and distance from the origin of a list of coordinates.

For more details see Culhane et al., 2002 and http://bioinf.ucd.ie/research/BGA.

**Value**

A list with a class bga containing:

- ord: Results of initial ordination. A list of class "dudi" (see dudi)
- bet: Results of between group analysis. A list of class "dudi" (see dudi), "between" (see bca)
- fac: The input classvec, the factor or vector which described the classes in the input dataset

**Author(s)**

Aedin Culhane

**References**


**See Also**

See Also bga, suppl, suppl.bga, bca, bga.jackknife

**Examples**

data(khan)

if (require(ade4, quiet = TRUE)) {
  khan.bga<-bga(khan$train, classvec=khan$train.classes)
}

khan.bga
plot(khan.bga, genelabels=khan$annotation$Symbol)
# Provide a view of the principal components (axes) of the bga

```r
heatplot(khan.bga$bet$ls, dend="none")
```

---

**bga.jackknife**  
*Jackknife between group analysis*

**Description**

Performs one-leave-out jackknife analysis of a between group analysis as described by Culhane et al., 20002

**Usage**

```r
bga.jackknife(data, classvec, ...)
```

**Arguments**

- `data`: Input dataset. A matrix, data.frame If the input is gene expression data in a matrix or data.frame. The columns contain the cases (array samples) which will be jackknifed.
- `classvec`: A factor or vector which describes the classes in the training dataset
- `...`: further arguments passed to or from other methods

**Details**

Performs a one-leave-out cross validation of between group analysis bga. Input is a training dataset. This can take 5-10 minutes to compute on standard data gene expression matrix.

In jackknife one leave out analysis, one case (column) is removed. The remaining dataset is subjected to **bga**. Then the class of the case that was removed is predicted using **suppl**. This analysis is repeated until all samples have been removed and predicted.

**Value**

A list containing

- results The projected co-ordinates of each sample
- summary A summary of number and percentage of correctly assigned samples

**Author(s)**

Aedin Culhane

**References**


**See Also**

- **bga**, **bga.suppl**, **suppl.bga**, **bca**, **plot.bga**
**Examples**
```r
data(khan)
# NOTE using a very reduced dataset (first 5 genes) to speed up results
# hence expect poor prediction accuracy
dim(khan$train)
print("using only small subset of data")
if (require(ade4, quiet = TRUE)) {
  bga.jackknife(khan$train[1:5,], khan$train.classes) }
```

---

**bga.suppl**  
*Between group analysis with supplementary data projection*

**Description**

*bga.suppl* performs a *bga* between group analysis with projection of supplementary points using *suppl*

**Usage**

```r
bga.suppl(dataset, supdata, classvec, supvec = NULL, suponly = FALSE, type="coa", ...)
```

**Arguments**

- **dataset**  
  Training dataset. A *matrix*, *data.frame*, *ExpressionSet* or *marrayRaw-class*. If the input is gene expression data in a *matrix* or *data.frame*. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.

- **supdata**  
  Test or blind dataset. A *matrix*, *data.frame*, *ExpressionSet* or *marrayRaw-class*. If the input is gene expression data in a *matrix* or *data.frame*. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively. The test dataset *supdata* and the training dataset *dataset* must contain the same number of variables (genes).

- **classvec**  
  A *factor* or *vector* which describes the classes in the training data *dataset*.

- **supvec**  
  A *factor* or *vector* which describes the classes in the test dataset *supdata*.

- **suponly**  
  Logical indicating whether the returned output should contain the test class assignment results only. The default value is FALSE, that is the training coordinates, test coordinates and class assignments will all be returned.

- **type**  
  Character, "coa", "pca" or "nsc" indicating which data transformation is required. The default value is type="coa".

- **...**  
  Further arguments passed to or from other methods.

**Details**

*bga.suppl* calls *bga* to perform between group analysis (*bga*) on the training dataset, then it calls *suppl* to project the test dataset onto the *bga* axes. It returns the coordinates and class assignment of the cases (microarray samples) in the test dataset as described by Culhane et al., 2002.

The test dataset must contain the same number of variables (genes) as the training dataset.

The input format of both the training dataset and test dataset are verified using *array2ade4*. Use *plot.bga* to plot results from *bga*. 
Coinertia analysis: Explore the covariance between two datasets

Description

Performs CIA on two datasets as described by Culhane et al., 2003. Used for meta-analysis of two or more datasets.
Usage

cia(df1, df2, cia.nf=2, cia.scan=FALSE, nsc=TRUE,...)
## S3 method for class 'cia'
plot(x, nlab = 10, axis1 = 1, axis2 = 2, genecol = "gray25",
genelabels1 = rownames(ciares$co), genelabels2 = rownames(ciares$li), ...)

Arguments

df1 The first dataset. A matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
df2 The second dataset. A matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
cia.nf Integer indicating the number of coinertia analysis axes to be saved. Default value is 2.
cia.scan Logical indicating whether the coinertia analysis eigenvalue (scree) plot should be shown so that the number of axes, cia.nf can be selected interactively. Default value is FALSE.
nsc A logical indicating whether coinertia analysis should be performed using two non-symmetric correspondence analyses dudi.nsc. The default=TRUE is highly recommended. If FALSE, COA dudi.coa will be performed on df1, and row weighted COA dudi.rwcoa will be performed on df2 using the row weights from df1.
x An object of class cia, containing the CIA projected coordinates to be plotted.
nlab Numeric. An integer indicating the number of variables (genes) to be labelled on plots.
axis1 Integer, the column number for the x-axis. The default is 1.
axis2 Integer, the column number for the y-axis. The default is 2.
genecol Character, the colour of genes (variables). The default is "gray25". genelabels1, genelabels2 A vector of variables labels, by default the row.names of each input matrix df1, and df2 are used.
...
... further arguments passed to or from other methods.

Details

CIA has been successfully applied to the cross-platform comparison (meta-analysis) of microarray gene expression datasets (Culhane et al., 2003). Please refer to this paper and the vignette for help in interpretation of the output from CIA.

Co-inertia analysis (CIA) is a multivariate method that identifies trends or co-relationships in multiple datasets which contain the same samples. That is the rows or columns of the matrix have to be weighted similarly and thus must be "matchable". In cia, it is assumed that the analysis is being performed on the microarray cases, and thus the columns will be matched between the 2 datasets. Thus please ensure that the order of cases (the columns) in df1 and df2 are equivalent before performing CIA.
CIA simultaneously finds ordinations (dimension reduction diagrams) from the datasets that are most similar. It does this by finding successive axes from the two datasets with maximum covariance. CIA can be applied to datasets where the number of variables (genes) far exceeds the number of samples (arrays) such is the case with microarray analyses.

cia calls coinertia in the ADE4 package. For more information on coinertia analysis please refer to coinertia and several recent reviews (see below).

In the paper by Culhane et al., 2003, the datasets df1 and df2 are transformed using COA and Row weighted COA respectively, before coinertia analysis. It is now recommended to perform non symmetric correspondence analysis (NSC) rather than correspondence analysis (COA) on both datasets.

**The RV coefficient**

In the results, in the object cia returned by the analysis, \$coinertia\$RV gives the RV coefficient. This is a measure of global similarity between the datasets, and is a number between 0 and 1. The closer it is to 1 the greater the global similarity between the two datasets.

**Plotting and visualising cia results**

plot.cia draws 3 plots.

The first plot uses S.match.col to plots the projection (normalised scores \$mY and \$mX) of the samples from each dataset onto the one space. Cases (microarray samples) from one dataset are represented by circles, and cases from the second dataset are represented by arrow tips. Each circle and arrow is joined by a line, where the length of the line is proportional to the divergence between the gene expression profiles of that sample in the two datasets. A short line shows good agreement between the two datasets.

The second two plots call plot.genes are show the projection of the variables (genes, \$li and \$co) from each dataset in the new space. It is important to note both the direction of project of Variables (genes) and cases (microarray samples). Variables and cases that are projected in the same direction from the origin have a positive correlation (ie those genes are upregulated in those microarray samples)

*Please refer to the help on bga for further discussion on graphing and visualisation functions in MADE4.*

**Value**

An object of the class cia which contains a list of length 4.

call list of input arguments, df1 and df2

coinertia A object of class "coinertia", sub-class dudi. See coinertia

coa1 Returns an object of class "coa" or "nsc", with sub-class dudi. See dudi.coa or dudi.nsc

coa2 Returns an object of class "coa" or "nsc", with sub-class dudi. See dudi.coa or dudi.nsc

**Author(s)**

Aedin Culhane

**References**

Culhane AC, et al., 2003 Cross platform comparison and visualisation of gene expression data using co-inertia analysis. BMC Bioinformatics. 4:59
See Also

See also coinertia, plot.cia

Examples

data(NCI60)
print("This will take a few minutes, please wait...")

if (require(ade4, quiet = TRUE)) {
  # Example data are "G1_Ross_1375.txt" and "G5_Affy_1517.txt"
  coin <- cia(NCI60$Ross, NCI60$Affy)
}

attach(coin)
summary(coin)
summary(coin$coinertia)

# $coinertia$RV will give the RV-coefficient, the greater (scale 0-1) the better

cat(paste("The RV coefficient is a measure of global similarity between the datasets.\n", "The two datasets analysed are very similar. ",
  "The RV coefficient of this coinertia analysis is: ", coin$coinertia$RV,\n  "sep= ""))

plot(coin)
plot(coin, classvec=NCI60$classes[,2], clab=0, cpoint=3)

---

commonMap

Highlight common points between two 1D plots

Description

CommonMap draws two 1D plots, and links the common points between the two.

Usage

commonMap(x, y, hor=TRUE, cex=1.5, scaled=TRUE, ...)

Arguments

x The coordinates of the first axis
y The coordinates of the second axis
hor Logical, whether a horizontal line should be drawn on plot. Default is TRUE.
cex Numeric. The amount by which plotting text and symbols should be scaled
  relative to the default
scaled Logical, whether the data in x and y are scaled. Scaling is useful for visualising
  small or large data values. Set to FALSE if actually or true values should be
  visualised. The default is TRUE.
... further arguments passed to or from other method

Details

Useful for mapping the genes in common from coinertia analysis This graphs a 1D graph. x and y
are the coordinates from two different analyses but the rows of each vectors correspond (ie common
genes)
Note

This is useful for examining common points in axes from coinertia analysis, or comparing results from two different analysis.

Author(s)

Ailis Fagan and Aedin Culhane

See Also

See also between.graph, graph1D

Examples

```r
a <- rnorm(20)
b <- rnorm(20)
par(mfrow=c(2,2))
commonMap(a, b)
commonMap(a, b, hor=FALSE, col="red", pch=19)
commonMap(a, b, col="blue", cex=2, pch=19)

# If the vectors contain different variables, the rows should define the variables that correspond
a[15:20] <- NA
b[10:15] <- NA
cbind(a, b)
commonMap(a, b, col="dark green", pch=18)
```

`comparelists`  
Return the intersect, difference and union between 2 vectors

Description

This is a very simple function which compares two vectors, x and y. It returns the intersection and unique lists. It is useful for comparing two genelists.

Usage

```r
comparelists(dx, dy, ...)
## S3 method for class 'comparelists'
print(x, ...)
```

Arguments

- `dx, dy`: A vector.
- `x`: An object from comparelists.
- `...`: Further arguments passed to or from other methods.

Details

reports on the intersect, difference and union between two lists.
do3d

Value

An object of class comparelists:

- `intersect`: Vector containing the intersect between x and y
- `Set.Diff`: Vector containing the elements unique to X obtained using `setdiff`
- `XinY`: Numeric, indicating the number of elements of x in y
- `YinX`: Numeric, indicating the number of elements of y in x
- `Length.X`: Numeric, the number of elements in x
- `Length.Y`: Numeric, the number of elements in y
- `...`: Further arguments passed to or from other methods

Author(s)

Aedin Culhane

See Also

See also `intersect`, `setdiff`

Examples

```r
a <- sample(LETTERS, 20)
b <- sample(LETTERS, 10)
z <- comparelists(a, b)
z$Set.Diff
z$intersect
```

---

**do3d**

Generate 3D graph(s) using scatterplot3d

Description

do3d is a wrapper for scatterplot3d. do3d will draw a single 3D xyz plot and will plot each group of points in a different colour, given a factor.

rotate3d calls do3d to draw multiple 3D plots in which each plot is marginally rotated on the x-y axis.

Usage

```r
do3d(dataset, x = 1, y = 2, z = 3, angle = 40, classvec = NULL, classcol = NULL, col = NULL, cex.lab = 0.3, pch = 19, cex.symbols = 1, ...)
rotate3d(dataset, x = 1, y = 2, z = 3, beg = 180, end = 360, step = 12, savefiles = FALSE, classvec = NULL, classcol = NULL, col = NULL, ...)```

Arguments

dataset  XYZ coordinates to be plotted. A matrix or data.frame with 3 or more columns. Usually results from multivariate analysis, such as the \$co or \$li coordinates from a PCA dudi.pca, or COA dudi.coa or the \$Sis, \$co coordinates from bga.

x  Numeric, the column number for the x-axis, the default is 1 (that is dataset[,1])
y  Numeric, the column number for the y-axis, the default is 2 (that is dataset[,2])
z  Numeric, the column number for the z-axis, the default is 3 (that is dataset[,3])
angle  Numeric, the angle between x and y axis. Note the result depends on scaling. See scatterplot3d

classvec  A factor or vector which describes the classes in dataset

classcol  A factor or vector which list the colours for each of the classes in the dataset. By default NULL. When NULL, getcol is used to obtain an optimum set of colours of the classes in classvec.

cex.lab  Numeric. The magnification to be used for the axis annotation relative to the current default text and symbol size. Default is 0.3

pch  Integer specifying a symbol or single character to be used when plotting points. The default is pch= 19

cex.symbols  Numeric. The magnification to be used for the symbols relative to the current default text size. Default is 1

col  A character indicating a colour. To be used if all points are to be one colour. If classvec, classcol and col are all NULL. all points will be drawn in red by default.

beg  Numeric. The starting angle between the x and y axis for rotate3d. Rotate3d will draw plots in which they are rotated from angle beg to angle end

der  Numeric. The final angle between the x and y axis for rotate3d. Rotate3d will draw plots in which they are rotated from angle beg to angle end

step  Numeric. Increment of the sequence between the starting angle beg and the final angle end.

savefiles  Logical, indicating whether the plot should be saved as a pdf file. The default is FALSE

...  further arguments passed to or from other methods

Details

This calls scatterplot3d to plot a 3d representation of results.

It is also worth exploring the package rgl which enables dynamic 3d plot (that can be rotated)

library(rgl) plot3d(khan.coa$co[,1], khan.coa$co[,2],khan.coa$co[,3], size=4, col=khan$train.classes) rgl.snapshot(file="test.png", top=TRUE) rgl.close()

Value

Produces plots of the xyz coordinates.

Author(s)

Aedin Culhane
getcol

Specialised colour palette with set of 21 maximally contrasting colours

description

Special colour palette developed to maximise the contrast between colours. Colours were selected for visualising groups of points on xy or xyz plots on a white background. Because of this, there are few pastel colours are in this palette. getcol contains 2 palettes of 12 and 21 colours.

Usage

getcol(nc = c(1:3), palette = NULL, test = FALSE)

Arguments

nc Numeric. Integer or vector in range 1 to 21. This selects colours from palette

palette A character to select either palette "colours1" or "colours2". colours1 contains 12 colours, colours2 contains 21 colours

test A logical, if TRUE a plot will be drawn to display the palettes colours1, colours2 and any selected colours.

Details

Colours1 contains the 12 colours,"red","blue","green","cyan","magenta","yellow","grey","black","brown", "orange","violet","purple"). These were choosen, as these are compatible with rasmol and chime, that are used in html3D. Colours2 contains 21 colours. These were selected so as to maximise the contrast between groups.

For other colour palettes in R, see colors, palette, rainbow, heat.colors, terrain.colors, topo.colors or cm.colors.

Also see the library RColorBrewer

Value

A vector containing a list of colours.
Author(s)
Aedin Culhane

See Also
See also colors, palette, rainbow, heat.colors, terrain.colors, topo.colors or cm.colors, RColorBrewer

Examples

getcol(3)
getcol(c(1:7))
getcol(10, test=TRUE)
getcol(c(1:5, 7, 15, 16), palette="colours2", test=TRUE)

graph1D

Plot 1D graph of axis from multivariate analysis

Description

Draw 1D plot of an axis from multivariate analysis. Useful for visualising an individual axis from analyses such as PCA dudi.pca or COA dudi.coa. It accepts a factor so that groups of points can be coloured. It can also be used for graphing genes, and will only label n genes at the ends of the axis.

Usage

graph1D(dfx, classvec=NULL, ax = 1, hor=FALSE, s.nam=row.names(dfx), n=NULL, scaled=TRUE, col="red", width=NULL, ...)

Arguments

dfx  vector, matrix, or data.frame, which contains a column with axis coordinates
ax   Numeric, indicating column of matrix, or data.frame to be plotted. The default is 1.
classvec Factor, indicating sub-groupings or classes in dfx or dfx[,ax]
hor   Logical, indicating whether the graph should be drawn horizontal or vertically. The default is vertically.
s.nam Vector. labels of dfx, The default is row.names(dfx)
n   Numeric. Whether all rows should be plotted, n=10 would label only the 10 variables at the end of the axis. By default all variables (row of dfx) are labelled
scaled A logical indicating whether the plot should be scaled to fit. The default is TRUE
col   A character or vector indicating the colour(s) for points or groups of points. If points are to be coloured according to a factor, length(col) should equal length(levels(classvec))
width A vector of length 2, which is the width (of a vertical plot) or height (of a horizontal plot). This can be increased if variable labels are unreadable. The default is c(-2,1)
...  further arguments passed to or from other methods
heatplot

Draws heatmap with dendrograms.

Description

heatplot calls `heatmap.2` using a red-green colour scheme by default. It also draws dendrograms of the cases and variables using correlation similarity metric and average linkage clustering as described by Eisen. heatplot is useful for a quick overview or exploratory analysis of data.

Usage

heatplot(dataset, dend = c("both", "row", "column", "none"), cols.default = TRUE, lowcol = "green", highcol = "red", ...)

Arguments

dataset	a matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
dend	A character indicating whether dendrograms should be drawn for both rows and columns "both", just rows "row" or column "column" or no dendrogram "none". Default is both.
cols.default	Logical. Default is TRUE. Use blue-brown color scheme.
lowcol, highcol	Character indicating colours to be used for down and upregulated genes when drawing heatmap if the default colors are not used, that is cols.default = FALSE.
scale	Default is row. Scale and center either "none", "row", or "column".
classvec, classvec2	A factor or vector which describes the classes in columns or rows of the dataset. Default is NULL. If included, a color bar including the class of each column (array sample) or row (gene) will be drawn. It will automatically add to either the columns or row, depending if the length(as.character(classvec)) == nrow(dataset) or ncol(dataset).
classvecCol A vector of length the number of levels in the factor classvec. These are the colors to be used for the row or column colorbar. Colors should be in the same order, as the levels(factor(classvec))

distfun A character, indicating function used to compute the distance between both rows and columns. Defaults to 1- Pearson Correlation coefficient

method The agglomeration method to be used. This should be one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". See hclust for more details. Default is "ave"

dualScale A logical indicating whether to scale the data by row and columns. Default is TRUE

zlim A vector of length 2, with lower and upper limits using for scaling data. Default is c(-3,3)

scaleKey A logical indicating whether to draw a heatmap color-key bar. Default is TRUE

returnSampleTree A logical indicating whether to return the sample (column) tree. If TRUE it will return an object of class dendrogram. Default is FALSE.

... further arguments passed to or from other methods.

Details

The hierarchical plot is produced using average linkage cluster analysis with a correlation metric distance. heatplot calls heatmap.2 in the R package gplots.

NOTE: We have changed heatplot scaling in made4 (v 1.19.1) in Bioconductor v2.5. Heatplot by default dual scales the data to limits of -3,3. To reproduce older version of heatplot, use the parameters dualScale=FALSE, scale="row".

Value

Plots a heatmap with dendrogram of hierarchical cluster analysis. If returnSampleTree is TRUE, it returns an object dendrogram which can be manipulated using

Note

Because Eisen et al., 1998 use green-red colours for the heatmap heatplot uses these by default however a blue-red or yellow-blue are easily obtained by changing lowcol and highcol

Author(s)

Aedin Culhane

References


See Also

See also as hclust, heatmap and dendrogram
Examples

data(khan)

## Change color scheme
heatplot(khan$train[1:30,],
heatplot(khan$train[1:30,], cols.default=FALSE, lowcol="white", highcol="red")

## Add labels to rows, columns
heatplot(khan$train[1:26,], labCol = c(64:1), labRow=LETTERS[1:26])

## Add a color bar
heatplot(khan$train[1:26,], classvec=khan$train.classes)
heatplot(khan$train[1:26,], classvec=khan$train.classes, classvecCol=c("magenta", "yellow", "cyan", "orange"))

## Change the scaling to the older made4 version (pre Bioconductor 2.5)
heatplot(khan$train[1:26,], classvec=khan$train.classes, dualScale=FALSE, scale="row")

## Getting the members of a cluster and manuipulating the tree
sTree<-heatplot(khan$train, classvec=khan$train.classes, returnSampleTree=TRUE)
class(sTree)
plot(sTree)

## Cut the tree at the height=1.0
lapply(cut(sTree,h=1)$lower, labels)

## Zoom in on the first cluster
plot(cut(sTree,1)$lower[[1]])
str(cut(sTree,1.0)$lower[[1]])

## Visualizing results from an ordination using heatplot
if (require(ade4, quiet = TRUE)) {
  res<-$ord(khan$train, ord.nf=5) # save 5 components from correspondence analysis
  khan.coa = res$ord
}

# Provides a view of the components of the Correspondence analysis (gene projection)
heatplot(khan.coa$li, dend="row", dualScale=FALSE) # first 5 components, do not cluster columns, only rows

# Provides a view of the components of the Correspondence analysis (sample projection)
# The difference between tissues and cell line samples are defined in the first axis.
heatplot(khan.coa$co, margins=c(4,20), dend="row") # Change the margin size. The default is c(5,5)

# Add a colorbar, change the heatmap color scheme and no scaling of data
heatplot(khan.coa$co,classvec2=khan$train.classes, cols.default=FALSE, lowcol="blue", dend="row", dualScale=FALSE)
apply(khan.coa$co,2, range)

html3D

Produce web page with a 3D graph that can be viewed using Chime web browser plug-in, and/or a pdb file that can be viewed using Ras-mol
**Description**

html3D produces a pdb file that can be viewed using the freeware protein structure viewer Rasmol and a html web page with a 3D graph that can be rotated and manipulated in a web browser that supports the chime web browser plug-in.

**Usage**

```r
html3D(df, classvec = NULL, writepdb = FALSE, filenamebase = "output",
       writehtml = FALSE, title = NULL, scaled=TRUE,xyz.axes=c(1:3), ...)
```

**Arguments**

- `df`: A `matrix` or `data.frame` containing the x,y,z coordinates. Typically the output from `bga` such as the \$ls or \$co files, or other xyz coordinates (\$li or \$co) produced by PCA, COA or other `dudi`
- `classvec`: factor or vector which describes classes in the df. Default is NULL. If specified each group will be coloured in contrasting colours
- `writepdb`: Logical. The default is FALSE. If TRUE a file will be saved which can be read into Rasmol.
- `writehtml`: Logical. The default is FALSE, If TRUE a web html file will be saved which can be viewed in any web browser than supports chime.
- `filenamebase`: Character. The basename of the html or pdb file(s) to be saved. The default is "output", which will save files output.pdb, output.html, if writepdb or writehtml are TRUE respectively.
- `title`: Character, the title (header) of the web page saved if writehtml is TRUE. The default is NULL.
- `scaled`: Logical indicating whether the data should be scaled for best fit. The default is TRUE
- `xyz.axes`: vector indicating which axes to use for x, y and z axes. By default, the first 3 columns of df.
- `...`: further arguments passed to or from other methods

**Details**

Produces a html file, of a 3D graph which can be rotated using the FREEWARE chime (win, MacOS). Chime can be downloaded from [http://www.mdlchime.com/](http://www.mdlchime.com/).

html3D will colour samples by classvec if given one, and will produce chime script to highlight groups, spin on/off, and include button for restore for example see [http://bioinf.ucd.ie/research/BGA/supplement.html](http://bioinf.ucd.ie/research/BGA/supplement.html)

html3d calls `chime3D` to produce the html web page with a 3D graph.

**Value**

html3D produces the pdb output file which can be read in Rasmol or other molecular structure viewers. html3D produces a html file with a 3D graph that can be rotated and manipulated in a web browser that supports the chime web browser plug-in.
Note

Note chime is only available on windows or Mac OS currently. Using the chime plug-in on Linux is slightly complicated but is available if the CrossOver Plug-in is installed. Instructions on installing this and chime on Linux are available at http://mirrors.rcsb.org/SMS/STINGm/help/chime_linux.html

If you wish to view a 3D graph in Rasmol, you will need to execute a Rasmol script similar to

```r
load pdbfilename.pdb;
set axes on; select off;
connect; set ambient 40;
rotate x 180; select *;
spacefill 40
```

html3D calls chime3D to produce the html file from the pdb file.

The author would like to thank Willie Taylor, The National Institute for Medical Research, London, UK for help with the awk command on which this function is based.

Author(s)

Aedin Culhane

Examples

```r
data(khan)
if (require(ade4, quiet = TRUE)) {
  khan.bga<-bga(khan$train, khan$train.classes)
}

out.3D <-html3D(khan.bga$bet$ls, khan.bga$fac, writepdb=TRUE, filenamebase ="Khan", writehtml=TRUE)

## Not run:
browseURL(paste("file://", file.path(paste(getwd(),"/khan.html", sep="")), sep=""))

## End(Not run)
```

khan


Description


Usage

data(khan)
Khan is dataset containing the following:

- **\$train**: `data.frame` of 306 rows and 64 columns. The training dataset of 64 arrays and 306 gene expression values
- **\$test**: `data.frame`, of 306 rows and 25 columns. The test dataset of 25 arrays and 306 genes expression values
- **\$gene.labels.imagesID**: `vector` of 306 Image clone identifiers corresponding to the rownames of \$train and \$test.
- **\$strain.classes**: `factor` with 4 levels "EWS", "BL-NHL", "NB" and "RMS", which correspond to the four groups in the \$train dataset
- **\$test.classes**: `factor` with 5 levels "EWS", "BL-NHL", "NB", "RMS" and "Norm" which correspond to the five groups in the \$test dataset
- **\$annotation**: `data.frame` of 306 rows and 8 columns. This table contains further gene annotation retrieved from SOURCE [http://SOURCE.stanford.edu](http://SOURCE.stanford.edu) in May 2004. For each of the 306 genes, it contains:
  - **\$CloneID**: Image Clone ID
  - **\$UGCluster**: The Unigene cluster to which the gene is assigned
  - **\$Symbol**: The HUGO gene symbol
  - **\$LLID**: The locus ID
  - **\$UGRepAcc**: The nucleotide sequence accession number
  - **\$LLRepProtAcc**: The protein sequence accession number
  - **\$Chromosome**: Chromosome location
  - **\$Cytoband**: Cytoband location

Details

Khan et al., 2001 used cDNA microarrays containing 6567 clones of which 3789 were known genes and 2778 were ESTs to study the expression of genes in four types of small round blue cell tumours of childhood (SRBCT). These were neuroblastoma (NB), rhabdomyosarcoma (RMS), Burkitt lymphoma, a subset of non-Hodgkin lymphoma (BL), and the Ewing family of tumours (EWS). Gene expression profiles from both tumour biopsy and cell line samples were obtained and are contained in this dataset. The dataset downloaded from the website contained the filtered dataset of 2308 gene expression profiles as described by Khan et al., 2001. This dataset is available from the [http://bioinf.ucd.ie/people/aedin/R/](http://bioinf.ucd.ie/people/aedin/R/).

In order to reduce the size of the MADE4 package, and produce small example datasets, the top 50 genes from the ends of 3 axes following bga were selected. This produced a reduced datasets of 306 genes.

Source


References


Examples

```r
data(khan)
summary(khan)
```

**Description**

NCI60 is a dataset of gene expression profiles of 60 National Cancer Institute (NCI) cell lines. These 60 human tumour cell lines are derived from patients with leukaemia, melanoma, along with, lung, colon, central nervous system, ovarian, renal, breast and prostate cancers. This panel of cell lines have been subjected to several different DNA microarray studies using both Affymetrix and spotted cDNA array technology. This dataset contains subsets from one cDNA spotted (Ross et al., 2000) and one Affymetrix (Staunton et al., 2001) study, and are pre-processed as described by Culhane et al., 2003.

**Usage**

```r
data(NCI60)
```

**Format**

The format is: List of 3

- `\$Ross: data.frame` containing 144 rows and 60 columns. 144 gene expression log ratio measurements of the NCI60 cell lines.
- `\$Affy: data.frame` containing 144 rows and 60 columns. 144 Affymetrix gene expression average difference measurements of the NCI60 cell lines.
- `\$classes: Data matrix` of 60 rows and 2 columns. The first column contains the names of the 60 cell line which were analysed. The second column lists the 9 phenotypes of the cell lines, which are BREAST, CNS, COLON, LEUK, MELAN, NSCLC, OVAR, PROSTATE, RENAL.
- `\$Annot: Data matrix` of 144 rows and 4 columns. The 144 rows contain the 144 genes in the \$Ross and \$Affy datasets, together with their Unigene IDs, and HUGO Gene Symbols. The Gene Symbols obtained for the \$Ross and \$Affy datasets differed (see note below), hence both are given. The columns of the matrix are the IMAGE ID of the clones of the \$Ross dataset, the HUGO Gene Symbols of these IMAGE clone ID obtained from SOURCE, the Affymetrix ID of the \$Affy dataset, and the HUGO Gene Symbols of these Affymetrix IDs obtained using annaffy.
Details

The datasets were processed as described by Culhane et al., 2003.

The Ross data frame contains gene expression profiles of each cell line in the NCI-60 panel, which were determined using spotted cDNA arrays containing 9,703 human cDNAs (Ross et al., 2000). The data were downloaded from The NCI Genomics and Bioinformatics Group Datasets resource http://discover.nci.nih.gov/datasetsNature2000.jsp. The updated version of this dataset (updated 12/19/01) was retrieved. Data were provided as log ratio values.

In this study, rows (genes) with greater than 15 and were removed from analysis, reducing the dataset to 5643 spot values per cell line. Remaining missing values were imputed using a K nearest neighbour method, with 16 neighbours and a Euclidean distance metric (Troyanskaya et al., 2001). The dataset $\$Ross$ contains a subset of the 144 genes of the 1375 genes set described by Scherf et al., 2000. This datasets is available for download from http://bioinf.ucd.ie/people/aedin/R/.

In order to reduce the size of the example datasets, the Unigene ID's for each of the 1375 IMAGE ID's for these genes were obtained using SOURCE http://source.stanford.edu. These were compared with the Unigene ID's of the 1517 gene subset of the $\$Affy$ dataset. 144 genes were common between the two datasets and these are contained in $\$Ross$.

The Affy data were derived using high density Hu6800 Affymetrix microarrays containing 7129 probe sets (Staunton et al., 2001). The dataset was downloaded from the Whitehead Institute Cancer Genomics supplemental data to the paper from Staunton et al., http://www-genome.wi.mit.edu/mpr/NCI60/, where the data were provided as average difference (perfect match-mismatch) values. As described by Staunton et al., an expression value of 100 units was assigned to all average difference values less than 100. Genes whose expression was invariant across all 60 cell lines were not considered, reducing the dataset to 4515 probe sets. This dataset NCI60$\$Affy$ of 1517 probe set, contains genes in which the minimum change in gene expression across all 60 cell lines was greater than 500 average difference units. Data were logged (base 2) and median centred. This datasets is available for download from http://bioinf.ucd.ie/people/aedin/R/.

In order to reduce the size of the example datasets, the Unigene ID’s for each of the 1517 Affymetrix ID of these genes were obtained using the function aafUniGene in the annaffy Bioconductor package. These 1517 Unigene IDs were compared with the Unigene ID's of the 1375 gene subset of the $\$Affy$ dataset. 144 genes were common between the two datasets and these are contained in $\$Affy$.

Source

These pre-processed datasets were available as a supplement to the paper:


References


Examples

data(NCI60)
summary(NCI60)

---

ord

**Ordination**

Description

Run principal component analysis, correspondence analysis or non-symmetric correspondence analysis on gene expression data.

Usage

```r
ord(dataset, type="coa", classvec=NULL, ord.nf=NULL, trans=FALSE, ...)
## S3 method for class 'ord'
plot(x, axis1=1, axis2=2, arraycol=NULL, genecol="gray25", nlab=10, genelabels= NULL, arraylabels=NULL, ...)
```

Arguments

dataset Training dataset. A matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.

classvec A factor or vector which describes the classes in the training dataset.

type Character, "coa", "pca" or "nsc" indicating which data transformation is required. The default value is type="coa".

ord.nf Numeric. Indicating the number of eigenvector to be saved, by default, if NULL, all eigenvectors will be saved.

trans Logical indicating whether 'dataset' should be transposed before ordination. Used by BGA Default is FALSE.

x An object of class ord. The output from ord. It contains the projection coordinates from ord, the \$co or \$li coordinates to be plotted.

arraycol, genecol Character, colour of points on plot. If arraycol is NULL, arraycol will obtain a set of contrasting colours using getcol, for each classes of cases (microarray samples) on the array (case) plot. genecol is the colour of the points for each variable (genes) on gene plot.
ord

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axis1 Integer, the column number for the x-axis. The default is 1.

axis2 Integer, the column number for the y-axis. The default is 2.

genelabels A vector of variables labels, if genelabels=NULL the row.names of input matrix dataset will be used.

arraylabels A vector of variables labels, if arraylabels=NULL the col.names of input matrix dataset will be used.

... further arguments passed to or from other methods.

Details

ord calls either dudi.pca, dudi.coa or dudi.nsc on the input dataset. The input format of the dataset is verified using array2ade4.

If the user defines microarray sample groupings, these are colours on plots produced by plot.ord.

Plotting and visualising bga results:

2D plots: plotarrays to draw an xy plot of cases (\$ls). plotgenes, is used to draw an xy plot of the variables (genes).

3D plots: 3D graphs can be generated using do3D and html3D. html3D produces a web page in which a 3D plot can be interactively rotated, zoomed, and in which classes or groups of cases can be easily highlighted.

1D plots, show one axis only: 1D graphs can be plotted using graph1D. graph1D can be used to plot either cases (microarrays) or variables (genes) and only requires a vector of coordinates (\$li, \$co)

Analysis of the distribution of variance among axes:

The number of axes or principal components from a ord will equal nrow the number of rows, or the ncol, number of columns of the dataset (whichever is less).

The distribution of variance among axes is described in the eigenvalues ($eig) of the ord analysis. These can be visualised using a scree plot, using scatterutil.eigen as it done in plot.ord.

It is also useful to visualise the principal components from a using a ord or principal components analysis dudi.pca, or correspondence analysis dudi.coa using a heatmap. In MADE4 the function heatplot will plot a heatmap with nicer default colours.

Extracting list of top variables (genes):

Use topgenes to get list of variables or cases at the ends of axes. It will return a list of the top n variables (by default n=5) at the positive, negative or both ends of an axes. sumstats can be used to return the angle (slope) and distance from the origin of a list of coordinates.

Value

A list with a class ord containing:

ord Results of initial ordination. A list of class "dudi" (see dudi)

fac The input classvec, the factor or vector which described the classes in the input dataset. Can be NULL.

Author(s)

Aedin Culhane
See Also

See Also \texttt{dudi.pca}, \texttt{dudi.coa} or \texttt{dudi.nsc}, \texttt{bga}.

Examples

data(khan)
if (require(ade4, quiet = TRUE)) {
  khan.coa <- ord(khan$train, classvec=khan$train.classes, type="coa")
}
khan.coa
plot(khan.coa, genelabels=khan$annotation$Symbol)
plotarrays(khan.coa)
# Provide a view of the first 5 principal components (axes) of the correspondence analysis
heatplot(khan.coa$ord$co[,1:5], dend="none", dualScale=FALSE)

overview

\textit{Draw boxplot, histogram and hierarchical tree of gene expression data}

Description

Very simple wrapper function that draws a boxplot, histogram and hierarchical tree of expression data

Usage

\texttt{overview(dataset, labels = NULL, title = \"\", classvec = NULL, hc = TRUE, boxplot = TRUE, hist = TRUE, returnTree = \FALSE, ...)}

Arguments

dataset \hspace{1cm} A \texttt{matrix}, \texttt{data.frame}, \texttt{ExpressionSet} or \texttt{marrayRaw-class}. If the input is gene expression data in a \texttt{matrix} or \texttt{data.frame}. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.

labels \hspace{1cm} Vector, labels to be placed on samples in plots. Default is \texttt{rownames(dataset)}.

title \hspace{1cm} Character, label to be placed on plots. Default is \texttt{NULL}.

classvec \hspace{1cm} A \texttt{factor} or \texttt{vector} which describes the classes in columns of the \texttt{dataset}. Default is \texttt{NULL}. If included columns (array samples) on the dendrogram will be coloured by class.

hc \hspace{1cm} Logical. Draw dendrogram of hierarchical cluster analysis of cases. Default is \texttt{TRUE}.

boxplot \hspace{1cm} Logical. Draw boxplot. Default is \texttt{TRUE}.

hist \hspace{1cm} Logical. Draw histogram. Default is \texttt{TRUE}.

returnTree \hspace{1cm} Logical. Return the hierarchical cluster analysis results. Default is \texttt{FALSE}.

Details

The hierarchical plot is produced using average linkage cluster analysis with Pearson’s correlation metric as described by Eisen et al.,1999.
Author(s)
Aedin Culhane

See Also
See also as boxplot, hclust, hist

Examples
data(khan)

logkhan<-log2(khan$train)
print(class(logkhan))
overview(logkhan, title="Subset of Khan Train")
overview(logkhan, classvec=khan$train.classes, labels=khan$train.classes,title="Subset of Khan Train")
overview(logkhan, classvec=khan$train.classes, labels=khan$train.classes,title="Subset of Khan Train", boxplot=FALSE, his=FALSE)

plotarrays

Graph xy plot of variable (array) projections from ordination, between group analysis or coinertia analysis.

Description
Graph xy plot of variables using s.var, s.groups or s.match.col. Useful for visualising array coordinates (\$li) resulting from ord, bga or cia of microarray data.

Usage

plotarrays(coord, axis1 = 1, axis2 = 2, arraylabels = NULL, classvec=NULL, graph = c("groups", "simple", "labels", "groups2", "coinertia", "coinertia2"), labelsize=1, star=1, ellipse=1, arraycol=NULL, ...)

Arguments

coord a data.frame or matrix or object from ord bga or cia analysis with at least two columns, containing x, y coordinates to be plotted

axis1 An integer, the column number for the x-axis. Default is 1, so axes 1 is dudi-var[,1]

axis2 An integer, the column number for the y-axis. Default is 2, so axes 2 is dudi-var[,2]

arraylabels A vector of variables labels. Default is row.names(coord)

classvec A factor or vector which describes the classes in coord. Default is NULL. If included variables will be coloured by class.

graph A character of type "groups", "simple", "labels", "groups2", "coinertia" or "coinertia2" which specifies the type of plot type or "graph" to be drawn. By default the graph will be selected depending on the class of coord, and whether a classvector is specified

labelsize Size of sample labels, by default=1

star If drawing groups, whether to join samples to centroid creating a "star"

ellipse If drawing groups, whether to draw an ellipse or ring around the samples

arraycol Character with length equal to the number of levels in the factor classvec. Colors for each of the levels in the factor classvec

... further arguments passed to or from other method
plotgenes

Details

plotarrays calls the function s.var, s.groups or s.match.col.

If you wish to return a table or list of the top array at the end of an axis, use the function topgenes.

Value

An xy plot

Note

plotarrays plots variables using s.var, s.groups, s.match.col which are modifieds version of s.label, s.class, and s.match.

Author(s)

Aedin Culhane

See Also

See Also as s.var and s.label

Examples

data(khan)
if (require(ade4, quiet = TRUE)) {
  khan.bga<-bga(khan$train, khan$train.classes)
}
attach(khan.bga)
par(mfrow=c(2,1))
plotarrays(khan.bga)
plotarrays(khan.bga, graph="simple")
plotarrays(khan.bga, graph="labels")
plotarrays(khan.bga, graph="groups")
plotarrays(khan.bga, graph="groups2")
plotgenes

Arguments

coord     a data.frame or matrix or object from ord bga or cia analysis with at least two columns, containing x, y coordinates to be plotted.
nlab      Numeric. An integer indicating the number of variables at ends of axes to be labelled.
axis1     An integer, the column number for the x-axis. Default is 1, so axis 1 is dudi-var[,1].
axis2     An integer, the column number for the y-axis. Default is 2, so axis 2 is dudi-var[,2].
genelabels A vector of gene (variable) labels. Default is row.names(coord)
boxes     A logical, indicating whether a box should be plotted surrounding each variable label. The default is TRUE.
colpoints The colour of the points on the plot. The default is "black".
...       further arguments passed to or from other method.

Details

plotgenes calls the function genes which return an index of the "top" variables at the ends of the x and y axes.
If you wish to return a table or list of the top genes at the end of an axis, use the function topgenes.

Value

An xy plot

Note

plotgenes plots variables using s.var, which is a modified version of s.label.

Author(s)

Aedin Culhane

See Also

See Also as s.var and s.label

Examples

data(khan)
if (require(ade4, quiet = TRUE)) {
  khan.ord<-ord(khan$train, classvec=khan$train.classes)
}
par(mfrow=c(2,2))
#s.var(khan.ord$co, col=as.numeric(khan$train.classes), clabel=0.8)
plotgenes(khan.ord, colpoints="red")
plotgenes(khan.ord, colpoints="red", genelabels=khan$annotation$Symbol)
plotgenes(khan.ord, colpoints="gray", genelabels=khan$annotation$Symbol,boxes=FALSE)
Function which performs a hierarchical cluster analysis of data, drawing a dendrogram, with colorbars for different sample covariate beneath the dendrogram.

**Arguments**

- `dataset`: a matrix, data.frame, ExpressionSet or marrayRaw-class. If the input is gene expression data in a matrix or data.frame. The rows and columns are expected to contain the variables (genes) and cases (array samples) respectively.
- `labels`: Vector, labels to be placed on samples in plots. Default is rownames(dataset).
- `title`: Character, label to be placed on plots. Default is NULL.
- `classvec`: A factor or vector or matrix or data.frame which describes the classes in columns of the dataset. Default is NULL.
- `covars`: Numeric. The columns of the data.frame classve to be used as class vectors. These will be displayed as color bars under the dendrogram. The default is 1 (column 1).
- `returnTree`: Logical. Return the hierarchical cluster analysis results. Default is FALSE.
- `getPalette`: Function, which generates a palette of colors. The default uses getcol function in made4. Other examples are provided below

**Details**

The hierarchical plot is produced using average linkage cluster analysis with 1- Pearson’s correlation metric. The default set of colors used to generate the color bars of the plots can be changed (see example below). By default, if there is only two levels in the factor, the colors will be black and grey.

**Author(s)**

Aedin Culhane

**See Also**

See also as overview, hclust
Examples

data(khan)
logkhan<-log2(khan$train)

# Get a character vector which defines which khan samples are cell lines or tissue sample
khanAnnot=cbind(as.character(khan$train.classes),khan$cellType)
print(khanAnnot[1:3,])

# Add 2 color bar, one for cancer subtype, another for cell type under dendrogram
prettyDend(logkhan, classvec=khanAnnot, covars = c(1,2), labels=khan$train.classes)

# To change the palette of colors
# Use topo.colors(), see colors() for more help on inbuilt palettes

prettyDend(logkhan, classvec=khanAnnot, covars = c(1,2), labels=khan$train.classes, getPalette=topo.colors)

# To use RColorBrewer Palettes

library(RColorBrewer)

# Use RColorBrewer Dark2 which contains 8 colors
prettyDend(logkhan, classvec=khanAnnot, covars = c(1,2), labels=khan$train.classes, getPalette=function(x)

# Use RColorBrewer Set1 which contains 9 colors
prettyDend(logkhan, classvec=khanAnnot, covars = c(1,2), labels=khan$train.classes, getPalette=function(x)

randomiser

Randomly reassign training and test samples

Description

This function is used to check for bias between a training and test data. It return a new index, which randomly re-assigns samples in the training data to the test dataset and vice versa.

Usage

randomiser(ntrain = 77, ntest = 19)

Arguments

ntrain Numeric. A integer indicating the number of cases in the training dataset
ntest Numeric. A integer indicating the number of cases in the test dataset

Details

Produces new indices that can be used for training/test datasets

Value

It returns a list, containing 2 vectors

train A vector of length ntrain, which can be used to index a new training dataset
test A vector of length ntest, which can be used to index a new test dataset
**sumstats**

**Author(s)**

Aedin Culhane

**Examples**

```r
randomiser(10,5)
train<-matrix(rnorm(400), ncol=20, nrow=20, dimnames=list(1:20, paste("train",letters[1:20], sep=".")))
test<-matrix(rnorm(200), ncol=10, nrow=20, dimnames=list(1:20, paste("test", LETTERS[1:10], sep=".")))
all<-cbind(train,test)

colnames(train)
colnames(test)
newInd<-randomiser(ntrain=20, ntest=10)

newtrain<-all[,newInd$train]
newtest<-all[,newInd$test]

colnames(newtrain)
colnames(newtest)
```

---

| sumstats | Summary statistics on xy co-ordinates, returns the slopes and distance from origin of each co-ordinate. |

---

**Description**

Given a `data.frame` or `matrix` containing xy coordinates, it returns the slope and distance from origin of each coordinate.

**Usage**

`sumstats(array, xax = 1, yax = 2)`

**Arguments**

- `array` A `data.frame` or `matrix` containing xy coordinates, normally a `$co`, `$li` from `dudi` such as PCA or COA, or `$ls` from `bga`
- `xax` Numeric, an integer indicating the column of the x axis coordinates. Default `xax=1`
- `yax` Numeric, an integer indicating the column of the x axis coordinates. Default `yax=2`

**Details**

In PCA or COA, the variables (upregulated genes) that are most associated with a case (microarray sample), are those that are projected in the same direction from the origin.
Variables or cases that have a greater contribution to the variance in the data are projected further from the origin in PCA. Equally variables and cases with the strong association have a high chi-square value, and are projected with greater distance from the origin in COA. See a description from Culhane et al., 2002 for more details.

Although the projection of co-ordinates are best visualised on an xy plot, sumstats returns the slope and distance from origin of each x,y coordinate in a matrix.

**Value**

A matrix (ncol=3) containing

- **slope**
- **angle (in degrees)**
- **distance from origin**

of each x,y coordinates in a matrix.

**Author(s)**

Aedin Culhane

**Examples**

```r
data(khan)
if (require(ade4, quiet = TRUE)) {
  khan.bga<-bga(khan$train, khan$train.classes)
  plotarrays(khan.bga$bet$ls, classvec=khan$train.classes)
  st.out<-sumstats(khan.bga$bet$ls)
  # Get stats on classes EWS and BL
  EWS<-khan$train.classes==levels(khan$train.classes)[1]
  st.out[EWS,]
  BL<-khan$train.classes==levels(khan$train.classes)[2]
  st.out[BL,]
  # Add dashed line to plot to highlight min and max slopes of class BL
  slope.BL.min<min(st.out[BL,1])
  slope.BL.max<max(st.out[BL,1])
  abline(c(0,slope.BL.min), col="red", lty=5)
  abline(c(0,slope.BL.max), col="red", lty=5)
```

---

**suppl**

Projection of supplementary data onto axes from a between group analysis

**Description**

Projection and class prediction of supplementary points onto axes from a between group analysis, `bga`.
Usage

```r
suppl(dudi.bga, supdata, supvec = NULL, assign=TRUE, ...)
## S3 method for class 'suppl'
plot(x, dudi.bga, axis1=1, axis2=2, supvec=sup$true.class, supvec.pred= sup$predicted, ...)
```

Arguments

- `dudi.bga`: An object returned by `bga`.
- `supdata`: Test or blind dataset. Accepted formats are a `matrix`, `data.frame`, `ExpressionSet` or `marrayRaw-class`.
- `supvec`: A factor or vector which describes the classes in the training dataset.
- `supvec.pred`: A factor or vector which describes the classes which were predicted by `suppl`.
- `assign`: A logical indicating whether class assignment should be calculated using the method described by Culhane et al., 2002. The default value is `TRUE`.
- `x`: An object returned by `suppl`.
- `axis1`: Integer, the column number for the x-axis. The default is 1.
- `axis2`: Integer, the column number for the y-axis. The default is 2.
- `...`: Further arguments passed to or from other methods.

Details

After performing a between group analysis on a training dataset using `bga`, a test dataset can be then projected onto `bga` axes using `suppl`. `suppl` returns the projected coordinates and assignment of each test case (array).

The test dataset must contain the same number of variables (genes) as the training dataset. The input format of both the training dataset and test dataset are verified using `array2ade4`. Use `plot.bga` to plot results from `bga`.

Value

A list containing:

- `suppl`: An object returned by `suppl`.

Author(s)

Aedin Culhane

References


See Also

See Also `bga`, `bca`, `plot.bga`, `bga.jackknife`
Examples

```r
data(khan)
# khan.bga <- bga(khan$train, khan$train.classes)
if (require(ade4, quiet = TRUE)) {
  khan.bga <- bga.suppl(khan$train, supdata = khan$test, classvec = khan$train.classes, supvec = khan$test.classes)
  khan.bga
  plot.bga(khan.bga, genelabels = khan$annotation$Symbol)
  plot.suppl(khan.bga$suppl, khan.bga)
  plot.suppl(khan.bga$suppl, khan.bga, supvec = NULL, supvec.pred = NULL)
  plot.suppl(khan.bga$suppl, khan.bga, axis1 = 2, axis2 = 3, supvec = NULL, supvec.pred = NULL)
}
```

```
topgenes
Topgenes, returns a list of variables at the ends (positive, negative or both) of an axis
```

Description

topgenes will return a list of the top N variables from the positive, negative or both ends of an axis. That is, it returns a list of variables that have the maximum and/or minimum values in a vector.

Usage

```
topgenes(x, n = 10, axis = 1, labels = row.names(x), ends = "both", ...)
```

Arguments

- **x**: A vector, matrix or data.frame. Typically a data frame \$co or \$li from dudi or \$s, \$l, \$co from bga.
- **n**: An integer indicating the number of variables to be returned. Default is 5.
- **axis**: An integer indicating the column of x. Default is 1 (first axis, of \$co or \$li file)
- **labels**: A vector of row names, for x[,axis]. Default values is row.names(x)
- **ends**: A string, "both", "neg", "pos", indicating whether variable label should be return from both, the negative or the positive end of an axis. The default is both.
- **...**: further arguments passed to or from other methods

Details

topgenes calls genes1d. genes1d is similar to genes, but returns an index of genes at the ends of one axis not two axes. Given a \$co or \$li file it will return that variables at the ends of the axis.

Value

Returns a vector or list of vectors.
Author(s)
Aedin Culhane

See Also
See Also as genes

Examples

# Simple example
a <- rnorm(50)
order(a)
topgenes(a, labels = c(1:length(a)), ends = "neg")

# Applied example
data(khan)
if (require(ade4, quiet = TRUE)) {
khan.coa <- ord(khan$train[1:100,])
ind <- topgenes(khan.coa, ends = "pos")
ind.ID <- topgenes(khan.coa, ends = "pos", labels = khan$gene.labels.imagesID)
ind.symbol <- topgenes(khan.coa, ends = "pos", labels = khan$annotationSymbol)
Top10.pos <- cbind("Gene Symbol" = ind.symbol,
                   "Clone ID" = ind.ID, "Coordinates" = khan.coa$ord$li[ind,], row.names = c(1:length(ind)))
Top10.pos
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