Package 'biplotEZ'

September 14, 2024

Title EZ-to-Use Biplots

Version 2.1

```
Description Provides users with an EZ-to-use platform for representing data
       with biplots. Currently principal component analysis (PCA), canonical variate
       analysis (CVA) and simple correspondence analysis (CA) biplots are included.
       This is accompanied by various formatting
       options for the samples and axes. Alpha-bags and concentration ellipses
       are included for visual enhancements and interpretation. For an extensive
       discussion on the topic, see Gower, J.C., Lubbe, S. and le Roux, N.J.
       (2011, ISBN: 978-0-470-01255-0) Understanding Biplots. Wiley: Chichester.
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```

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Description

This function produces α -bags, which is a useful graphical summary of the scatter plot. The alphabag refers to a contour which contains $\alpha\%$ of the observations.

Usage

```
alpha.bags(bp, alpha = 0.95, which = NULL, col = ez.col[which], lty = 1,
lwd = 1, max = 2500, trace = TRUE, opacity = 0.25, outlying=FALSE)
```

Arguments

bp	an object of class biplot.	
alpha	numeric vector between 0 and 1 to determine coverage of the bag (α) , with default 0.95.	
which	numeric vector indicating the selection of groups or classes to be fitted with $\alpha\text{-bags}.$	
col	vector of colours for the α -bags. Multiple α bags for one group will be displayed in the same colour.	
lty	vector of line types for the $\alpha\text{-bags}.$ The same line type will be used per value of $\alpha.$	
lwd	vector of line widths for the $\alpha\text{-bags}.$ The same line width will be used per value of $\alpha.$	
max	maximum number of samples to include in α -bag calculations, with default 2500. If more samples are in the group, a random sample of size max is taken for the computations.	
trace	logical, indicating progress of computation.	
opacity	level of opacity, with default 0.5.	
outlying	utlying logical indicating whether only outlying points should be plotted. Note which argument may be overwritten when TRUE	

Value

A list with the following components is available:

alpha.bags	list of coordinates for the α -bags for each group.
col	vector of colours for the α -bags.
lty	vector of line types for the α -bags.
lwd	vector of line widths for the α -bags.

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References

Gower, J., Gardner-Lubbe, S. & Le Roux, N. (2011, ISBN: 978-0-470-01255-0) *Understanding Biplots*. Chichester, England: John Wiley & Sons Ltd.

Examples

```
biplot (iris[,1:4]) |> PCA(group.aes=iris[,5]) |> alpha.bags(alpha=0.95) |> plot()
biplot (iris[,1:4],group.aes=iris[,5]) |> PCA() |> alpha.bags(alpha=0.95) |> plot()
```

AoD

Use the Analysis of Distance (AoD) method to construct the biplot

Description

This function appends the biplot object with elements resulting from using the AoD method.

Usage

```
AoD(bp, classes=bp$classes, dist.func=NULL, dist.func.cat=NULL, dim.biplot = c(2,1,3), e.vects = 1:ncol(bp$X), weighted = c("unweighted", "weighted"), show.class.means = TRUE, axes = c("regression", "splines"), ...)
```

more arguments to dist.func.

Arguments

	bp	an object of class biplot obtained from preceding function biplot().
	classes	a vector of the same length as the number of rows in the data matrix with the class indicator for the samples.
	dist.func	a character string indicating which distance function is used to compute the Euclidean embeddable distances between samples. One of NULL (default) which computes the Euclidean distance or other functions that can be used for the dist() function.
	dist.func.cat	a character string indicating which distance function is used to compute the Euclidean embeddable distances between samples. One of NULL (default) which computes the extended matching coefficient or other functions.
	dim.biplot	the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.
e.vects the vector indicating which eigenvectors (canonical variates) should be ploin the biplot, with default 1:dim.biplot.		the vector indicating which eigenvectors (canonical variates) should be plotted in the biplot, with default 1:dim.biplot.
	weighted	a character string indicating the weighting of the classes. One of "unweighted" for each class to receive equal weighting or "weighted" for each class to receive their class sizes as weights.
show.class.means		
		a logical value indicating whether to plot the class means on the biplot.
	axes	a character string indicating the type of biplot axes to be used in the biplot. One of "regression" or "splines".

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Value

Object of class biplot

Examples

```
biplot(iris[,1:4]) |> AoD(classes=iris[,5])
# create a CVA biplot
biplot(iris[,1:4]) |> AoD(classes=iris[,5]) |> plot()
```

AoD.biplot

Calculate elements for the Analysis of Distance (AoD) biplot

Description

This function is used to construct the AoD biplot

Usage

```
## $3 method for class 'biplot'
AoD(
    bp,
    classes = bp$classes,
    dist.func = NULL,
    dist.func.cat = NULL,
    dim.biplot = c(2, 1, 3),
    e.vects = 1:ncol(bp$X),
    weighted = c("unweighted", "weighted"),
    show.class.means = TRUE,
    axes = c("regression", "splines"),
    ...
)
```

Arguments

dim.biplot

bp	an object of class biplot obtained from preceding function biplot().	
classes	a vector of the same length as the number of rows in the data matrix with th class indicator for the samples.	
dist.func	a character string indicating which distance function is used to compute the Euclidean embeddable distances between samples. One of NULL (default) which computes the Euclidean distance or other functions that can be used for the dist() function.	
dist.func.cat	a character string indicating which distance function is used to compute the Euclidean embeddable distances between samples. One of NULL (default) which computes the extended matching coefficient or other functions.	

the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

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e.vects the vector indicating which eigenvectors (canonical variates) should be plotted

in the biplot, with default 1:dim.biplot.

a character string indicating the weighting of the classes. One of "unweighted" weighted

for each class to receive equal weighting or "weighted" for each class to receive

their class sizes as weights.

show.class.means

a logical value indicating whether to plot the class means on the biplot.

a character string indicating the type of biplot axes to be used in the biplot. One axes

of "regression" or "splines".

more arguments to dist.func.

Value

an object of class biplot.

Examples

```
biplot(iris) |> AoD(classes = iris[,5]) |> plot()
```

axes

Format aesthetics for the biplot axes

Description

This function allows the user to format the aesthetics for the biplot axes.

Usage

```
axes(bp, X.names=colnames(bp$X), which = 1:bp$p, col = grey(0.7), lwd = 1, lty = 1,
label.dir = "Orthog", label.col = col, label.cex = 0.75, label.line = 0.1,
label.offset=rep(0,4), ticks = 5, tick.col = col, tick.size = 1, tick.label = TRUE,
tick.label.side = "below", tick.label.col = tick.col, tick.label.cex = 0.6,
predict.col = col, predict.lwd = lwd, predict.lty = lty, ax.names = X.names,
orthogx = 0, orthogy = 0, vectors = FALSE, unit.circle=FALSE)
```

Arguments

bp	an object of class biplot.	
X.names	a vector of column names of bp to specify which axes should be labelled.	
which	a vector containing the columns or variables for which the axes should be displayed, with default $1:p$.	
col	the colour(s) for the axes, with default grey(0.7). Alternatively, provide a vector of colours corresponding to X .names.	
lwd	the line width(s) for the axes, with default 1.	

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lty	the line type(s) for the axes, with default 1.	
label.dir	a character string indicating the placement of the axis titles to the side of the figure. One of "Orthog" for axis titles to appear orthogonal to the side of the figure (default), "Hor" for axis titles to appear horizontally or "Paral" for axis titles to appear parallel to the side of the figure.	
label.col	the colour(s) for the axis labels, with default, col.	
label.cex	the label expansion for the axis labels, with default 0.75.	
label.line	the distance of the axis title from the side of the figure, with default $\emptyset.1$.	
label.offset	a four-component numeric vector controlling the distances axis titles are displayed from the side of the figure, with default $rep(0,4)$. Sides are numbered 1 to 4 according to R conventions.	
ticks	an integer-valued vector indicating the number of tickmarks for each axis, with default 5 for each axis.	
tick.col	the colour(s) for the tick marks, with default col.	
tick.size	a vector specifying the sizes of tick marks for each axis, with default 1 for each .	
tick.label	a logical value indicating whether the axes should be labelled, with default TRUE.	
tick.label.side		
	a character string indicating the position of the tick label. One of "below" for the label to appear below the tick mark (default) or "above" for the label to appear above the tick mark.	
tick.label.col	the colour(s) for the tick mark labels, with default tick.col.	
tick.label.cex	the label expansion for the tick mark labels, with default 0.6.	
predict.col	the colour(s) for the predicted samples, with default col.	
predict.lwd	<pre>lwd the line width(s) for the predicted samples, with default lwd.</pre>	
predict.lty	dict.lty the line type(s) for the predicted samples, with default lty.	
ax.names	a vector of size p containing user defined titles for the axes.	
orthogx	a numeric vector of size p specifying the x-coordinate of the parallel tran formation of each axis, with default 0 for each axis. This is only used who dim.biplot = 2.	
orthogy	a numeric vector of size p specifying the y-coordinate of the parallel transformation of each axis, with default 0 for each axis. This is only used when dim.biplot = 2.	
vectors	a logical value indicating whether vector representation (calibrated axes) should be displayed on the biplot. This is only used when a PCA biplot is produced.	
unit.circle	it.circle a logical value indicating whether a unit circle should be displayed on the biplo	

Value

The object of class biplot will be appended with a list called axes containing the following elements:

which a vector containing the columns for which the axes are displayed.

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the colour(s) of the axes. col lwd the line width(s) of the axes. 1ty the line type(s) of the axes. label.dir the placement of the axis titles to the side of the figure. label.col the colour(s) of the axis titles. label.cex the label expansion(s) of the axis titles. label.line the distance(s) of the axis titles from the side of the figure. ticks the number of tick marks per axis. tick.col the colour(s) of the tick marks. tick.size the size(s) of the tick marks. tick.label logical value(s) indicating whether axes are labelled. tick.label.side the position of the tick mark labels. tick.label.col the colour(s) of the tick mark labels. tick.label.cex the expansion(s) of the tick mark labels. the colour(s) of the predicted samples. predict.col predict.lty the line type(s) of the predicted samples. the line width(s) of the predicted samples. predict.lwd the user defined axis titles. names orthogx the horizontal translations for each axis. orthogy the vertical translations for each axis. vectors a logical value indicating whether calibrated axes are plotted.

See Also

biplot()

Examples

```
biplot(iris[,1:4]) |> PCA() |> axes(col="purple") |> plot()
biplot(iris[,1:4]) |> PCA() |> samples(col="purple",pch=15) |> axes() |> plot()
```

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axes_coordinates

Calibrate Biplot Axes

Description

Convenience function to obtain the coordinates of the calibrated ticks marks on the biplot

Usage

```
axes_coordinates(x)
```

Arguments

Х

an object of class biplot

Value

An ordered list containing the coordinates the of tick marks to plotted on the biplot

Examples

```
x<-biplot(iris) |> PCA()
coordinates<-axes_coordinates(x)</pre>
```

biplot

First step to create a new biplot with biplotEZ

Description

This function produces a list of elements to be used when producing a biplot, which provides a useful data analysis tool and allows the visual appraisal of the structure of large data matrices. Biplots are the multivariate analogue of scatter plots. They approximate the multivariate distribution of a sample in a few dimensions and they superimpose on this display representations of the variables on which the samples are measured.

Usage

```
biplot(data, classes = NULL, group.aes = NULL, center = TRUE, scaled = FALSE,
Title = NULL)
```

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Arguments

data a data frame or numeric matrix containing all variables the user wants to analyse.

classes a vector identifying class membership.

group.aes a vector identifying groups for aesthetic formatting.

center a logical value indicating whether data should be column centered, with default

TRUE.

scaled a logical value indicating whether data should be standardised to unit column

variances, with default FALSE.

Title the title of the biplot to be rendered, enter text in " ".

Details

This function is the entry-level function in biplotEZ to construct a biplot display. It initialises an object of class biplot which can then be piped to various other functions to build up the biplot display.

Value

means

A list with the following components is available:

X the matrix of the centered and scaled numeric variables.

Xcat the data frame of the categorical variables.

raw. X the original data.

classes the vector of category levels for the class variable. This is to be used for colour,

pch and cex specifications.

na.action the vector of observations that have been removed.

center a logical value indicating whether \mathbf{X} is centered.

scaled a logical value indicating whether X is scaled.

sd the vector of standard deviations for each numeric variable.

the vector of means for each numeric variable.

n the number of observations.

p the number of variables.

group.aes the vector of category levels for the grouping variable. This is to be used for

colour, pch and cex specifications.

g. names the descriptive names to be used for group labels.

g the number of groups.

Title the title of the biplot rendered

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Useful links

The biplot display can be built up in four broad steps depending on the needs for the display. Firstly, choose an appropriate method to construct the display; Secondly, change the aesthetics of the display; Thirdly, append the display with supplementary features such as axes, samples and means; Finally, superimpose shapes, characters or elements onto the display.

1. Different types of biplots:

- PCA(): Principal Component Analysis biplot of various dimensions
- CVA(): Canonical Variate Analysis biplot
- PCO(): Principal Coordinate Analysis biplot
- CA(): Correspondence Analysis biplot
- regress(): Regression biplot method

2. Customise the biplot display with aesthetic functions:

- samples(): Change the formatting of sample points on the biplot display
- axes(): Change the formatting of the biplot axes

3. Supplement the existing biplot with additional axes, samples and group means:

- newsamples(): Add and change formatting of additional samples
- newaxes(): Add and change formatting of additional axes
- means(): Insert class means to the display, and format appropriately

4. Append the biplot display:

- alpha.bags(): Add α -bags
- ellipses(): Add ellipses
- density2D(): Add 2D density regions

Other useful links:

- plot()
- fit.measures()
- legend.type()
- interpolate()
- prediction()
- classify()
- reflect()
- rotate()

References

Gabriel, K.R. (1971) The biplot graphic display of matrices with application to principal component analysis. *Biometrika*. 58(3):453–467. Gower, J., Gardner-Lubbe, S. & Le Roux, N. (2011, ISBN: 978-0-470-01255-0) *Understanding Biplots*. Chichester, England: John Wiley & Sons Ltd. Gower, J.C. & Hand, D.J.(1996, ISBN: 0-412-71630-5) *Biplots*. London: Chapman & Hall.

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Examples

```
biplot(data = iris)
# create a PCA biplot
biplot(data = iris) |> PCA() |> plot()
```

biplotEZ

biplotEZ: EZ-to-Use Biplots

Description

Details

The goal of biplotEZ is to provide users an EZ-to-use platform for visually representing their data with biplots. Currently, this package includes principal component analysis (PCA) and canonical variate analysis (CVA) biplots. This is accompanied by various formatting options for the samples and axes. Alpha-bags and concentration ellipses are included for visual enhancements and interpretation.

Details

Package: biplotEZ
Type: Package
Version: 2.0

Date: 05-04-2024 License: MIT LazyLoad: TRUE

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Core Functions

- biplot
- PCA
- CVA
- **C**A

Code Availability

The newest version of the package can be obtained on GitHub: https://github.com/MuViSU/biplotEZ

CA

Correspondence Analysis (CA) method

Description

This function produces a list of elements to be used for CA biplot construction by approximation of the Pearson residuals.

Usage

```
CA(bp, dim.biplot = c(2,1,3), e.vects = 1:ncol(bp$X), variant = "Princ",
lambda.scal = FALSE)
```

Arguments

bp	object of class biplot obtained from preceding function biplot(center = FALSE). In order to maintain the frequency table, the input should not be centered or scaled. For CA, bp should be a contingency table.
dim.biplot	dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.
e.vects	which eigenvectors (canonical variates) to extract, with default 1:dim.biplot.
variant	which correspondence analysis variant, with default "Princ", presents a biplot with rows in principal coordinates and columns in standard coordinates. variant = "Stand", presents a biplot with rows in standard coordinates and columns in principal coordinates. variant = "symmetric", presents a symmetric biplot with row and column standard coordinates scaled equally by the singular values.
lambda.scal	logical value to request lambda-scaling, default is FALSE.

Value

A list with the following components is available:

Z Combined data frame of the row and column coordinates.

r Numer of levels in the row factor.

c Numer of levels in the column factor.

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rowcoor Row coordinates based on the selected variant.

colcoor Column coordinates based on the selected variant.

P Correspondence Matrix

Smat Standardised Pearson residuals.

SVD Singular value decomposition solution: d, u, v.

qual Quality of the approximation.

lambda.val The computed lambda value if lambda-scaling is requested.

See Also

```
biplot()
```

Examples

```
# Creating a CA biplot with rows in principal coordinates:
biplot(HairEyeColor[,,2], center = FALSE) |> CA() |> plot()
# Creating a CA biplot with rows in standard coordinates:
biplot(HairEyeColor[,,2], center = FALSE) |> CA(variant = "Stand") |>
samples(col=c("magenta","purple"), pch=c(15,17), label.col="black") |> plot()
# Creating a CA biplot with rows and columns scaled equally:
biplot(HairEyeColor[,,2], center = FALSE) |> CA(variant = "Symmetric") |>
samples(col=c("magenta","purple"), pch=c(15,17), label.col="black") |> plot()
```

CA.biplot

CA biplot

Description

Performs calculations for a CA biplot.

Usage

```
## S3 method for class 'biplot'
CA(
   bp,
   dim.biplot = c(2, 1, 3),
   e.vects = 1:ncol(bp$X),
   variant = "Princ",
   lambda.scal = FALSE
)
```

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Arguments

bp object of class biplot obtained from preceding function biplot(center = FALSE).

In order to maintain the frequency table, the input should not be centered or

scaled. For CA, bp should be a contingency table.

dim.biplot dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

e.vects which eigenvectors (canonical variates) to extract, with default 1:dim.biplot.

variant which correspondence analysis variant, with default "Princ", presents a biplot with rows in principal coordinates and columns in standard coordinates. variant

= "Stand", presents a biplot with rows in standard coordinates and columns in principal coordinates. variant = "symmetric", presents a symmetric biplot with row and column standard coordinates scaled equally by the singular values.

lambda.scal logical value to request lambda-scaling, default is FALSE.

Value

an object of class CA, inherits from class biplot.

Examples

```
biplot(HairEyeColor[,,2], center = FALSE) |> CA() |> plot()
```

CATPCA

Categorical Principal Component Analysis

Description

Categorical Principal Component Analysis

Usage

```
CATPCA(bp, dim.biplot = c(2, 1, 3), e.vects = 1:ncol(bp$X), group.aes = NULL, show.class.means = FALSE)
```

Arguments

bp an object of class biplot obtained from preceding function biplot().

dim.biplot dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

e.vects which eigenvectors (principal components) to extract, with default 1:dim.biplot.

group.aes vector of the same length as the number of rows in the data matrix for differen-

tiated aesthetics for samples.

show.class.means

logical, indicating whether group means should be plotted in the biplot.

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Value

```
an object of class biplot
```

Examples

```
CATPCA (iris)
```

classification

Classification biplot method

Description

This function produces a list of elements to be used for constructing a classification biplot.

Usage

```
classification(bp, Pmat, dim.biplot = c(2, 1, 3), e.vects = 1:ncol(bp$X),
group.aes=NULL, axes = "regression", col=ez.col, opacity=0.4, borders = FALSE)
```

Arguments

bp an object of class biplot obtained from preceding function biplot().	an object of class biplot obtained from preceding function biplot().	
Pmat a matrix containing the posterior probability for the classes		
dim.biplot dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.		
e.vects which eigenvectors (principal components) to extract, with default 1:dim.bipl		
group.aes vector of the same length as the number of rows in the data matrix for differentiated aesthetics for samples.		
axes type of axes, defaults to "regression"		
col colour of the classification regions		
opacity opacity of classification regions		

logical, indicating whether borders should be added to classification regions

Value

Object of class biplot with the following elements:

References

borders

Gardner-Lubbe, S., 2016. A triplot for multiclass classification visualisation. *Computational Statistics & Data Analysis*, 94, pp.20-32.

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Examples

```
biplot(iris[,1:4]) |>
classification(predict(MASS::lda(Species ~ ., data = iris))$posterior)
# create a classification biplot
biplot(iris[,1:4]) |>
classification(predict(MASS::lda(Species ~ ., data = iris))$posterior) |>
plot()
```

classification.biplot classification biplot

Description

Performs calculations for a classification biplot.

Usage

```
## S3 method for class 'biplot'
classification(
   bp,
   Pmat,
   dim.biplot = c(2, 1, 3),
   e.vects = 1:ncol(bp$X),
   group.aes = NULL,
   axes = "regression",
   col = ez.col,
   opacity = 0.4,
   borders = FALSE
)
```

Arguments

bp	an object of class biplot obtained from preceding function biplot().	
Pmat	a matrix containing the posterior probability for the classes	
dim.biplot	dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.	
e.vects	which eigenvectors (principal components) to extract, with default 1: dim. biplot.	
group.aes	vector of the same length as the number of rows in the data matrix for differentiated aesthetics for samples.	
axes	type of axes, defaults to "regression"	
col	colour of the classification regions	
opacity	opacity of classification regions	
borders	logical, indicating whether borders should be added to classification regions	

Value

an object of class biplot.

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classify

Classify samples into classes

Description

Classify samples into classes

Usage

```
classify(
  bp,
  classify.regions = TRUE,
  col = ez.col,
  opacity = 0.4,
  borders = FALSE
)
```

Arguments

bp an object of class biplot

classify.regions

a logical value indicating whether classifications regions should be shown in the

biplot, with default TRUE.

col the colours of the classification regions

opacity the opacity levels of the classification regions borders the border colours of the classification regions

Value

A list object called classify appended to the object of class biplot with the following elements:

table the confusion matrix resulting from the classification into classes.

rate the classification accuracy rate.

classify.regions

a logical value indicating whether classification regions are shown in the biplot.

aes a list of chosen aesthetics for the colours, opacity levels and border colours of

the classification regions.

Examples

```
biplot(iris[,1:4],classes = iris[,5]) |> CVA() |> axes(col="black") |>
  classify(col=c("red","blue","orange"),opacity=0.1) |> plot()
```

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Format aesthetics for the category level points

Description

This function allows the user to format the aesthetics for the category level points (CLPs).

Usage

```
CLPs (bp, which = 1:ncol(bp$Xcat), col = "black", cex = 0.6)
```

Arguments

bp	an object of class biplot.
which	a vector containing the columns or variables for which the CLPs should be displayed, with default ncol(Xcat).
col	the colour(s) for the CLPs, with default black.
cex	the character expansion(s) for the CLPs, with default 0.6.

Value

The object of class biplot will be appended with a list called CLP.aes containing the following elements A list with the following components is available:

```
which a vector containing the columns or variables for which the CLPs are displayed.

col the colour(s) of the CLPs.

cex the character expansion(s) of the plotting characters of the CLPs.
```

See Also

```
biplot, CA, AoD
```

Examples

20 CVA

CVA

Perform Canonical Variate Analysis (CVA)

Description

This function appends the biplot object with elements resulting from performing CVA.

Usage

Arguments

bp an object of class biplot obtained from preceding function biplot().

classes a vector of the same length as the number of rows in the data matrix with the

class indicator for the samples.

dim.biplot the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

e.vects the vector indicating which eigenvectors (canonical variates) should be plotted

in the biplot, with default 1:dim.biplot.

weightedCVA a character string indicating which type of CVA to perform. One of "weighted"

(default) for a weighted CVA to be performed (The centring matrix will be a diagonal matrix with the class sizes ($\mathbf{C} = \mathbf{N}$), "unweightedCent" for unweighted CVA to be performed (The centring matrix is the usual centring matrix ($\mathbf{C} = \mathbf{I}_G - G^{-1} \mathbf{1}_G \mathbf{1}_G'$)) or "unweightedI" for unweighted CVA to be performed while retaining the weighted centroid (The centring matrix is an indicator matrix

 $(\mathbf{C} = \mathbf{I}_G)$).

show.class.means

a logical value indicating whether to plot the class means on the biplot.

low.dim a character string indicating which method to use to construct additional dimen-

sion(s) if the dimension of the canonical space is smaller than dim.biplot. One of "sample.opt" (default) for maximising the sample predictivity of the individual samples in the biplot or "Bhattacharyya.dist" which is based on the decomposition of the Bhattacharyya distance into a component for the sample means and a component for the dissimilarity between the sample covariance

matrices.

Value

Object of class CVA with the following elements:

X the matrix of the centered and scaled numeric variables.

Xcat the data frame of the categorical variables.

raw. X the original data.

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classes the vector of category levels for the class variable. This is to be used for colour,

pch and cex specifications.

na.action the vector of observations that have been removed.
center a logical value indicating whether **X** is centered.
scaled a logical value indicating whether **X** is scaled.
means the vector of means for each numerical variable.

sd the vector of standard deviations for each numerical variable.

n the number of observations.
p the number of variables.

group.aes the vector of category levels for the grouping variable. This is to be used for

colour, pch and cex specifications.

g.names the descriptive names to be used for group labels.

g the number of groups.

Title the title of the biplot rendered.

Lmat the matrix for transformation to the canonical space.

Linv the inverse of L.

eigenvalues the vector of eigenvalues of the two-sided eigenvalue problem.

Z the matrix with each row containing the details of the points to be plotted (i.e.

coordinates).

ax.one.unit one unit in the positive direction of each biplot axis.

Gmat the indicator matrix defining membership of the classes.

Xmeans the matrix of the class means.

Zmeans the matrix of the class mean coordinates that are plotted in the biplot.

e.vects the vector indicating which canonical variates are plotted in the biplot.

Cmat the centring matrix based on different choices of weighting described in argu-

ments.

Bmat the between class sums of squares and cross products matrix.

Wmat the within class sums of squares and cross products matrix.

Mrr the matrix used for prediction from the canonical space (the inverse of $\mathbf{M}=$

LV).

Mr the first r dimensions of the solution to be plotted.

Nmat the matrix with the class sizes on the diagonal.

lambda.mat the matrix with the eigenvalues of $\mathbf{W}^{-1/2}\mathbf{B}\mathbf{W}^{-1/2}$ on the diagonal.

class.means a logical value indicating whether the class means should be plotted in the biplot.

dim. biplot the dimension of the biplot.

low.dim the method used to construct additional dimension(s).

See Also

biplot()

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Examples

```
biplot(iris[,1:4]) |> CVA(classes=iris[,5])
# create a CVA biplot
biplot(iris[,1:4]) |> CVA(classes=iris[,5]) |> plot()
```

CVA.biplot

Calculate elements for the CVA biplot

Description

This function performs calculations for the construction of a CVA biplot.

Usage

```
## $3 method for class 'biplot'
CVA(
   bp,
   classes = bp$classes,
   dim.biplot = c(2, 1, 3),
   e.vects = 1:ncol(bp$X),
   weightedCVA = "weighted",
   show.class.means = TRUE,
   low.dim = "sample.opt"
)
```

Arguments

bp an object of class biplot obtained from preceding function biplot().

classes a vector of the same length as the number of rows in the data matrix with the

class indicator for the samples.

dim.biplot the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

e.vects the vector indicating which eigenvectors (canonical variates) should be plotted

in the biplot, with default 1:dim.biplot.

weightedCVA a character string indicating which type of CVA to perform. One of "weighted"

(default) for a weighted CVA to be performed (The centring matrix will be a diagonal matrix with the class sizes ($\mathbf{C}=\mathbf{N}$), "unweightedCent" for unweighted CVA to be performed (The centring matrix is the usual centring matrix ($\mathbf{C}=\mathbf{I}_G-G^{-1}\mathbf{1}_G\mathbf{1}_G'$)) or "unweightedI" for unweighted CVA to be performed while retaining the weighted centroid (The centring matrix is an indicator matrix

 $(\mathbf{C} = \mathbf{I}_G)).$

show.class.means

a logical value indicating whether to plot the class means on the biplot.

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low.dim

a character string indicating which method to use to construct additional dimension(s) if the dimension of the canonical space is smaller than dim.biplot. One of "sample.opt" (default) for maximising the sample predictivity of the individual samples in the biplot or "Bhattacharyya.dist" which is based on the decomposition of the Bhattacharyya distance into a component for the sample means and a component for the dissimilarity between the sample covariance matrices.

Value

an object of class CVA, inherits from class biplot.

Examples

```
biplot(iris[,1:4]) |> CVA(classes=iris[,5])
```

CVAlowdim

Construct additional dimensions when the dimension of the canonical space is smaller than the dimension of the biplot

Description

This function is used to add dimensions to the CVA biplot when the dimension of the canonical space K is smaller than the dimension of the biplot (dim.biplot). This function is already used in the CVA calculations, and will therefore not have to be used in isolation.

Usage

```
CVAlowdim(bp, G, W, Mmat, low.dim, K, e.vects)
```

Arguments

bp an object of class biplot.

G the indicator matrix defining membership of the classes.
W the within class sums of squares and cross products matrix.

Mmat the eigenvector matrix from CVA.

low.dim a character string indicating which method to use to construct additional di-

mension(s) if the dimension of the canonical space is smaller than dim.biplot. One of "sample.opt" (default) for maximising the sample predictivity of the individual samples in the biplot or Bhattacharyya.dist which is based on the decomposition of the Bhattacharyya distance into a component for the sample means and a component for the dissimilarity between the sample covariance

matrices.

K the dimension of the canonical space.

e.vects the vector indicating which canonical variates are plotted in the biplot, with

default 1:dim.biplot

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Value

A list with three components:

Mr the first r dimensions of the solution to be plotted.

Mrr the matrix used for prediction from the canonical space.

Lmat the matrix for transformation to the canonical space.

density1D

Creates a kernel density in 1-dimension

Description

Creates a kernel density in 1-dimension

Usage

```
density1D(
   bp,
   which = NULL,
   h = "nrd0",
   kernel = "gaussian",
   col = ez.col,
   lwd = 1.5,
   legend.mar = c(2, 5, 0, 5)
)
```

Arguments

bp object of class biplot

which which group.
h bandwidth.

kernel character string giving the smoothing kernel to be used.

col colours to be used for each of the density curves.

lwd linewidth of density curve.legend.mar The margin line of the legend.

Value

An object of class biplot.

Examples

```
biplot (iris,classes=iris[,5]) |> CVA(dim=1) |> density1D() |> plot()
```

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density2D

Create a density in 2-dimensions

Description

Create a density in 2-dimensions

Usage

```
density2D(
   bp,
   which = NULL,
   contours = F,
   h = NULL,
   n = 100,
   col = c("green", "yellow", "red"),
   contour.col = "black",
   cuts = 50,
   cex = 0.6,
   tcl = -0.2,
   mgp = c(0, -0.25, 0),
   layout.heights = c(100, 10),
   legend.mar = c(2, 5, 0, 5)
)
```

Arguments

bp	object of class biplot
which	which group.
contours	logical indicating whether contours are added to the density plot
h	vector of bandwidths for x and y directions, see kde2d.
n	number of grid points in each direction. Can be scalar or a length-2 integer vector.
col	vector of colours to use to form a 'continuous' sequence of colours.
contour.col	colour of the contours.
cuts	number of colours in col.
cex	character expansion.
tcl	The length of tick marks as a fraction of the height of a line of text.
mgp	The margin line.
layout.heights	A vector of values for the heights of rows.
legend.mar	The margin line of the legend.

26 ellipses

Value

An object of class biplot.

Examples

```
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |>
  density2D(which=3,col=c("white","purple","cyan","blue")) |> plot()
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |>
  density2D(which=3,col=c("white","purple","cyan","blue"),contours = TRUE,
  contour.col = "grey") |> plot()
```

ellipses

Concentration ellipses (κ -ellipses)

Description

This function produces κ -ellipses, which is a useful geometrical description of the data points about the sample mean.

Usage

```
ellipses(bp, df=2, kappa = NULL, which = NULL,
alpha = 0.95, col = bp$sample$col[which], lty = 1, lwd = 1,
opacity = 0.25, trace = TRUE)
```

Arguments

bp	an object of class biplot.
df	degrees of freedom, with default 2.
kappa	value to construct κ -ellipse (the value of κ).
which	the selection of the group for ellipse construction.
alpha	size of α -bag, with default 0.95.
col	colour of ellipse. Multiple $\kappa\text{-ellipse}$ for one group will be displayed in the same colour.
lty	line type of ellipse. The same line type will be used per value of κ .
lwd	line width of ellipse. The same line width will be used per value of κ .
opacity	level of opacity, with default 0.25.
trace	logical, indicating progress of computation.

Value

A list with the following components is available:

conc.ellipses list of coordinates for the κ -ellipses for each group.

vector of colours for the κ -ellipses.
1ty vector of line types for the κ -ellipses.
1wd vector of line widths for the κ -ellipses.

alpha vector of α values.

References

Gower, J., Gardner-Lubbe, S. & Le Roux, N. (2011, ISBN: 978-0-470-01255-0) *Understanding Biplots*. Chichester, England: John Wiley & Sons Ltd.

Examples

```
biplot (iris[,1:4]) |> PCA(group.aes=iris[,5]) |> ellipses(kappa=2) |> plot()
```

extended.matching.coefficient

Extended matching coefficient

Description

Extended matching coefficient

Usage

```
extended.matching.coefficient(X)
```

Arguments

X a data frame containing the categorical variables used for computing the EMC

distance

Value

a dist object

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Examples

```
mtdf <- as.data.frame(mtcars)
mtdf$cyl <- factor(mtdf$cyl)
mtdf$vs <- factor(mtdf$vs)
mtdf$am <- factor(mtdf$am)
mtdf$gear <- factor(mtdf$gear)
mtdf$carb <- factor(mtdf$carb)
extended.matching.coefficient(mtdf[,8:11])</pre>
```

fit.measures

Compute measures of fit for the biplot.

Description

This function computes the measures of fit for the biplot. The biplot object is augmented with additional items, which can differ depending on the type of biplot. The measures provide information on the overall quality of fit and the adequacy of representation of variables.

Usage

```
fit.measures(bp)
```

Arguments

bp an object of class biplot.

Value

An object of class biplot. The object is augmented with additional items, depending on the type of biplot object.

quality the overall quality of fit.

adequacy the adequacy of representation of variables.

For an object of class PCA:

axis.predictivity

the fit measure of each individual axis.

sample.predictivity

the fit measure for each individual sample.

For an object of class CVA:

axis.predictivity

the fit measure of each individual axis.

class.predictivity

the fit measure for each class mean.

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```
within.class.axis.predictivity
```

the fit measure for each axis based on values expressed as deviations from their class means.

```
within.class.sample.predictivity
```

the fit measure for each sample expressed as deviation from its class mean.

Examples

```
out <- biplot (iris[,1:4]) |> PCA() |> fit.measures()
summary(out)
```

interpolate

Interpolate supplementary points and variables to add to the biplot

Description

This function adds supplementary points and variables to the plot from a new data set.

Usage

```
interpolate(bp, newdata = NULL, newvariable = NULL)
```

Arguments

bp an object of class biplot obtained from preceding function biplot().

newdata a new data set, similar in structure to the data set supplied to biplot() contain-

ing supplementary data points to be added onto the biplot.

newvariable a new data set, similar in structure to the data set supplied to biplot() contain-

ing supplementary variables to be added onto the biplot.

Value

The object of class biplot will be appended with the following elements:

Xnew.raw the new data.

Xnew the matrix of the centered and scaled new numeric variables of new data.

Xnew.cat the matrix of the categorical variables of new data.

Znew the matrix of the coordinates of the new data in the biplot.

Examples

```
biplot(data = iris[1:145,]) |> PCA() |> interpolate(newdata = iris[146:150,]) |> plot()
```

30 legend.type

legend.type	Format the legend for the biplot	
-------------	----------------------------------	--

Description

This function enables the user to format the legend and make a required selection to display.

Usage

```
legend.type(bp, samples = FALSE, means = FALSE, bags = FALSE,
                   ellipses=FALSE, regions=FALSE, new=FALSE, ...)
```

Arguments

bp	an object of class biplot.
samples	a logical value indicating whether a legend should be printed for samples, with default FALSE.
means	a logical value indicating whether a legend should be printed for means, with default FALSE.
bags	a logical value indicating whether a legend should be printed for bags, with default FALSE.
ellipses	a logical value indicating whether a legend should be printed for concentration ellipses, with default FALSE.
regions	a logical value indicating whether a legend should be printed for classification regions, with default FALSE.
new	a logical value indicating whether the legend should appear in a new window, with default FALSE.
	additional arguments to be sent to legend().

Value

A list with the following components is available:

samples a logical value indicating whether a legend for samples are provided. a logical value indicating whether a legend for class means are provided. means a logical value indicating whether a legend for α -bags are provided. bags ellipses a logical value indicating whether a legend for κ -ellipses are provided. a logical value indicating whether a legend for classification regions are proregions vided. new

a logical value indicating whether the legend appears on new plot.

Examples

```
biplot (iris[,1:4], Title="Test biplot") |> PCA(group.aes = iris[,5]) |>
   legend.type(samples=TRUE) |> plot()
```

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means	Format aesthetics for the class or group means	

Description

This function allows the user to format the aesthetics for the class means or group means.

Usage

```
means (bp, which = NULL, col = NULL, pch = 15, cex = 1, label = FALSE,
label.col = NULL, label.cex = 0.75, label.side = "bottom", label.offset = 0.5,
opacity = 1, shade.darker = TRUE)
```

Arguments

bp	an object of class biplot.
which	a vector containing the groups or classes for which the means should be displayed, with default bp\$g.
col	the colour(s) for the means, with default as the colour of the samples.
pch	the plotting character(s) for the means, with default 15.
cex	the character expansion(s) for the means, with default 1.
label	a logical value indicating whether the means should be labelled, with default TRUE.
label.col	a vector of the same length as which with label colours for the means, with default as the colour of the means.
label.cex	a vector of the same length as which with label text expansions for the means, with default 0.75 .
label.side	the side at which the label of the plotted mean point appears, with default bottom. Note that unlike the argument pos in text(), options are "bottom", "left", "top", "right" and not 1, 2, 3, 4.
label.offset	the offset of the label from the plotted mean point. See ?text for a detailed explanation of the argument offset.
opacity	transparency of means.
shade.darker	a logical value indicating whether the colour of the mean points should be made a shade darker than the default or specified colour, with default TRUE.

Details

The number of classes or groups (defined by group.aes) is indicated as g. If an argument is not of length g, recycling is used.

newaxes

Value

The object of class biplot will be appended with a list called means containing the following elements:

which a vector containing the groups or classes for which the means are displayed.

col the colour(s) of the means.

pch the plotting character(s) of the means.

cex the character expansion(s) of the plotting character(s) of the means.

label a logical value indicating whether means are labelled.

label.col the label colours of the means.

label.cex the label text expansions of the samples.label.side the side at which the label of the plotted mean point appears.

label.offset the offset of the label from the plotted mean point.

opacity the opacity level of the plotted points.

See Also

biplot()

Examples

newaxes

Format aesthetics for the supplementary (new) biplot axes

Description

This function allows the user to format the aesthetics for the supplementary (new) biplot axes.

Usage

```
newaxes(bp, X.new.names=bp$var.names, which = 1:bp$num.vars, col = "orange", lwd = 1,
lty = 1, label.dir = "Orthog", label.col = col, label.cex = 0.75, label.line = 0.1,
ticks = 5, tick.col = col, tick.size = 1, tick.label = TRUE, tick.label.col = tick.col,
tick.label.cex = 0.6, tick.label.side = "below", predict.col = col, predict.lwd = lwd,
predict.lty = lty, ax.names = X.new.names, orthogx = 0, orthogy = 0)
```

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Arguments

bp	an object of class biplot.
X.new.names	a vector of the new column names of bp to specify which axes should be labelled.
which	a vector containing the new columns or variables for which the axes should be displayed, with default 1:num.vars.
col	the colour(s) for the axes, with default grey(0.7). Alternatively, provide a vector of colours corresponding to X .names.
lwd	the line width(s) for the axes, with default 1.
lty	the line type(s) for the axes, with default 1.
label.dir	a character string indicating the placement of the axis titles to the side of the figure. One of "Orthog" for axis titles to appear orthogonal to the side of the figure (default), "Hor" for axis titles to appear horizontally or "Paral" for axis titles to appear parallel to the side of the figure.
label.col	the colour(s) for the axis labels, with default, col.
label.cex	the label expansion for the axis labels, with default 0.75.
label.line	the distance of the axis title from the side of the figure, with default 0.1 .
ticks	an integer-valued vector indicating the number of tickmarks for each axis, with default 5 for each axis.
tick.col	the colour(s) for the tick marks, with default col.
tick.size	a vector specifying the sizes of tick marks for each axis, with default 1 for each $\boldsymbol{\cdot}$
tick.label	a logical value indicating whether the axes should be labelled, with default TRUE.
tick.label.col	the colour(s) for the tick mark labels, with default tick.col.
tick.label.cex tick.label.side	the label expansion for the tick mark labels, with default 0.6.
	a character string indicating the position of the tick label. One of "below" for the label to appear below the tick mark (default) or "above" for the label to appear above the tick mark.
predict.col	the colour(s) for the predicted samples, with default col.
predict.lwd	the line width(s) for the predicted samples, with default lwd.
predict.lty	the line type(s) for the predicted samples, with default 1ty.
ax.names	a vector of size p containing user defined titles for the axes.
orthogx	a numeric vector of size p specifying the x-coordinate of the parallel transformation of each axis, with default \emptyset for each axis. This is only used when dim.biplot = 2.
orthogy	a numeric vector of size p specifying the y-coordinate of the parallel transformation of each axis, with default 0 for each axis. This is only used when $dim.biplot = 2$.

Value

The object of class biplot will be appended with a list called newaxes containing elements similar to that of axes.

newsamples

See Also

```
biplot, axes
```

Examples

```
biplot(data = iris[,1:2]) |> PCA() |> interpolate(newvariable = iris[3:4]) |>
newaxes(col="gold") |> plot()
```

newsamples

Format aesthetics for the supplementary (new) biplot samples

Description

This function allows formatting changes to new samples.

Usage

```
newsamples (bp, col = "darkorange1", pch = 1, cex = 1, label = FALSE,
label.name = NULL, label.col = NULL, label.cex = 0.75, label.side = "bottom",
label.offset = 0.5, connected = FALSE, connect.col = "black", connect.lty=1,
connect.lwd=1)
```

Arguments

bp	an object of class biplot.
col	the colour(s) for the new samples, with default darkorange1.
pch	the plotting character(s) for the new samples, with default 1.
cex	the character expansion(s) for the new samples, with default 1.
label	a logical value indicating whether new samples should be labelled or not, with default FALSE.
label.name	the label names for the new samples.
label.col	a vector of the same length as the number of new samples containing the colour(s) for the labels of the new samples, with default the colour of the sample points.
label.cex	the label text expansion(s) for the new samples, with default 0.75.
label.side	the side at which the label of the plotted point appears, with default bottom. Note that unlike the argument pos in text(), options are "bottom", "left", "top", "right" and not 1, 2, 3, 4.
label.offset	the offset of the label from the plotted point. See ?text for a detailed explanation of the argument offset.
connected	a logical value indicating whether samples are connected in order of rows of the data matrix, with default FALSE.
connect.col	the colour of the connecting line, with default black.
connect.lty	the line type of the connecting line, with default 1.
connect.lwd	the line width of the connecting line, with default 1.

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Value

The object of class biplot will be appended with a list called newsamples containing the following elements:

col the colour(s) of the new samples.

pch the plotting character(s) of the new samples.

the character expansion(s) of the plotting character(s) of the new samples.

label a logical value indicating whether new samples are labelled.

label.col the label colours of the new samples.

label.cex the label text expansions of the new samples.

label.side the side at which the label of the plotted point appears.

label.offset the offset of the label from the plotted point.

connected a logical value indicating whether new samples are connected.

connect.col the colour of the connecting line.

connect.lty the line type of the connecting line.

connect.lwd the line width of the connecting line.

See Also

biplot, samples

Examples

```
biplot(data = iris[1:145,]) |> PCA() |> samples(col = "grey") |>
interpolate(newdata = iris[146:150,]) |> newsamples(col = rainbow(6), pch=15) |> plot()
```

PCA

Perform Principal Components Analysis (PCA)

Description

This function appends the biplot object with elements resulting from performing PCA.

Usage

```
PCA(bp, dim.biplot = c(2, 1, 3), e.vects = 1:ncol(bp$X),
group.aes = NULL, show.class.means = FALSE, correlation.biplot = FALSE)
```

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Arguments

bp an object of class biplot obtained from preceding function biplot().

the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2. dim.biplot e.vects

the vector indicating which eigenvectors (principal components) should be plotted in the biplot, with default 1:dim.biplot.

group.aes

a vector of the same length as the number of rows in the data matrix for differ-

entiated aesthetics for samples.

show.class.means

a logical value indicating whether group means should be plotted in the biplot.

correlation.biplot

a logical value. If FALSE, the distances between sample points are optimally approximated in the biplot. If TRUE, the correlations between variables are optimally approximated by the cosine of the angles between axes. Default is FALSE.

Value

An object of class PCA with the following elements:

Χ the matrix of the centered and scaled numeric variables.

Xcat the data frame of the categorical variables.

the original data. raw.X

classes the vector of category levels for the class variable. This is to be used for colour,

pch and cex specifications.

the vector of observations that have been removed. na.action center a logical value indicating whether X is centered. scaled a logical value indicating whether X is scaled. means the vector of means for each numerical variable.

sd the vector of standard deviations for each numerical variable.

the number of observations. n the number of variables.

the vector of category levels for the grouping variable. This is to be used for group.aes

colour, pch and cex specification.

g.names the descriptive names to be used for group labels.

the number of groups. g

Title the title of the biplot rendered.

Ζ the matrix with each row containing the details of the points that are plotted (i.e.

coordinates).

the matrix for transformation to the principal components. Lmat

Linv the inverse of L.

eigenvalues the vector of eigenvalues of the covariance matrix of X. one unit in the positive direction of each biplot axis. ax.one.unit

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e.vects the vector indicating which principal components are plotted in the biplot. Vr the 1:dim.biplot columns of ${\bf V}$.

dim.biplot the dimension of the biplot.

class.means a logical value indicating whether group means are plotted in the biplot.

Zmeans the matrix of class mean coordinates that are plotted in the biplot.

References

Gabriel, K.R. (1971) The biplot graphic display of matrices with application to principal component analysis. *Biometrika*. 58(3):453–467.

See Also

```
biplot()
```

Examples

```
biplot(iris[,1:4]) |> PCA()
# create a PCA biplot
biplot(data = iris) |> PCA() |> plot()
```

PCA.biplot

Calculate elements for the PCA biplot

Description

This function performs calculations for the construction of a PCA biplot.

```
## S3 method for class 'biplot'
PCA(
   bp,
   dim.biplot = c(2, 1, 3),
   e.vects = 1:ncol(bp$X),
   group.aes = NULL,
   show.class.means = FALSE,
   correlation.biplot = FALSE
)
```

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Arguments

bp an object of class biplot obtained from preceding function biplot().

dim.biplot the dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2.

e.vects the vector indicating which eigenvectors (principal components) should be plot-

ted in the biplot, with default 1:dim.biplot.

group.aes a vector of the same length as the number of rows in the data matrix for differ-

entiated aesthetics for samples.

show.class.means

a logical value indicating whether group means should be plotted in the biplot.

correlation.biplot

a logical value. If FALSE, the distances between sample points are optimally approximated in the biplot. If TRUE, the correlations between variables are optimally approximated by the cosine of the angles between axes. Default is FALSE.

Value

an object of class PCA, inherits from class biplot.

Examples

```
biplot(iris[,1:4]) |> PCA()
# create a PCA biplot
biplot(data = iris) |> PCA() |> plot()
```

PC0

Principal Coordinate Analysis (PCO) biplot method

Description

Principal Coordinate Analysis (PCO) biplot method

Usage

Arguments

bp an object of class biplot obtained from preceding function biplot().

Dmat nxn matrix of Euclidean embeddable distances between samples

dist.func function to compute Euclidean embeddable distances between samples. The

default NULL computes Euclidean distance.

PCO.biplot 39

function to compute Euclidean embeddable distance between categorical varidist.func.cat ables for the samples. The default NULL computes the extended matching coefficient. dim.biplot dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2. e.vects which eigenvectors (canonical variates) to extract, with default 1:dim.biplot. e.vects group.aes vector of the same length as the number of rows in the data matrix for differentiated aesthetics for samples. show.class.means logical, indicating whether to plot the class means on the biplot. axes type of biplot axes, currently only regression axes are implemented more arguments to dist.func

Value

Object of class biplot

Examples

```
biplot(iris[,1:4]) |> PCO(dist.func = sqrtManhattan)
# create a CVA biplot
biplot(iris[,1:4]) |> PCO(dist.func = sqrtManhattan) |> plot()
```

PCO.biplot

PCO biplot

Description

Computes Principal Coordinate Analysis biplot

```
## S3 method for class 'biplot'
PCO(
   bp,
   Dmat = NULL,
   dist.func = NULL,
   dist.func.cat = NULL,
   dim.biplot = c(2, 1, 3),
   e.vects = NULL,
   group.aes = NULL,
   show.class.means = FALSE,
   axes = c("regression", "splines"),
   ...
)
```

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Arguments

bp an object of class biplot obtained from preceding function biplot(). Dmat nxn matrix of Euclidean embeddable distances between samples dist.func function to compute Euclidean embeddable distances between samples. The default NULL computes Euclidean distance. dist.func.cat function to compute Euclidean embeddable distance between categorical variables for the samples. The default NULL computes the extended matching coefficient. dim.biplot dimension of the biplot. Only values 1, 2 and 3 are accepted, with default 2. e.vects which eigenvectors (canonical variates) to extract, with default 1:dim.biplot. e.vects vector of the same length as the number of rows in the data matrix for differengroup.aes tiated aesthetics for samples. show.class.means logical, indicating whether to plot the class means on the biplot.

type of biplot axes, currently only regression axes are implemented axes

more arguments to dist.func

Value

an object of class biplot.

Examples

```
biplot(iris) |> PCO(dist.func=sqrtManhattan) |> plot()
```

plot.biplot

Generic Plotting function of objects of class biplot

Description

Generic Plotting function of objects of class biplot

```
## S3 method for class 'biplot'
plot(
  х,
  exp.factor = 1.2,
  axis.predictivity = NULL,
  sample.predictivity = NULL,
  zoom = FALSE,
  xlim = NULL,
 ylim = NULL,
)
```

prediction 41

Arguments

x An object of class biplot.

exp.factor a numeric value with default axes of the biplot. Larger values are specified

for zooming out with respect to sample points in the biplot display and smaller values are specified for zooming in with respect to sample points in the biplot

display.

axis.predictivity

either a logical or a numeric value between 0 and 1. If it is a numeric value, this value is used as threshold so that only axes with axis predictivity larger than the threshold is displayed. If axis.predictivity = TRUE, the axis colour is

'diluted' in proportion with the axis predictivity.

sample.predictivity

either a logical or a numeric value between 0 and 1. If it is a numeric value, this value is used as threshold so that only samples with sample predictivity larger than the threshold is displayed. If sample.predictivity = TRUE, the sample

size is shrinked in proportion with the sample predictivity.

zoom a logical value allowing the user to select an area to zoom into.

xlim the horizontal limits of the plot.

ylim the vertical limits of the plot.

... additional arguments.

Value

An object of class biplot.

Examples

```
biplot (iris[,1:4]) |> PCA() |> plot()
```

prediction

Predict samples to display on the biplot

Description

This function makes predictions of sample points, variables and means and displays them on the biplot.

```
prediction(bp, predict.samples = NULL, predict.means = NULL, which = 1:bp$p)
```

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Arguments

```
bp an object of class biplot obtained from preceding function biplot().

predict.samples
a vector specifying which samples to predict.

predict.means a vector specifying which group means to predict.

which a vector specifying which variable to do the prediction.
```

Value

A list object called predict appended to the object of class biplot with the following elements:

samples a vector of indices of samples which are being predicted.

predict.means a vector of group names of groups for which the means are being predicted.

which the vector of indices variables which are being predicted.

predict.mat the matrix of predicted samples.

predict.means.mat

the matrix of predicted group means.

Examples

```
biplot(data = iris[,1:4]) |> PCA(group.aes=iris[,5], show.class.means = TRUE) |>
prediction(141:145,1:3) |> plot()
```

print.biplot

Generic print function for objects of class biplot

Description

This function is used to print output when the biplot object is created.

Usage

```
## S3 method for class 'biplot' print(x, ...)
```

Arguments

x an object of class biplot.... additional arguments.

Value

This function will not produce a return value, it is called for side effects.

print.CA 43

Examples

```
out <- biplot (iris[,1:4]) |> PCA()
out
```

print.CA

Generic print function of objects of class CA

Description

Generic print function of objects of class CA

Usage

```
## S3 method for class 'CA'
print(x, ...)
```

Arguments

x an object of class CA ... additional arguments

Value

no return value, called for side effects

Examples

```
out <- biplot(HairEyeColor[,,1], center=FALSE) |> CA()
out
```

reflect

Reflect the biplot about a chosen axis

Description

This function provides the user with an option to reflect the biplot horizontally, vertically or diagonally.

```
reflect(bp, reflect.axis = c("FALSE", "x", "y", "xy"))
```

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Arguments

bp an object of class biplot

reflect.axis a character string indicating which axis about to reflect. One of FALSE (default),

"x" for reflection about the x-axis, "y" for reflection about the y-axis and "xy"

for reflection about both axes.

Value

An object of class biplot

Examples

```
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |> reflect("x") |> plot()
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |> reflect("y") |> plot()
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |> reflect("xy") |> plot()
```

regress

Regression biplot method

Description

Regression biplot method

Usage

Arguments

bp an object of class biplot obtained from preceding function biplot().

Z the matrix of coordinates of the samples

group.aes vector of the same length as the number of rows in the data matrix for differen-

tiated aesthetics for samples.

 $\verb|show.group.means||$

logical, indicating whether group means should be plotted in the biplot.

axes the type of axes to be fitted to the biplot. Options are 'regression' for linear

regression axes (default) and 'splines' for B-spline axes.

Value

Object of class biplot

Examples

```
biplot(iris[,1:4]) |> regress(Z=cmdscale(dist(iris[,1:4]))) |> plot()
```

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regress.biplot

Regression biplot

Description

Computes regression biplot axes

Usage

```
## $3 method for class 'biplot'
regress(
   bp,
   Z,
   group.aes = NULL,
   show.group.means = TRUE,
   axes = c("regression", "splines")
)
```

Arguments

bp an object of class biplot obtained from preceding function biplot().

Z the matrix of coordinates of the samples

group.aes vector of the same length as the number of rows in the data matrix for differen-

tiated aesthetics for samples.

show.group.means

logical, indicating whether group means should be plotted in the biplot.

axes

the type of axes to be fitted to the biplot. Options are 'regression' for linear

regression axes (default) and 'splines' for B-spline axes.

Value

an object of class biplot.

Examples

```
biplot(iris) |> regress(Z = cmdscale(dist(iris[,1:4]))) |> plot()
```

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rotate

Rotate the biplot a chosen amount of degrees

Description

This function provides the user with an option to rotate the biplot anti-clockwise or clockwise.

Usage

```
rotate(bp, rotate.degrees = 0)
```

Arguments

```
bp an object of class biplot

rotate.degrees a value specifying the degrees the biplot should be rotated, with default 0. A positive value results in anti-clockwise rotation and a negative value in clockwise rotation.
```

Value

An object of class biplot.

Examples

```
biplot(iris[,1:4],group.aes = iris[,5]) |> PCA() |> rotate(200) |> plot()
```

samples

Format aesthetics for the biplot samples

Description

This function allows the user to format the aesthetics for the samples.

```
samples (bp, which = 1:bp$g, col = ez.col, pch = 16, cex = 1,
label = FALSE, label.name = NULL, label.col=NULL, label.cex = 0.75,
label.side = "bottom", label.offset = 0.5,
connected=FALSE, connect.col = "black", connect.lty = 1,
connect.lwd = 1, opacity = 1)
```

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Arguments

bp	an object of class biplot.
which	a vector containing the groups or classes for which the samples should be displayed, with default bp\$g.
col	the colour(s) for the samples, with default blue.
pch	the plotting character(s) for the samples, with default 16.
cex	the character expansion(s) for the samples, with default 1.
label	a logical value indicating whether the samples should be labelled, with default FALSE. Alternatively, specify "ggrepel" for non-overlapping placement of labels.
label.name	a vector of the same length as which with label names for the samples, with default NULL. If NULL, the rownames(bp) are used. Alternatively, a custom vector of length n should be used.
label.col	a vector of the same length as which with label colours for the samples, with default as the same colour of the sample points.
label.cex	a vector of the same length as which with label text expansions for the samples, with default 0.75 .
label.side	the side at which the label of the plotted point appears, with default bottom. Note that unlike the argument pos in text(), options are "bottom", "left", "top", "right" and not 1, 2, 3, 4.
label.offset	the offset of the label from the plotted point. See ?text for a detailed explanation of the argument offset.
connected	a logical value indicating whether samples are connected in order of rows of the data matrix, with default FALSE.
connect.col	the colour of the connecting line, with default black.
connect.lty	the line type of the connecting line, with default 1.
connect.lwd	the line width of the connecting line, with default 1.
opacity	the opacity level of the plotted points, with default 1 for an opaque point.

Details

The arguments which, col, pch and cex are based on the specification of group. aes or classes. If no groups are specified, a single colour, plotting character and / or character expansion is expected. If g groups are specified, vectors of length g is expected, or values are recycled to length g.

The arguments label, label.cex, label.side and label.offset are based on the sample size n. A single value will be recycled n times or a vector of length n is expected.

Value

The object of class biplot will be appended with a list called samples containing the following elements:

which a vector containing the groups or classes for which the samples (and means) are displayed.

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col	the colour(s) of the samples.
pch	the plotting character(s) of the samples.
cex	the character expansion(s) of the plotting character(s) of the samples.
label	a logical value indicating whether samples are labelled.
label.name	the label names of the samples.
label.col	the label colours of the samples.
label.cex	the label text expansions of the samples.
label.side	the side at which the label of the plotted point appears
label.offset	the offset of the label from the plotted point.
connected	a logical value indicating whether samples are connected in order of the rows of the data matrix.
connect.col	the colour of the connecting line.
connect.lty	the line type of the connecting line.
connect.lwd	the line width of the connecting line.
opacity	the opacity level of the plotted points.

See Also

biplot()

Examples

sqrtManhattan Computes the square root of the Manhattan distance An example of a Euclidean embeddable distance metric

Description

Computes the square root of the Manhattan distance An example of a Euclidean embeddable distance metric

```
sqrtManhattan(X)
```

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Arguments

Χ

matrix of samples x variables for computation of samples x samples distance matrix

Value

a dist object

Examples

```
sqrtManhattan(iris[,1:4])
```

summary.biplot

Generic summary function for objects of class biplot

Description

This function is used to print summary output of the biplot. These summary outputs are related to measures of fit.

Usage

```
## S3 method for class 'biplot'
summary(
  object,
  adequacy = TRUE,
  axis.predictivity = TRUE,
  sample.predictivity = TRUE,
  class.predictivity = TRUE,
  within.class.axis.predictivity = TRUE,
  within.class.sample.predictivity = TRUE,
  ...
)
```

Arguments

object an object of class biplot.

adequacy a logical value indicating whether variable adequacies should be reported, with

default TRUE.

axis.predictivity

a logical value indicating whether axis predictivities should be reported, with

default TRUE.

sample.predictivity

a logical value indicating whether sample predictivities should be reported, with

default TRUE.

50 translate_axes

```
class.predictivity
```

a logical value indicating whether class predictivities should be reported, with default TRUE (only applicable to objects of class CVA).

within.class.axis.predictivity

a logical value indicating whether within class axis predictivity should be reported, with default TRUE (only applicable to objects of class CVA).

within.class.sample.predictivity

a logical value indicating whether within class sample predictivity should be reported, with default TRUE (only applicable to objects of class CVA).

... additional arguments.

Value

This function will not produce a return value, it is called for side effects.

Examples

```
out <- biplot (iris[,1:4]) |> PCA() |> fit.measures()
summary(out)
```

translate_axes

Translate biplot axes

Description

Automatically or manually translate the axes away from the center of the plot

Usage

```
translate_axes(bp, delta = 0, swop = FALSE, distances = NULL)
```

Arguments

bp An object of class biplot

delta numeric value indicating distance between axes

swop logical. Change the direction in which axes are translated

distances numeric vector of distances. Used to manually parallel translate the axes.

Details

This function uses the same algorithm implemented in TDAbiplot in the bipl5 package. It translates the axes out of the center of the plot. Correlated axes generally gets translated in the same direction.

This function calculates the orthogy and orthogy paramaters in axes()

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Value

An object of class biplot with the translated distances appended under bp\$axes

Examples

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