# Package 'corregp' 

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Description A collection of tools for correspondence regression, i.e. the correspondence analysis of the crosstabulation of a categorical variable Y in function of another one X , where X can in turn be made up of the combination of various categorical variables. Consequently, correspondence regression can be used to analyze the effects for a polytomous or multinomial outcome variable.
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## Description

This package provides functions and methods for performing correspondence regression, i.e. the correspondence analysis of the crosstabulation of a categorical variable Y in function of another one X , where X can in turn be made up of the combination of various categorical variables. Consequently, correspondence regression can be used to analyze the effects for a polytomous or multinomial outcome variable. The central function in the package is corregp, which enables methods for printing, summarizing and plotting the output. Additionally, there are functions for computing confidence intervals, ellipses or 3D ellipsoids (by means of bootstrapping).

## Contents

This package consists of the following datasets, functions, generics and methods (some internal functions are no longer exported in version 2):

## Datasets:

- HairEye Hair and eye color of statistics students (data frame).
- COMURE The use of linguistic variants in translations vs. non-translations and in six different registers.
- AVT The use of linguistic variants in audio-visual translation (subtitles).
- TSS The use of inflected or uninflected determiners in vernacular Belgian Dutch.


## Functions:

- ci A helper function to compute confidence intervals on the basis of a numeric vector.
- corregp The basic function to perform correspondence regression. Typically, one starts here, and then one uses print, summary, anova, screeplot or plot methods.
- corregplicate A function for repeated correspondence regressions with bootstrapping in order to handle large data sets.


## Generics:

- cint Compute confidence intervals.
- cell Compute confidence ellipses.
- cell3d Compute 3D confidence ellipsoids.
- ciplot Plot confidence intervals.
- pcplot Plot parallel coordinates.
- agplot Plot an association graph.
- plotag Plot an association graph.


## Methods:

- print. corregp Print the output of a correspondence regression.
- summary. corregp Give a summary of a correspondence regression.
- print. summary. corregp Print the summary of a correspondence regression.
- screeplot.corregp Make a scree plot on the basis of the output of a correspondence regression.
- anova. corregp Give an anova table on the basis of a correspondence regression.
- print. anova. corregp Print an anova table on the basis of a correspondence regression.
- coef. corregp Give the coefficients on the basis of a correspondence regression.
- coefficients.corregp Give the coefficients on the basis of a correspondence regression.
- fitted. corregp Give the fitted values on the basis of a correspondence regression.
- fitted.values.corregp Give the fitted values on the basis of a correspondence regression.
- residuals.corregp Give the residuals on the basis of a correspondence regression.
- resid. corregp Give the residuals on the basis of a correspondence regression.
- cint.corregp Compute confidence intervals on the basis of the output of a correspondence regression. Typically, this function is not so much used directly as it is called by a ciplot. corregp command.
- ciplot.corregp Plot confidence intervals on the basis of the output of a correspondence regression.
- pcplot.corregp Make a parallel coordinate plot on the basis of the output of a correspondence regression.
- cell.corregp Compute confidence ellipses on the basis of the output of a correspondence regression. Typically, this function is not so much used directly as it is called by a plot.corregp command.
- plot.corregp Plot the output (and the optional confidence ellipses) of a correspondence regression.
- cell3d. corregp Compute 3D confidence ellipsoids on the basis of a correspondence regression. Typically, this function is not so much used directly as it is called by a plot3d.corregp command.
- plot3d.corregp Plot the 3D output (and the optional confidence ellipsoids) of a correspondence regression.
- agplot.corregp Make an association graph on the basis of the output of a correspondence regression.
- plotag. corregp Make an association graph on the basis of the output of a correspondence regression.


## Future prospects

- Specify a predict method for a.o. supplementary points.
- Specify a plot method for an anova table.
- Enable scale transformations for all plots (and corresponding confidence regions).
- Provide the possibility for so-called "calibration lines".


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## Acknowledgements

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```
agplot.corregp Plotting an Association Graph for Correspondence Regression
```


## Description

Function to make an association graph of the (significant) coordinate scores in correspondence regression.

## Usage

```
## S3 method for class 'corregp'
agplot(x, axes = NULL, ysub = NULL, xsub = NULL,
    sort = NULL, na.rm = FALSE, col = "black", cex = par("cex"),
    font = par("font"), family = par("family"), lwd = par("lwd"),
    lty = par("lty"), ycol = col, xcol = col, ncol = c("white",
    "lightgray"), nwid = lwd, lcol = col, lwid = lwd, pcol = lcol,
    ppos = NULL, ptyp = "simple", zoom = 1, hshft = 0, vshft = 0,
    main = NULL, cl = 0.95, nq = TRUE, digits = 2, ...)
    ## S3 method for class 'corregp'
    plotag(x, axes = NULL, ysub = NULL, xsub = NULL,
    sort = NULL, na.rm = FALSE, col = "black", cex = par("cex"),
    font = par("font"), family = par("family"), lwd = par("lwd"),
```

```
    lty = par("lty"), ycol = col, xcol = col, ncol = c("white",
    "lightgray"), nwid = lwd, lcol = col, lwid = lwd, pcol = lcol,
    ppos = NULL, ptyp = "simple", zoom = 1, hshft = 0, vshft = 0,
    main = NULL, cl = 0.95, nq = TRUE, digits = 2, ...)
agplot(x, ...)
plotag(x, ...)
```


## Arguments

X
axes
ysub Vector of indices to select a subset of the Y levels.
xsub Vector of indices to select a subset of the X levels. Can also be "all" or "both" (or abbreviations).
sort Vector of axes for which to sort the coordinate scores. The default (NULL) plots all levels in the order in which they appear in the correspondence regression $x$.
na.rm Logical specifying whether to omit NA coordinates from the plot. Defaults to FALSE.
col Color of the association graph: either numeric or see colors.
cex Character expansion factor: a number to specify the size of the text labels.
font Font of the text labels (levels): 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. Defaults to 1 .
family Font family of the text labels (levels): can be "serif", "sans", "mono" or one of the Hershey fonts.
lwd Line width of the association graph: a number to specify the line width.
lty Line type of the association graph (i.e. linking edges): 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1.
ycol Color of the levels in Y: either numeric or see colors.
xcol Color of the levels in X: either numeric or see colors.
ncol Fill color of the nodes: either numeric or see colors. Defaults to c ("white", "lightgray"): the first value is for the nodes of the axes and the second value is for the nodes of the $X$ and $Y$ levels.
nwid Line width of the nodes: a number to specify the line width. If a vector of two values is specified, then the first width is for the nodes of the axes and the second width is for the nodes of the X and Y levels.
lcol
The output of a call to corregp (i.e. an object of class "corregp").
The axes for which to plot the association graph: a vector of indices. Defaults to all the axes. of the Wershey fonts.

Color of the links (edges): either numeric or see colors. If a vector of two values is specified, then the first color is for the scores $>0$ and the second color is for the scores $<0$.

| lwid | Line width of the links (edges): a number to specify the line width. If a vector of two values is specified, then the first width is for the scores $>0$ and the second width is for the scores $<0$. |
| :---: | :---: |
| pcol | Color of the pointer (arrow head): either numeric or see colors. If a vector of two values is specified, then the first color is for the scores $>0$ and the second color is for the scores $<0$. |
| ppos | Relative position of the pointer (arrow head): a vector of values between 0 and 1 for each axis. |
| ptyp | Type of of the pointer (arrow head): can be "simple", "curved", "triangle", "circle", "ellipse" or "T". Defaults to "simple". |
| zoom | Zoom factor of the association graph. Defaults to 1. |
| hshft | Horizontal shift of the association graph. Defaults to 0 . |
| vshft | Vertical shift of the association graph. Defaults to 0 . |
| main | The main title of the association graph. |
| cl | The confidence level for the confidence intervals. Defaults to 0.95. |
| nq | Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence interval. Defaults to TRUE. If FALSE, then the confidence interval is computed directly with the quantile function. |
| digits | Integer specifying the number of decimals for the scores as labels of the links (edges). Defauls to 2. |
|  | Further arguments passed to or from other methods. |

## Details

Association graphs (of a corregp output) in the corregp package make use of various functionalities of the package diagram.

## Value

A plot window containing the association graph.

## See Also

```
corregp, cint.corregp, pcplot.corregp, plot3d.corregp.
```


## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
agplot(haireye.crg, axes = 1:2, xsub = c("Hair", "Sex"))
plotag(haireye.crg, axes = 1:2, xsub = c("Hair", "Sex"))
```


## Description

Method to construct an ANOVA table for correspondence regression, i.e. a table with the Chisquared deviation for each term in the formula of the corregp call (or of each individual level in X in case xep $=$ FALSE).

## Usage

\#\# S3 method for class 'corregp'
anova(object, $n f=$ NULL, $c l=0.95, n q=$ TRUE,... )

## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
nf The number of dimensions to be retained in the reduced space. Defaults to all dimensions (no reduction).
cl The confidence level for the confidence intervals. Defaults to 0.95.
nq Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence interval. Defaults to TRUE. If FALSE, then the confidence interval is computed directly with the quantile function.
.. Further arguments passed to or from other methods.

## Details

If object was made with bootstrap replications, then anova.corregp will automatically compute confidence intervals for the Chi-squared deviations by means of the ci function.

## Value

A matrix with the Chi-squared deviations for all the terms in the formula of object, based on the selected number of dimensions. If object was made with the argument xep $=$ FALSE, then the output contains the Chi-squared deviation for every individual level in X .

## See Also

print. anova.corregp, ci, summary.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
anova(haireye.crg, nf = 2)
```


## Description

This data set was a follow-up study to the COMURE project and was conducted at the Department of Translation, Interpreting and Communication of Ghent University between 2014 and 2018.

## Format

A data frame with 3302 rows and 7 variables.

- Variant The linguistic variant used in a set of alternatives (27 levels).
- Variable The linguistic variable specifying a set of alternatives (13 levels).
- Variety The dichotomization of Variant into standard and non-standard.
- Speaker The role of the speaker in the data (2 levels).
- Language The language (and source language) of the data (3 levels).
- Language2 The same as Language but with the observations of level intra.nl set to NA.
- Genre The genre or register of the data (2 levels).


## Source

Prieels, L., I. Delaere, K. Plevoets and G. De Sutter (2015) A corpus-based multivariate analysis of linguistic norm-adherence in audiovisual and written translation. Across Languages and Cultures 16 (2), 209-231.

## Examples

```
data(AVT)
# The execution of corregp may be slow, due to bootstrapping:
avt.crg <- corregp(Variant ~ Speaker * Language * Genre, data = AVT, part = "Variable", b = 3000)
avt.crg
summary(avt.crg, parm = "b", add_ci = TRUE)
screeplot(avt.crg, add_ci = TRUE)
anova(avt.crg, nf = 2)
avt.col <- ifelse( xtabs(~ Variant + Variety, data = AVT)[, "Standard"] > 0, "blue", "red")
plot(avt.crg, x_ell = TRUE, xsub = c("Speaker", "Language", "Genre"), col_btm = avt.col,
    col_top = "black")
```

```
cell.corregp Confidence Ellipses for Correspondence Regression
```


## Description

Method to compute confidence ellipses for coordinates in correspondence regression.

## Usage

```
## S3 method for class 'corregp'
    cell(object, parm = "x", axes = 1:2, cl = 0.95,
        np = 100, ...)
    cell(object, ...)
```


## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to compute the confidence ellipses. Can be either " y ", " $x$ ", or any vector of term names in $X$, level names in $X$ or level names in $Y$. Defaults to " $x$ ".
axes The axes for which to compute the confidence ellipses: a vector of two values. Defaults to the first two axes.
cl The confidence level for the confidence ellipses. Defaults to 0.95.
$\mathrm{np} \quad$ The number of points to represent the confidence ellipses. Defaults to 100.
... Further arguments passed to or from other methods.

## Details

cell (of a corregp output) makes use of ellipse from the package ellipse.
Typically, cell is not so much used directly as it is called by a plot. corregp command.

## Value

A list containing np points for each confidence ellipse of interest.

## See Also

```
plot.corregp.
```


## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
cell(haireye.crg, parm = "y")
cell(haireye.crg, parm = c("Hair", "Sex"))
```

```
cell3d.corregp 3D Confidence Ellipsoids for Correspondence Regression
```


## Description

Method to compute 3D confidence ellipsoids for coordinates in correspondence regression.

## Usage

\#\# S3 method for class 'corregp'
cell3d(object, parm = "x", axes $=1: 3$, cl $=0.95, \ldots$ )
cell3d(object, ...)

## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to compute the confidence ellipsoids. Can be either " $y$ ", " $x$ ", or any vector of term names in $X$, level names in $X$ or level names in Y. Defaults to " $x$ ".
axes The axes for which to compute the confidence ellipsoids: a vector of three values. Defaults to the first three axes.
cl The confidence level for the confidence ellipsoids. Defaults to 0.95.
... Further arguments passed to or from other methods.

## Details

cell3d (of a corregp output) makes use of ellipse3d from the package rgl.
Typically, cell3d is not so much used directly as it is called by a plot3d.corregp command.

## Value

A list containing coordinate points for each confidence ellipsoid of interest.

## See Also

plot3d.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
cell3d(haireye.crg, parm = "y")
cell3d(haireye.crg, parm = c("Hair", "Sex"))
```


## Description

This is the basic function for computing a confidence interval on the basis of a sample of data values.

## Usage

ci(x, cl = 0.95, nq = TRUE)

## Arguments

x
cl The confidence level for the confidence interval. Defaults to 0.95.
nq Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence interval. Defaults to TRUE. If FALSE, then the confidence interval is computed directly with the quantile function.

## Value

A vector with two components Lower and Upper giving the lower and upper confidence limits respectively.

## See Also

ciplot. corregp, anova.corregp, agplot.corregp, confint.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
ci(haireye.crg$conf$eigen[, 1])
ci(haireye.crg$conf$eigen[, 2])
```


## Description

Method to compute confidence intervals for coordinates in correspondence regression.

```
Usage
    ## S3 method for class 'corregp'
    cint(object, parm = "x", axis, cl = 0.95, nq = TRUE,
        ...)
    cint(object, ...)
```


## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to compute the confidence intervals. Can be either " y ", " $x$ ", or any vector of term names in $X$, level names in $X$ or level names in $Y$. Defaults to " $x$ ".
axis The axis for which to compute the confidence intervals.
cl The confidence level for the confidence interval. Defaults to 0.95.
nq Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence intervals. Defaults to TRUE. If FALSE, then the confidence intervals are computed directly with the quantile function.
.. Further arguments passed to or from other methods.

## Details

cint (of a corregp output) makes use of ci.
Typically, cint is not so much used directly as it is called by a ciplot. corregp command.

## Value

A matrix with Lower and Upper confidence limits for the coordinates of interest.

## See Also

ci, ciplot. corregp, agplot.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
cint(haireye.crg, parm = "y", axis = 1)
cint(haireye.crg, parm = c("Hair", "Sex"), axis = 1)
```

ciplot.corregp Plotting Confidence Intervals for Correspondence Regression

## Description

Method to plot confidence intervals for coordinates in correspondence regression.

## Usage

```
## S3 method for class 'corregp'
ciplot(x, parm = "x", axis, cl = 0.95, nq = TRUE,
    horiz = FALSE, na.rm = FALSE, type = "p", col = "darkgrey",
        cex = par("cex"), font = par("font"), family = par("family"),
        alim = NULL, adir = 1, ecol = "darkgrey", ewid = par("lwd"),
        etyp = par("lty"), psym = 16, pcol = par("col"), pcex = cex,
        pbgc = par("bg"), lwd = ewid, lty = etyp, sfrac = 0.01, gap = 0,
        main = NULL, sub = NULL, ...)
    ciplot(x, ...)
```


## Arguments

horiz Logical specifying whether the confidence intervals should be plotted horizon-

X
parm
axis
cl
nq
na.rm
type

The output of a call to corregp (i.e. an object of class "corregp").
The parameter for which to plot the confidence intervals. Can be either " $y$ ", " $x$ ", or any vector of term names in $X$, level names in $X$ or level names in $Y$. Defaults to " $x$ ".
The axis for which to plot the confidence intervals.
The confidence level for the confidence intervals. Defaults to 0.95.
Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence intervals. Defaults to TRUE. If FALSE, then the confidence intervals are computed directly with the quantile function. tally or not. Defaults to FALSE.
Logical specifying whether to omit NA coordinates from the plot. Defaults to FALSE.
The type of plot: see plot. default. For correspondence regression, there is an additional option "labs" which plots the text labels at the centers of the confidence intervals. Defaults to " p ".

| col | Color of the text labels: either numeric or see colors. |
| :---: | :---: |
| cex | Character expansion factor: a number to specify the size of the text labels. |
| font | Font of the text labels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. Defaults to 1 . |
| family | Font family of the text labels: can be "serif", "sans", "mono" or one of the Hershey fonts. |
| alim | Vector of two values specifying the lower and upper limit between which to plot the axis. |
| adir | Reading direction of the text labels on the (horizontal) axis: either a numeric value between 0 and 3 (see the las argument in the graphical parameters par) or a character value matching either "horizontal" or "vertical". Defaults to 1 (horizontal). |
| ecol | Color of the error bars: either numeric or see colors. |
| ewid | Width of the error bars: a number to specify the line width. |
| etyp | Line type of the error bars: 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1. |
| psym | The symbol (or "plotting character") to use for the centers of the confidence intervals. |
| pcol | Color of the center symbol: either numeric or see colors. |
| pcex | Character expansion factor of the center symbol. |
| pbgc | Background color of the center symbol: either numeric or see colors. |
| lwd | Width of all lines except for the error bars, e.g. the connecting lines: a number to specify the line width. |
| 1 ty | Line type of all lines except for the error bars, e.g. the connecting lines: 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1. |
| sfrac | Width of "crossbar" at the end of error bar as a fraction of the x plotting region. Defaults to 0.01 . |
| gap | Space left between the center of the error bar and the lines marking the error bar in units of the height (width) of the letter "O". Defaults to 0 . |
| main | The main title of the plot. |
| sub | The subtitle of the plot. |
|  | Further arguments passed to or from other methods. |

## Details

ciplot (of a corregp output) makes use of plotCI from the package gplots.

## Value

A plot window containing the confidence intervals.

## See Also

ci, plotCI.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
ciplot(haireye.crg, parm = "y", axis = 1)
ciplot(haireye.crg, parm = c("Hair", "Sex"), axis = 1)
```

coef.corregp Extracting Coefficients from Correspondence Regression

## Description

Method to extract the coefficients (i.e. scores) of a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
coef(object, parm = "x", axes = NULL, ...)
    ## S3 method for class 'corregp'
    coefficients(object, parm = "x", axes = NULL, ...)
```


## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to extract the coefficients. Can be either " $y$ ", " $x$ ", or any vector of term names in X , level names in X or level names in Y . Defaults to " $x$ ".
axes The axes for which to extract the coefficients: a vector of indices. Defaults to all the axes.
... Further arguments passed to or from other methods.

## Details

The coefficients in correspondence regression are the same as the coordinate scores.

## Value

A matrix or vector with coefficients (i.e. scores) for the parameters and axes of interest.

## See Also

fitted.corregp, residuals.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
coef(haireye.crg, parm = c("Hair", "Sex"), axes = 1:2)
coefficients(haireye.crg, parm = c("Hair", "Sex"), axes = 1:2)
```


## COMURE

The Use of Linguistic Variants in Translations vs. Non-translations and in Six Different Registers

## Description

This data set was a case study in the COMURE project ("corpus-based, multivariate research of register variation in translated and non-translated Belgian Dutch") which was conducted at the Department of Translation, Interpreting and Communication of Ghent University between 2010 and 2014.

## Format

A data frame with 3762 rows and 5 variables.

- Variant The linguistic variant used in a set of alternatives ( 27 levels).
- Variable The linguistic variable specifying a set of alternatives (13 levels).
- Variety The dichotomization of Variant into standard and non-standard.
- Register The register or "Text type" of the data (6 levels).
- Language The language (and source language) of the data (3 levels).


## Source

Delaere, I., G. De Sutter and K. Plevoets (2012) Is translated language more standardized than non-translated language? Target 24 (2), 203-224.

## Examples

```
data(COMURE)
# The execution of corregp may be slow, due to bootstrapping:
comure.crg <- corregp(Variant ~ Register * Language, data = COMURE, part = "Variable", b = 3000)
comure.crg
summary(comure.crg, parm = "b", add_ci = TRUE)
screeplot(comure.crg, add_ci = TRUE)
anova(comure.crg, nf = 2)
comure.col <- ifelse( xtabs(~ Variant + Variety, data = COMURE)[, "Standard"] > 0, "blue", "red")
plot(comure.crg, x_ell = TRUE, xsub = c("Register", "Language"), col_btm = comure.col,
    col_top = "black")
```

```
confGet Getting conf Components from corregp Objects
```


## Description

Internal function for retrieving the conf component(s) in a corregp object.

## Usage

confGet(crg, parm)

## Arguments

crg The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to retrieve the conf components. Can be either " $y$ ", " $x$ ", or any vector of term names in $X$, level names in $X$ or level names in $Y$.

## Details

confGet is an internal function to be called by cint. corregp, cell. corregp or cell3d.corregp, but not by users.

## Value

A list of components selected with parm.
corregp Correspondence Regression

## Description

This is the basic function for correspondence regression, i.e. the correspondence analysis of a contingency table formed by the categorical variables $Y$ and $X$, where $X$ can be in turn made up of the combinations of various categorical variables.

## Usage

corregp(formula, data, part $=$ NULL, $b=0$, xep $=$ TRUE, std $=$ FALSE, rel = TRUE, phi = FALSE, chr = ".", b_scheme = "multinomial")

## Arguments

| formula | A formula specification of which factors to cross with each other. The left-hand <br> (y) side must be a single factor. The right-hand side (x) can involve all the usual <br> specifications of interactions and/or nested analyses. |
| :--- | :--- |
| data | The data frame containing the variables specified in the formula. <br> Character vector specifying the names of conditional factors (e.g. a factor par- <br> tioning the levels of the left-hand side y into groups). This argument is relevant <br> for analyses in which one wants to remove between-item variation. <br> Number of the bootstrap replications (simulations). If 0 (i.e. the default), then <br> the analysis is exploratory. |
| b | Logical specifying whether to output the separate terms in the right-hand side <br> (x) as components in a list. If FALSE, then all x output is collected in a matrix. |
| xep | Logical specifying whether to output the standardized coordinates. Defaults to <br> FALSE. |
| rel | Logical specifying whether to divide the coordinates by the sqrt of their totals, <br> so that one obtains coordinates for the relative frequencies (as is customary in <br> correspondence analysis). Defaults to TRUE. |
| phi | Logical specifying whether to compute the output on the scale of the Chi-squared <br> value of the contingency table or of the Phi-squared value (which is Chi-squared <br> divided by $N$ ). Reminiscent of corresp in package MASS, defaults to FALSE. |
| chr | Character specifying the separator string for constructing the interaction terms. <br> b_scheme$\quad$Character specifying the sampling scheme for bootstrapping. Must match either <br> "multinomial" (the default) or "product-multinomial". |

## Details

Correspondence regression rests on the idea, described by Gilula and Haberman (1988), of using a correspondence analysis to model a polytomous or multinomial (i.e. 'multi-category') response variable $(\mathrm{Y})$ in terms of other (possibly interacting) factors ( X ) (see also 3.2 in Van der Heijden et al. 1989). These are specified in the argument formula, which can be constructed in all the usual ways of specifying a model formula: e.g.

- $\mathrm{Y} \sim \mathrm{X} 1$ + X2 + X1 : X2 or Y ~ X1 * X2
- $Y \sim(X 1+X 2+X 3)^{\wedge} 2$
- Y ~ X1 * X2 * X3 - X1 : X2 : X3
- ...

Correspondence regression then crosstabulates the $Y$ factor with all the combinations in $X$, thus producing a typical contingency table, on which a simple correspondence analysis is performed (see Greenacre 2017: 121-128 for the outline of this approach). The more general effects in $X$ are obtained by aggregating the combinations.
Correspondence regression also allows for inferential validation of the effects, which is done by means of the bootstrap (in fact, Monte Carlo simulation). Setting the argument $b$ to a number $>0, \mathrm{~b}$ replicates of the contingency table are generated with multinomial sampling. From these, $b$ new values are derived for the coordinates in both $Y$ and $X$ as well as for the eigenvalues (also
called the "principal inertias"). On the basis of the replicate/simulated values, confidence intervals, ellipses or ellipsoids can be computed. CAUTION: bootstrapping/simulation is computationally quite intensive, so it can take a while to reach results, especially with a large $b$.
The argument parm can be used when one wants to perform a correspondence regression of $Y$ onto $X$ conditional on other factors. These conditioning factors are therefore equivalent to random factors, and corregp always conditions on the joint variable of all the specified factors. One such use of conditioning factors is a so-called lectometric analysis in linguistics, where the levels of $Y$ are grouped/partitioned/nested into clusters and one wants to exclude the heterogeneity between the clusters.

## Value

An object of class "corregp", i.e. a list with components:
eigen A vector of eigenvalues of the correpondence regression.
$y \quad$ The coordinates (matrix) of the Y levels.
$x \quad$ The coordinates of the X levels. If xep is TRUE, then this is a list with a component for each term name.
freq A list of the frequencies of every Y and X level.
conf If $b>0$. A list of bootstrap replicates for the eigenvalues, the coordinates of Y levels, the coordinates of X levels and the frequencies of both the Y levels and the X levels.
aux A list of auxiliary information (such as the U and V matrices of the SVD, the specified values for all the arguments) to be passed to other functions and methods.

## References

Gilula, Z. and S.J. Haberman (1988) The analysis of multivariate contingency tables by restricted canonical and restricted association models. Journal of the American Statistical Association 83 (403), 760-771.

Greenacre, M. (2017) Correspondence analysis in practice, Third edition. Boca Raton: Chapman and Hall/CRC.
Van der Heijden, P.G.M., A. de Falguerolles and J. de Leeuw (1989) A combined approach to contingency table analysis using correspondence analysis and log-linear analysis. Applied Statistics 38 (2), 249-292.

## See Also

print.corregp, summary.corregp, screeplot.corregp, anova.corregp, plot.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
haireye.crg
```


## corregplicate Repeated Correspondence Regression

## Description

A function for repeated correspondence regressions with bootstrapping in order to handle large data sets. This is essentially a wrapper replicate $(n=r$, expr $=\operatorname{corregp}(\ldots)$, simplify = FALSE), so it may dissappear in the future.

## Usage

corregplicate(formula, data, part $=$ NULL, $b=100, r=10$, xep $=$ TRUE, std = FALSE, rel = TRUE, phi = FALSE, chr = ".", b_scheme = "multinomial")

## Arguments

| formula | A formula specification of which factors to cross with each other. The left-hand <br> (y) side must be a single factor. The right-hand side (x) can involve all the usual <br> specifications of interactions and/or nested analyses. |
| :--- | :--- |
| data | The data frame containing the variables specified in the formula. <br> Character vector specifying the names of conditional factors (e.g. a factor par- <br> tioning the levels of the left-hand side y into groups). This argument is relevant <br> for analyses in which one wants to remove between-item variation. <br> number of the bootstrap replications (simulations). |
| b | Number of repeated calls to corregp. |
| rep | Logical specifying whether to output the separate terms in the right-hand side <br> (x) as components in a list. If FALSE, then all x output is collected in a matrix. |
| std | Logical specifying whether to output the standardized coordinates. Defaults to <br> FALSE. |
| rel | Logical specifying whether to divide the coordinates by the sqrt of their totals, <br> so that one obtains coordinates for the relative frequencies (as is customary in <br> correspondence analysis). Defaults to TRUE. |
| phi | Logical specifying whether to compute the output on the scale of the Chi-squared <br> value of the contingency table or of the Phi-squared value (which is Chi-squared <br> divided by $N$ ). Reminiscent of corresp in package MASS, defaults to FALSE. |
| chr | Character specifying the separator string for constructing the interaction terms. <br> b_scheme |
| Character specifying the sampling scheme for bootstrapping. Must match either |  |
| "multinomial" (the default) or "product-multinomial". |  |

## Value

An object of class "corregp" in which the bootstrap replications of all the repeated calls to corregp are put together.

## See Also

corregp.
fitted.corregp Extracting Fitted Values from Correspondence Regression

## Description

Method to extract the fitted values of a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
fitted(object, parm = "all", nf = NULL, ...)
## S3 method for class 'corregp'
fitted.values(object, parm = "all", nf = NULL, ...)
```


## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to extract the fitted values. Can be "all", "both" (or abbreviations), " $y$ " or " $x$ " for the fitted values of every cell in the data, but it can also be any vector of term names in X or level names in X. Defaults to "all".
$\mathrm{nf} \quad$ The number of dimensions to be retained in the reduced space. Defaults to all dimensions (no reduction).
... Further arguments passed to or from other methods.

## Details

If all dimensions are retained, then the fitted values will only be equal to the observed counts if no conditioning factors were specified with the argument "part" in the corregp call. This is because the associations with the conditioning factors (in "part") are not taken into account.

## Value

A matrix or vector with the fitted values for the parameters of interest, based on the selected number of dimensions.

## See Also

coef.corregp, residuals.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
fitted(haireye.crg, parm = c("Hair", "Sex"), nf = 2)
fitted.values(haireye.crg, parm = c("Hair", "Sex"), nf = 2)
```

HairEye Hair and Eye Color of Statistics Students (Data Frame)

## Description

The distribution of hair color, eye color and sex among 592 statistics students (from Snee 1974 and Friendly 1992).

## Format

A data frame with 592 rows and 3 variables.

## Source

This is simply a data frame version of the in-built data set HairEyeColor.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
haireye.crg
summary(haireye.crg, parm = "b", add_ci = TRUE)
screeplot(haireye.crg, add_ci = TRUE)
anova(haireye.crg, nf = 2)
plot(haireye.crg, x_ell = TRUE, xsub = c("Hair", "Sex"))
```

pcplot.corregp Parallel Coordinate Plotting for Correspondence Regression

## Description

Method to produce a parallel coordinate plot of the output of a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
pcplot(x, parm = "x", axes, add_ci = FALSE, cl = 0.95,
    nq = TRUE, col = "darkgrey", cex = par("cex"), font = par("font"),
    family = par("family"), lwd = par("lwd"), lty = par("lty"),
    lcol = col, psym = NULL, pcol = col, pcex = cex, ecol = "red",
    ewid = 1, etyp = 2, acol = "black", awid = 1, atyp = 1,
    acex = cex, afnt = font, adir = 1, add_scale = FALSE, main = NULL,
    sub = NULL, ...)
pcplot(x, ...)
```


## Arguments

X
parm
axes
add_ci Logical specifying whether to include the confidence intervals. Defaults to FALSE.
cl The confidence level for the confidence intervals. Defaults to 0.95.
nq Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence intervals. Defaults to TRUE. If FALSE, then the confidence intervals are computed directly with the quantile function.
col Color of the text labels: either numeric or see colors.
cex Character expansion factor: a number to specify the size of the text labels.
font Font of the text labels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. Defaults to 1 .
family Font family of the text labels: can be "serif", "sans", "mono" or one of the Hershey fonts.
lwd Width of the connecting lines: a number to specify the line width.
lty Line type of the connecting lines: 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1.
lcol Color of the connecting lines: either numeric or see colors.
psym The symbol (or "plotting character") for the values of the coordinates on the axes.
pcol Color of the symbol for the values on the axes: either numeric or see colors.
pcex Character expansion factor of the symbol for the values on the axes.
ecol Color of the error lines (connecting the confidence intervals on each axis): either numeric or see colors.
$\left.\begin{array}{ll}\text { ewid } & \begin{array}{l}\text { Width of the error lines (connecting the confidence intervals on each axis): a } \\ \text { number to specify the line width. }\end{array} \\ \text { etyp } & \begin{array}{l}\text { Line type of the error lines (connecting the confidence intervals on each axis): } 0 \\ \text { or "blank", } 1 \text { or "solid", } 2 \text { or "dashed", } 3 \text { or "dotted", } 4 \text { or "dotdash", } 5 \text { or } \\ \text { "longdash", } 6 \text { or "twodash". Defaults to } 2 .\end{array} \\ \text { acol } & \text { Color of the parallel axes: either numeric or see colors. } \\ \text { awid } & \text { Width of the parallel axes: a number to specify the line width. } \\ \text { atyp } & \begin{array}{l}\text { Line type of the parallel axes: } 0 \text { or "blank", } 1 \text { or "solid", } 2 \text { or "dashed", } 3 \text { or } \\ \text { "dotted", } 4 \text { or "dotdash", } 5 \text { or "longdash", } 6 \text { or "twodash". Defaults to } 1 .\end{array} \\ \text { acex } & \begin{array}{l}\text { Character expansion factor for the labels of the parallel axes. }\end{array} \\ \text { afnt } & \begin{array}{l}\text { Font for the labels of the parallel axes: } 1 \text { for plain, } 2 \text { for bold, } 3 \text { for italic, and } 4 \\ \text { for bold italic. }\end{array} \\ \text { Reading direction of the labels on the parallel axes: either a numeric value } \\ \text { between } 0 \text { and } 3 \text { (see the las argument in the graphical parameters par) or a } \\ \text { character value matching either "horizontal" or "vertical". Defaults to } 1\end{array}\right]$ (horizontal).

## Details

Although adding lines for confidence intervals is possible, it is not recommended, as it typically leads to an unreadable plot.

## Value

A parallel coordinate plot containing the output of a correspondence regression.

## See Also

```
ciplot.corregp, plot.corregp, plot3d.corregp, agplot.corregp.
```


## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
pcplot(haireye.crg, parm = "y", axes = 1:3)
pcplot(haireye.crg, parm = c("Hair", "Sex"), axes = 1:3)
```


## Description

Basic method to plot the output of a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
plot(x, axes = 1:2, y_btm = TRUE, y_ell = FALSE,
    x_ell = FALSE, ysub = NULL, xsub = NULL, hlim = NULL, vlim = NULL,
    expa_btm = 1, expa_top = 1, asp = 1, asp_btm = asp, asp_top = asp,
    col_btm = "darkgrey", col_top = "red", cex_btm = par("cex"),
    cex_top = cex_btm, font_btm = par("font"), font_top = font_btm,
    fam_btm = par("family"), fam_top = fam_btm, col_ell = par("col"),
    lwd_ell = par("lwd"), lty_ell = par("lty"), col_ori = par("col"),
    lwd_ori = par("lwd"), lty_ori = 1, main = NULL, sub = NULL,
    hlab = NULL, vlab = NULL, cl = 0.95, np = 100, add_ori = TRUE, ...)
```


## Arguments

X
axes
y_btm
y_ell Logical specifying whether the confidence ellipses of the Y levels should be plotted. Defaults to FALSE.
x_ell Logical specifying whether the confidence ellipses of the $X$ levels should be plotted. Defaults to FALSE.
ysub Vector of indices to select a subset of the Y levels.
xsub Vector of indices to select a subset of the $X$ levels.
hlim Vector of two values specifying the lower and upper limit between which to plot the horizontal axis.
vlim Vector of two values specifying the lower and upper limit between which to plot the vertical axis.
expa_btm Expansion factor for the bottom coordinates: a number to rescale the axes.
expa_top Expansion factor for the top coordinates: a number to rescale the axes.
asp The aspect ratio for the whole plot. See plot.window.
asp_btm
asp_top The aspect ratio for the top coordinates. See plot.window.
col_btm Color of the bottom levels: either numeric or see colors. Defaults to "darkgrey".

| col_top | Color of the top levels: either numeric or see colors. Defaults to "red". |
| :---: | :---: |
| cex_btm | Character expansion factor of the bottom levels: a number to specify the size of the text labels. |
| cex_top | Character expansion factor of the top levels: a number to specify the size of the text labels. |
| font_btm | Font of the bottom levels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. Defaults to 1. |
| font_top | Font of the top levels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. Defaults to 1. |
| fam_btm | Font family of the bottom levels: can be "serif", "sans", "mono" or one of the Hershey fonts. |
| fam_top | Font family of the top levels: can be "serif", "sans", "mono" or one of the Hershey fonts. |
| col_ell | Color of the confidence ellipses: either a number or see colors. |
| lwd_ell | Width of the confidence ellipses: a number to specify the line width. |
| lty_ell | Line type of the confidence ellipses: 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1. |
| col_ori | Color of the lines through the origin: either numeric or see colors. |
| lwd_ori | Width of the lines through the origin: a number to specify the line width. |
| lty_ori | Line type of the lines through the origin: 0 or "blank", 1 or "solid", 2 or "dashed", 3 or "dotted", 4 or "dotdash", 5 or "longdash", 6 or "twodash". Defaults to 1 . |
| main | The main title of the plot. |
| sub | The subtitle of the plot. |
| hlab | The title of the horizontal axis. |
| vlab | The title of the vertical axis. |
| cl | The confidence level for the confidence ellipses. Defaults to 0.95. |
| np | The number of points to represent the confidence ellipses. Defaults to 100. |
| add_ori | Logical specifying whether to add lines through the origin. Defaults to TRUE. |
|  | Further arguments passed to or from other methods. |

## Details

The plot of a correspondence regression is by definition a biplot.

## Value

A plot window containing the output of a correspondence regression.

## References

Gower, J., S. Lubbe and N. Le Roux (2011) Understanding biplots. Chichester: Wiley.
Greenacre, M. (2010) Biplots in practice. Bilbao: Fundacion BBVA.

## See Also

corregp, summary.corregp, screeplot.corregp, anova.corregp, biplot.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
plot(haireye.crg, x_ell = TRUE, xsub = c("Hair", "Sex"))
```

plot3d.corregp 3D Plotting for Correspondence Regression

## Description

Method to produce a 3D plot for a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
plot3d(x, axes = 1:3, y_btm = TRUE, y_ell = FALSE,
    x_ell = FALSE, ysub = NULL, xsub = NULL, hlim = NULL, vlim = NULL,
    dlim = NULL, asp = par3d("scale"), col_btm = "darkgrey",
    col_top = "red", cex_btm = par3d("cex"), cex_top = cex_btm,
    font_btm = par3d("font"), font_top = font_btm,
    fam_btm = par3d("family"), fam_top = fam_btm, col_ell = "black",
    lwd_ell = 1, lty_ell = "shade", opa_ell = 0.2, col_ori = "grey",
    lwd_ori = 1, main = NULL, sub = NULL, hlab = NULL, vlab = NULL,
    dlab = NULL, cl = 0.95, add_ori = TRUE, ...)
```


## Arguments

x
axes
y_btm
y_ell Logical specifying whether the confidence ellipsoids of the Y levels should be plotted. Defaults to FALSE.
x_ell Logical specifying whether the confidence ellipsoids of the X levels should be plotted. Defaults to FALSE.
ysub Vector of indices to select a subset of the Y levels.
xsub Vector of indices to select a subset of the $X$ levels.
hlim Vector of two values specifying the lower and upper limit between which to plot the horizontal axis.

| vlim | Vector of two values specifying the lower and upper limit between which to plot the vertical axis. |
| :---: | :---: |
| dlim | Vector of two values specifying the lower and upper limit between which to plot the "depth" axis. |
| asp | The aspect ratio for the whole plot. See aspect3d. |
| col_btm | Color of the bottom levels: either numeric or see colors. Defaults to "darkgrey". |
| col_top | Color of the top levels: either numeric or see colors. Defaults to "red". |
| cex_btm | Character expansion factor of the bottom levels: a number to specify the size of the text labels. |
| cex_top | Character expansion factor of the top levels: a number to specify the size of the text labels. |
| font_btm | Font of the bottom levels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. |
| font_top | Font of the top levels: 1 for plain, 2 for bold, 3 for italic, and 4 for bold italic. |
| fam_btm | Font family of the bottom levels: can be "serif", "sans", "mono" or code"symbol". |
| fam_top | Font family of the top levels: can be "serif", "sans", "mono" or "symbol". |
| col_ell | Color of the confidence ellipsoids: either a number or see colors. Defaults to "black". |
| lwd_ell | Width of the confidence ellipsoids: a number to specify the line width. |
| lty_ell | Line type of the confidence ellipsoids: either "shade", "wire", or "dots". Defaults to "shade". |
| opa_ell | Opaqueness of the confidence ellipsoids: a number between 0 for fully transparent and 1 for fully opaque. Defaults to 0.2. |
| col_ori | Color of the lines through the origin: either a number or see colors. Defaults to "grey". |
| lwd_ori | Width of the lines through the origin: a number to specify the line width. Defaults to 1 . |
| main | The main title of the plot. |
| sub | The subtitle of the plot. |
| hlab | The title of the horizontal axis. |
| vlab | The title of the vertical axis. |
| dlab | The title of the "depth" axis. |
| cl | The confidence level for the confidence ellipsoids. Defaults to 0.95. |
| add_ori | Logical specifying whether to add lines through the origin. Defaults to TRUE. |
|  | Further arguments passed to or from other methods. |

## Details

plot3d (of a corregp output) makes use of plot3d (and text3d and abclines3d) from the package rgl.

## Value

A 3D plot window containing the output of a correspondence regression.

## See Also

corregp, pcplot.corregp, agplot.corregp, plot3d.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
plot3d(haireye.crg, x_ell = TRUE, xsub = c("Hair", "Sex"))
```

print.anova.corregp Printing the ANOVA Table of Correspondence Regression

## Description

Method to print the output of anova.corregp.

## Usage

\#\# S3 method for class 'anova.corregp' print(x, ...)

## Arguments

x
The output of a call to anova on a "corregp" object (i.e. an object of class "anova.corregp").
$\ldots \quad$ Further arguments passed to or from other methods.

## Value

The output of a call to anova on a "corregp" object.

## See Also

anova. corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
anova(haireye.crg, nf = 2)
```

```
print.corregp Printing Correspondence Regression
```


## Description

Method to print the output of corregp.

## Usage

\#\# S3 method for class 'corregp'
print(x, nf = 2, ...)

## Arguments

| x | The output of a call to corregp (i.e. an object of class "corregp"). |
| :--- | :--- |
| nf | The number of dimensions to print. Defaults to the first two dimensions. |
| $\ldots$ | Further arguments passed to or from other methods. |

## Value

The output of a call to corregp.

## See Also

corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
haireye.crg
print(haireye.crg, nf = 3)
```

print.summary.corregp Printing the Summary of Correspondence Regression

## Description

Method to print the output of summary. corregp.

## Usage

\#\# S3 method for class 'summary.corregp'
print(x, ...)

## Arguments

X
The output of a call to summary on a "corregp" object (i.e. an object of class "summary.corregp").
... Further arguments passed to or from other methods.

## Value

The output of a call to summary on a "corregp" object. The eigenvalues and contributions are printed with TOTALs.

## See Also

summary. corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
summary(haireye.crg, add_ci = TRUE)
summary(haireye.crg, parm = "y", contrib = "pts_axs", nf = 2)
```

```
residuals.corregp Extracting Residuals from Correspondence Regression
```


## Description

Method to extract the residuals of a correspondence regression.

## Usage

```
## S3 method for class 'corregp'
residuals(object, parm = "all", nf = NULL, ...)
## S3 method for class 'corregp'
resid(object, parm = "all", nf = NULL, ...)
```


## Arguments

object The output of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to extract the residuals. Can be "all", "both" (or abbreviations), " $y$ " or " $x$ " for the residuals of every cell in the data, but it can also be any vector of term names in X or level names in X . Defaults to "all".
$\mathrm{nf} \quad$ The number of dimensions to be retained in the reduced space. Defaults to all dimensions (no reduction).
... Further arguments passed to or from other methods.

## Details

If all dimensions are retained, then the residuals will only be exactly zero to the observed counts if no conditioning factors were specified with the argument "part" in the corregp call. This is because the associations with the conditioning factors (in "part") are not taken into account.

## Value

A matrix or vector with the residuals for the parameters of interest, based on the selected number of dimensions.

## See Also

```
coef.corregp, fitted.corregp.
```


## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
residuals(haireye.crg, parm = c("Hair", "Sex"), nf = 2)
resid(haireye.crg, parm = c("Hair", "Sex"), nf = 2)
```

```
screeplot.corregp Scree Plotting
```


## Description

Method to produce a scree plot, i.e. a bar chart of the eigenvalues.

## Usage

```
## S3 method for class 'corregp'
screeplot(x, type = "value", add_ci = FALSE, cl = 0.95,
    nq = TRUE, ...)
```


## Arguments

add_ci Logical specifying whether to include the confidence intervals. Defaults to
x
type
cl

The output of a call to corregp (i.e. an object of class "corregp").
A character specification of which type of values to plot: either "value" for the actual eigenvalues, "\%" for percentages or "cum_\%" for cumulative percentages. Defaults to "value". FALSE.
The confidence level for the confidence intervals. Defaults to 0.95.
nq Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the computation of the confidence intervals. Defaults to TRUE. If FALSE, then the confidence intervals are computed directly with the quantile function.
.. Further arguments passed to or from other methods.

## Details

screeplot (of a corregp output) makes use of barplot2 from the package gplots.

## Value

A plot window containing the scree plot.

## See Also

corregp, summary.corregp, anova.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
screeplot(haireye.crg, add_ci = TRUE)
```

```
summary.corregp Summarizing Correspondence Regression
```


## Description

Method to produce a summary of a correspondence regression.

## Usage

\#\# S3 method for class 'corregp'
summary (object, parm = NULL, contrib = NULL, nf = NULL, add_ci = FALSE, cl = 0.95, nq = TRUE, ...)

## Arguments

object The outout of a call to corregp (i.e. an object of class "corregp").
parm The parameter for which to compute the contributions contrib. Can be either " $y$ " for the Y contributions, " $x$ " for the $X$ contributions, "both" which can be abbreviated to " $b$ ", or a vector of term names in X. Defaults to " $b$ ".
contrib The type of contributions to be computed: either from points to axes (absolute contributions) or from axes to points (squared correlations). The specification can be "pnts_to_axes" or "axes_to_pnts", "pts2axs" or "axs2pts", "p_a" or "a_p", or any other reasonable abbreviation.

| nf | The number of dimensions to be retained in the reduced space. Defaults to all <br> dimensions (no reduction). |
| :--- | :--- |
| add_ci | Logical specifying whether to compute confidence intervals for the eigenvalues <br> (and eigenvalues only). Defaults to FALSE. |
| cl | The confidence level for the confidence intervals. Defaults to 0.95. |
| nq | Logical specifying whether to use a normal quantile (i.e. apply qnorm) in the <br> computation of the confidence intervals. Defaults to TRUE. If FALSE, then the <br> confidence intervals are computed directly with the quantile function. |
| $\ldots$ | Further arguments passed to or from other methods. |

## Value

An object of class "summary.corregp", providing a summary of a correspondence regression, i.e. a list with components:
\(\left.$$
\begin{array}{ll}\begin{array}{l}\text { formula } \\
\text { data }\end{array} & \begin{array}{l}\text { The formula specified to the formula argument in the call to corregp. } \\
\text { part }\end{array}
$$ <br>
che name of the data frame specified to the data argument in the call to corregp. <br>
phi_squared \& The name of the factor specified to the part argument in the call to corregp. <br>
N The chi-squared value of the correspondence regression. <br>
The phi-squared value of the correspondence regression, i.e. the chi-squared <br>

value divided by N .\end{array}\right]\)| The total number of observations. |
| :--- |
| eigen |
| Depending on add_ci: if FALSE, a matrix of the actual eigenvalues, their per- |
| centages and cumulative percentages; if TRUE, a list of the actual eigenvalues, |
| their percentages and cumulative percentages together with the lower and upper |
| confidence limits for each. |

## See Also

corregp, print. summary. corregp, anova.corregp.

## Examples

```
data(HairEye)
haireye.crg <- corregp(Eye ~ Hair * Sex, data = HairEye, b = 3000)
summary(haireye.crg, add_ci = TRUE)
summary(haireye.crg, parm = "y", contrib = "pts_axs", nf = 2)
```

The Use of Inflected or Uninflected Determiners in the Belgian Dutch Vernacular

## Description

The distribution of the Belgian Dutch -e(n)-suffix with 14 determiners in 14 registers and for several speaker characteristics.

## Format

A data frame with 40778 rows and 13 variables.

- Variant The linguistic variant used in a set of alternatives (35 levels).
- Variable The linguistic variable specifying a set of alternatives (14 levels).
- Inflected Numeric variable specifying whether the linguistic variant is inflected (1) or not (0).
- Register The register of the data in the Spoken Dutch Corpus (14 levels: see here for their definition).
- Register2 The dichotomization of Register into private and public.
- SpeakerID The ID of the speaker in the Spoken Dutch Corpus (1144 levels).
- Region The region in which the speaker lived until the age of 18 (4 levels).
- Sex The sex of the speaker (2 levels).
- BirthYear The year in which the speaker was born (63 levels).
- Decade The decade in which the speaker was born (7 levels).
- Generation The generation cohort in which the speaker was born (5 levels).
- Education The level of education of the speaker (3 levels).
- Occupation The level of occupation of the speaker (10 levels: see here for their definition).


## Source

Plevoets, K. (2008) Tussen spreek- en standaardtaal. Leuven, Doctoral dissertation. Available online here.

## Examples

```
data(TSS)
# The execution of corregp may be slow, due to bootstrapping:
tss.crg <- corregp(Variant ~ Register2 * Region, data = TSS, part = "Variable", b = 3000)
tss.crg
summary(tss.crg, parm = "b", add_ci = TRUE)
screeplot(tss.crg, add_ci = TRUE)
anova(tss.crg, nf = 2)
tss.col <- ifelse( xtabs(~ Variant + Inflected, data = TSS)[, 1] > 0, "blue", "red")
plot(tss.crg, x_ell = TRUE, xsub = c("Register2", "Region"), col_btm = tss.col, col_top = "black")
```


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