# Package 'OpenRepGrid'

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Title Tools to Analyze Repertory Grid Data

LazyData yes

Type Package

LazyLoad yes

**Description** Analyze repertory grids, a qualitative-quantitative

data collection technique devised by George A. Kelly in the 1950s. Today, grids are used across various domains ranging from clinical psychology to marketing. The package contains functions to quantitatively analyze and visualize repertory grid data (see e.g. Bell, 2005, <doi:10.1002/0470013370.ch9>;

Fransella, Bell, & Bannister, 2004, ISBN: 978-0-470-09080-0).

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URL http://openrepgrid.org,

https://github.com/markheckmann/OpenRepGrid

**Imports** methods, graphics, grid, utils, stats, grDevices, plyr, stringr, abind, rgl, colorspace, GPArotation, psych, XML, tcltk, pvclust, openxlsx

Collate 'bertin.r' 'calc.r' 'data-openrepgrid.r' 'dev-functions.r'
 'distance.R' 'double-entry.R' 'export.r' 'gmMain.r' 'import.r'
 'measures.r' 'onair.r' 'openrepgrid.r' 'repgrid-basicops.r' 'repgrid-constructs.r'
 'repgrid-elements.r' 'repgrid-output.r' 'repgrid-plots.r'
 'repgrid-ratings.r' 'rgl-3d.r' 'settings.r' 'utils-import.r'
 'utils.r' 'zzz.r'

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Concatenate repgrid objects.

# Description

Simple concatenation of repgrid objects or list containing repgrid objects using the '+' operator.

<sup>+,</sup>repgrid,repgrid-method

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# Usage

```
## S4 method for signature 'repgrid,repgrid'
e1 + e2
## S4 method for signature 'list,repgrid'
e1 + e2
## S4 method for signature 'repgrid,list'
e1 + e2
```

# **Arguments**

e1, e2 A repgrid object.

# **Details**

Methods for "+" function.

# Author(s)

Mark heckmann

# **Examples**

```
x <- bell2010
x + x
x + list(x,x)
list(x,x) + x</pre>
```

addConstruct

Add a new construct to an existing grid object.

# **Description**

Add a new construct to an existing grid object.

# Usage

```
addConstruct(x, 1.name = NA, r.name = NA, scores = NA, 1.preferred = NA,
    r.preferred = NA, 1.emerged = NA, r.emerged = NA, position = NA,
    side = "pre")
```

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# Arguments

X	repgrid object.
1.name	Name of the left pole (character string).
r.name	Name of the right pole (character string).
scores	Numerical ratings for the new construct row (length must match number of elements in the grid).
1.preferred	Is the left one the preferred pole? (logical).
r.preferred	Is the right one the preferred pole? (logical).
1.emerged	Is the left one the emergent pole? (logical).
r.emerged	Is the right one the emergent pole? (logical).
position	An integer at which row the construct will be added. TODO. Does not work properly.
side	Not yet in use.

# Value

repgrid object.

# Author(s)

Mark Heckmann

# See Also

addElement

```
## Not run:

# show grid
bell2010
addConstruct(bell2010, "left pole", "pole right", c(3,1,3,2,5,4,6,3,7,1))

## End(Not run)
```

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addElement	Add an element to an	existing grid.
------------	----------------------	----------------

# **Description**

Add an element to an existing grid.

# Usage

```
addElement(x, name = NA, scores = NA, abbreviation = NA, status = NA,
    position = NA, side = "pre")
```

# **Arguments**

x repgrid object.

name Name of the new element (character string).

scores Numerical ratings for the new element column (length must match number of

constructs in the grid).

abbreviation Abbreviation for element name. status Element status (not yet in use).

position An integer at which column the element will be added. TODO: Does not work

properly yet.

side Not yet in use.

# Value

repgrid object

## Author(s)

Mark Heckmann

# See Also

addConstruct

```
## Not run:
bell2010
addElement(bell2010, "new element", c(1,2,5,4,3,6,5,2,7))
## End(Not run)
```

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alignByIdeal

Align constructs using the ideal element to gain pole preferences.

# **Description**

The direction of the constructs in a grid is arbitrary and a reflection of a scale does not affect the information contained in the grid. Nonetheless, the direction of a scale has an effect on interelement correlations (Mackay, 1992) and on the spatial representation and clustering of the grid (Bell, 2010). Hence, it is desirable to follow a protocol to align constructs that will render unique results. A common approach is to align constucts by pole preference, i. e. alignining all positive and negative poles. This can e. g. be achieved using swapPoles. If an ideal element is present, this element can be used to identify the positive and negative pole. The function alignByIdeal will align the constructs accordingly. Note that this approach does not always yield definite results as sometimes ratings do not show a clear preference for one pole (Winter, Bell & Watson, 2010). If a preference cannot be determined definitely, the construct direction remains unchanged (a warning is issued in that case).

## Usage

```
alignByIdeal(x, ideal, high = TRUE)
```

#### **Arguments**

x repgrid object

ideal Number of the element that is used for alignment (the ideal).

high Logical. Whether to align the constructs so the ideal will have high ratings on

the constructs (i.e. TRUE, default) or low ratings (FALSE). High scores will lead to the preference pole on the right side, low scores will align the preference pole

on the left side.

#### Value

repgrid object with aligned constructs.

#### Author(s)

Mark Heckmann

#### References

Bell, R. C. (2010). A note on aligning constructs. Personal Construct Theory & Practice, 7, 42-48.

Mackay, N. (1992). Identification, Reflection, and Correlation: Problems in the bases of repertory grid measures. *International Journal of Personal Construct Psychology*, *5*(1), 57-75.

Winter, D. A., Bell, R. C., & Watson, S. (2010). Midpoint ratings on personal constructs: Constriction or the middle way? *Journal of Constructivist Psychology*, 23(4), 337-356.

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# See Also

```
alignByLoadings
```

# **Examples**

alignByLoadings

Align constructs by loadings on first pricipal component.

# Description

In case a construct loads negatively on the first principal component, the function alignByLoadings will reverse it so that all constructs have positive loadings on the first principal component (see deatil section for more).

# Usage

```
alignByLoadings(x, trim = 20, index = TRUE)
```

# **Arguments**

x repgrid object.

trim The number of characters a construct is trimmed to (default is 10). If NA no

trimming is done. Trimming simply saves space when displaying the output.

index Whether to print the number of the construct (e.g. for correltion matrices). The

default is TRUE.

## **Details**

The direction of the constructs in a grid is arbitrary and a reflection of a scale does not affect the information contained in the grid. Nonetheless, the direction of a scale has an effect on interelement correlations (Mackay, 1992) and on the spatial representation and clustering of the grid (Bell, 2010). Hence, it is desirable to follow a protocol to align constructs that will render unique results. A common approach is to align constructs by pole preference, but this information is not always accessible. Bell (2010) proposed another solution for the problem of construct alignment. As a unique protocol he suggests to align constructs in a way so they all have positive loadings on the first component of a grid PCA.

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#### Value

An object of class alignByLoadings containing a list of calculations with the following entries:

cor.before Construct correlation matrix before reversal

loadings.before

Loadings on PCs before reversal

reversed Constructs that have been reversed

cor.after Construct correlation matrix after reversal

loadings.after Loadings on PCs after reversal

#### Note

Bell (2010) proposed a solution for the problem of construct alignment. As construct reversal has an effect on element correlation and thus on any measure that based on element correlation (Mackay, 1992), it is desireable to have a standard method for construct alignment independently from its semantics (preferred pole etc.). Bell (2010) proposes to align constructs in a way so they all have positive loadings on the first component of a grid PCA.

#### Author(s)

Mark Heckmann

#### References

Bell, R. C. (2010). A note on aligning constructs. *Personal Construct Theory & Practice*, 7, 42-48. Mackay, N. (1992). Identification, Reflection, and Correlation: Problems in the bases of repertory grid measures. *International Journal of Personal Construct Psychology*, 5(1), 57-75.

# See Also

```
alignByIdeal
```

```
# reproduction of the example in the Bell (2010)
# constructs aligned by loadings on PC 1
bell2010
alignByLoadings(bell2010)

# save results
a <- alignByLoadings(bell2010)

# modify printing of resukts
print(a, digits=5)

# access results for further processing
names(a)
a$cor.before</pre>
```

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```
a$loadings.before
a$reversed
a$cor.after
a$loadings.after
```

bertin

Make Bertin display of grid data.

# Description

One of the most popular ways of displaying grid data has been adopted from Bertin's (1974) graphical proposals, which have had an immense influence onto data visualization. One of the most appealing ideas presented by Bertin is the concept of the reordable matrix. It is comprised of graphical displays for each cell, allowing to identify structures by eye-balling reordered versions of the data matrix (see Bertin, 1974). In the context of repertory grids, the display is made up of a simple colored rectangle where the color denotes the corresponding score. Bright values correspond to low, dark to high scores. For an example of how to analyze a Bertin display see e.g. Dick (2000) and Raeithel (1998).

# Usage

```
bertin(x, colors = c("white", "black"), showvalues = TRUE, xlim = c(0.2, 0.8), ylim = c(0, 0.6), margins = c(0, 1, 1), cex.elements = 0.7, cex.constructs = 0.7, cex.text = 0.6, col.text = NA, border = "white", lheight = 0.75, id = c(T, T), cc = 0, cr = 0, cc.old = 0, cr.old = 0, col.mark.fill = "#FCF5A4", print = TRUE, ...)
```

# **Arguments**

X	repgrid object.
colors	Vector. Two or more colors defining the color ramp for the bertin (default c("white", "black")).
showvalues	Logical. Wether scores are shown in bertin
xlim	Vector. Left and right limits inner bertin (default c(.2, .8)).
ylim	Vector. Lower and upper limits of inner bertin default(c(.0, .6)).
margins	Vector of length three (default margins= $c(0,1,1)$ ). 1st element denotes the left, 2nd the upper and 3rd the right margin in npc coordinates (i.e. 0 to zero).
cex.elements	Numeric. Text size of element labels (default .7).
cex.constructs	Numeric. Text size of construct labels (default .7).
cex.text	Numeric. Text size of scores in bertin cells (default .7).
col.text	Color of scores in bertin (default NA). By default the color of the text is chosen according to the background color. If the background ist bright the text will be black and vice versa. When a color is specified the color is set independent of background.

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border	Border color of the bertin cells (default white).
lheight	Line height for constructs.
id	Logical. Wheteher to print id number for constructs and elements respectively (default $c(T,T)$ ).
сс	Numeric. Current column to mark.
cr	Numeric. Current row to mark.
cc.old	Numeric. Column to unmark.
cr.old	Numeric. Row to unmark.
col.mark.fill	Color of marked row or column (default "#FCF5A4").
print	Print whole bertin. If FALSE only current and old row and column are printed.
	Optional arguments to be passed on to bertinBase.

#### Value

NULL just for the side effects, i.e. printing.

## References

Bertin, J. (1974). *Graphische Semiologie: Diagramme, Netze, Karten.* Berlin, New York: de Gruyter.

Dick, M. (2000). The Use of Narrative Grid Interviews in Psychological Mobility Research. *Forum Qualitative Social forschung / Forum: Qualitative Social Research*, 1(2).

Raeithel, A. (1998). Kooperative Modellproduktion von Professionellen und Klienten - erlauetert am Beispiel des Repertory Grid. *Selbstorganisation, Kooperation, Zeichenprozess: Arbeiten zu einer kulturwissenschaftlichen, anwendungsbezogenen Psychologie* (pp. 209-254). Opladen: Westdeutscher Verlag.

```
## Not run:
bertin(feixas2004)
bertin(feixas2004, c("white", "darkblue"))
bertin(feixas2004, showvalues=F)
bertin(feixas2004, border="grey")
bertin(feixas2004, cex.text=.9)
bertin(feixas2004, id=c(F, F))

bertin(feixas2004, cc=3, cr=4)
bertin(feixas2004, cc=3, cr=4, col.mark.fill="#e6e6e6")

## End(Not run)
```

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bertinCluster

Bertin display with corresponding cluster anaylsis.

#### Description

Element columns and constructs rows are ordered according to cluster criterion. Various distance measures as well as cluster methods are supported.

#### **Usage**

```
bertinCluster(x, dmethod = c("euclidean", "euclidean"), cmethod = c("ward",
    "ward"), p = c(2, 2), align = TRUE, trim = NA, type = c("triangle"),
    xsegs = c(0, 0.2, 0.7, 0.9, 1), ysegs = c(0, 0.1, 0.7, 1), x.off = 0.01,
    y.off = 0.01, cex.axis = 0.6, col.axis = grey(0.4), draw.axis = TRUE,
    ...)
```

#### **Arguments**

rid	object.
	rid

The distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary", or "minkowski". Default is "euclidean". Any unambiguous substring can be given (e.g. "euc" for "euclidean"). A vector of length two can be passed if a different distance measure for constructs and elements is wanted (e.g.c("euclidean", "manhattan")). This will apply euclidean distance to the constructs and manhattan distance to the elements. For

additional information on the different types see ?dist.

The agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". Default is "ward". A vector of length two can be passed if a different cluster method for constructs and elements is wanted (e.g.c("ward", "euclidean")). This will apply ward clustering to the constructs and single linkage clustering to the elements. If only one of either constructs or elements is to be clustered the value NA can be supplied. E.g. to cluster

elements only use c(NA, "ward").

The power of the Minkowski distance, in case "minkowski" is used as argument for dmethod. p can be a vector of length two if different powers are wanted for constructs and elements respectively (e.g. c(2,1)).

Whether the constructs should be aligned before clustering (default is TRUE). If not, the grid matrix is clustered as is. See Details section in function cluster

for more information.

The number of characters a construct is trimmed to (default is 10). If NA no trimming is done. Trimming simply saves space when displaying the output.

Type of dendrogram. Either or "triangle" (default) or "rectangle" form.

Numeric vector of normal device coordinates (ndc i.e. 0 to 1) to mark the widths of the regions for the left labels, for the bertin display, for the right labels and

for the vertical dendrogram (i.e. for the constructs).

cmethod

dmethod

p

align

trim

type xsegs bertinCluster 13

ysegs	Numeric vector of normal device coordinates (ndc i.e. 0 to 1) to mark the heights of the regions for the horizontal dendrogram (i.e. for the elements), for the bertin display and for the element names.
x.off	Horizontal offset between construct labels and construct dendrogram and (default is 0.01 in normal device coordinates).
y.off	Vertical offset between bertin display and element dendrogram and (default is 0.01 in normal device coordinates).
cex.axis	cex for axis labels, default is .6.
col.axis	Color for axis and axis labels, default is grey(.4).
draw.axis	Whether to draw axis showing the distance metric for the dendrograms (default is TRUE).
	additional parameters to be passed to function bertin.

#### Value

A list of two hclust object, for elements and constructs respectively.

## Author(s)

Mark Heckmann

### See Also

cluster

```
## Not run:
  # default is euclidean distance and ward clustering
  bertinCluster(bell2010)
  ### applying different distance measures and cluster methods
  # euclidean distance and single linkage clustering
  bertinCluster(bell2010, cmethod="single")
  # manhattan distance and single linkage clustering
  bertinCluster(bell2010, dmethod="manhattan", cm="single")
  # minkowksi distance with power of 2 = euclidean distance
  bertinCluster(bell2010, dm="mink", p=2)
  ### using different methods for constructs and elements
  # ward clustering for constructs, single linkage for elements
  bertinCluster(bell2010, cmethod=c("ward", "single"))
   # euclidean distance measure for constructs, manhatten
   # distance for elements
  bertinCluster(bell2010, dmethod=c("euclidean", "man"))
  # minkowski metric with different powers for constructs and elements
```

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```
bertinCluster(bell2010, dmethod="mink", p=c(2,1)))
  ### clustering either constructs or elements only
  # euclidean distance and ward clustering for constructs no
  # clustering for elements
  bertinCluster(bell2010, cmethod=c("ward", NA))
  # euclidean distance and single linkage clustering for elements
  # no clustering for constructs
  bertinCluster(bell2010, cm=c(NA, "single"))
  ### changing the appearance
  # different dendrogram type
  bertinCluster(bell2010, type="rectangle")
  # no axis drawn for dendrogram
  bertinCluster(bell2010, draw.axis=F)
  ### passing on arguments to bertin function via \dots
   # grey cell borders in bertin display
  bertinCluster(bell2010, border="grey")
  # omit printing of grid scores, i.e. colors only
  bertinCluster(bell2010, showvalues=FALSE)
  ### changing the layout
  # making the vertical dendrogram bigger
  bertinCluster(bell2010, xsegs=c(0, .2, .5, .7, 1))
  # making the horizontal dendrogram bigger
  bertinCluster(bell2010, ysegs=c(0, .3, .8, 1))
## End(Not run)
```

bindConstructs

Concatenate the constructs of two or more grids.

## **Description**

I.e. the constructs are combined to form one long grid. The girds must have the same set of elements and an identical scale range. The order of the elements may differ.

# Usage

```
bindConstructs(..., index = FALSE)
```

# Arguments

. . . One or more repgrid objects or a list containing repgrid object.

index TODO. Logical (default TRUE). Whether to add an index at the end of each construct name so it remains clear from which grid each construct came.

## **Details**

This function can be used in order to analyse multiple grids as one 'big grid' (eg. Slater, 1977, chap. 11).

#### Value

repgrid object with concatenated constructs.

## Author(s)

Mark Heckmann

#### References

Slater, P. (1977). The measurement of intrapersonal space by grid technique. London: Wiley.

#### **Examples**

```
a <- randomGrid()
b <- randomGrid()
b@elements <- rev(a@elements)  # reverse elements
bindConstructs(a, b)
bindConstructs(a, b, a)

# using lists of repgrid objects
bindConstructs(a, list(a, b))</pre>
```

biplot2d

Draw a two-dimensional biplot.

# Description

The biplot is the central way to create a joint plot of elements and constructs. Depending on te parameters chosen it contains information on the distances between elements and constructs. Also the relative values the elements have on a construct can be read off by projetion the element onto the construct vector. A lot of parameters can be changed rendering different types of biplots (ESA, Slater's) and different looks (colors, text size). See the example section below to get started.

# Usage

```
biplot2d(x, dim = c(1, 2), map.dim = 3, center = 1, normalize = 0,
    g = 0, h = 1 - g, col.active = NA, col.passive = NA,
    e.point.col = "black", e.point.cex = 0.9, e.label.col = "black",
    e.label.cex = 0.7, e.color.map = c(0.4, 1), c.point.col = "black",
    c.point.cex = 0.8, c.label.col = "black", c.label.cex = 0.7,
    c.color.map = c(0.4, 1), c.points.devangle = 91, c.labels.devangle = 91,
```

```
c.points.show = TRUE, c.labels.show = TRUE, e.points.show = TRUE,
e.labels.show = TRUE, inner.positioning = TRUE,
outer.positioning = TRUE, c.labels.inside = FALSE, c.lines = TRUE,
col.c.lines = grey(0.9), flipaxes = c(FALSE, FALSE), strokes.x = 0.1,
strokes.y = 0.1, offsetting = TRUE, offset.labels = 0, offset.e = 1,
axis.ext = 0.1, mai = c(0.2, 1.5, 0.2, 1.5), rect.margins = c(0.01, 1.5)
0.01), srt = 45, cex.pos = 0.7, xpd = TRUE, unity = FALSE,
unity3d = FALSE, scale.e = 0.9, zoom = 1, var.show = TRUE,
var.cex = 0.7, var.col = grey(0.1), ...)
```

#### **Arguments**

repgrid object. Х

dim Dimensions (i.e. principal components) to be used for biplot (default is c(1,2)).

map.dim Third dimension (depth) used to map aesthetic attributes to (default is 3).

center Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= doublecentering (construct and element means), 4= midpoint centering of rows (con-

structs). The default is 1 (row centering).

normalize A numeric value indicating along what direction (rows, columns) to normalize

by standard deviations. 0 = none, 1 = rows, 2 = columns (default is 0).

Power of the singular value matrix assigned to the left singular vectors, i.e. the g

constructs.

h Power of the singular value matrix assigned to the right singular vectors, i.e. the

elements.

col.active Columns (elements) that are no supplementary points, i.e. they are used in the

SVD to find principal components. default is to use all elements.

Columns (elements) that are supplementary points, i.e. they are NOT used in the col.passive

SVD but projecte into the component space afterwards. They do not determine

the solution. Default is NA, i.e. no elements are set supplementary.

Color of the element symbols. The default is "black". Two values can be e.point.col entered that will create a color ramp. The values of map.dim are mapped onto

the ramp. If only one color color value is supplied (e.g. "black") no mapping occurs and all elements will have the same color irrespective of their value on

the map. dim dimension.

e.point.cex Size of the element symbols. The default is .9. Two values can be entered that will create a size ramp. The values of map. dim are mapped onto the ramp.

If only one color size value is supplied (e.g. .8) no mapping occurs and all elements will have the same size irrespective of their value on the map. dim di-

mension.

e.label.col

Color of the element label. The default is "black". Two values can be entered that will create a color ramp. The values of map, dim are mapped onto the ramp.

If only one color color value is supplied (e.g. "black") no mapping occurs and all labels will have the same color irrespective of their value on the map.dim

dimension.

e.label.cex Size of the element labels. The default is .7. Two values can be entered that will create a size ramp. The values of map.dim are mapped onto the ramp. If only one color size value is supplied (e.g. .7) no mapping occurs and all labels will have the same size irrespective of their value on the map.dim dimension.

- e.color.map Value range to determine what range of the color ramp defined in e.color will be used for mapping the colors. Default is c(.4, ,1). Usually not important for the user.
- c.point.col Color of the construct symbols. The default is "black". Two values can be entered that will create a color ramp. The values of map.dim are mapped onto the ramp. If only one color color value is supplied (e.g. "black") no mapping occurs and all construct will have the same color irrespective of their value on the map.dim dimension.
- c.point.cex Size of the construct symbols. The default is .8. Two values can be entered that will create a size ramp. The values of map.dim are mapped onto the ramp. If only one color size value is supplied (e.g. .8) no mapping occurs and all construct will have the same size irrespective of their value on the map.dim dimension.
- c.label.col Color of the construct label. The default is "black". Two values can be entered that will create a color ramp. The values of map. dim are mapped onto the ramp. If only one color color value is supplied (e.g. "black") no mapping occurs and all labels will have the same color irrespective of their value on the map.dim dimension.
- c.label.cex Size of the construct labels. The default is .7. Two values can be entered that will create a size ramp. The values of map.dim are mapped onto the ramp. If only one color size value is supplied (e.g. .7) no mapping occurs and all labels will have the same size irrespective of their value on the map.dim dimension.
- c.color.map Value range to determine what range of the color ramp defined in c.color will be used for mapping. Default is c(.4, ,1). Usually not important for the user.
- c.points.devangle

The deviation angle from the x-y plane in degrees. These can only be calculated if a third dimension map.dim is specified. Only the constructs that do not depart more than the specified degrees from the x-y plane will be printed. This facilitates the visual interpretation, as only vectors represented near the current plane are shown. Set the value to 91 (default) to show all vectors.

c.labels.devangle

The deviation angle from the x-y plane in degrees. These can only be calculated if a third dimension map.dim is specified. Only the labels of constructs that do not depart more than the specified degrees from the x-y plane will be printed. Set the value to 91 (default) to show all construct labels.

- c.points.show Whether the constructs are printed (default is TRUE). FALSE will surpress the printing of the constructs. To only print certain constructs a numeric vector can be provided (e.g. c(1:10)).
- c.labels.show Whether the construct labels are printed (default is TRUE). FALSE will surpress the printing of the labels. To only print certain construct labels a numeric vector can be provided (e.g. c(1:10)).

e.points.show Whether the elements are printed (default is TRUE). FALSE will surpress the printing of the elements. To only print certain elements a numeric vector can be provided (e.g. c(1:10)).

e.labels.show Whether the element labels are printed (default is TRUE). FALSE will surpress the printing of the labels. To only print certain element labels a numeric vector can be provided (e.g. c(1:10)).

inner.positioning

Logical. Whether to calculate positions to minimize overplotting of elements and construct labels (default isTRUE). Note that the positioning may slow down the plotting.

outer.positioning

Logical. Whether to calculate positions to minimize overplotting of of construct labels on the outer borders (default isTRUE). Note that the positioning may slow down the plotting.

c.labels.inside

Logical. Whether to print construct labels next to the points. Can be useful during inspection of the plot (default FALSE).

c.lines Logical. Whether construct lines from the center of the biplot to the sourrounding box are drawn (default is FALSE).

col.c.lines The color of the construct lines from the center to the borders of the plot (default is gray(.9)).

flipaxes Logical vector of length two. Whether x and y axes are reversed (default is c(F,F)).

strokes.x Length of outer strokes in x direction in NDC. strokes.y Length of outer strokes in y direction in NDC.

offsetting Do offsetting? (TODO)

offset.labels Offsetting parameter for labels (TODO).
offset.e offsetting parameter for elements (TODO).

axis.ext Axis extension factor (default is .1). A bigger value will zoom out the plot.

mai Margins available for plotting the labels in inch (default is c(.2, 1.5, .2, 1.5)).

rect.margins Vector of length two (default is c(.07, .07)). Two values specifying the additional horizontal and vertical margin around each label.

srt Angle to rotate construct label text. Only used in case offsetting=FALSE.

cex.pos Cex parameter used during positioning of labels if prompted. Does usually not

have to be changed by user.

xpd Logical (default is TRUE). Wether to extend text labels over figure region. Usu-

ally not needed by the user.

unity Scale elements and constructs coordinates to unit scale in 2D (maximum of 1)

so they are printed more neatly (default TRUE).

unity3d Scale elements and constructs coordinates to unit scale in 3D (maximum of 1)

so they are printed more neatly (default TRUE).

scale.e	Scaling factor for element vectors. Will cause element points to move a bit more to the center. (but only if unity or unity3d is TRUE). This argument is for visual appeal only.
zoom	Scaling factor for all vectors. Can be used to zoom the plot in and out (default 1).
var.show	Show explained sum-of-squares in biplot? (default TRUE).
var.cex	The cex value for the percentages shown in the plot.
var.col	The color value of the percentages shown in the plot.
	parameters passed on to come.

# **Details**

For the construction of a biplot the grid matrix is first centered and normalized according to the prompted options.

Next, the matrix is decomposed by singular value decomposition (SVD) into

$$X = UDV^T$$

The biplot is made up of two matrices

$$X = GH^T$$

These matrices are construed on the basis of the SVD results.

$$\hat{X} = UD^g D^h V^T$$

Note that the grid matrix values are only recovered and the projection property is only given if g+h=1

# Author(s)

Mark Heckmann

#### See Also

```
Unsophisticated biplot: biplotSimple;
```

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

```
## Not run:
```

```
biplot2d(boeker)  # biplot of boeker data
biplot2d(boeker, c.lines=T)  # add construct lines
biplot2d(boeker, center=2)  # with column centering
biplot2d(boeker, center=4)  # midpoint centering
biplot2d(boeker, normalize=1)  # normalization of constructs
```

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```
biplot2d(boeker, dim=2:3)
                                  # plot 2nd and 3rd dimension
  biplot2d(boeker, dim=c(1,4))
                                  # plot 1st and 4th dimension
  biplot2d(boeker, g=1, h=1)
                                        # assign singular values to con. & elem.
  biplot2d(boeker, g=1, h=1, center=1) # row centering (Slater)
  biplot2d(boeker, g=1, h=1, center=4) # midpoint centering (ESA)
  biplot2d(boeker, e.color="red", c.color="blue")
                                                    # change colors
                                                    # mapped onto color range
  biplot2d(boeker, c.color=c("white", "darkred"))
  biplot2d(boeker, unity=T)
                                            # scale con. & elem. to equal length
  biplot2d(boeker, unity=T, scale.e=.5)
                                            # scaling factor for element vectors
  biplot2d(boeker, e.labels.show=F)
                                            # do not show element labels
  biplot2d(boeker, e.labels.show=c(1,2,4)) # show labels for elements 1, 2 and 4
  biplot2d(boeker, e.points.show=c(1,2,4)) # only show elements 1, 2 and 4
  biplot2d(boeker, c.labels.show=c(1:4))
                                            # show constructs labels 1 to 4
  biplot2d(boeker, c.labels.show=c(1:4))
                                            # show constructs labels except 1 to 4
  biplot2d(boeker, e.cex.map=1)
                                  # change size of texts for elements
  biplot2d(boeker, c.cex.map=1) # change size of texts for constructs
  biplot2d(boeker, g=1, h=1, c.labels.inside=T) # constructs inside the plot
  biplot2d(boeker, g=1, h=1, c.labels.inside=T, # different margins and elem. color
           mai=c(0,0,0,0), e.color="red")
  biplot2d(boeker, strokes.x=.3, strokes.y=.05) # change length of strokes
  biplot2d(boeker, flipaxes=c(T, F))
                                          # flip x axis
  biplot2d(boeker, flipaxes=c(T, T))
                                          # flip x and y axis
  biplot2d(boeker, outer.positioning=F)
                                          # no positioning of con.-labels
  biplot2d(boeker, c.labels.devangle=20) # only con. within 20 degree angle
## End(Not run)
```

biplot3d

Draw grid in rgl (3D device).

#### **Description**

The 3D biplot opens an interactive 3D device that can be rotated and zoomed using the mouse. A 3D device facilitates the exploration of grid data as significant proportions of the sum-of-squares are often represented beyond the first two dimensions. Also, in a lot of cases it may be of interest to explore the grid space from a certain angle, e.g. to gain an optimal view onto the set of elements under investigation (e.g. Raeithel, 1998).

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# Usage

```
biplot3d(x, dim = 1:3, labels.e = TRUE, labels.c = TRUE, lines.c = TRUE,
  lef = 1.3, center = 1, normalize = 0, g = 0, h = 1,
  col.active = NA, col.passive = NA, c.sphere.col = grey(0.4),
  c.cex = 0.6, c.text.col = grey(0.4), e.sphere.col = grey(0),
  e.cex = 0.6, e.text.col = grey(0), alpha.sphere = 0.05,
  col.sphere = "black", unity = FALSE, unity3d = FALSE, scale.e = 0.9,
  zoom = 1, ...)
```

# Arguments

guments	
Х	repgrid object.
dim	Dimensions to display.
labels.e	Logical. whether element labels are displayed.
labels.c	Logical. whether construct labels are displayed.
lines.c	Numeric. The way lines are drawn through the construct vectors. $\emptyset$ = no lines, 1 = lines from constructs to outer frame, 2 = lines from the center to outer frame.
lef	Construct lines extension factor
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). Default is 1 (row centering).
normalize	A numeric value indicating along what direction (rows, columns) to normalize by standard deviations. $0 = \text{none}$ , $1 = \text{rows}$ , $2 = \text{columns}$ (default is $0$ ).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
col.active	Columns (elements) that are no supplementary points, i.e. they are used in the SVD to find principal components. default is to use all elements.
col.passive	Columns (elements) that are supplementary points, i.e. they are NOT used in the SVD but projecte into the component space afterwards. They do not determine the solution. Default is NA, i.e. no elements are set supplementary.
c.sphere.col	Color of construct spheres.
c.cex	Size of construct text.
c.text.col	Color for construct text.
e.sphere.col	Color of elements.
e.cex	Size of element labels.
e.text.col	Color of element labels.
alpha.sphere	Numeric. alpha blending of the sourrounding sphere (default".05").
col.sphere	Color of sourrouding sphere (default"black").

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unity	Scale elements and constructs coordinates to unit scale (maximum of 1) so they are printed more neatly (default TRUE).
unity3d	To come.
scale.e	Scaling factor for element vectors. Will cause element points to move a bit more to the center (but only if unity or unity3d is TRUE). This argument is for visual appeal only.
zoom	Not yet used. Scaling factor for all vectors. Can be used to zoom the plot in and out (default 1).
	Parameters to be passed on.

## Author(s)

Mark Heckmann

#### References

Raeithel, A. (1998). Kooperative Modellproduktion von Professionellen und Klienten - erlauetert am Beispiel des Repertory Grid. *Selbstorganisation, Kooperation, Zeichenprozess: Arbeiten zu einer kulturwissenschaftlichen, anwendungsbezogenen Psychologie* (pp. 209-254). Opladen: Westdeutscher Verlag.

# See Also

```
Unsophisticated biplot: biplotSimple;
2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;
Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;
Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;
Function to set view in 3D: home.
```

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biplotEsa2d	Plot an eigenstructure analysis (ESA) biplot in 2D.	

## **Description**

The ESA is a special type of biplot suggested by Raeithel (e.g. 1998). It uses midpoint centering as a default. Note that the eigenstructure analysis is just a special case of a biplot that can also be produced using the biplot2d function with the arguments center=4, g=1, h=1. Here, only the arguments that are modified for the ESA biplot are described. To see all the parameters that can be changed see biplot2d.

#### Usage

```
biplotEsa2d(x, center = 4, g = 1, h = 1, ...)
```

# **Arguments**

x	repgrid object.
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). Eigenstructure analysis uses midpoint centering (4).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs. Eigenstructure analysis uses g=1.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements. Eigenstructure analyis uses h=1.
• • •	Additional parameters for be passed to biplot2d.

#### Author(s)

Mark Heckmann

# References

Raeithel, A. (1998). Kooperative Modellproduktion von Professionellen und Klienten. Erlaeutert am Beispiel des Repertory Grid. In A. Raeithel (1998). Selbstorganisation, Kooperation, Zeichenprozess. Arbeiten zu einer kulturwissenschaftlichen, anwendungsbezogenen Psychologie (p. 209-254). Opladen: Westdeutscher Verlag.

#### See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

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# **Examples**

```
## Not run:
    # See examples in \code{\link{biplot2d}} as the same arguments
    # can used for this function.
## End(Not run)
```

biplotEsa3d

Draw the eigenstructure analysis (ESA) biplot in rgl (3D device).

# **Description**

The 3D biplot opens an interactive 3D device that can be rotated and zoomed using the mouse. A 3D device facilitates the exploration of grid data as significant proportions of the sum-of-squares are often represented beyond the first two dimensions. Also, in a lot of cases it may be of interest to explore the grid space from a certain angle, e.g. to gain an optimal view onto the set of elements under investigation (e.g. Raeithel, 1998). Note that the eigenstructure analysisis just a special case of a biplot that can also be produced using the biplot3d function with the arguments center=4, g=1, h=1.

## Usage

```
biplotEsa3d(x, center = 1, g = 1, h = 1, ...)
```

# **Arguments**

Х	repgrid object.
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). Default is 4 (scale midpoint centering).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
	Additional arguments to be passed to biplot3d.

# Author(s)

Mark Heckmann

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## See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

# **Examples**

biplotEsaPseudo3d

Plot an eigenstructure analysis (ESA) in 2D grid with 3D impression (pseudo 3D).

# **Description**

The ESA is a special type of biplot suggested by Raeithel (e.g. 1998). It uses midpoint centering as a default. Note that the eigenstructure analysis is just a special case of a biplot that can also be produced using the biplot2d function with the arguments center=4, g=1, h=1. Here, only the arguments that are modified for the ESA biplot are described. To see all the parameters that can be changed see biplot2d and biplotPseudo3d.

# Usage

```
biplotEsaPseudo3d(x, center = 4, g = 1, h = 1, ...)
```

# **Arguments**

X	repgrid object.
---	-----------------

center Numeric. The type of centering to be performed. 0= no centering, 1= row

mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (con-

structs). Eigenstructure analyis uses midpoint centering (4).

g Power of the singular value matrix assigned to the left singular vectors, i.e. the

constructs. Eigenstructure analyis uses g=1.

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h Power of the singular value matrix assigned to the right singular vectors, i.e. the elements. Eigenstructure analysi uses h=1.

... Additional parameters for be passed to biplotPseudo3d.

## Author(s)

Mark Heckmann

#### See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

## **Examples**

```
## Not run:
    # See examples in \code{\link{biplotPseudo3d}} as the same arguments
    # can used for this function.
## End(Not run)
```

biplotPseudo3d

See biplotPseudo3d for its use. Draws a biplot of the grid in 2D with depth impression (pseudo 3D).

# Description

This version is basically a 2D biplot. It only modifies color and size of the symbols in order to create a 3D impression of the data points. This function will call the standard biplot2d function with some modified arguments. For the whole set of arguments that can be used see biplot2d. Here only the arguments special to biplotPseudo3d are outlined.

# Usage

```
biplotPseudo3d(x, dim = 1:2, map.dim = 3, e.point.col = c("white",
   "black"), e.point.cex = c(0.6, 1.2), e.label.col = c("white", "black"),
   e.label.cex = c(0.6, 0.8), e.color.map = c(0.4, 1),
   c.point.col = c("white", "darkred"), c.point.cex = c(0.6, 1.2),
   c.label.col = c("white", "darkred"), c.label.cex = c(0.6, 0.8),
   c.color.map = c(0.4, 1), ...)
```

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#### **Arguments**

x repgrid object.

dim Dimensions (i.e. principal components) to be used for biplot (default is c(1,2)).

map.dim Third dimension (depth) used to map aesthetic attributes to (default is 3).

Color(s) of the element symbols. Two values can be entered that will create a color ramp. The values of map.dim are mapped onto the ramp. The default is c("white", "black"). If only one color color value is supplied (e.g. "black") no mapping occurs and all elements will have the same color irrespective of their value on the map.dim dimension.

Size of the element symbols. Two values can be entered that will represents the lower and upper size of a range of cex the values of map.dim are mapped onto. The default is c(.6, 1.2). If only one cex value is supplied (e.g. .7) no mapping occurs and all elements will have the same size irrespective of their value on the map.dim dimension.

Color(s) of the element labels. Two values can be entered that will create a color ramp. The values of map.dim are mapped onto the ramp. The default is c("white", "black"). If only one color color value is supplied (e.g. "black") no mapping occurs and all element labels will have the same color irrespective of their value on the map.dim dimension.

Size of the element labels. Two values can be entered that will represents the lower and upper size of a range of cex the values of map.dim are mapped onto. The default is c(.6, .8). If only one cex value is supplied (e.g. .7) no mapping occurs and all element labels will have the same size irrespective of their value on the map.dim dimension.

Value range to determine what range of the color ramp defined in e.color will be used for mapping the colors. Default is c(.4, ,1). Usually not important for the user.

Color(s) of the construct symbols. Two values can be entered that will create a color ramp. The values of map.dim are mapped onto the ramp. The default is c("white", "darkred"). If only one color color value is supplied (e.g. "black") no mapping occurs and all elements will have the same color irrespective of their value on the map.dim dimension.

Size of the construct symbols. Two values can be entered that will represents the lower and upper size of a range of cex the values of map.dim are mapped onto. The default is  $c(.6,\ 1.2)$ . If only one cex value is supplied (e.g. .7) no mapping occurs and all elements will have the same size irrespective of their value on the map.dim dimension.

Color(s) of the construct labels. Two values can be entered that will create a color ramp. The values of map.dim are mapped onto the ramp. The default is c("white", "black"). If only one color color value is supplied (e.g. "black") no mapping occurs and all construct labels will have the same color irrespective of their value on the map.dim dimension.

Size of the construct labels. Two values can be entered that will represents the lower and upper size of a range of cex the values of map.dim are mapped onto. The default is c(.6, .9). If only one cex value is supplied (e.g. .7) no mapping

e.point.col

e.point.cex

e.label.col

e.label.cex

e.color.map

c.point.col

c.point.cex

c.label.col

c.label.cex

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occurs and all construct labels will have the same size irrespective of their value on the map.dim dimension.

c.color.map

Value range to determine what range of the color ramp defined in c.color will be used for mapping. Default is c(.4, ,1). Usually not important for the user.

... Additional parameters passed to biplot2d.

#### Author(s)

Mark Heckmann

# See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

```
## Not run:
  # biplot with 3D impression
  biplotPseudo3d(boeker)
   # Slater's biplot with 3D impression
  biplotPseudo3d(boeker, g=1, h=1, center=1)
  # show 2nd and 3rd dim. and map 4th
  biplotPseudo3d(boeker, dim=2:3, map.dim=4)
   # change elem. colors
  biplotPseudo3d(boeker, e.color=c("white", "darkgreen"))
   # change con. colors
  biplotPseudo3d(boeker, c.color=c("white", "darkgreen"))
  # change color mapping range
  biplotPseudo3d(boeker, c.colors.map=c(0, 1))
   # set uniform con. text size
  biplotPseudo3d(boeker, c.cex=1)
  # change text size mapping range
  biplotPseudo3d(boeker, c.cex=c(.4, 1.2))
## End(Not run)
```

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# **Description**

It will draw elements and constructs vectors using similar arguments as biplot2d. It is a version for quick exploration used during development.

# Usage

```
biplotSimple(x, dim = 1:2, center = 1, normalize = 0, g = 0, h = 1 -
g, unity = T, col.active = NA, col.passive = NA, scale.e = 0.9,
zoom = 1, e.point.col = "black", e.point.cex = 1,
e.label.col = "black", e.label.cex = 0.7, c.point.col = grey(0.6),
c.label.col = grey(0.6), c.label.cex = 0.6, ...)
```

# **Arguments**

rguments	
х	repgrid object.
dim	Dimensions (i.e. principal components) to be used for biplot (default is $c(1,2)$ ).
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). The default is 1 (row centering).
normalize	A numeric value indicating along what direction (rows, columns) to normalize by standard deviations. $0 = \text{none}$ , $1 = \text{rows}$ , $2 = \text{columns}$ (default is $0$ ).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
unity	Scale elements and constructs coordinates to unit scale in 2D (maximum of 1) so they are printed more neatly (default TRUE).
col.active	Columns (elements) that are no supplementary points, i.e. they are used in the SVD to find principal components. default is to use all elements.
col.passive	Columns (elements) that are supplementary points, i.e. they are NOT used in the SVD but projecte into the component space afterwards. They do not determine the solution. Default is NA, i.e. no elements are set supplementary.
scale.e	Scaling factor for element vectors. Will cause element points to move a bit more to the center. This argument is for visual appeal only.
ZOOM	Scaling factor for all vectors. Can be used to zoom the plot in and out (default 1).
e.point.col	Color of the element symbols (default is "black".
e.point.cex	Size of the element symbol (default is 1.
e.label.col	Color of the element labels (default is "black".
e.label.cex	Size of the element labels (default is .7.
c.point.col	Color of the construct lines (default is grey(.6).
c.label.col	Color of the construct labels (default is grey(.6).
c.label.cex	Size of the costruct labels (default is . 6.
• • •	Parameters to be passed on to center() and normalize.

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## Value

```
repgrid object.
```

#### Author(s)

Mark Heckmann

#### See Also

```
Unsophisticated biplot: biplotSimple;
```

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

### **Examples**

```
## Not run:
  biplotSimple(boeker)
  biplotSimple(boeker, unity=F)
  biplotSimple(boeker, g=1, h=1)
                                               # INGRID biplot
  biplotSimple(boeker, g=1, h=1, center=4)
                                               # ESA biplot
  biplotSimple(boeker, zoom=.9)
                                               # zooming out
                                               # scale element vectors
  biplotSimple(boeker, scale.e=.6)
  biplotSimple(boeker, e.point.col="brown")
                                               # change colors
  biplotSimple(boeker, e.point.col="brown",
                c.label.col="darkblue")
## End(Not run)
```

biplotSlater2d

Draws Slater's INGRID biplot in 2D.

## **Description**

The default is to use row centering and no normalization. Note that Slater's biplot is just a special case of a biplot that can be produced using the biplot2d function with the arguments center=1, g=1, h=1. The arguments that can be used in this function are the same as in biplot2d. Here, only the arguments that are set for Slater's biplot are described. To see all the parameters that can be changed see biplot2d.

#### Usage

```
biplotSlater2d(x, center = 1, g = 1, h = 1, ...)
```

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## **Arguments**

x	repgrid object.
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). Slater's biplot uses 1 (row centering).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
	Additional parameters for be passed to biplot2d.

## Author(s)

Mark Heckmann

#### See Also

```
Unsophisticated biplot: biplotSimple;
```

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

# Examples

```
## Not run:
    # See examples in \code{\link{biplot2d}} as the same arguments
    # can used for this function.
## End(Not run)
```

biplotSlater3d

*Draw the Slater's INGRID biplot in rgl (3D device).* 

# Description

The 3D biplot opens an interactive 3D device that can be rotated and zoomed using the mouse. A 3D device facilitates the exploration of grid data as significant proportions of the sum-of-squares are often represented beyond the first two dimensions. Also, in a lot of cases it may be of interest to explore the grid space from a certain angle, e.g. to gain an optimal view onto the set of elements under investigation (e.g. Raeithel, 1998). Note that Slater's biplot is just a special case of a biplot that can be produced using the biplot3d function with the arguments center=1, g=1, h=1.

32 biplotSlaterPseudo3d

## Usage

```
biplotSlater3d(x, center = 1, g = 1, h = 1, ...)
```

# Arguments

x repgrid object.

center Numeric. The type of centering to be performed. 0= no centering, 1= row

mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (con-

structs). Default is 1 (row i.e. construct centering).

g Power of the singular value matrix assigned to the left singular vectors, i.e. the

constructs.

h Power of the singular value matrix assigned to the right singular vectors, i.e. the

elements.

.. Additional arguments to be passed to biplot3d.

#### See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

biplotSlaterPseudo3d 33

# **Description**

The default is to use row centering and no normalization. Note that Slater's biplot is just a special case of a biplot that can be produced using the biplotPseudo3d function with the arguments center=1, g=1, h=1. Here, only the arguments that are modified for Slater's biplot are described. To see all the parameters that can be changed see biplot2d and biplotPseudo3d.

# Usage

```
biplotSlaterPseudo3d(x, center = 1, g = 1, h = 1, ...)
```

# **Arguments**

x	repgrid object.
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). Slater's biplot uses 1 (row centering).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
	Additional parameters for be passed to biplotPseudo3d.

# Author(s)

Mark Heckmann

# See Also

Unsophisticated biplot: biplotSimple;

2D biplots: biplot2d, biplotEsa2d, biplotSlater2d;

Pseudo 3D biplots: biplotPseudo3d, biplotEsaPseudo3d, biplotSlaterPseudo3d;

Interactive 3D biplots: biplot3d, biplotEsa3d, biplotSlater3d;

Function to set view in 3D: home.

```
## Not run:
    # See examples in \code{\link{biplotPseudo3d}} as the same arguments
    # can used for this function.
## End(Not run)
```

34 center

center

Centering of rows (constructs) and/or columns (elements).

# Description

Centering of rows (constructs) and/or columns (elements).

# Usage

```
center(x, center = 1, ...)
```

# Arguments

x repgrid object.

center Numeric. The type of centering to be performed. 0= no centering, 1= row

mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (con-

structs). of the scale(default FALSE). Default is 1 (row centering).

... Not evaluated.

### Value

matrix containing the transformed values.

## Note

If scale midpoint centering is applied no row or column centering can be applied simultaneously. TODO: After centering the standard representation mode does not work any more as it remains unclear what color values to attach to the centered values.

# Author(s)

Mark Heckmann

```
## Not run:

center(bell2010)  # no centering
center(bell2010, rows=T)  # row centering of grid
center(bell2010, cols=T)  # column centering of grid
center(bell2010, rows=T, cols=T)  # row and column centering
## End(Not run)
```

cluster 35

cluster	Cluster analysis (of constructs or elements).

## **Description**

cluster is a preliminary implementation of a cluster function. It supports various distance measures as well as cluster methods. More is to come.

**align**: Aligning will reverse constructs if necessary to yield a maximal similarity between constructs. In a first step the constructs are clustered including both directions. In a second step the direction of a construct that yields smaller distances to the adjacent constructs is preserved and used for the final clustering. As a result, every construct is included once but with an orientation that guarantees optimal clustering. This approach is akin to the procedure used in FOCUS (Jankowicz & Thomas, 1982).

# Usage

```
cluster(x, along = 0, dmethod = "euclidean", cmethod = "ward", p = 2,
    align = TRUE, trim = NA, main = NULL, mar = c(4, 2, 3, 15), cex = 0,
    lab.cex = 0.8, cex.main = 0.9, print = TRUE, ...)
```

# **Arguments**

x	repgrid object.
along	Along which dimension to cluster. $1 = \text{constructs only}$ , $2 = \text{elements only}$ , $0 = \text{both (default)}$ .
dmethod	The distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". Any unambiguous substring can be given. For additional information on the different types type ?dist.
cmethod	The agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid".
p	The power of the Minkowski distance, in case "minkowski" is used as argument for dmethod.
align	Whether the constructs should be aligned before clustering (default is TRUE). If not, the grid matrix is clustered as is. See Details section for more information.
trim	the number of characters a construct is trimmed to (default is 10). If NA no trimming is done. Trimming simply saves space when displaying the output.
main	Title of plot. The default is a name indicating the distance function and cluster method.
mar	Define the plot region (bottom, left, upper, right).
cex	Size parameter for the nodes. Usually not needed.
lab.cex	Size parameter for the constructs on the right side.
cex.main	Size parameter for the plot title (default is .9).

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 $\label{eq:continuous} \mbox{ Logical. We ther to print the dendrogram (default is TRUE)}.$ 

. . . Additional parameters to be passed to plotting function from as . dendrogram.

Type ?as . dendrogram for further information. This option is usually not needed,

except if special designs are needed.

#### Value

Reordered repgrid object.

#### Author(s)

Mark Heckmann

#### References

Jankowicz, D., & Thomas, L. (1982). An Algorithm for the Cluster Analysis of Repertory Grids in Human Resource Development. *Personnel Review*, 11(4), 15-22. doi:10.1108/eb055464.

#### See Also

bertinCluster

# **Examples**

```
## Not run:
 cluster(bell2010)
 cluster(bell2010, main="My cluster analysis")
                                                  # new title
 cluster(bell2010, type="t")
                                                  # different drawing style
 cluster(bell2010, dmethod="manhattan")
                                                  # using manhattan metric
 cluster(bell2010, cmethod="single")
                                                  # do single linkage clustering
 cluster(bell2010, cex=1, lab.cex=1)
                                                  # change appearance
 cluster(bell2010, lab.cex=.7,
                                                  # advanced appearance changes
          edgePar = list(lty=1:2, col=2:1))
## End(Not run)
```

clusterBoot

Multiscale bootstrap cluster analysis.

# **Description**

p-values are calculated for each branch of the cluster dendrogram to indicate the stability of a specific partition. clusterBoot will yield the same clusters as the cluster function (i.e. standard hierarchical clustering) with additional p-values. Two kindes of p-values are reported: bootstrap probabilities (BP) and approximately unbiased (AU) probabilities (see Details section for more information).

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### Usage

```
clusterBoot(x, along = 1, align = TRUE, dmethod = "euclidean",
  cmethod = "ward", p = 2, nboot = 1000, r = seq(0.8, 1.4, by = 0.1),
  seed = NULL, ...)
```

#### **Arguments**

Х	grid object
along	Along which dimension to cluster. $1 = constructs$ , $2 = elements$ .
align	Whether the constructs should be aligned before clustering (default is TRUE). If not, the grid matrix is clustered as is. See Details section for more information.
dmethod	The distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". Any unambiguous substring can be given. For additional information on the different types type ?dist.
cmethod	The agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid".
p	Power of the Minkowski metric. Not yet passed on to pvclust!
nboot	the number of bootstrap replications. The default is 1000.
r	numeric vector which specifies the relative sample sizes of bootstrap replications. For original sample size $n$ and bootstrap sample size $n'$ , this is defined as $r=n'/n$ .
seed	Random seed for bootstrapping. Can be set for reproducibility (see set.seed). Usually not needed.
	Arguments to pass on to pvclust.

#### **Details**

In standard (hierarchical) cluster analysis the question arises which of the idientified structures are significant or just emerged by chance. Over the last decade several methods have been developed to test structures for robustness. One line of research in this area is based on resampling. The idea is to resample the rows or columns of the data matrix and to build the dendrogram for each bootstrap sample (Felsenstein, 1985). The p-values indicates the pecentage of times a specific structure is identified across the bootstrap samples. It was shown that the p-value is biased (Hillis & Bull, 1993; Zharkikh & Li, 1995). In the literature several methods for bias correction have been proposed. In clusterBoot a method based on the *multiscale bootstrap* is used to derive corrected (approximately unbiased) p-values (Shimodaira, 2002, 2004). In conventional bootstrap analysis the size of the bootstrap sample is identical to the original sample size. Multiscale bootstrap varies the bootstrap sample size in order to infer a correction formula for the biased p-value on the basis of the variation of the results for the different sample sizes (Suzuki & Shimodaira, 2006).

align: Aligning will reverse constructs if necessary to yield a maximal similarity between constructs. In a first step the constructs are clustered including both directions. In a second step the direction of a construct that yields smaller distances to the adjacent constructs is preserved and used for the final clustering. As a result, every construct is included once but with an orientation that guarantees optimal clustering. This approach is akin to the procedure used in FOCUS (Jankowicz & Thomas, 1982).

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#### Value

A pyclust object as returned by the function pyclust

#### Author(s)

Mark Heckmann

#### References

Felsenstein, J. (1985). Confidence Limits on Phylogenies: An Approach Using the Bootstrap. *Evolution*, 39(4), 783. doi:10.2307/2408678

Hillis, D. M., & Bull, J. J. (1993). An Empirical Test of Bootstrapping as a Method for Assessing Confidence in Phylogenetic Analysis. *Systematic Biology*, 42(2), 182-192.

Jankowicz, D., & Thomas, L. (1982). An Algorithm for the Cluster Analysis of Repertory Grids in Human Resource Development. *Personnel Review, 11*(4), 15-22. doi:10.1108/eb055464.

Shimodaira, H. (2002) An approximately unbiased test of phylogenetic tree selection. *Syst, Biol.*, 51, 492-508.

Shimodaira,H. (2004) Approximately unbiased tests of regions using multistep- multiscale bootstrap resampling. *Ann. Stat.*, *32*, 2616-2614.

Suzuki, R., & Shimodaira, H. (2006). Pvclust: an R package for assessing the uncertainty in hierarchical clustering. *Bioinformatics*, 22(12), 1540-1542. doi:10.1093/bioinformatics/btl117

Zharkikh, A., & Li, W.-H. (1995). Estimation of confidence in phylogeny: the complete-and-partial bootstrap technique. *Molecular Phylogenetic Evolution*, *4*(1), 44-63.

```
## Not run:

# pvclust must be loaded
library(pvclust)

# p-values for construct dendrogram
s <- clusterBoot(boeker)
plot(s)
pvrect(s, max.only=FALSE)

# p-values for element dendrogram
s <- clusterBoot(boeker, along=2)
plot(s)
pvrect(s, max.only=FALSE)

## End(Not run)</pre>
```

constructCor 39

constructCor	Calculate correlations between constructs.	
--------------	--	--

## **Description**

Different types of correlations can be requested: PMC, Kendall tau rank correlation, Spearman rank correlation.

### Usage

```
constructCor(x, method = c("pearson", "kendall", "spearman"), trim = 20,
  index = FALSE)
```

#### **Arguments**

x repgrid object.

method A character string indicating which correlation coefficient is to be computed.

One of "pearson" (default), "kendall" or "spearman", can be abbreviated.

The default is "pearson".

trim The number of characters a construct is trimmed to (default is 20). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs with long names.

index Whether to print the number of the construct.

### Value

Returns a matrix of construct correlations.

### Author(s)

Mark Heckmann

### See Also

elementCor

```
# three different types of correlations
constructCor(mackay1992)
constructCor(mackay1992, method="kendall")
constructCor(mackay1992, method="spearman")
# format output
constructCor(mackay1992, trim=6)
constructCor(mackay1992, index=TRUE, trim=6)
```

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```
# save correlation matrix for further processing
r <- constructCor(mackay1992)
r
print(r, digits=5)
# accessing the correlation matrix
r[1, 3]</pre>
```

constructD

Calculate Somers' d for the constructs.

# **Description**

Somer'ss d is an assymetric association measure as it depends on which variable is set as dependent and independent. The direction of dependency needs to be specified.

## Usage

```
constructD(x, dependent = "columns", trim = 30, index = TRUE)
```

### **Arguments**

x repgrid object

dependent A string denoting the direction of dependency in the output table (as d is assy-

metrical). Possible values are "columns" (the default) for setting the columns as dependent, "rows" for setting the rows as the dependent variable and "symmetric" for the symmetrical Somers' d measure (the mean of the two directional values

for code"columns" and "rows").

trim The number of characters a construct is trimmed to (default is 30). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs with long names.

index Whether to print the number of the construct (default is TRUE).

#### Value

matrix of construct correlations.

### Note

Thanks to Marc Schwartz for supplying the code to calculate Somers' d.

### Author(s)

Mark Heckmann

constructPca 41

### References

Somers, R. H. (1962). A New Asymmetric Measure of Association for Ordinal Variables. *American Sociological Review*, 27(6), 799-811.

# Examples

```
## Not run:
    constructD(fbb2003)  # columns as dependent (default)
    constructD(fbb2003, "c")  # row as dependent
    constructD(fbb2003, "s")  # symmetrical index

# surpress printing
    d <- constructD(fbb2003, out=0, trim=5)
    d

# more digits
    constructD(fbb2003, dig=3)

# add index column, no trimming
    constructD(fbb2003, col.index=TRUE, index=F, trim=NA)

## End(Not run)</pre>
```

constructPca

Principal component analysis (PCA) of inter-construct correlations.

### **Description**

Various methods for rotation and methods for the calculation of the correlations are available. Note that the number of factors has to be specified. For more information on the PCA function itself type ?principal.

### Usage

```
constructPca(x, nfactors = 3, rotate = "varimax", method = "pearson",
    trim = NA)
```

## **Arguments**

```
x repgrid object.
nfactors Number of components to extract (default is 3).
rotate "none", "varimax", "promax" and "cluster" are possible rotations (default is none).
```

42 constructPca

method A character string indicating which correlation coefficient is to be computed.

One of "pearson" (default), "kendall" or "spearman", can be abbreviated.

The default is "pearson".

trim The number of characters a construct is trimmed to (default is 7). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs with long names.

#### Value

Returns an object of class constructPca.

#### Author(s)

Mark Heckmann

#### References

Fransella, F., Bell, R. & Bannister, D. (2003). *A Manual for Repertory Grid Technique* (2. Ed.). Chichester: John Wiley & Sons.

## See Also

To extract the PCA loadings for further processing see constructPcaLoadings.

```
## Not run:
    constructPca(bell2010)

# data from grid manual by Fransella et al. (2003, p. 87)
# note that the construct order is different
    constructPca(fbb2003, nfactors=2)

# no rotation
    constructPca(fbb2003, rotate="none")

# use a different type of correlation (Spearman)
    constructPca(fbb2003, method="spearman")

# save output to object
    m <- constructPca(fbb2003, nfactors=2)
    m

# different printing options
    print(m, digits=5)
    print(m, cutoff=.3)

## End(Not run)</pre>
```

constructPcaLoadings 43

## **Description**

Extract loadings from PCA of constructs.

# Usage

```
constructPcaLoadings(x)
```

## **Arguments**

Х

repgrid object. This object is returned by the function constructPca.

## Value

A matrix containing the factor loadings.

# **Examples**

```
p <- constructPca(bell2010)
l <- constructPcaLoadings(p)
l[1, ]
l[, 1]
l[1,1]</pre>
```

constructRmsCor

Root mean square (RMS) of inter-construct correlations.

## **Description**

The RMS is also known as 'quadratic mean' of the inter-construct correlations. The RMS serves as a simplification of the correlation table. It reflects the average relation of one construct to all other constructs. Note that as the correlations are squared during its calculation, the RMS is not affected by the sign of the correlation (cf. Fransella, Bell & Bannister, 2003, p. 86).

## Usage

```
constructRmsCor(x, method = "pearson", trim = NA)
```

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### **Arguments**

x repgrid object

method A character string indicating which correlation coefficient is to be computed.

One of "pearson" (default), "kendall" or "spearman", can be abbreviated.

The default is "pearson".

trim The number of characters a construct is trimmed to (default is NA). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs with long names.

#### Value

dataframe of the RMS of inter-construct correlations

### Author(s)

Mark Heckmann

### References

Fransella, F., Bell, R. C., & Bannister, D. (2003). A Manual for Repertory Grid Technique (2. Ed.). Chichester: John Wiley & Sons.

#### See Also

elementRmsCor, constructCor

### **Examples**

```
# data from grid manual by Fransella, Bell and Bannister
constructRmsCor(fbb2003)
constructRmsCor(fbb2003, trim=20)

# modify output
r <- constructRmsCor(fbb2003)
print(r, digits=5)
# access calculation results
r[2, 1]</pre>
```

data-bell2010

Grid data from Bell (2010).

## **Description**

Grid data orginated (but is not shown in the paper) from a study by Haritos, Gindinis, Doan and Bell (2004) on element role titles. It was used to demonstrate the effects of construct alignment in Bell (2010, p. 46).

data-bellmcgorry1992 45

#### References

Bell, R. C. (2010). A note on aligning constructs. *Personal Construct Theory and Practice*, 7, 43-48.

Haritos, A., Gindidis, A., Doan, C., & Bell, R. C. (2004). The effect of element role titles on construct structure and content. *Journal of constructivist psychology*, 17(3), 221-236.

data-bellmcgorry1992 Grid data from Bell and McGorry (1992).

# **Description**

The grid data set is used in Bell's technical report "Using SPSS to Analyse Repertory Grid Data" (1997, p. 6). Originally, the data comes from a study by Bell and McGorry (1992).

#### References

Bell, R. C. (1977). *Using SPSS to Analyse Repertory Grid Data*. Technical Report, University of Melbourne.

Bell, R. C., & McGorry, P. (1992). The analysis of repertory grids used to monitor the perceptions of recovering psychotic patients. In A. Thomson & P. Cummins (Eds.), *European Perspectives in Personal Construct Psychology* (p. 137-150). Lincoln, UK: European Personal Construct Association.

data-boeker

Grid data from Boeker (1996).

### **Description**

Grid data from a schizophrenic patient undergoing psychoanalytically oriented psychotherapy. The data was taken during the last stage of therapy (Boeker, 1996, p. 163).

#### References

Boeker, H. (1996). The reconstruction of the self in the psychotherapy of chronic schizophrenia: a case study with the Repertory Grid Technique. In: Scheer, J. W., Catina, A. (Eds.): *Empirical Constructivism in Europe - The Personal Construct Approach* (p. 160-167). Giessen: Psychosozial-Verlag.

46 data-leach2001

data-fbb2003

Grid data from Fransella, Bell and Bannister (2003).

# **Description**

A dataset used throughout the book "A Manual for Repertory Grid Technique" (Fransella, Bell and Bannister, 2003, p. 60).

#### References

Fransella, F., Bell, R. & Bannister, D. (2003). A Manual for Repertory Grid Technique (2. Ed.). Chichester: John Wiley & Sons.

data-feixas2004

Grid data from Feixas and Saul (2004).

### **Description**

A desription by the authors: "When Teresa, 22 years old, was seen by the second author (LAS) at the psychological services of the University of Salamanca, she was in the final year of her studies in chemical sciences. Although Teresa proves to be an excellent student, she reveals serious doubts about her self worth. She cries frequently, and has great difficulty in meeting others, even though she has a boyfriend who is extremely supportive. Teresa is anxiously hesitant about accepting a new job which would involve moving to another city 600 Km away from home." (Feixas & Saul, 2004, p. 77).

## References

Feixas, G., & Saul, L. A. (2004). The Multi-Center Dilemma Project: an investigation on the role of cognitive conflicts in health. *The Spanish Journal of Psychology*, 7(1), 69-78.

data-leach2001

Pre- and post therapy dataset from Leach et al. (2001).

## **Description**

Case as described by the authors: "Sarah, aged 32, was referred with problems of depression and sexual difficulties relating to childhood sexual abuse. She had three children and was living with her male partner. From the age of 9, her brother, an adult, had sexually abused Sarah. She attended a group for survivors of child sexual abuse and completed repertory grids prior to the group, immediately after the group and at 3- and 6-month follow-up." (Leach et al. 2001, p. 230).

data-mackay1992 47

### **Details**

leach2001a is the pre-therapy, leach2001b is the post-therapy therapy datset. The construct and elements are identical.

#### References

Leach, C., Freshwater, K., Aldridge, J., & Sunderland, J. (2001). Analysis of repertory grids in clinical practice. *The British Journal of Clinical Psychology*, 40, 225-248.

data-mackay1992

Grid data from Mackay (1992). Data set 'Grid C'-

#### **Description**

used in Mackay's paper on inter-element correlation (1992, p. 65).

### References

Mackay, N. (1992). Identification, reflection, and correlation: Problems in the bases of repertory grid measures. *International Journal of Personal Construct Psychology*, 5(1), 57-75.

data-raeithel

Grid data from Raeithel (1998).

### **Description**

Grid data to demonstrate the use of Bertin diagrams (Raeithel, 1998, p. 223). The context of its administration is unknown.

#### References

Raeithel, A. (1998). Kooperative Modellproduktion von Professionellen und Klienten. Erlaeutert am Beispiel des Repertory Grid. In A. Raeithel (1998). Selbstorganisation, Kooperation, Zeichenprozess. Arbeiten zu einer kulturwissenschaftlichen, anwendungsbezogenen Psychologie (p. 209-254). Opladen: Westdeutscher Verlag.

data-slater1977a

Drug addict's grid data set from Slater (1977, p. 32).

## **Description**

Drug addict's grid data set from Slater (1977, p. 32).

#### References

Slater, P. (1977). The measurement of intrapersonal space by grid technique. London: Wiley.

48 distance

data-slater1977b Grid data from Slater (1977).

# **Description**

Grid data (ranked) from a seventeen year old female psychiatric patient (Slater, 1977, p. 110). She was depressed, anxious and took to cutting herself. The data was originally reported by Watson (1970).

### References

Slater, P. (1977). The measurement of intrapersonal space by grid technique. London: Wiley. Watson, J. P. (1970). The relationship between a self-mutilating patient and her doctor. *Psychotherapy and Psychosomatics*, 18(1), 67-73.

distance

Distance measures (between constructs or elements).

# Description

Various distance measures between elements or constructs are calculated.

## Usage

```
distance(x, along = 1, dmethod = "euclidean", p = 2, trim = 20,
  index = TRUE, ...)
```

## **Arguments**

X	repgrid object.
along	Whether to calculate distance for $1 = \text{constructs}$ (default) or for $2 = \text{elements}$ .
dmethod	The distance measure to be used. This must be one of "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". Any unambiguous substring can be given. For additional information on the different types type ?dist.
p	The power of the Minkowski distance, in case "minkowski" is used as argument for dmethod.
trim	The number of characters a construct or element is trimmed to (default is 20). If NA no trimming occurs. Trimming simply saves space when displaying correlation of constructs with long names.
index	Whether to print the number of the construct or element in front of the name (default is TRUE). This is useful to avoid identical row names, which may cause an error.
	Additional parameters to be passed to function dist. Type dist for further information.

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## Value

matrix object.

# Author(s)

Mark Heckmann

## **Examples**

```
## Not run:
   # between constructs
  distance(bell2010, along=1)
  # between elements
  distance(bell2010, along=2)
  # several distance methods
  distance(bell2010, dm="man")
                                        # manhattan distance
  distance(bell2010, dm="mink", p=3) # minkowski metric to the power of 3
  # to save the results without printing to the console
  d <- distance(bell2010, trim=7)</pre>
  d
  # some more options when printing the distance matrix
  print(d, digits=5)
  print(d, col.index=FALSE)
  print(d, upper=FALSE)
  # accessing entries from the matrix
  d[1,3]
## End(Not run)
```

distanceHartmann

'Hartmann distance' (standardized Slater distances).

# Description

Calculate Hartmann distance

## Usage

```
distanceHartmann(x, method = "paper", reps = 10000, prob = NULL,
    progress = TRUE, distributions = FALSE)
```

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#### **Arguments**

x repgrid object.

method The method used for distance calculation, on of "paper", "simulate", "new".

"paper" uses the parameters as given in Hartmann (1992) for caclulation. "simulate" (default) simulates a Slater distribution for the caclulation. In a future version the time consuming simulation will be replaced by more accurate parameters for

Hartmann distances than used in Hartmann (1992).

reps Number of random grids to generate sample distribution for Slater distances

(default is 10000). Note that a lot of samples may take a while to calculate.

prob The probability of each rating value to occur. If NULL (default) the distribution

is uniform. The number of values must match the length of the rating scale.

progress Whether to show a progress bar during simulation (default is TRUE) (for method="simulate").

May be useful when the distribution is estimated on the basis of many quasis.

distributions Wether to additionally return the values of the simulated distributions (Slater

etc.) The default is FALSE as it will quickly boost the object size.

#### **Details**

Hartmann (1992) showed in a simulation study that Slater distances (see distanceSlater) based on random grids, for which Slater coined the expression quasis, have a skewed distribution, a mean and a standard deviation depending on the number of constructs elicited. He suggested a linear transformation (z-transformation) which takes into account the estimated (or expected) mean and the standard deviation of the derived distribution to standardize Slater distance scores across different grid sizes. 'Hartmann distances' represent a more accurate version of 'Slater distances'. Note that Hartmann distances are multiplied by -1. Hence, negative Hartmann values represent dissimilarity, i.e. a big Slater distance.

There are two ways to use this function. Hartmann distances can either be calculated based on the reference values (i.e. means and standard deviations of Slater distance distributions) as given by Hartmann in his paper. The second option is to conduct an instant simulation for the supplied grid size for each calculation. The second option will be more accurate when a big number of quasis is used in the simulation.

It is also possible to return the quantiles of the sample distribution and only the element distances consideres 'significant' according to the quantiles defined.

### Value

A matrix containing Hartmann distances.

In the attributes several additional parameters can be found:

"arguments" A list of several parameters including mean and sd of Slater distribution.

"quantiles" Quantiles for Slater and Hartmann distance distribition.

"distributions"

List with values of the simulated distributions.

distanceHartmann 51

### Calculation

The 'Hartmann distance' is calculated as follows (Hartmann 1992, p. 49).

$$D = -1(\frac{D_{slater} - M_c}{sd_c})$$

Where  $D_{slater}$  denotes the Slater distances of the grid,  $M_c$  the sample distribution's mean value and  $sd_c$  the sample distributions's standard deviation.

## Author(s)

Mark Heckmann

#### References

Hartmann, A. (1992). Element comparisons in repertory grid technique: Results and consequences of a Monte Carlo study. *International Journal of Personal Construct Psychology*, 5(1), 41-56.

# See Also

distanceSlater

```
## Not run:
  ### basics ###
  distanceHartmann(bell2010)
  distanceHartmann(bell2010, method="simulate")
  h <- distanceHartmann(bell2010, method="simulate")</pre>
  # printing options
  print(h)
  print(h, digits=6)
  # 'significant' distances only
  print(h, p=c(.05, .95))
  # access cells of distance matrix
  h[1,2]
   ### advanced ###
  # histogram of Slater distances and indifference region
  h <- distanceHartmann(bell2010, distributions=TRUE)</pre>
  1 <- attr(h, "distributions")</pre>
  hist(l$slater, breaks=100)
  hist(l$hartmann, breaks=100)
## End(Not run)
```

52 distanceNormalized

distanceNormalized	Standardized inter-element distances' (power transformed Hartmann distances).

### Description

Calculate power-transformed Hartmann distances.

### Usage

```
distanceNormalized(x, reps = 1000, prob = NULL, progress = TRUE,
    distributions = TRUE)
```

# Arguments

_	
x	repgrid object.
reps	Number of random grids to generate to produce sample distribution for Hartmann distances (default is 1000). Note that a lot of samples may take a while to calculate.
prob	The probability of each rating value to occur. If NULL (default) the distribution is uniform. The number of values must match the length of the rating scale.
progress	Whether to show a progress bar during simulation (default is TRUE) (for method="simulate"). May be useful when the distribution is estimated on the basis of many quasis.
distributions	Wether to additionally return the values of the simulated distributions (Slater etc.) The default is FALSE as it will quickly boost the object size.

### **Details**

Hartmann (1992) suggested a transformation of Slater (1977) distances to make them independent from the size of a grid. Hartmann distances are supposed to yield stable cutoff values used to determine 'significance' of inter-element distances. It can be shown that Hartmann distances are still affected by grid parameters like size and the range of the rating scale used (Heckmann, 2012). The function distanceNormalize applies a Box-Cox (1964) transformation to the Hartmann distances in order to remove the skew of the Hartmann distance distribution. The normalized values show to have more stable cutoffs (quantiles) and better properties for comparison across grids of different size and scale range.

The function distanceNormalize can also return the quantiles of the sample distribution and only the element distances consideres 'significant' according to the quantiles defined.

### Value

A matrix containing the standardized distances. Further data is contained in the object's attributes:

distanceNormalized 53

```
"arguments" A list of several parameters including mean and sd of Slater distribution.

"quantiles" Quantiles for Slater, Hartmann and power transformed distance distribitions.

"distributions"
```

List with values of the simulated distributions, if distributions=TRUE.

#### **Calculations**

The 'power tranformed Hartmann distance' are calulated as follows: The simulated Hartmann distribution is added a constant as the Box-Cox transformation can only be applied to positive values. Then a range of values for lambda in the Box-Cox transformation (Box & Cox, 1964) are tried out. The best lambda is the one maximizing the correlation of the quantiles with the standard normal distribution. The lambda value maximizing normality is used to transform Hartmann distances. As the resulting scale of the power transformation depends on lambda, the resulting values are z-transformed to derive a common scaling.

The code for the calculation of the optimal lambda was written by Ioannis Kosmidis.

## Author(s)

Mark Heckmann

#### References

Box, G. E. P., & Cox, D. R. (1964). An Analysis of Transformations. *Journal of the Royal Statistical Society. Series B (Methodological)*, 26(2), 211-252.

Hartmann, A. (1992). Element comparisons in repertory grid technique: Results and consequences of a Monte Carlo study. *International Journal of Personal Construct Psychology*, 5(1), 41-56.

Heckmann, M. (2012). Standardizing inter-element distances in grids - A revision of Hartmann's distances, 11th Biennal Conference of the European Personal Construct Association (EPCA), Dublin, Ireland, Paper presentation, July 2012.

Slater, P. (1977). The measurement of intrapersonal space by Grid technique. London: Wiley.

### See Also

distanceHartmann and distanceSlater.

```
## Not run:
    ### basics ###
    distanceNormalized(bell2010)
    n <- distanceNormalized(bell2010)
    n

# printing options
    print(n)
    print(n, digits=4)
    # 'significant' distances only</pre>
```

54 distanceSlater

```
print(n, p=c(.05, .95))

# access cells of distance matrix
n[1,2]

### advanced ###

# histogram of Slater distances and indifference region
n <- distanceNormalized(bell2010, distributions=TRUE)
l <- attr(n, "distributions")
hist(l$bc, breaks=100)</pre>

## End(Not run)
```

distanceSlater

'Slater distances' (standardized Euclidean distances).

## **Description**

Calculate Slater distance.

#### Usage

```
distanceSlater(x, trim = 20, index = TRUE)
```

# **Arguments**

x repgrid object.

trim The number of characters a construct or element is trimmed to (default is 20). If

NA no trimming occurs. Trimming simply saves space when displaying correla-

tion of constructs with long names.

index Whether to print the number of the construct or element in front of the name

(default is TRUE). This is useful to avoid identical row names, which may cause

an error.

### **Details**

The euclidean distance is often used as a measure of similarity between elements (see distance. A drawback of this measure is that it depends on the range of the rating scale and the number of constructs used, i. e. on the size of a grid.

An approach to standardize the euclidean distance to make it independent from size and range of ratings and was proposed by Slater (1977, pp. 94). The 'Slater distance' is the Euclidean distance divided by the expected distance. Slater distances bigger than 1 are greater than expected, lesser than 1 are smaller than expected. The minimum value is 0 and values bigger than 2 are rarely found. Slater distances have been be used to compare inter-element distances between different grids, where the grids do not need to have the same constructs or elements. Hartmann (1992) showed that Slater

distanceSlater 55

distance is not independent of grid size. Also the distribution of the Slater distances is asymmetric. Hence, the upper and lower limit to infer 'significance' of distance is not symmetric. The practical relevance of Hartmann's findings have been demonstrated by Schoeneich and Klapp (1998). To calculate Hartmann's version of the standardized distances see distanceHartmann

#### Value

A matrix with Slater distances.

#### Calculation

The Slater distance is calculated as follows. For a derivation see Slater (1977, p.94). Let matrix D contain the row centered ratings. Then

$$P = D^T D$$

and

$$S = tr(P)$$

The expected 'unit of expected distance' results as

$$U = (2S/(m-1))^{1/2}$$

where m denotes the number of elements of the grid. The standardized Slater distances is the matrix of Euclidean distances E devided by the expected distance U.

#### Author(s)

Mark Heckmann

#### References

Hartmann, A. (1992). Element comparisons in repertory grid technique: Results and consequences of a Monte Carlo study. *International Journal of Personal Construct Psychology*, *5*(1), 41-56.

Schoeneich, F., & Klapp, B. F. (1998). Standardization of interelement distances in repertory grid technique and its consequences for psychological interpretation of self-identity plots: An empirical study. *Journal of Constructivist Psychology*, 11(1), 49-58.

Slater, P. (1977). The measurement of intrapersonal space by Grid technique. Vol. II. London: Wiley.

### See Also

distanceHartmann

56 elementCor

# **Examples**

```
distanceSlater(bell2010)
distanceSlater(bell2010, trim=40)

d <- distanceSlater(bell2010)
print(d)
print(d, digits=4)

# using Norris and Makhlouf-Norris (problematic) cutoffs
print(d, cutoffs=c(.8, 1.2))</pre>
```

elementCor

Calculate the correlations between elements.

# Description

Note that simple element correlations as a measure of similarity are flawed as they are not invariant to construct reflection (Mackay, 1992; Bell, 2010). A correlation index invariant to construct reflection is Cohen's rc measure (1969), which can be calculated using the argument rc=TRUE which is the default option.

## Usage

```
elementCor(x, rc = TRUE, method = "pearson", trim = 20, index = TRUE)
```

# Arguments

X	repgrid object.
rc	Use Cohen's rc which is invariant to construct reflection (see desciption above). It is used as the default.
method	A character string indicating which correlation coefficient is to be computed. One of "pearson" (default), "kendall" or "spearman", can be abbreviated. The default is "pearson".
trim	The number of characters a construct is trimmed to (default is 20). If NA no trimming occurs. Trimming simply saves space when displaying correlation of constructs with long names.
index	Whether to print the number of the construct.

#### Value

matrix of element correlations

# Author(s)

Mark Heckmann

elementRmsCor 57

#### References

Bell, R. C. (2010). A note on aligning constructs. *Personal Construct Theory & Practice*, (7), 42-48.

Cohen, J. (1969). rc: A profile similarity coefficient invariant over variable reflection. *Psychological Bulletin*, 71(4), 281-284.

Mackay, N. (1992). Identification, Reflection, and Correlation: Problems In The Bases Of Repertory Grid Measures. *International Journal of Personal Construct Psychology*, *5*(1), 57-75.

#### See Also

constructCor

#### **Examples**

```
elementCor(mackay1992)
                                             # Cohen's rc
elementCor(mackay1992, rc=FALSE)
                                             # PM correlation
elementCor(mackay1992, rc=FALSE, method="spearman") # Spearman correlation
# format output
elementCor(mackay1992, trim=6)
elementCor(mackay1992, index=FALSE, trim=6)
# save as object for further processing
r <- elementCor(mackay1992)</pre>
# change output of object
print(r, digits=5)
print(r, col.index=FALSE)
print(r, upper=FALSE)
# accessing elements of the correlation matrix
r[1,3]
```

Description

elementRmsCor

The RMS is also known as 'quadratic mean' of the inter-element correlations. The RMS serves as a simplification of the correlation table. It reflects the average relation of one element with all other elements. Note that as the correlations are squared during its calculation, the RMS is not affected by the sign of the correlation (cf. Fransella, Bell & Bannister, 2003, p. 86).

Root mean square (RMS) of inter-element correlations.

#### Usage

```
elementRmsCor(x, rc = TRUE, method = "pearson", trim = NA)
```

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### **Arguments**

x repgrid object.

rc Whether to use Cohen's rc which is invariant to construct reflection (see descip-

tion above). It is used as the default.

method A character string indicating which correlation coefficient to be computed. One

of "pearson" (default), "kendall" or "spearman", can be abbreviated. The

default is "pearson".

trim The number of characters an element is trimmed to (default is NA). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs with long names.

#### **Details**

Note that simple element correlations as a measure of similarity are flawed as they are not invariant to construct reflection (Mackay, 1992; Bell, 2010). A correlation index invariant to construct reflection is Cohen's rc measure (1969), which can be calculated using the argument rc=TRUE which is the default option in this function.

#### Value

dataframe of the RMS of inter-element correlations

#### Author(s)

Mark Heckmann

#### References

Fransella, F., Bell, R. C., & Bannister, D. (2003). A Manual for Repertory Grid Technique (2. Ed.). Chichester: John Wiley & Sons.

## See Also

```
constructRmsCor, elementCor
```

```
# data from grid manual by Fransella, Bell and Bannister
elementRmsCor(fbb2003)
elementRmsCor(fbb2003, trim=10)

# modify output
r <- elementRmsCor(fbb2003)
print(r, digits=5)

# access second row of calculation results
r[2, "RMS"]</pre>
```

home 59

home

Rotate the interactive 3D device to default views.

# Description

Rotate the interactive 3D device to a default viewpoint or to a position defined by theta and phi in Euler angles. Three default viewpoints are implemented rendering a view so that two axes span a plane and the third axis is poiting out of the screen.

## Usage

```
home(view = 1, theta = NULL, phi = NULL)
```

# Arguments

view Numeric. Specifying one of three default views. 1 = XY, 2=XZ and 3=YZ-

plane.

theta Numeric. Euler angle. Overrides view setting.

phi Numeric. Euler angle. Overrides view setting.

return NULL.

## Author(s)

Mark Heckmann

## See Also

Interactive 3D biplots: biplot3d, biplotSlater3d, biplotEsa3d.

```
## Not run:
biplot3d(boeker)
home(2)
home(3)
home(1)
home(theta=45, phi=45)
## End(Not run)
```

60 importExcel

importExcel Import grid data from an Excel file.
--

## **Description**

You can define a grid using Microsoft Excel and by saving it as a .xlsx file. The .xlsx file has to be in a specified fixed format (see section Details).

#### **Usage**

```
importExcel(file, dir = NULL, sheetIndex = 1, min = NULL, max = NULL)
```

## **Arguments**

file	A vector of filenames including the full path if file is not in current working directory. The file suffix has to be .xlsx (used since Excel 2007).
dir	Alternative way to supply the directory where the file is located (default NULL).
sheetIndex	The number of the Excel sheet that contains the grid data.
min	Optional argument (numeric, default NULL) for minimum rating value in grid.
max	Optional argument (numeric, default NULL) for maximum rating value in grid.

### **Details**

Excel file structure: The first row contains the minimum of the rating scale, the names of the elements and the maximum of the rating scale. Below every row contains the left construct pole, the ratings and the right construct pole.

```
E2
                     E3 E4
                                         5
left pole 1
                 5
                      3
                          4
                              right pole 1
left pole 2
                      1
                              right pole 2
             3
                 1
                          3
left pole 3
                 2
                      5
                              right pole 3
```

Note that the maximum and minimum value has to be defined using the min and max arguments if no values are supplied at the beginning and end of the first row. Otherwise the scaling range is inferred from the available data and a warning is issued as the range may be erroneous. This may effect other functions that depend on knowing the correct range and it is thus strongly recommended to set the scale range correctly.

# Value

A single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

importGridcor 61

### Author(s)

Mark Heckmann

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt

# **Examples**

```
## Not run:

# Open Excel file delivered along with the package
file <- system.file("extdata", "grid_01.xlsx", package = "OpenRepGrid")
rg <- importExcel(file)

# To see the structure of the Excel file try to open it as follows.
# Requires Excel to be installed.
system2("open", file)

# Import more than one Excel file
files <- system.file("extdata", c("grid_01.xlsx", "grid_02.xlsx") , package = "OpenRepGrid")
rg <- importExcel(files)

## End(Not run)</pre>
```

importGridcor

Import GRIDCOR data files.

### **Description**

Reads the file format that is used by the grid program GRIDCOR (Feixas & Cornejo, 2002).

### Usage

```
importGridcor(file, dir = NULL)
```

### **Arguments**

file filename including path if file is not in current working directory. File can also

be a complete URL. The fileformat is .dat.

dir alternative way to supply the directory where the file is located (default NULL).

### Value

a single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

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#### Note

Note that the GRIDCOR data sets the minimum ratings scale range to 1. The maximum value can differ and is defined in the data file.

Also note that both Gridcor and Gridstat data files do have the same suffix .dat. Make sure not to mix themn up.

### Author(s)

Mark Heckmann

### References

Feixas, G., & Cornejo, J. M. (2002). GRIDCOR: Correspondence Analysis for Grid Data (version 4.0). Barcelona: Centro de Terapia Cognitiva. Retrieved from http://www.terapiacognitiva.net/record/gridcor.htm.

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt, importExcel

```
## Not run:
# using the pop-up selection menu
rg <- importGridcor()

# supposing that the data file gridcor.dat is in the current directory
file <- "gridcor.dat"
rg <- importGridcor(file)

# specifying a directory (arbitrary example directory)
dir <- "/Users/markheckmann/data"
rg <- importGridcor(file, dir)

# using a full path
rg <- importGridcor("/Users/markheckmann/data/gridcor.dat")

# load GRIDCOR data from URL
rg <- importGridcor("http://www.openrepgrid.uni-bremen.de/data/gridcor.dat")

## End(Not run)</pre>
```

importGridstat 63

|--|

### **Description**

Reads the file format that is used by the latest version of the grid program gridstat (Bell, 1998).

### Usage

```
importGridstat(file, dir = NULL, min = NULL, max = NULL)
```

#### **Arguments**

file	Filename including path if file is not in current working directory. File can also be a complete URL. The fileformat is .dat. If no file is supplied a selection pop up menu is opened to select the files.
dir	Alternative way to supply the directory where the file is located (default NULL).
min	Optional argument (numeric, default NULL) for minimum rating value in grid.
max	Optional argument (numeric, default NULL) for maximum rating value in grid.

#### Value

A single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

#### Note

Note that the gridstat data format does not contain explicit information about the range of the rating scale used (minimum and maximum). By default the range is inferred by scanning the ratings and picking the minimal and maximal values as rating range. You can set the minimal and maximal value by hand using the min and max arguments or by using the setScale() function. Note that if the rating range is not set, it may cause several functions to not work properly. A warning will be issued if the range is not set explicitly when using the importing function.

The function only reads data from the latest GridStat version. The latest version allows the seperation of the left and right pole by using on of the following symbols /: - (hyphene, colon and dash). Older versions may not seperate the left and right pole. This will cause all labels to be assigned to the left pole only when importing. You may fix this by simply entering one of the construct seperator symbols into the GridStat file between each left and right construct pole.

The third line of a GridStat file may contain a no labels statement (i.e. a line containing any string of 'NOLA', 'NO L', 'NoLa', 'No L', 'Nola', 'No l', 'nola' or 'no l'). In this case only ratings are supplied, hence, default names are assigned to elements and constructs.

### Author(s)

Mark Heckmann

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### References

Bell, R. C. (1998) GRIDSTAT: A program for analysing the data of a repertory grid. Melbourne: Author

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt, importExcel

## **Examples**

```
## Not run:
# using the pop-up selection menu
rg <- importGridstat()</pre>
# supposing that the data file gridstat.dat is in the current working directory
file <- "gridstat.dat"
rg <- importGridstat(file)</pre>
# specifying a directory (example)
dir <- "/Users/markheckmann/data"</pre>
rg <- importGridstat(file, dir)</pre>
# using a full path (example)
rg <- importGridstat("/Users/markheckmann/data/gridstat.dat")</pre>
# load gridstat data from URL
rg <- importGridstat("http://www.openrepgrid.uni-bremen.de/data/gridstat.dat")</pre>
# setting rating scale range
rg <- importGridstat(file, dir, min=1, max=6)</pre>
## End(Not run)
```

importGridsuite

Import Gridsuite data files.

### **Description**

Import Gridsuite data files.

## Usage

```
importGridsuite(file, dir = NULL)
```

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## **Arguments**

file	Filename including path if file is not in current working directory. File can also
	be a complete URL. The fileformat is .dat.

dir Alternative way to supply the directory where the file is located (default NULL).

### Value

A single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

#### Note

The developers of Gridsuite have proposed to use an XML scheme as a standard exchange format for repertory grid data (Walter, Bacher & Fromm, 2004). This approach is also embraced by the OpenRepGrid package.

TODO: The element and construct IDs are not used yet. Thus, if the output should be in different order the current mechanism will cause false assignments.

### Author(s)

Mark Heckmann

#### References

```
http://www.gridsuite.de/
```

Walter, O. B., Bacher, A., & Fromm, M. (2004). A proposal for a common data exchange format for repertory grid data. *Journal of Constructivist Psychology*, 17(3), 247. doi:10.1080/10720530490447167

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt, importExcel

```
## Not run:
# using the pop-up selection menu
rg <- importGridsuite()
# supposing that the data file gridsuite.xml is in the current directory
file <- "gridsuite.xml"
rg <- importGridsuite(file)
# specifying a directory (arbitrary example directory)
dir <- "/Users/markheckmann/data"
rg <- importGridsuite(file, dir)
# using a full path
rg <- importGridsuite("/Users/markheckmann/data/gridsuite.xml")</pre>
```

66 importScivesco

```
# load Gridsuite data from URL
rg <- importGridsuite("http://www.openrepgrid.uni-bremen.de/data/gridsuite.xml")
## End(Not run)</pre>
```

importScivesco

Import sci:vesco data files.

## **Description**

Import sci:vesco data files.

### Usage

```
importScivesco(file, dir = NULL)
```

### **Arguments**

file Filename including path if file is not in current working directory. File can also

be a complete URL. The fileformat is .dat.

dir Alternative way to supply the directory where the file is located (default NULL).

#### Value

A single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

### Note

Sci:Vesco offers the options to rate the construct poles seperately or using a bipolar scale. The seperated rating is done using the "tetralemma" field. The field is a bivariate plane on which each of the four (tetra) corners has a different meaning in terms of rating. Using this approach also allows ratings like: "both poles apply", "none of the poles apply" and all intermediate ratings can be chosen. This relaxes the bipolarity assumption often assumed in grid theory and allows for deviation from a strict bipolar rating if the constructs are not applied in a bipolar way. Using the tetralemma field for rating requires to analyze each construct seperately though. This means we get a double entry grid where the emergent and constrast pole ratings might not simply be a reflection of on another. The tetralemma field is not yet supported and importing will fail. Currently only bipolar ratings are suppoerted.

If a tetralemma field has been used for rating, OpenRepGrid will offer the option to transform the scores into "normal" grid ratings (i.e. restricted to bipolarity) by projecting the ratings from the bivariate tetralemma field onto the diagonal of the tetralemma field and thus forcing a bipolar rating type. This option is not recommended due to the fact that the conversion is susceptible to error when both ratings are near to zero.

TODO: For developers: The element IDs are not used yet. This might cause wrong assignments.

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#### Author(s)

Mark Heckmann

#### References

Menzel, F., Rosenberger, M., Buve, J. (2007). Emotionale, intuitive und rationale Konstrukte verstehen. *Personalfuehrung*, 4(7), 91-99.

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt, importExcel

### **Examples**

```
## Not run:

# supposing that the data file scivesco.scires is in the current directory
file <- "scivesco.scires"
rg <- importScivesco(file)

# specifying a directory (arbitrary example directory)
dir <- "/Users/markheckmann/data"
rg <- importScivesco(file, dir)

# using a full path
rg <- importScivesco("/Users/markheckmann/data/scivesco.scires")

# load Gridsuite data from URL
rg <- importScivesco("http://www.openrepgrid.uni-bremen.de/data/scivesco.scires")

## End(Not run)</pre>
```

importTxt

Import grid data from a text file.

### **Description**

If you do not have a grid program at hand you can define a grid using a standard text editor and by saving it as a .txt file. The .txt file has to be in a fixed format. There are three mandatory blocks each starting and ending with a predefined tag in uppercase letters. The first block starts with ELEMENTS and ends with END ELEMENTS and contains one element in each line. The other mandatory blocks contain the constructs and ratings (see below). In the block containing the constructs the left and right pole are seperated by a colon (:). To define missing values use NA like in the example below. One optional block contains the range of the rating scale used defined by two numbers. The order of the blocks is arbitrary. All text not contained within the blocks is discarded and can thus be used for comments.

68 importTxt

### Usage

```
importTxt(file, dir = NULL, min = NULL, max = NULL)
```

#### **Arguments**

file	A vector of filenames including the full path if file is not in current working directory. File can also be a complete URL. The file suffix has to be .txt. If no file is supplied a selection pop up menu is opened to select the files.
dir	Alternative way to supply the directory where the file is located (default NULL).
min	Optional argument (numeric, default NULL) for minimum rating value in grid.
max	Optional argument (numeric, default NULL) for maximum rating value in grid.

#### **Details**

```
----- .txt file -----
anything not contained within the tags will be discarded
ELEMENTS
element 1
element 2
element 3
END ELEMENTS
CONSTRUCTS
left pole 1 : right pole 1
left pole 2 : right pole 2
left pole 3 : right pole 3
left pole 4 : right pole 4
END CONSTRUCTS
RATINGS
1 3 2
4 1 1
1 4 4
3 1 1
END RATINGS
RANGE
1 4
END RANGE
----- end of file -----
```

Note that the maximum and minimum value has to be defined using the min and max arguments if no RANGE block is contained in the data file. Otherwise the scaling range is inferred from the available data and a warning is issued as the range may be erroneous. This may effect other functions that depend on knowing the correct range and it is thus strongly recommended to set the scale range

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correctly.

### Value

A single repgrid object in case one file and a list of repgrid objects in case multiple files are imported.

### Author(s)

Mark Heckmann

#### See Also

importGridcor, importGridstat, importScivesco, importGridsuite, importTxt, importExcel

## **Examples**

```
## Not run:
# using the pop-up selection menu
rg <- importTxt()</pre>
# supposing that the data file sample.txt is in the current directory
file <- "sample.txt"</pre>
rg <- importTxt(file)</pre>
# specifying a directory (arbitrary example directory)
dir <- "/Users/markheckmann/data"</pre>
rg <- importTxt(file, dir)</pre>
# using a full path
rg <- importTxt("/Users/markheckmann/data/sample.txt")</pre>
# load .txt data from URL
rg <- importTxt("http://www.openrepgrid.uni-bremen.de/data/sample.txt")</pre>
# importing more than one .txt file via R code
files <- c("sample.txt", "sample_2.txt")</pre>
rg <- importTxt(files)</pre>
## End(Not run)
```

indexBias

Calculate 'bias' of grid as defined by Slater (1977).

### **Description**

"Bias records a tendency for reponses to accumulate at one end of the grading scale" (Slater, 1977, p.88).

70 indexConflict1

### Usage

```
indexBias(x, min, max, digits = 2)
```

## **Arguments**

x repgrid object.

min Minimum grid scale value.
max Maximum grid scale value.

digits Numeric. Number of digits to round to (default is 2).

### Value

Numeric.

## Note

STATUS: Working and checked against example in Slater, 1977, p. 87.

## Author(s)

Mark Heckmann

## References

Slater, P. (1977). The measurement of intrapersonal space by Grid technique. London: Wiley.

### See Also

indexVariability

indexConflict1	Conflict measure for grids (Slade & Sheehan, 1979) based on correla-
	tions.

# Description

Conflict measure as proposed by Slade and Sheehan (1979)

### Usage

```
indexConflict1(x)
```

# Arguments

x repgrid object.

indexConflict1 71

#### **Details**

The first approach to mathematically derive a conflict measure based on grid data was presented by Slade and Sheehan (1979). Their operationalization is based on an approach by Lauterbach (1975) who applied the *balance theory* (Heider, 1958) for a quantitative assessment of psychological conflict. It is based on a count of balanced and imbalanced triads of construct correlations. A triad is imbalanced if one or all three of the correlations are negative, i. e. leading to contrary implications. This approach was shown by Winter (1982) to be flawed. An improved version was proposed by Bassler et al. (1992) and has been implemented in the function indexConflict2.

The table below shows when a triad made up of the constructs A, B, and C is balanced and imbalanced.

cor(A,B)	cor(A,C)	cor(B,C)	Triad characteristic
+	+	+	balanced
+	+	-	imbalanced
+	-	+	imbalanced
+	-	-	balanced
-	+	+	imbalanced
-	+	-	balanced
-	-	+	balanced
-	-	-	imbalanced

#### Value

A list with the following elements:

total Total number of triads

imbalanced Number of imbalanced triads

prop.balanced Proportion of balanced triads

prop.imbalanced

Proportion of imbalanced triads

#### Author(s)

Mark Heckmann

#### References

Bassler, M., Krauthauser, H., & Hoffmann, S. O. (1992). A new approach to the identification of cognitive conflicts in the repertory grid: An illustrative case study. *Journal of Constructivist Psychology*, 5(1), 95-111.

Heider, F. (1958). The Psychology of Interpersonal Relation. John Wiley & Sons.

Lauterbach, W. (1975). Assessing psychological conflict. *The British Journal of Social and Clinical Psychology*, 14(1), 43-47.

Slade, P. D., & Sheehan, M. J. (1979). The measurement of 'conflict' in repertory grids. *British Journal of Psychology*, 70(4), 519-524.

72 indexConflict2

Winter, D. A. (1982). Construct relationships, psychological disorder and therapeutic change. *The British Journal of Medical Psychology*, 55 (Pt 3), 257-269.

### See Also

indexConflict2 for an improved version of this measure; see indexConflict3 for a measure based on distances.

## **Examples**

```
## Not run:
    indexConflict1(feixas2004)
    indexConflict1(boeker)
## End(Not run)
```

indexConflict2

Conflict measure for grids (Bassler et al., 1992) based on correlations.

# Description

Conflict measure as proposed by Bassler et al. (1992).

#### Usage

```
indexConflict2(x, crit = 0.03)
```

### Arguments

x repgrid object.

crit Sensitivity criterion with which triads are marked as unbalanced. A bigger val-

ues will lead to less imbalanced triads. The default is 0.03. The value should

be adjusted with regard to the researchers interest.

#### **Details**

The function calculates the conflict measure as devised by Bassler et al. (1992). It is an improved version of the ideas by Slade and Sheehan (1979) that have been implemented in the function indexConflict1. The new approach also takes into account the magnitude of the correlations in a traid to assess whether it is balanced or imbalanced. As a result, small correlations that are psychologically meaningless are considered accordingly. Also, correlations with a small magnitude, i. e. near zero, which may be positive or negative due to chance alone will no longer distort the measure (Bassler et al., 1992).

Description of the balance / imbalance assessment:

indexConflict2 73

- 1. Order correlations of the triad by absolute magnitude, so that  $r_{max} > r_{mdn} > r_{min}$ .
- 2. Apply Fisher's Z-transformation and devision by 3 to yield values between 1 and -1
- 3. Check whether the triad is balanced by assessing if the following relation holds:
  - If  $Z_{max}Z_{mdn}>0$ , the triad is balanced if  $Z_{max}Z_{mdn}-Z_{min}<=crit$  Z\_max x Z\_mdn Z\_min <= crit .
  - If  $Z_{max}Z_{mdn} < 0$ , the triad is balanced if  $Z_{min} Z_{max}Z_{mdn} <= crit Z_{min} Z_{max} \times Z_{mdn} <= crit Z_{min} Z_{max} \times Z_{min} <= crit Z_{min} Z_{min} Z_{min} \times Z_{min} <= crit Z_{min} <= crit Z_{min} <= crit Z_{min} Z_{min} <= crit Z_{min} Z_{min} <= crit Z_{min} Z_{min} <= crit Z_{min} <= crit Z_{min} Z_{min} <= crit Z_{mi$

#### Personal remarks (MH)

I am a bit suspicious about step 2 from above. To devide by 3 appears pretty arbitrary. The r for a z-values of 3 is 0.9950548 and not 1. The r for 4 is 0.9993293. Hence, why not a value of 4, 5, or 6? Denoting the value to devide by with a, the relation for the first case translates into  $aZ_{max}Z_{mdn} <= \frac{crit}{a} + Z_{min}$  a x Z\_max x Z\_mdn =< crit/a + Z\_min. This shows that a bigger value of a will make it more improbabale that the relation will hold.

#### Author(s)

Mark Heckmann

#### References

Bassler, M., Krauthauser, H., & Hoffmann, S. O. (1992). A new approach to the identification of cognitive conflicts in the repertory grid: An illustrative case study. *Journal of Constructivist Psychology*, 5(1), 95-111.

Slade, P. D., & Sheehan, M. J. (1979). The measurement of 'conflict' in repertory grids. *British Journal of Psychology*, 70(4), 519-524.

#### See Also

See indexConflict1 for the older version of this measure; see indexConflict3 for a measure based on distances instead of correlations.

```
## Not run:
indexConflict2(bell2010)

x <- indexConflict2(bell2010)
print(x)

# show conflictive triads
print(x, output=2)

# accessing the calculations for further use
x$total
x$imbalanced
x$prop.balanced</pre>
```

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```
x$prop.imbalanced
x$triads.imbalanced
## End(Not run)
```

indexConflict3

Conflict or inconsistenciy measure for grids (Bell, 2004) based on distances.

## Description

Conflict measure as proposed by Bell (2004).

## Usage

```
indexConflict3(x, p = 2, e.out = NA, e.threshold = NA, c.out = NA,
    c.threshold = NA, trim = 20)
```

## **Arguments**

x	repgrid object.
р	The power of the Minkowski distance. p=2 (default) will result in euclidean distances, p=1 in city block distances.
e.out	Numeric. A vector giving the indexes of the elements for which detailed stats (number of conflicts per element, discrepancies for triangles etc.) are promted (default NA, i.e. no detailed stats for any element).
e.threshold	Numeric. Detailed stats are prompted for those elements with a an attributable percentage to the overall conflicts higher than the supplied threshold (default NA).
c.out	Numeric. A vector giving the indexes of the constructs for which detailed stats (discrepancies for triangles etc.) are promted (default NA, i. e. no detailed stats).
c.threshold	Numeric. Detailed stats are prompted for those constructs with a an attributable percentage to the overall conflicts higher than the supplied threshold (default NA).
trim	The number of characters a construct (element) is trimmed to (default is 10). If NA no trimming is done. Trimming simply saves space when displaying the output.

## **Details**

Measure of conflict or inconsistency as proposed by Bell (2004). The identification of conflict is based on distances rather than correlations as in other measures of conflict indexConflict1 and indexConflict2. It assesses if the distances between all components of a triad, made up of one element and two constructs, satisfies the "triangle inequality" (cf. Bell, 2004). If not, a triad is regarded as conflictive. An advantage of the measure is that it can be interpreted not only as a

indexConflict3 75

global measure for a grid but also on an element, construct, and element by construct level making it valuable for detailed feedback. Also, differences in conflict can be submitted to statistical testing procedures.

Status: working; output for euclidean and manhattan distance checked against Gridstat output. TODO: standardization and z-test for discrepancies; Index of Conflict Variation.

#### Value

A list (invisibly) containing containing:

potential	number of potential conflicts
actual	count of actual conflicts
overall	percentage of conflictive relations
e.count	number of involvements of each element in conflictive relations
e.perc	percentage of involvement of each element in total of conflictive relations
e.count	number of involvements of each construct in conflictive relation
c.perc	percentage of involvement of each construct in total of conflictive relations
e.stats	detailed statistics for prompted elements
c.stats	detailed statistics for prompted constructs
e.threshold	threshold percentage. Used by print method
c.threshold	threshold percentage. Used by print method
enames	trimmed element names. Used by print method
cnames	trimmed construct names. Used by print method

## output

For further control over the output see print.indexConflict3.

#### Author(s)

Mark Heckmann

#### References

Bell, R. C. (2004). A new approach to measuring inconsistency or conflict in grids. Personal Construct Theory & Practice, (1), 53-59.

#### See Also

See indexConflict1 and indexConflict2 for conflict measures based on triads of correlations.

#### **Examples**

```
## Not run:

# calculate conflicts
indexConflict3(bell2010)

# show additional stats for elements 1 to 3
indexConflict3(bell2010, e.out=1:3)

# show additional stats for constructs 1 and 5
indexConflict3(bell2010, c.out=c(1,5))

# finetune output

## change number of digits
x <- indexConflict3(bell2010)
print(x, digits=4)

## omit discrepancy matrices for constructs
x <- indexConflict3(bell2010, c.out=5:6)
print(x, discrepancies=FALSE)

## End(Not run)</pre>
```

indexDilemma

Detect implicative dilemmas (conflicts).

#### Description

Implicative Dilemmas

#### Usage

```
indexDilemma(x, self = 1, ideal = ncol(x), diff.mode = 1,
  diff.congruent = NA, diff.discrepant = NA, diff.poles = 1,
  r.min = 0.2, exclude = FALSE, digits = 2, show = F, output = 1,
  index = T, trim = 20)
```

#### Arguments

diff.mode

x repgrid object.

self Numeric. Index of self element. ideal Numeric. Index of ideal self element.

Numeric. Mode to classify construct pairs into congruent and discrepant. diff.mode=1

will use the difference in ratings between the self and the ideal element to determine if the construct is congruent or discrepant. No other modes have yet been

implemented.

diff.congruent Is used if diff.mode=1. Maximal difference between element ratings to define

construct as congruent (default diff.congruent=1). Note that the value needs

to be adjusted by the user according to the rating scale used.

diff.discrepant

Is used if diff.mode=1. Minimal difference between element ratings to define construct as discrepant (default diff.discrepant=3). Note that the value needs

to be adjusted by the user according to the rating scale used.

diff.poles Not yet implemented.

r.min Minimal correlation to determine implications between constructs.

exclude Whether to exclude the elements self and ideal self during the calculation of the

inter-construct correlations. (default is FALSE).

digits Numeric. Number of digits to round to (default is 2).

show Whether to additionally plot the distribution of correlations to help the user as-

sess what level is adequate for r.min.

output The type of output printed to the console. output=1 prints classification of the

construct into congruent and discrepant and the detected dilemmas. output=1 only prints the latter. output=0 will surpress printing. Note that the type of output does not affect the object that is returned invisibly which will be the

same in any case (see value).

index Whether to print index numbers in front of each construct (default is TRUE).

trim The number of characters a construct (element) is trimmed to (default is 20).

If NA no trimming is done. Trimming simply saves space when displaying the

output.

#### **Details**

Implicative dilemmas are closely related to the notion of conflict. An implicative dilemma arises when a desired change on one construct is associated with an undesired implication on another construct. E. g. a timid subject may want to become more socially skilled but associates being socially skilled with different negative characteristics (selfish, insensitive etc.). Hence, he may anticipate that becoming less timid will also make him more selfish (cf. Winter, 1982). As a consequence the subject will resist to the change if the negative presumed implications will threaten the patients identity and the predictive power of his construct system. From this stance the resistance to change is a logical consequence coherent with the subjects construct system (Feixas, Saul, & Sanchez, 2000). The investigation of the role of cognitive dilemma in different disorders in the context of PCP is a current field of research (e.g. Feixas & Saul, 2004, Dorough et al. 2007).

The detection of implicative dilemmas happens in two steps. First the constructs are classified as being 'congruent' or 'discrepant'. Second the correlation between a congruent and discrepant construct pair is assessed if it is big enough to indicate an implication.

#### Classifying the construct

To detect implicit dilemmas the construct pairs are first identified as 'congruent' or 'discrepant'. The assessment is based on the rating differences between the elements 'self' and 'ideal self'. A construct is 'congruent' if the construction of the 'self' and the preferred state (i.e. ideal self) are the same or similar. A construct is discrepant if the construction of the 'self' and the 'ideal' is dissimilar. Suppose the element 'self' is rated 2 and 'ideal self' 5 on a scale from 1 to 6. The ratings

differences are 5-2 = 3. If this difference is smaller than e.g. 1 the construct is 'congruent', if it is bigger than 3 it is 'discrepant'.

The values used to classify the constructs 'congruent' or 'discrepant' can be determined in several ways (cf. Bell, 2009):

- 1. They are set 'a priori'.
- 2. They are implicitly derived by taking into account the rating differences to the other constructs. Not yet implemented.

The value mode is determined via the argument diff.mode.

If no 'a priori' criteria to determine if the construct is congruent or discrepant is supplied as an argument, the values are chosen acording to the range of the rating scale used. For the following scales the defaults are chosen as:

Scale	'A priori' criteria
1 2	-> con: <=0 disc: >=1
1 2 3	-> con: <=0 disc: >=2
1 2 3 4	-> con: <=0 disc: >=2
1 2 3 4 5	-> con: <=1 disc: >=3
1 2 3 4 5 6	-> con: <=1 disc: >=3
1234567	-> con: <=1 disc: >=4
12345678	-> con: <=1 disc: >=5
123456789	-> con: <=2 disc: >=5
12345678910	-> con: <=2 disc: >=6

#### **Defining the correlations**

As the implications between constructs cannot be derived from a rating grid directly, the correlation between two constructs is used as an indicator for implication. A large correlation means that one construct pole implies the other. A small correlation indicates a lack of implication. The minimum criterion for a correlation to indicate implication is set to .35 (cf. Feixas & Saul, 2004). The user may also chose another value. To get a an impression of the distribution of correlations in the grid, a visualization can be prompted via the argument show. When calculating the correlation used to assess if an implication is given or not, the elements under consideration (i. e. self and ideal self) can be included (default) or excluded. The options will cause different correlations (see argument exclude).

#### Example of an implicative dilemma

A depressive person considers herself as timid and wished to change to the opposite pole she defines as extraverted. This construct is called discrepant as the construction of the 'self' and the desired state (e.g. described by the 'ideal self') on this construct differ. The person also considers herself as sensitive (preferred pole) for which the opposite pole is selfish. This construct is congruent, as the person construes herself as she would like to be. If the person now changed on the discrepant construct from the undesired to the desired pole, i.e. from timid to extraverted, the question can be asked what consequences such a change has. If the person construes being timid and being sensitive as related and that someone who is extraverted will not be timid, a change on the first construct

will imply a change on the congruent construct as well. Hence, the positive shift from timid to extraverted is presumed to have a undesired effect in moving from sensitive towards selflish. This relation is called an implicative dilemma. As the implications of change on a construct cannot be derived from a rating grid directly, the correlation between two constructs is used as an indicator for implication.

#### Value

Called for console output. Invisbly returns a list containing the result dataframes and all results from the calculations.

#### Author(s)

Mark Heckmann, Alejandro García

#### References

Bell, R. C. (2009). *Gridstat version 5 - A Program for Analyzing the Data of A Repertory Grid* (manual). University of Melbourne, Australia: Department of Psychology.

Dorough, S., Grice, J. W., & Parker, J. (2007). Implicative dilemmas and psychological well-being. *Personal Construct Theory & Practice*, (4), 83-101.

Feixas, G., & Saul, L. A. (2004). The Multi-Center Dilemma Project: an investigation on the role of cognitive conflicts in health. *The Spanish Journal of Psychology*, 7(1), 69-78.

Feixas, G., Saul, L. A., & Sanchez, V. (2000). Detection and analysis of implicative dilemmas: implications for the therapeutic process. In J. W. Scheer (Ed.), *The Person in Society: Challenges to a Constructivist Theory*. Giessen: Psychosozial-Verlag.

Winter, D. A. (1982). Construct relationships, psychological disorder and therapeutic change. *British Journal of Medical Psychology*, *55* (Pt 3), 257-269.

```
## Not run:
indexDilemma(boeker, self=1, ideal=2)
indexDilemma(boeker, self=1, ideal=2, out=2)

# additionally show correlation distribution
indexDilemma(boeker, self=1, ideal=2, show=T)

# adjust minimal correlation
indexDilemma(boeker, 1, 2, r.min=.25)

# adjust congruence and discrepance ranges
indexDilemma(boeker, 1, 2, diff.con=0, diff.disc=4)

## End(Not run)
```

80 indexIntensity

indexIntensity Intensity in	ndex
-----------------------------	------

#### **Description**

Calculate intensity index.

## Usage

```
indexIntensity(x, rc = FALSE, trim = 30)
```

#### **Arguments**

x repgrid object.

rc Whether to use Cohen's rc for the calculation of inter-element correlations. See

elementCor for further explanations of this measure.

trim The number of characters a construct is trimmed to (default is 30). If NA no

trimming occurs. Trimming simply saves space when displaying correlation of

constructs or elements with long names.

#### **Details**

The Intensity index has been suggested by Bannister (1960) as a measure of the amount of construct linkage. Bannister suggested that the score reflects the degree of organization of the construct system under investigation (Bannister & Mair, 1968). The index resulted from his and his colleagues work on construction systems of patient suffering schizophrenic thought disorder. The concept of intensity has a theoretical connection to the notion of "tight" and "loose" construing as proposed by Kelly (1991). While tight constructs lead to unvarying prediction, loose constructs allow for varying predictions. Bannister hypothesized that schizophrenic thought disorder is liked to a process of extremely loose construing leading to a loss of predictive power of the subject's construct system. The Intensity score as a structural measure is thought to reflect this type of system disintegration (Bannister, 1960).

Implementation as in the Gridcor programme and explained on the correspoding help pages: "... the sum of the squared values of the correlations of each construct with the rest of the constructs, averaged by the total number of constructs minus one. This process is repeated with each element, and the overall Intensity is calculated by averaging the intensity scores of constructs and elements." <a href="http://www.terapiacognitiva.net/record/pag/man11.htm">http://www.terapiacognitiva.net/record/pag/man11.htm</a>. Currently the total is calculated as the unweighted average of all single scores (for elements and construct).

#### Value

An object of class indexIntensity containing a list with the following elements:

c.int Intensity scores by construct.e.int Intensity scores by element.

indexPvaff 81

```
c.int.mean Average intensity score for constructs.e.int.mean Average intensity score for elements.total.int Total intensity score.
```

## Development

TODO: Results have not been tested against other programs' results.

## Author(s)

Mark Heckmann

## References

Bannister, D. (1960). Conceptual structure in thought-disordered schizophrenics. *The Journal of mental science*, 106, 1230-49.

```
indexIntensity(bell2010)
indexIntensity(bell2010, trim=NA)

# using Cohen's rc for element correlations
indexIntensity(bell2010, rc=TRUE)

# save output
x <- indexIntensity(bell2010)
x

# printing options
print(x, digits=4)

# accessing the objects' content
x$c.int
x$e.int
x$c.int.mean
x$e.int.mean
x$total.int</pre>
```

82 indexPvaff

#### **Description**

The PVAFF is used as a measure of cognitive complexity. It was introduced in an unpublished PhD thesis by Jones (1954, cit. Bonarius, 1965). To calculate it, the 'first factor' is extracted from the construct correlation matrix by principal component analysis. The PVAFF reflects the amount of variation that is accounted for by a single linear component. If a single latent component is able to explain the variation in the grid, the cognitive complexity is said to be low. In this case the construct system is regarded as 'simple' (Bell, 2003).

#### Usage

```
indexPvaff(x)
```

## **Arguments**

x repgrid object.

#### **Details**

The percentage of variance is calculated using the corelation matrix of te constructs submitted to svd.

## **Development**

TODO: Results have not yet been checked against other grid programs.

#### Author(s)

Mark Heckmann

#### References

Bell, R. C. (2003). An evaluation of indices used to represent construct structure. In G. Chiari & M. L. Nuzzo (Eds.), *Psychological Constructivism and the Social World* (pp. 297-305). Milan: FrancoAngeli.

Bonarius, J. C. J. (1965). Research in the personal construct theory of George A. Kelly: role construct repertory test and basic theory. In B. A. Maher (Ed.), *Progress in experimental personality research* (Vol. 2). New York: Academic Press.

James, R. E. (1954). *Identification in terms of personal constructs* (Unpublished doctoral thesis). Ohio State University, Columbus, OH.

```
indexPvaff(bell2010)
indexPvaff(feixas2004)

# save results to object
p <- indexPvaff(bell2010)
p</pre>
```

index Variability 83

index Variability

Calculate 'variability' of a grid as defined by Slater (1977).

## **Description**

Variability records a tendency for the responses to gravitate towards both end of the gradings scale. (Slater, 1977, p.88).

## Usage

```
indexVariability(x, min, max, digits = 2)
```

## **Arguments**

x repgrid object.

min Minimum grid scale value.

max Maximum grdi scale value.

digits Numeric. Number of digits to round to (default is 2).

## Value

Numeric.

## Note

STATUS: working and checked against example in Slater, 1977, p.88.

## Author(s)

Mark Heckmann

## References

Slater, P. (1977). The measurement of intrapersonal space by Grid technique. London: Wiley.

#### See Also

indexBias

84 left

left

Move construct or element in grid to the left, right, up or down.

## Description

Move construct or element in grid to the left, right, up or down.

Move element in grid to the right.

Move construct in grid upwards.

Move construct in grid downwards.

## Usage

```
left(x, pos = \emptyset)
right(x, pos = \emptyset)
up(x, pos = \emptyset)
down(x, pos = \emptyset)
```

## **Arguments**

x repgrid object.

pos

Row (column) number of construct (element) to be moved leftwards, rightwards, upwards or downwards. The default is  $\emptyset$ . For indexes outside the range of the grid no moving is done.

## Value

```
repgrid object
repgrid object
repgrid object
repgrid object
```

## Author(s)

Mark Heckmann

```
## Not run:
    x <- randomGrid()
    left(x, 2)  # 2nd element to the left
    right(x, 1)  # 1st element to the right
    up(x, 2)  # 2nd construct upwards
    down(x, 1)  # 1st construct downwards</pre>
```

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```
## End(Not run)
```

makeRepgrid

Make a new repgrid object.

## Description

The function creates a repgrid object from scratch. A number of paramters have to be defined in order to make a new grid (see parameters).

## Usage

```
makeRepgrid(args)
```

## Arguments

args

Arguments needed for the construction of the grid (list). These include name followed by a vector containing the element names. 1. name followed by a vector with the left construct poles. r.name followed by a vector with the right construct poles. scores followed by a vector containing the rating scores row wise.

#### Value

NULL

#### Author(s)

Mark Heckmann

86 normalize

normalize

Normalize rows or columns by its standard deviation.

## Description

Normalize rows or columns by its standard deviation.

Not evaluated.

## Usage

```
normalize(x, normalize = 0, ...)
```

## **Arguments**

```
x matrix

normalize A numeric value indicating along what direction (rows, columns) to normalize by standard deviations. 0 = none, 1= rows, 2 = columns (default is 0).
```

#### Value

Not yet definde TODO!

## Author(s)

Mark Heckmann

```
## Not run:

x <- matrix(sample(1:5, 20, rep=T), 4)
normalize(x, 1)  # normalizing rows
normalize(x, 2)  # normalizing columns
## End(Not run)</pre>
```

OpenRepGrid 87

OpenRepGrid

**OpenRepGrid**: an R package for the analysis of repertory grids.

#### **Description**

The **OpenRepGrid** package provides tools for the analysis of repertory grid data. The repertory grid is a method devised by George Alexander Kelly in his seminal work "The Psychology of Personal Constructs" published in 1955. The repertory grid has been used in and outside the context of Personal Construct Psychology (PCP) in a broad range of fields. For an introduction into the technique see e.g. Fransella, Bell and Bannister (2003).

#### Note

To get started with **OpenRepGrid** visit the project's home under www.openrepgrid.org. On this site you will find tutorials, explanation about the theory, methods of analysis and the according R code.

To see the preferable citation of the **OpenRepGrid** package, type citation("OpenRepGrid") into the R console.

Disclaimer: Note that the package is distributed under the GPL 2 license. It is work in progress and is continuously being improved by hopefully numerous contributors. It may contain bugs and errors. There is no warranty whatsoever for the program.

#### Author(s)

**OpenRepGrid** was initiated by Mark Heckmann. Current contributors are: Mark Heckmann, Alejandro García. Researchers interested in developing the package are invited to join.

The **OpenRepGrid** package development is hosted on github (http://github.com/markheckmann/OpenRepGrid). The github site provides information and allows to file bug reports or feature requests. Bug reports can also be emailed to the package maintainer or issued on http://www.openrepgrid.org under section *Suggestions/Issues*. The package maintainer is Mark Heckmann <heckmann(at)uni-bremen.de>.

#### References

Fransella, F., Bell, R. C., & Bannister, D. (2003). *A Manual for Repertory Grid Technique* (2. *Ed.*). Chichester: John Wiley & Sons.

Kelly, G. A. (1955). *The psychology of personal constructs. Vol. I, II.* New York: Norton, (2nd printing: 1991, Routledge, London, New York).

OpenRepGrid-overview **OpenRepGrid**: Annotated overview of package functions.

**Description** 

# This documentation page contains an overview over the package functions ordered by topics. The best place to start learning OpenRepGrid will be the package website <a href="http://www.openrepgrid.org">http://www.openrepgrid.org</a> though.

## Functions sorted by topic

#### Manipulating grids

left Move construct(s) to the left
right Move construct(s) to the right
up Move construct(s) upwards
down Move construct(s) downwards

#### Loading and saving data

importGridcor
importGridstat
importGridsuite
importScivesco
importTxt
Import GRIDCOR data files
Import Gridstat data files
Import Gridsuite data files
Import sci:vesco data files
Import grid data from a text file

## **Analyzing constructs**

Descriptive statistics of constructs Construct correlations distance Root mean square of inter-construct correlations Somers' D Principal component analysis (PCA) of construct correlation matrix Cluster analysis of constructs

Save grid in a text file (txt)

## Analyzing elements

#### Visual representation

Bertin plots

bertin Make Bertin display of grid data

saveAsTxt

bertinCluster Bertin display with corresponding cluster analysis

**Biplots** 

biplot2d Draw a two-dimensional biplot

biplotEsa2d Plot an eigenstructure analysis (ESA) biplot in 2D

biplotSlater2d Draws Slater's INGRID biplot in 2D

biplotPseudo3d See 'biplotPseudo3d' for its use. Draws a biplot of the grid in 2D with depth impression (pseudo 3

biplotEsaPseudo3d Plot an eigenstructure analysis (ESA) in 2D grid with 3D impression (pseudo 3D)

biplotSlaterPseudo3d Draws Slater's biplot in 2D with depth impression (pseudo 3D)

biplot3d Draw grid in rgl (3D device)

biplotEsa3d Draw the eigenstructure analysis (ESA) biplot in rgl (3D device)

biplotSlater3d Draw the Slater's INGRID biplot in rgl (3D device)

biplotSimple A graphically unsophisticated version of a biplot

#### **Index measures**

indexConflict1 Conflict measure for grids (Slade & Sheehan, 1979) based on correlations Conflict measure for grids (Bassler et al., 1992) based on correlations indexConflict3 Conflict or inconsistencity measure for grids (Bell, 2004) based on distances

indexDilemma Detect implicative dilemmas (conflicts)

indexIntensity Intensity index

indexPvaff Percentage of Variance Accounted for by the First Factor (PVAFF)

indexBias Calculate 'bias' of grid as defined by Slater (1977)

indexVariability Calculate 'variability' of a grid as defined by Slater (1977)

#### Special features

alignByIdeal Align constructs using the ideal element to gain pole preferences alignByLoadings Align constructs by loadings on first pricipal component

reorder2d Order grid by angles between construct and/or elements in 2D

#### **Settings**

**OpenRepGrid** uses several default settings e.g. to determine how many construct characters to display by default when displaying a grid. The function settings can be used to show and change these settings. Also it is possible to store the settings to a file and load the settings file to restore the

settings.

settings Show and modify global settings for OpenRepGrid Save OpenRepGrid settings to file Load OpenRepGrid settings from file

#### **Grid datasets**

**OpenRepGrid** already contains some ready to use grid data sets. Most of the datasets are taken from the literature. To output the data simply type Type the name of the dataset to the console and press enter.

Single grids

bel12010 Grid data from a study by Haritos et al. (2004) on role titles; used for demonstration of construct Grid from a psychotic patient used in Bell (1997, p. 6). Data originated from a study by Bell and bellmcgorry1992 boeker Grid from seventeen year old female schizophrenic patient undergoing last stage of psychoanalyt fbb2003 Dataset used in A manual for Repertory Grid Technique (Fransella, Bell, & Bannister, 2003b, p. 6 Grid from a 22 year old Spanish girl suffering self-worth problems (Feixas & Saul, 2004, p. 77). feixas2004 mackay1992 Dataset Grid C used in Mackay's paper on inter-element correlation (1992, p. 65). leach2001a, leach2001b Pre- (a) and post-therapy (b) dataset from sexual child abuse survivor (Leach, Freshwater, Aldrid Grid data to demonstrate the use of Bertin diagrams (Raeithel, 1998, p. 223). The context of its a raeithel slater1977a Drug addict grid dataset from (Slater, 1977, p. 32). Grid dataset (ranked) from a seventeen year old female psychiatric patient (Slater, 1977, p. 110): slater1977b

Multiple grids

NOT YET AVAILABLE

#### **Functions for developers**

OpenRepGrid: internal functions overview for developers.

Below you find a guide for developers: these functions are usually not needed by the casual user. The internal functions have a twofold goal 1) to provide means for advanced numerical grid analysis and 2) to facilitate function development. The function for these purposes are internal, i.e. they are not visible in the package documentation. Nonetheless they do have a documentation that can be accesses in the same way as for other functions. More in the details section.

#### Functions for advanced grid analysis

The package provides functions to facilitate numerical research for grids. These comprise the generation of random data, permutation of grids etc. to facilitate Monte Carlo simulations, batch analysis of grids and other methods. With R as an underlying framework, the results of grid analysis easily

lend themselves to further statistical processing and analysis within R. This is one of the central advantages for researchers compared to other standard grid software. The following table lists several functions for these purposes.

randomGrid
randomGrids
permuteConstructs
permuteGrid
quasiDistributionDistanceSlater

#### **Modules for function development**

Beside the advanced analysis feature the developer's functions comprise low-level modules to create new functions for grid analysis. Though the internal structure of a repgrid object in R is simple (type e.g. str(bell2010, 2) to get an impression), it is convenient to not have to deal with access on this level. Several function like e.g. getElementNames are convenient wrappers that perform standard tasks needed when implementing new functions. The following table lists several functions for these purposes.

getRatingLayer Retrieve grid scores from grid object. Get the number of constructs in a grid object. getNoOfConstructs getNoOfElements Get the number of elements in a grid object. Get grid dimensions, i.e. contructs x elements. dim getScale Get minimum and maximum scale value used in grid. Get midpoint of the grid rating scale. getScaleMidpoint Get construct names. getConstructNames getConstructNames2 Get construct names (another newer version). Retrieve element names of repgrid object. getElementNames Concatenate the constructs of two grids. bindConstructs doubleEntry Join the constructs of a grid with the same reversed constructs.

#### Other internal functions

#### importTxtInternal

#### Author(s)

Current members of the **OpenRepGrid** development team: Mark Heckmann. Everyone who is interested in developing the package is invited to join.

The **OpenRepGrid** package development is hosted on github (http://github.com/markheckmann/OpenRepGrid). The github site provides information and allows to file bug reports or feature requests. Bug reports can also be emailed to the package maintainer or issued on http://www.openrepgrid.org under section *Suggestions/Issues*. The package maintainer is Mark Heckmann <heckmann(at)uni-bremen.de>.

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permuteConstructs

Generate a list with all possible construct reflections of a grid.

## Description

Generate a list with all possible construct reflections of a grid.

## Usage

```
permuteConstructs(x, progress = TRUE)
```

## **Arguments**

x repgrid object.

progress

Whether to show a progress bar (default is TRUE). This may be sensible for a

larger number of elements.

#### Value

A list of repgrid objects with all possible permutations of the grid.

#### Author(s)

Mark Heckmann

## **Examples**

```
## Not run:
    1 <- permuteConstructs(mackay1992)
    1
## End(Not run)</pre>
```

randomGrid

Generate a random grid (quasis) of prompted size.

## **Description**

This feature is useful for research purposes like exploring distributions of indexes etc.

## Usage

```
randomGrid(nc = 10, ne = 15, nwc = 8, nwe = 5, range = c(1, 5), prob = NULL, options = 1)
```

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#### **Arguments**

nc	Number of constructs (default 10).
ne	Number of elements (default 15).
nwc	Number of random words per construct.
nwe	Number of random words per element.
range	Minimal and maximal scale value (default c(1, 5)).
prob	The probability of each rating value to occur. If NULL (default) the distribution is uniform.
options	Use random sentences as constructs and elements (1) or not (0). If not, the

elements and constructs are given default names and are numbered.

#### Value

repgrid object.

## Author(s)

Mark Heckmann

## **Examples**

```
## Not run:
    x <- randomGrid()
    x
    x <- randomGrid(10, 25)
    x
    x <- randomGrid(10, 25, options=0)
    x

## End(Not run)</pre>
```

randomGrids

Generate a list of random grids (quasis) of prompted size.

## Description

This feature is useful for research purposes like exploring distributions of indexes etc. The function is a simple wrapper around randomGrid.

## Usage

```
randomGrids(rep = 3, nc = 10, ne = 15, nwc = 8, nwe = 5, range = c(1, 5), prob = NULL, options = 1)
```

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## **Arguments**

rep	Number of grids to be produced (default is 3).
nc	Number of constructs (default 10).
ne	Number of elements (default 15).
nwc	Number of random words per construct.
nwe	Number of random words per element.
range	Minimal and maximal scale value (default c(1, 5)).
prob	The probability of each rating value to occur. If NULL (default) the distribution is uniform.
options	Use random sentences as constructs and elements (1) or not (0). If not, the

elements and constructs are given default names and are numbered.

## Value

A list of repgrid objects.

#### Author(s)

Mark Heckmann

## **Examples**

```
## Not run:
    x <- randomGrids()
    x
    x <- randomGrids(5, 3, 3)
    x
    x <- randomGrids(5, 3, 3, options=0)
    x

## End(Not run)</pre>
```

reorder2d

Order grid by angles between construct and/or elements in 2D.

## Description

The approach is to reorder the grid matrix by their polar angles on the first two principal components from a data reduction technique (here the biplot, i.e. SVD). The function reorder2d reorders the grid according to the angles between the x-axis and the element (construct) vectors derived from a 2D biplot solution. This approach is apt to identify circumplex structures in data indicated by the diagonal stripe in the display (see examples).

reorder2d 95

## Usage

```
reorder2d(x, dim = c(1, 2), center = 1, normalize = 0, g = 0, h = 1 - g, rc = TRUE, re = TRUE, ...)
```

## Arguments

Х	repgrid object.
dim	Dimension of 2D solution used to calculate angles (default c(1,2)).
center	Numeric. The type of centering to be performed. 0= no centering, 1= row mean centering (construct), 2= column mean centering (elements), 3= double-centering (construct and element means), 4= midpoint centering of rows (constructs). The default is 1 (row centering).
normalize	A numeric value indicating along what direction (rows, columns) to normalize by standard deviations. $\emptyset$ = none, 1= rows, 2 = columns (default is $\emptyset$ ).
g	Power of the singular value matrix assigned to the left singular vectors, i.e. the constructs.
h	Power of the singular value matrix assigned to the right singular vectors, i.e. the elements.
rc	Logical. Reorder constructs by similarity (default TRUE).
re	Logical. Reorder elements by similarity (default TRUE).
	Not evaluated.

## Value

Reordered repgrid object.

## Author(s)

Mark Heckmann

```
## Not run:

x <- feixas2004
reorder2d(x)  # reorder grid by angles in first two dimensions
reorder2d(x, rc=F)  # reorder elements only
reorder2d(x, re=F)  # reorder constructs only

## End(Not run)</pre>
```

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saveAsExcel

Save grid in a Microsoft Excel file (.xlsx)

## Description

```
saveAsExcel will save the grid as a Microsoft Excel file (.xlsx).
```

## Usage

```
saveAsExcel(x, file, sheet = 1)
```

## Arguments

x A repgrid object.

file Filename to save the grid to. The name should have the suffix .xlsx.

sheet Index of the sheet to write to.

## Value

Invisibly returns the name of the file.

## Author(s)

Mark Heckmann

## See Also

```
importExcel
```

```
## Not run:

x <- randomGrid(options=0)
 saveAsExcel(x, "grid.xlsx")

## End(Not run)</pre>
```

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saveAsTxt

Save grid in a text file (txt).

#### Description

saveAsTxt will save the grid as a .txt file in format used by **OpenRepGrid**. This file format can also easily be edited by hand (see importTxt for a description). The funtion will open an interactive dialog box to let the user enter a filename if no file argument is supplied in the function call.

## Usage

```
saveAsTxt(x, file = NA)
```

## **Arguments**

x repgrid object.

file Filename to save the grid to. The name should have the suffix .txt. If the function

is called without specifying this argumnet a dialog box is opened.

#### Value

Invisibly returns the name of the file.

#### Note

```
Structure of a txt file that can be read by importTxt.
----- .txt file -----
anything not contained within the tags will be discarded
                        ELEMENTS
                        element 1
                        element 2
                        element 3
                        END ELEMENTS
                        CONSTRUCTS
                        left pole 1 : right pole 1
                        left pole 2 : right pole 2
                        left pole 3 : right pole 3
                        left pole 4 : right pole 4
                        END CONSTRUCTS
                        RATINGS
                        1 3 2
                        4 1 1
                        1 4 4
```

3 1 1

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END RATINGS

RANGE 1 4

END RANGE

```
----- end of file -----
```

## Author(s)

Mark Heckmann

#### See Also

```
importTxt
```

#### **Examples**

```
## Not run:
x <- randomGrid()
saveAsTxt(x, "random.txt")
## End(Not run)</pre>
```

setConstructAttr

Set the attributes of a construct

## **Description**

Set the attributes of a construct i.e. name, abbreviation, status etc.

## Usage

```
setConstructAttr(x, pos, l.name, r.name, l.preferred, r.preferred, l.emerged,
    r.emerged)
```

## Arguments

x repgrid object.

pos Row number of construct in the grid to be changed

1. name Name of the left pole (string) (optional).r. name Name of the right pole (string) (optional).

1. preferred Logical. Is the left one the preferred pole? (optional).

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```
r.preferred Logical. Is the right one the preferred pole? (optional).
l.emerged Logical. Is the left one the emergent pole? (optional).
r.emerged Logical. Is the right one the emergent pole? (optional).
```

## Value

repgrid object

## Author(s)

Mark Heckmann

## See Also

```
setElementAttr
```

## **Examples**

setElementAttr

Set the attributes of an element

## **Description**

Set the attributes of an element i.e. name, abbreviation, status etc.

## Usage

```
setElementAttr(x, pos, name, abb, status)
```

## **Arguments**

X	repgrid object.
pos	Column number of element in the grid whose attributes are changed.
name	New element name (optional).
abb	Abbreviation of element name (optional).
status	Status of element (e.g. ideal etc.) (optional).

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## Value

```
repgrid object
```

#### Note

Currently the main purpose is to change element names. Future implementations will allow to set further attributes.

#### Author(s)

Mark Heckmann

#### See Also

```
setConstructAttr
```

## **Examples**

```
## Not run:
    x <- setElementAttr(boeker, 1, "new name")  # change name of first element
    x
## End(Not run)</pre>
```

setScale

Set the scale range of a grid.

## **Description**

The scale must be known for certain operations, e.g. to swap the construct poles. If the user construes a grid he should make sure that the scale range is set correctly.

## Usage

```
setScale(x, min, max, step, ...)
```

## **Arguments**

```
x repgrid object.
min Minimal possible scale value for ratings.
max Maximal possible scale value for ratings.
step Steps the scales uses (not yet in use).
... Not evaluated.
```

settings 101

#### Value

```
repgrid object
```

#### Author(s)

Mark Heckmann

#### **Examples**

```
## Not run:
    x <- bell2010
    x <- setScale(x, 0, 8)  # not set correctly
    x
    x <- setScale(x, 1, 7)  # set correctly
    x
## End(Not run)</pre>
```

settings

global settings for OpenRepGrid

## **Description**

global settings for OpenRepGrid

## Usage

```
settings(...)
```

#### **Arguments**

Use parameter value pairs (par1=val1, par2=val2) to change a parameter. Use parameter names to request parameter's value ("par1", "par2").

#### Note

Currently the following parameters can be changed, ordered by topic. The default value is shown in the brackets at the end of a line.

Printing grid to the console

- show. scale Show grid scale info? (TRUE)
- show.meta Show grid meta data? (TRUE)
- show.trim Number of chars to trim strings to (30)
- show. cut Maximum number of characters printed on the sides of a grid (20)
- c.no Print construct ID number? (TRUE)
- e.no Print element ID number? (TRUE)

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#### **Examples**

```
## Not run:
# get current settings
settings()

# get some parameters
settings("show.scale", "show.meta")

# change parameters
bell2010

settings(show.meta=F)
bell2010

settings(show.scale=F, show.cut=30)
bell2010

## End(Not run)
```

settingsLoad

Load OpenRepGrid settings

## **Description**

OpenRepGrid settings saved in an a settings file with the extension .orgset can be loaded to restore the settings.

## Usage

```
settingsLoad(file)
```

## Arguments

file

Path of the file to be loaded. If a path is not supplied an interactive file chooser dialog is opened.

settingsSave

Save OpenRepGrid settings

## **Description**

The current settings of OpenRepGrid can be saved into a file with the extension .orgset.

## Usage

```
settingsSave(file)
```

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## **Arguments**

file

Path of the file to be saved to. If a path is not supplied an interactive file saver dialog is opened.

shift

Shift construct or element to first position.

## Description

Shifts the whole grid vertically or horizontally so that the order remains the same but the prompted element or construct appears in first position.

## Usage

```
shift(x, c = 1, e = 1)
```

## Arguments

x repgrid object.

c Index of construct to be shifted to first position.

e Index of element to be shifted to first position.

## Value

repgrid object.

## Author(s)

Mark Heckmann

```
## Not run:

# shift element 13: 'Ideal self' to first position
shift(feixas2004, 13)

x <- randomGrid(5,10)
shift(x, 3, 5)

## End(Not run)</pre>
```

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show, repgrid-method

Show method for repgrid

## **Description**

Show method for repgrid

## Usage

```
## S4 method for signature 'repgrid'
show(object)
```

## Arguments

object

A repgrid object.

statsElements

Descriptive statistics for constructs and elements of a grid.

## Description

Descriptive statistics for constructs and elements of a grid.

Descriptive statistics for constructs and elements of a grid.

## Usage

```
statsElements(x, index = TRUE, trim = 20)
statsConstructs(x, index = T, trim = 20)
```

## **Arguments**

x repgrid object.

index Whether to print the number of the element.

trim The number of characters an element or a construct is trimmed to (default is

20). If NA no trimming occurs. Trimming simply saves space when displaying

correlation of constructs or elements with long names.

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#### Value

```
A dataframe containing the following measures is returned invisibly (see describe): item name item number number of valid cases mean standard deviation trimmed mean (with trim defaulting to .1) median (standard or interpolated) mad: median absolute deviation (from the median) minimum maximum skew kurtosis standard error
```

#### Note

Note that standard deviation and variance are estimations, i.e. including Bessel's correction. For more info type ?describe.

Note that standard deviation and variance are estimated ones, i.e. including Bessel's correction. For more info type ?describe.

#### Author(s)

Mark Heckmann
Mark Heckmann

```
## Not run:
    statsConstructs(fbb2003)
    statsConstructs(fbb2003, trim=10)
    statsConstructs(fbb2003, trim=10, index=F)

    statsElements(fbb2003)
    statsElements(fbb2003, trim=10)
    statsElements(fbb2003, trim=10, index=F)

    # save the access the results
    d <- statsElements(fbb2003)
    d
    d["mean"]
    d[2, "mean"] # mean rating of 2nd element

    d <- statsConstructs(fbb2003)
    d
    d["sd"]
    d[1, "sd"] # sd of ratings on first construct</pre>
```

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```
## End(Not run)
```

swapConstructs

Swap the position of two constructs in a grid.

## Description

Swap the position of two constructs in a grid.

## Usage

```
swapConstructs(x, pos1 = 1, pos2 = 1)
```

## **Arguments**

x repgrid object.

pos1 Row number of first construct to be swapped (default=1).

pos2 Row number of second construct to be swapped (default=1).

#### Value

repgrid object

## Author(s)

Mark Heckmann

```
## Not run:
    x <- randomGrid()
    swapConstructs(x, 1, 3)  # swap constructs 1 and 3
    swapConstructs(x, 1:2, 3:4)  # swap construct 1 with 3 and 2 with 4
## End(Not run)</pre>
```

swapElements 107

swa	n⊦ I	emer	าปร

Swap the position of two elements in a grid.

## Description

Swap the position of two elements in a grid.

## Usage

```
swapElements(x, pos1 = 1, pos2 = 1)
```

## **Arguments**

```
x repgrid object.
```

pos1 Column number of first element to be swapped (default=1).

pos2 Column number of second element to be swapped (default=1).

#### Value

repgrid object.

#### Author(s)

Mark Heckmann

## **Examples**

```
## Not run:
    x <- randomGrid()
    swapElements(x, 1, 3)  # swap elements 1 and 3
    swapElements(x, 1:2, 3:4)  # swap element 1 with 3 and 2 with 4
## End(Not run)</pre>
```

swapPoles

Swaps the construct poles.

## Description

Swaps the constructs poles and re-adjusts ratings accordingly.

#### Usage

```
swapPoles(x, pos)
```

#### **Arguments**

x repgrid object.

pos Row number of construct whose poles are swapped

## Value

repgrid object.

#### Note

Please note that the scale of the rating grid has to be set in order to swap poles. If the scale is unknown no swapping occurs and a warning is issued on the console.

#### Author(s)

Mark Heckmann

#### **Examples**

```
## Not run:

x <- randomGrid()
swapPoles(x, 1)  # swap construct poles of construct
swapPoles(x, 1:2)  # swap construct poles of construct 1 and 2
swapPoles(x)  # swap all construct poles
## End(Not run)</pre>
```

```
[,repgrid,ANY,ANY-method
```

Extract parts of the repgrid object.

## Description

Methods for "[", i.e., subsetting of repgrid objects.

## Usage

```
## S4 method for signature 'repgrid,ANY,ANY'
x[i, j, ..., drop = TRUE]
```

## **Arguments**

```
x A repgrid object.i, j Row and column indices.... Not evaluated.drop Not used.
```

[<-,repgrid-method 109

## Author(s)

Mark heckmann

## **Examples**

```
x <- randomGrid()
x[1:4, ]
x[ , 1:3]
x[1:4,1:3]
x[1,1]</pre>
```

[<-,repgrid-method

Method for "<-" assignment of the repgrid ratings.

## Description

It should be possible to use it for ratings on all layers.

## Usage

```
## S4 replacement method for signature 'repgrid' x[i, j, ...] \leftarrow value
```

## **Arguments**

x A repgrid object.
i, j Row and column indices.
... Not evaluated.
value Numeric replacement value(s).

#### Author(s)

Mark Heckmann

```
## Not run:
    x <- randomGrid()
    x[1,1] <- 2
    x[1, ] <- 4
    x[ ,2] <- 3
## End(Not run)</pre>
```

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