

Package ‘SCVA’

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Title Single-Case Visual Analysis

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Description Make graphical representations of single case data and transform graphical displays back to raw data, as discussed in Bulte and Onghena (2013) <[doi:10.22237/jmasm/1383280020](https://doi.org/10.22237/jmasm/1383280020)>. The package also includes tools for visually analyzing single-case data, by displaying central location, variability and trend.

License GPL (>= 2)

Imports ggExtra, ggplot2, graphics, plotly, scales, stats, utils

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SCVA-package

Single-Case Visual Analysis

Description

Make graphical representations of single case data and transform graphical displays back to raw data. The package also includes tools for visually analyzing single-case data, by displaying central location, variability and trend.

Details

Package: SCVA
Type: Package
Version: 1.3.1
Date: 2019-12-20
License: GPL version 2 or newer

Author(s)

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AB

Hypothetical AB data

Description

Hypothetical data collected in an AB phase design with 27 measurement times.

Usage

```
data(AB)
```

Format

A data frame with 27 observations.

- The first column contains the condition/phase labels ("A" and "B").
- The second column contains the obtained scores.
- The rows and columns are not labeled.

References

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

Examples

```
data(AB)
```

graph	<i>Graphical representation of single-case data</i>
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Description

The observed single-case data are plotted.

Usage

```
graph(design,data=read.table(file.choose(new=FALSE)),xlab="Measurement Times",
ylab="Scores",ylim=NULL,legendxy=NULL,labels=c("A","B","A","B"))
```

Arguments

design	Type of single-case design: "AB", "ABA", "ABAB", "CRD"(completely randomized design), "RBD" (randomized block design), "ATD" (alternating treatments design), "MBD" (multiple-baseline AB design) or "Custom" (user specified design).
data	File in which the data can be found. Default: a window pops up in which the file can be selected.
xlab	Label x axis.
ylab	Label y axis.
ylim	Y axis limits in the form c(min, max).
legendxy	Optional legend location x and y coordinates in the form c(x coordinate, y coordinate). Only used when design is "CRD", "RBD", "ATD" or "Custom".
labels	Optional labels for treatment levels in the form c("A", "B").

Details

When using the default data argument, a window will pop up to ask in what file the data can be found. This text file containing the data should consist of two columns for single-case phase and alternation designs: the first with the condition labels and the second with the obtained scores. For multiple-baseline designs it should consist of these two columns for EACH unit. This way, each row represents one measurement occasion. It is important not to label the rows or columns.

Missing data should be indicated as NA. For calculations, missing data are omitted.

For alternation designs, after the plot is drawn, the location of the legend should be indicated by a left mouse click.

References

Bulte, I., & Onghena, P. (2008). An R package for single-case randomization tests. *Behavior Research Methods*, 40, 467-478.

Bulte, I., & Onghena, P. (2009). Randomization tests for multiple baseline designs: An extension of the SCRT-R package. *Behavior Research Methods*, 41, 477-485.

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

See Also

[graph.CL](#) to plot a measure of central tendency as a line parallel to the abscissa.

[graph.VAR](#) to display variability information.

[graph.TREND](#) to display a possible trend in the data.

[graphly](#) to display an interactive plot.

Examples

```
data(AB)
graph(design = "AB", data = AB)
```

graph.CL

Plot a measure of central tendency

Description

A measure of central tendency ((trimmed) mean, (broadened) median, M-estimator) is plotted as a horizontal reference line superimposed on the raw time series data.

Usage

```
graph.CL(design,CL,tr,data=read.table(file.choose(new=FALSE)),xlab="Measurement Times",
ylab="Scores",ylim=NULL,legendxy=NULL,labels=c("A","B","A","B"))
```

Arguments

design	Type of single-case design: "AB", "ABA", "ABAB", "CRD"(completely randomized design), "RBD" (randomized block design), "ATD" (alternating treatments design), "MBD" (multiple-baseline AB design) or "Custom" (user specified design).
CL	Measure of central tendency: "mean", "median", "bmed" (broadened median), "trimmean" (trimmed mean), or "mest" (M-estimator of location).

tr	If CL="trimmean": the percentage of observations that has to be removed from the end of the distribution before computing the mean. It can be any value from 0 (regular arithmetic mean) to 0.5. Usually 20 percent of the observations is trimmed (so tr=0.2). If CL="mest": the desired value for the constant K. Usually a percentile of the standard normal distribution is chosen. Wilcox (2005) suggests using K=1.28, which corresponds to the 90th percentile of the standard normal distribution and covers 80 percent of the underlying distribution.
data	File in which the data can be found. Default: a window pops up in which the file can be selected.
xlab	Label x axis.
ylab	Label y axis.
ylim	Y axis limits in the form c(min, max).
legendxy	Optional legend location x and y coordinates in the form c(x coordinate, y coordinate). Only used when design is "CRD", "RBD", "ATD" or "Custom".
labels	Optional labels for treatment levels in the form c("A", "B").

Details

When using the default data argument, a window will pop up to ask in what file the data can be found. This text file containing the data should consist of two columns for single-case phase and alternation designs: the first with the condition labels and the second with the obtained scores. For multiple-baseline designs it should consist of these two columns for EACH unit. This way, each row represents one measurement occasion. It is important not to label the rows or columns.

Missing data should be indicated as NA. For calculations, missing data are omitted. Please note that some of the complicated plots may not work if there is missing data.

For alternation designs, after the plot is drawn, the location of the legend should be indicated by a left mouse click.

Note

For the calculation of the M-estimator of location, the function `mest(x,bend=1.28)` from Wilcox (2005) is used.

Wilcox, R.R. (2005). Introduction to robust estimation and hypothesis testing (2nd ed.). San Diego, CA: Elsevier Academic Press.

References

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

See Also

[graph](#) to simply plot raw single-case data.

[graph.VAR](#) to display variability information.

[graph.TREND](#) to display a possible trend in the data.

[graphly](#) to display an interactive plot.

Examples

```
data(AB)
graph.CL(design = "AB", CL = "mean", data = AB)
```

graph.extract	<i>Extract raw data from graphs</i>
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Description

Extract data values from existing graphs in jpeg format.

Usage

```
graph.extract(MT, refX, refY, save = "no", image = read.jpeg(file.choose()))
```

Arguments

MT	Measurement times: number of observations
refX	2 reference values for the X scale (e.g., refX=c(2, 14)).
refY	2 reference values for the Y scale (e.g., refY=c(10, 16)).
save	Save the extracted data points to a file (save="yes") or just see them as output in the R console (default: save="no").
image	Jpeg file in which the graph can be found. Default: a window pops up in which the file can be selected.

Details

When using the default image argument, a window will pop up to ask in what jpeg file the graph can be found.

Then the reference values should be located on the graph by clicking with the left mouse button: first the X values and then the Y values. The marked reference points are indicated with a blue cross.

Then each data point should be located sequentially. The marked data points are indicated with a red circle.

When finished, the calibrated true data points are displayed as output in the R console and plotted in a new graph, as a quick visual check.

When choosing to save these data values to a file (save="yes"), another window will pop up to ask where to save them. Here one can create a new txt file in the folder of choice.

Note

This function is an adaptation of: Timothee. (2010, March 5). Data visualization (in R). Getting data from an image (introductory post). [Web log message]. Retrieved from <http://rdataviz.wordpress.com/2010/03/05/getting-data-from-an-image-introductory-post/>.

We would like to thank David Howell for pointing us to this function.

The function based on functions adopted from the ReadImages package, which has been orphaned and archived by CRAN.

References

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

graph.TREND

Plot a trend in central location

Description

Visualizes systematic shifts in central location over time using several methods. A vertical line graph plots the deviations from each data point to a measure of central tendency against time. Regression lines superimpose a linear function on the raw data by means of least squares regression, the split-middle method or the resistant trend line fitting method. The presence of a nonlinear trend can be displayed with running medians.

Usage

```
graph.TREND(design, TREND, CL, tr, data = read.table(file.choose(new = FALSE)),
  xlab = NULL, ylab = NULL, ylim = NULL, legendxy = NULL, labels = c("A", "B", "A", "B"))
```

Arguments

design	Type of single-case design: "AB", "ABA", "ABAB", "CRD"(completely randomized design), "RBD" (randomized block design), "ATD" (alternating treatments design), "MBD" (multiple-baseline AB design) or "Custom" (user specified design).
TREND	Trend visualization: vertical line plot ("VLP"); trend lines by means of least squares regression ("LSR"), split-middle method ("SM"), resistant trend line fitting ("RTL"), or running medians ("RM3", "RM5" or "RM42", depending on the desired batch size).
CL	Measure of central tendency: "mean", "median", "bmed" (broadened median), "trimmean" (trimmed mean), or "mest" (M-estimator of location).
tr	If CL="trimmean": the percentage of observations that has to be removed from the end of the distribution before computing the mean. It can be any value from 0 (regular arithmetic mean) to 0.5. Usually 20 percent of the observations is trimmed (so tr=0.2). If CL="mest": the desired value for the constant K. Usually a percentile of the standard normal distribution is chosen. Wilcox (2005) suggests using K=1.28, which corresponds to the 90th percentile of the standard normal distribution and covers 80 percent of the underlying distribution.
data	File in which the data can be found. Default: a window pops up in which the file can be selected.
xlab	Label x axis.
ylab	Label y axis.

ylim	Y axis limits in the form c(min, max).
legendxy	Optional legend location x and y coordinates in the form c(x coordinate, y coordinate). Only used when design is "CRD", "RBD", "ATD" or "Custom".
labels	Optional labels for treatment levels in the form c("A", "B").

Details

When using the default data argument, a window will pop up to ask in what file the data can be found. This text file containing the data should consist of two columns for single-case phase and alternation designs: the first with the condition labels and the second with the obtained scores. For multiple-baseline designs it should consist of these two columns for EACH unit. This way, each row represents one measurement occasion. It is important not to label the rows or columns.

Missing data should be indicated as NA. For calculations, missing data are omitted. Please note that some of the complicated plots may not work if there is missing data.

For alternation designs, after the plot is drawn, the location of the legend should be indicated by a left mouse click.

Note

For the calculation of the M-estimator of location, the function `mest(x,bend=1.28)` from Wilcox (2005) is used.

Wilcox, R.R. (2005). Introduction to robust estimation and hypothesis testing (2nd ed.). San Diego, CA: Elsevier Academic Press.

References

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

See Also

[graph](#) to simply plot raw single-case data.

[graph.CL](#) to plot a measure of central tendency as a line parallel to the abscissa.

[graph.VAR](#) to display variability information.

[graphly](#) to display an interactive plot.

Examples

```
data(AB)
graph.TREND(design = "AB", TREND = "VLP", CL = "mean", data = AB)
graph.TREND(design = "AB", TREND = "LSR", data = AB)
graph.TREND(design = "AB", TREND = "SM", data = AB)
graph.TREND(design = "AB", TREND = "RTL", data = AB)
graph.TREND(design = "AB", TREND = "RM3", data = AB)
graph.TREND(design = "AB", TREND = "RM5", data = AB)
graph.TREND(design = "AB", TREND = "RM42", data = AB)
```


graph.VAR

*Plot an estimate of variability***Description**

Information about variability in the data is displayed by three methods. Range bar graphs consist of a vertical line for each phase, created by connecting three points: an estimate of central tendency ((trimmed) mean, (broadened) median, M-estimator), the minimum and the maximum. Range lines consist of a pair of lines parallel to the X-axis, passing through the lowest and highest values for each phase, and superimposed on the raw data. Trended ranges display changes in variability within phases. For all these methods, the influence of outliers may be lessened by using a trimmed range, in which only a sample of the data set is used.

Usage

```
graph.VAR(design, VAR, dataset = "regular", CL, tr,
data = read.table(file.choose(new = FALSE)), xlab = NULL, ylab = NULL,
ylim = NULL, legendxy = NULL, labels = c("A", "B", "A", "B"))
```

Arguments

design	Type of single-case design: "AB", "ABA", "ABAB", "CRD"(completely randomized design), "RBD" (randomized block design), "ATD" (alternating treatments design), "MBD" (multiple-baseline AB design) or "Custom" (user specified design).
VAR	Estimate of variability: range bar graph ("RB"), range lines ("RL"), or trended range ("TR").
dataset	Use the whole dataset ("regular") or reduce the influence of outliers by removing the 10-20 percent extreme values from each phase ("trimmed"). Default: the whole dataset is used.
CL	Measure of central tendency: "mean", "median", "bmed" (broadened median), "trimmean" (trimmed mean), or "mest" (M-estimator of location).
tr	If CL="trimmean": the percentage of observations that has to be removed from the end of the distribution before computing the mean. It can be any value from 0 (regular arithmetic mean) to 0.5. Usually 20 percent of the observations is trimmed (so tr=0.2). If CL="mest": the desired value for the constant K. Usually a percentile of the standard normal distribution is chosen. Wilcox (2005) suggests using K=1.28, which corresponds to the 90th percentile of the standard normal distribution and covers 80 percent of the underlying distribution.
data	File in which the data can be found. Default: a window pops up in which the file can be selected.
xlab	Label x axis.
ylab	Label y axis.
ylim	Y axis limits in the form c(min, max).

legendxy	Optional legend location x and y coordinates in the form c(x coordinate, y coordinate). Only used when design is "CRD", "RBD", "ATD" or "Custom".
labels	Optional labels for treatment levels in the form c("A", "B").

Details

When using the default data argument, a window will pop up to ask in what file the data can be found. This text file containing the data should consist of two columns for single-case phase and alternation designs: the first with the condition labels and the second with the obtained scores. For multiple-baseline designs it should consist of these two columns for EACH unit. This way, each row represents one measurement occasion. It is important not to label the rows or columns.

Missing data should be indicated as NA. For calculations, missing data are omitted. Please note that some of the complicated plots may not work if there is missing data.

For alternation designs, after the plot is drawn, the location of the legend should be indicated by a left mouse click.

Note

For the calculation of the M-estimator of location, the function `mest(x,bend=1.28)` from Wilcox (2005) is used.

Wilcox, R.R. (2005). Introduction to robust estimation and hypothesis testing (2nd ed.). San Diego, CA: Elsevier Academic Press.

References

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

See Also

[graph](#) to simply plot raw single-case data.

[graph.CL](#) to plot a measure of central tendency as a line parallel to the abscissa.

[graph.TREND](#) to display a possible trend in the data.

[graphly](#) to display an interactive plot.

Examples

```
data(AB)
graph.VAR(design = "AB", VAR = "RB", CL = "mean", data = AB)
graph.VAR(design = "AB", VAR = "RL", data = AB)
graph.VAR(design = "AB", VAR = "TR", data = AB)
graph.VAR(design = "AB", VAR = "RL", dataset = "trimmed", data = AB)
```

graphly

Interactive plot of single-case data

Description

The observed single-case data are plotted using plotly, a web-based interactive charting library.

Usage

```
graphly(design,data=read.table(file.choose(new=FALSE)),xlab="Measurement Times",
ylab="Scores",ylim=NULL,labels=c("A","B","A","B"))
```

Arguments

design	Type of single-case design: "AB", "ABA", "ABAB", "CRD"(completely randomized design), "RBD" (randomized block design), "ATD" (alternating treatments design), "MBD" (multiple-baseline AB design) or "Custom" (user specified design).
data	File in which the data can be found. Default: a window pops up in which the file can be selected.
xlab	Label x axis.
ylab	Label y axis.
ylim	Y axis limits in the form c(min, max).
labels	Optional labels for treatment levels in the form c("A", "B").

Details

When using the default data argument, a window will pop up to ask in what file the data can be found. This text file containing the data should consist of two columns for single-case phase and alternation designs: the first with the condition labels and the second with the obtained scores. For multiple-baseline designs it should consist of these two columns for EACH unit. This way, each row represents one measurement occasion. It is important not to label the rows or columns.

Missing data should be indicated as NA. For calculations, missing data are omitted.

References

Bulte, I., & Onghena, P. (2008). An R package for single-case randomization tests. *Behavior Research Methods*, 40, 467-478.

Bulte, I., & Onghena, P. (2009). Randomization tests for multiple baseline designs: An extension of the SCRT-R package. *Behavior Research Methods*, 41, 477-485.

Bulte, I., & Onghena, P. (2012). When the truth hits you between the eyes: A software tool for the visual analysis of single-case experimental data. *Methodology*, 8, 104-114.

<http://ppw.kuleuven.be/home/english/research/mesrg>

<https://plotly-r.com/overview.html>

See Also

[graph](#) to simply plot raw single-case data.

[graph.CL](#) to plot a measure of central tendency as a line parallel to the abscissa.

[graph.VAR](#) to display variability information.

[graph.TREND](#) to display a possible trend in the data.

Examples

```
data(AB)
graphly(design = "AB", data = AB)
```

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