

Package ‘blandr’

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Title Bland-Altman Method Comparison

Version 0.6.0

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Description Carries out Bland Altman analyses (also known as a Tukey mean-difference plot) as described by JM Bland and DG Altman in 1986 <[doi:10.1016/S0140-6736\(86\)90837-8](https://doi.org/10.1016/S0140-6736(86)90837-8)>. This package was created in 2015 as existing Bland-Altman analysis functions did not calculate confidence intervals. This package was created to rectify this, and create reproducible plots. This package is also available as a module for the 'jamovi' statistical spreadsheet (see <<https://www.jamovi.org>> for more information).

Depends R (>= 3.2.0)

License GPL-3

Encoding UTF-8

LazyData true

URL <https://github.com/deepankardatta/blandr/>

BugReports <https://github.com/deepankardatta/blandr/issues>

Imports glue, ggplot2, knitr, stringr, jmvcore (>= 0.8.5), markdown, rmarkdown

Suggests testthat

SystemRequirements pandoc (>=1.12.3)

VignetteBuilder knitr

Collate 'blandr.data.preparation.r' 'blandr.dataset.fibre.r'
'blandr.dataset.sbp.r' 'blandr.dataset.o2sats.r'
'blandr.dataset.pefr.r' 'blandr.dataset.load.r'
'blandr.plot.ggplot.r' 'blandr.plot.rplot.r'
'blandr.plot.limits.r' 'blandr.statistics.r' 'blandr.draw.r'
'blandr.output.text.r' 'blandr.display.and.draw.r'
'blandr.display.and.plot.r' 'blandr.method.comparison.r'
'blandr.output.report.r' 'blandr.plot.normality.r'
'blandr.plot.qq.r' 'blandr.plot.r'

```
'blandr.statistics.formula.R' 'jamovibaanalysis.b.R'
'jamovibaanalysis.h.R' 'jamovibaplothistogram.b.R'
'jamovibaplothistogram.h.R' 'jamovibaplotqq.b.R'
'jamovibaplotqq.h.R' 'jamovibastats.b.R' 'jamovibastats.h.R'
'print.blandr.R'
```

RoxygenNote 7.3.1

Language en-GB

NeedsCompilation no

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`bland.altman.PEFR.1986`*Sample PEFr comparison data from Bland-Altman (1986)*

Description

This is the sample PEFr data set from the 1986 Lancet paper written by Bland and Altman. I do not claim any copyright on the data - this is meant to allow testing of the function. I encourage future package authors to use the .rda file if they so wish.

Usage

```
data("bland.altman.PEFR.1986")
```

Format

A data frame with 17 observations on the following 4 variables.

WrightFirst a numeric vector

WrightSecond a numeric vector

MiniWrightFirst a numeric vector

MiniWrightSecond a numeric vector

References

Bland, J. M., & Altman, D. (1986). Statistical methods for assessing agreement between two methods of clinical measurement. *The Lancet*, 327(8476), 307-310. PMID:2868172. [http://dx.doi.org/10.1016/S0140-6736\(86\)90837-8](http://dx.doi.org/10.1016/S0140-6736(86)90837-8)

Examples

```
data(bland.altman.PEFR.1986)
```

`blandr.data.preparation`*Data preparation for method comparison analysis*

Description

Prepares the data and runs error checks before the calling function runs whatever method analysis mode is wants.

Usage

```
blandr.data.preparation(method1, method2, sig.level)
```

Arguments

method1	A list of numbers.
method2	A list of numbers.
sig.level	Significance level. Is not optional in this function, as the calling package should have a default value to pass if needed

Value

method.comparison A data frame of paired values. These have been data checked, and empty rows omitted, from the originally supplied data.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Calls the function - do note that this function was really
# meant to be called from other functions and not a stand-alone funtion
blandr.data.preparation( measurement1 , measurement2, sig.level=0.95 )
```

blandr.dataset.fibre	<i>Function to load D'arbela mean velocity of circumferential fibre shortening dataset from internet</i>
----------------------	--

Description

Loads the D'arbela mean velocity of circumferential fibre shortening dataset from Martin Bland's website.

Usage

```
blandr.dataset.fibre()
```

Value

converted.from.dct A data frame containing the dataset

Note

The function converts the STATA DCT data format into a data frame that R can process.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

<https://www-users.york.ac.uk/~mb55/datasets/pefr.dct> - The D'arbela mean velocity of circumferential fibre shortening dataset from Martin Bland's website

<https://www-users.york.ac.uk/~mb55/datasets/datasets.htm> - Martin Bland's example data-set webpage

Bland JM, Altman DG. (1986) Statistical methods for assessing agreement between two methods of clinical measurement. *Lancet* i, 307-310.

Examples

```
blandr.dataset.fibre()
pefr.data <- blandr.dataset.fibre()
```

`blandr.dataset.load` *Function to load example data sets*

Description

Loads example data sets from the internet.

Usage

```
blandr.dataset.load(dataset.name)
```

Arguments

`dataset.name` Loads the requisite data set. See the description for further details.

Value

example.dataset A data frame containing the requisite dataset

Note

Dataset 1 ("1", "PEFR", "pefr") - Bland Altman PEFR dataset (from `blandr.dataset.pefr`)

Dataset 2 ("2", "o2sats", "sealey") - Selaey oxygen saturations dataset (from `blandr.dataset.o2sats`)

Dataset 3 ("3", "fibre", "darbela") - D'arbela mean velocity of circumferential fibre shortening dataset (from `blandr.dataset.fibre`)

Dataset 4 ("4", "sbp", "close") - Close systolic blood pressure dataset (from `blandr.dataset.sbp`)

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
blandr.dataset.load( "pefr" )  
pefr.data <- blandr.dataset.load( "pefr" )
```

`blandr.dataset.o2sats` *Function to load Bland-Altman oxygen saturation dataset from internet*

Description

Loads the Bland-Altman oxygen saturation dataset from Martin Bland's website.

Usage

```
blandr.dataset.o2sats()
```

Value

converted.from.dct A data frame containing the dataset

Note

The function converts the STATA DCT data format into a data frame that R can process.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

<https://www-users.york.ac.uk/~mb55/datasets/sealey.dct> - The oxygen saturation dataset from Martin Bland's website

<https://www-users.york.ac.uk/~mb55/datasets/datasets.htm> - Martin Bland's example data-set webpage

Bland JM, Altman DG. (1986) Statistical methods for assessing agreement between two methods of clinical measurement. *Lancet* i, 307-310.

Examples

```
blandr.dataset.o2sats()  
pefr.data <- blandr.dataset.o2sats()
```

blandr.dataset.pefr	<i>Function to load Bland-Altman PEFR dataset from internet</i>
---------------------	---

Description

Loads the Bland-Altman PEFR dataset from Martin Bland's website.

Usage

```
blandr.dataset.pefr()
```

Value

converted.from.dct A data frame containing the dataset

Note

The function converts the STATA DCT data format into a data frame that R can process.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

<https://www-users.york.ac.uk/~mb55/datasets/pefr.dct> - The PEFR dataset from Martin Bland's website

<https://www-users.york.ac.uk/~mb55/datasets/datasets.htm> - Martin Bland's example data-set webpage

Bland JM, Altman DG. (1986) Statistical methods for assessing agreement between two methods of clinical measurement. Lancet i, 307-310.

Examples

```
blandr.dataset.pefr()
pefr.data <- blandr.dataset.pefr()
```

blandr.dataset.sbp	<i>Function to load Close systolic blood pressure dataset from internet</i>
--------------------	---

Description

Loads the Close systolic blood pressure dataset from Martin Bland's website.

Usage

```
blandr.dataset.sbp()
```

Value

converted.from.dct A data frame containing the dataset

Note

The function converts the STATA DCT data format into a data frame that R can process.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

<https://www-users.york.ac.uk/~mb55/datasets/pefr.dct> - The Close systolic blood pressure dataset from Martin Bland's website

<https://www-users.york.ac.uk/~mb55/datasets/datasets.htm> - Martin Bland's example data-set web-page

Bland JM, Altman DG. (1995) Comparing methods of measurement: why plotting difference against standard method is misleading. Lancet, 346, 1085-7.

Examples

```
blandr.dataset.sbp()
pefr.data <- blandr.dataset.sbp()
```

blandr.display.and.draw
Bland-Altman display and draw for R

Description

Stub function: calls both the display and plots functions (in that order). Uses the same parameters as the plot and display functions to allow easy all-in-one use.

This function may be deprecated in future, as you really can use the functions easily separately.

Usage

```
blandr.display.and.draw(
  method1,
  method2,
  plotter = "ggplot",
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  sig.level = 0.95,
  annotate = FALSE,
  ciDisplay = TRUE,
  ciShading = FALSE,
  normalLow = FALSE,
  normalHigh = FALSE,
  lowest_y_axis = FALSE,
  highest_y_axis = FALSE,
  point_size = 0.8
)
```

Arguments

method1	A list of numbers.
method2	A list of numbers.
plotter	(Optional- default='ggplot') Selects which graphics engine to use to plot the Bland-Altman charts. 2 options are 'ggplot' or 'rplot'. If unknown parameter sent, will default to 'ggplot'
method1name	(Optional) Plotting name for 1st method, default 'Method 1'
method2name	(Optional) Plotting name for 2nd method, default 'Method 2'
plotTitle	(Optional) Title name, default 'Bland-Altman plot for comparison of 2 methods'
sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.
annotate	(Optional) TRUE/FALSE switch to provides annotations to plot, default=FALSE

ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default=TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default=TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
lowest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the lower boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
highest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the upper boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
point_size	(Optional) Size of marker for each dot. Default is cex=0.8

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates a plot, with no optional arguments
blandr.display.and.draw( measurement1 , measurement2 )

# Generates a plot, with title
blandr.display.and.draw( measurement1 , measurement2 , plotTitle = 'Bland-Altman example plot' )
```

blandr.display.and.plot

(DEPRECATED) Bland-Altman display and plot for R

Description

(DEPRECATED) Re-directs to blandr.display.and.draw

Usage

```

blandr.display.and.plot(
  method1,
  method2,
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  sig.level = 0.95,
  annotate = FALSE,
  ciDisplay = TRUE,
  ciShading = FALSE,
  normalLow = FALSE,
  normalHigh = FALSE,
  lowest_y_axis = FALSE,
  highest_y_axis = FALSE,
  point_size = 0.8
)

```

Arguments

method1	A list of numbers.
method2	A list of numbers.
method1name	(Optional) Plotting name for 1st method, default 'Method 1'
method2name	(Optional) Plotting name for 2nd method, default 'Method 2'
plotTitle	(Optional) Title name, default 'Bland-Altman plot for comparison of 2 methods'
sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.
annotate	(Optional) TRUE/FALSE switch to provides annotations to plot, default=FALSE
ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default=TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default=TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
lowest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the lower boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
highest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the upper boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
point_size	(Optional) Size of marker for each dot. Default is cex=0.8

Author(s)

Deepankar Datta deepankar.datta@gmail.com

`blandr.draw`

Bland-Altman drawing function for R

Description

Bland-Altman drawing function. Depends on the `blandr.statistics` function in the package. Will generate a plot via the standard R plotting functions.

Usage

```
blandr.draw(
  method1,
  method2,
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  sig.level = 0.95,
  LoA.mode = 1,
  annotate = FALSE,
  ciDisplay = TRUE,
  ciShading = TRUE,
  normalLow = FALSE,
  normalHigh = FALSE,
  lowest_y_axis = FALSE,
  highest_y_axis = FALSE,
  point_size = 0.8,
  overlapping = FALSE,
  plotter = "ggplot",
  x.plot.mode = "means",
  y.plot.mode = "difference",
  plotProportionalBias = FALSE,
  plotProportionalBias.se = TRUE,
  assume.differences.are.normal = TRUE
)
```

Arguments

<code>method1</code>	A vector of numbers corresponding to the results from method 1.
<code>method2</code>	A vector of numbers corresponding to the results from method 2.
<code>method1name</code>	(Optional) Plotting name for 1st method, default 'Method 1'
<code>method2name</code>	(Optional) Plotting name for 2nd method, default 'Method 2'
<code>plotTitle</code>	(Optional) Title name, default 'Bland-Altman plot for comparison of 2 methods'

sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.
LoA.mode	(Optional) Switch to change how accurately the limits of agreement (LoA) are calculated from the bias and its standard deviation. The default is LoA.mode=1 which calculates LoA with the more accurate 1.96x multiplier. LoA.mode=2 uses the 2x multiplier which was used in the original papers. This should really be kept at default, except to double check calculations in older papers.
annotate	(Optional) TRUE/FALSE switch to provides annotations to plot, default=FALSE
ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default=TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default=TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
lowest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the lower boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
highest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the upper boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
point_size	(Optional) Size of marker for each dot. Default is cex=0.8
overlapping	(Optional) TRUE/FALSE switch to increase size of plotted point if multiple values using ggplot's geom_count, default=FALSE. Not currently recommend until I can tweak the graphics to make them better
plotter	(Optional- default='ggplot') Selects which graphics engine to use to plot the Bland-Altman charts. 2 options are 'ggplot' or 'rplot'. If unknown parameter sent, will default to 'ggplot'
x.plot.mode	(Optional) Switch to change x-axis from being plotted by means ("means") or by either 1st method ("method1") or 2nd method ("method2"). Default is "means". Anything other than "means" will switch to default mode.
y.plot.mode	(Optional) Switch to change y-axis from being plotted by difference ("difference") or by proportion magnitude of measurements ("proportion"). Default is "difference". Anything other than "proportional" will switch to default mode.
plotProportionalBias	(Optional) TRUE/FALSE switch. Plots a proportional bias line. Default is FALSE.
plotProportionalBias.se	(Optional) TRUE/FALSE switch. If proportional bias line is drawn, switch to plot standard errors. See stat_smooth for details. Default is TRUE.
assume.differences.are.normal	(Optional, not operationally used currently) Assume the difference of means has a normal distribution. Will be used to build further analyses

Note

Started 2015-11-14

Last update 2015-11-19

Originally designed for LAVAS and CVLA

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates a plot, with no optional arguments
blandr.draw( measurement1 , measurement2 )

# Generates a plot, using the in-built R graphics
blandr.draw( measurement1 , measurement2 , plotter = 'rplot' )

# Generates a plot, with title changed
blandr.draw( measurement1 , measurement2 , plotTitle = 'Bland-Altman example plot' )

# Generates a plot, with title changed, and confidence intervals off
blandr.draw( measurement1 , measurement2 , plotTitle = 'Bland-Altman example plot' ,
ciDisplay = FALSE , ciShading = FALSE )
```

blandr.method.comparison

Bland-Altman method comparison

Description

Everyone likes graphs, lines and T-tests. This uses the data provided to generate simple tests whilst trying to explain why they should be treated with caution in method comparison studies. This is hopefully the first step in getting people to use the Bland-Altman functions as I suspect everyone will try to do these tests anyway.

Usage

```
blandr.method.comparison(method1, method2, sig.level = 0.95)
```

Arguments

method1	A list of numbers.
method2	A list of numbers.
sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

Based on: (1) Bland, J. M., & Altman, D. (1986). Statistical methods for assessing agreement between two methods of clinical measurement. The Lancet, 327(8476), 307-310. [http://dx.doi.org/10.1016/S0140-6736\(86\)90837-8](http://dx.doi.org/10.1016/S0140-6736(86)90837-8)

Linnet K., Limitations of the paired t-test for evaluation of method comparison data. Clin Chem. 1999 Feb;45(2):314-5. PMID: 9931067

Zaki R, Bulgiba A, Ismail R, Ismail NA. Statistical Methods Used to Test for Agreement of Medical Instruments Measuring Continuous Variables in Method Comparison Studies: A Systematic Review PLoS ONE 2012 7(5): e37908. doi: 10.1371/journal.pone.0037908

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Call the function
blandr.method.comparison( measurement1 , measurement2 )
```

blandr.output.report *Bland-Altman report generator*

Description

Generates a report for the Bland-Altman statistics using rMarkdown and Shiny.

Usage

```
blandr.output.report(method1, method2)
```

Arguments

method1	A list of numbers for the first method
method2	A list of numbers for the second method

Note

Use the function to generate a report. You can also take the .Rmd file to customise it and create your own report. Or use rMarkdown to save the contents. I couldn't add this to the function as it's not allowed in CRAN. On the other hand a full Shiny app would take too long. So this is a stop-gap way of creating this function. Hopefully I can improve it in the future

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# NOT RUN
# Generates two random measurements
# measurement1 <- rnorm(100)
# measurement2 <- rnorm(100)

# blandr.output.report( measurement1 , measurement2 )
#
# Use this to manually run the rmarkdown template
# However specify where the template is
# Also define your methods as method1 and method2 exactly
# For a reason I can't fathom (or how the list of parameters is constructed)
# not naming them method1 and method2 makes them invisible to the rMarkdown document
#
# rmarkdown::run( file = "blandr_report_template.Rmd" ,
#   render_args = list( runtime = "shiny" ,
#     params = list( method1 = method1 ,
#       method2 = method2 ) ) )
# END OF NOT RUN
```

blandr.output.text *(DEPRECATED) Bland-Altman summary statistics display function*

Description

(DEPRECATED) Displays results of Bland-Altman analysis in a nicer text format. Relies on the blandr.statistics function in the package.

Usage

```
blandr.output.text(method1, method2, sig.level = 0.95)
```

Arguments

method1	A list of numbers.
method2	A list of numbers.
sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Displays basic statistics for the two measurements in a readable form
blandr.output.text( measurement1 , measurement2 )
```

blandr.plot

(DEPRECATED) Bland-Altman drawing function for R

Description

(DEPRECATED) Re-directs to blandr.draw.r

Usage

```
blandr.plot(
  method1,
  method2,
  plotter = "ggplot",
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  sig.level = 0.95,
  annotate = FALSE,
  ciDisplay = TRUE,
  ciShading = TRUE,
  normalLow = FALSE,
  normalHigh = FALSE,
  lowest_y_axis = FALSE,
  highest_y_axis = FALSE,
  point_size = 0.8
)
```

Arguments

method1	A vector of numbers corresponding to the results from method 1.
method2	A vector of numbers corresponding to the results from method 2.
plotter	(Optional- default='ggplot') Selects which graphics engine to use to plot the Bland-Altman charts. 2 options are 'ggplot' or 'rplot'. If unknown parameter sent, will default to 'ggplot'
method1name	(Optional) Plotting name for 1st method, default 'Method 1'
method2name	(Optional) Plotting name for 2nd method, default 'Method 2'
plotTitle	(Optional) Title name, default 'Bland-Altman plot for comparison of 2 methods'
sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.
annotate	(Optional) TRUE/FALSE switch to provides annotations to plot, default=FALSE
ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default=TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default=TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
lowest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the lower boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
highest_y_axis	(Optional) Defaults to NULL If given a continuous variable will use this as the upper boundary of the y axis. Useful if need multiple plots with equivalent y-axes.
point_size	(Optional) Size of marker for each dot. Default is cex=0.8

Author(s)

Deepankar Datta deepankar.datta@gmail.com

`blandr.plot.ggplot` *Bland-Altman plotting function, using ggplot2*

Description

Draws a Bland-Altman plot using data calculated using the other functions, using ggplot2

Usage

```

blandr.plot.ggplot(
  statistics.results,
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  ciDisplay = TRUE,
  ciShading = TRUE,
  normalLow = FALSE,
  normalHigh = FALSE,
  overlapping = FALSE,
  x.plot.mode = "means",
  y.plot.mode = "difference",
  plotProportionalBias = FALSE,
  plotProportionalBias.se = TRUE,
  assume.differences.are.normal = TRUE
)

```

Arguments

statistics.results	A list of statistics generated by the blandr.statistics function: see the function's return list to see what variables are passed to this function
method1name	(Optional) Plotting name for 1st method, default "Method 1"
method2name	(Optional) Plotting name for 2nd method, default "Method 2"
plotTitle	(Optional) Title name, default "Bland-Altman plot for comparison of 2 methods"
ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default is TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default is TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
overlapping	(Optional) TRUE/FALSE switch to increase size of plotted point if multiple values using ggplot's geom_count, default=FALSE. Not currently recommend until I can tweak the graphics to make them better
x.plot.mode	(Optional) Switch to change x-axis from being plotted by means (="means") or by either 1st method (="method1") or 2nd method (="method2"). Default is "means". Anything other than "means" will switch to default mode.
y.plot.mode	(Optional) Switch to change y-axis from being plotted by difference (="difference") or by proportion magnitude of measurements (="proportion"). Default is "difference". Anything other than "proportional" will switch to default mode.
plotProportionalBias	(Optional) TRUE/FALSE switch. Plots a proportional bias line. Default is FALSE.

`plotProportionalBias.se`
 (Optional) TRUE/FALSE switch. If proportional bias line is drawn, switch to plot standard errors. See `stat_smooth` for details. Default is TRUE.

`assume.differences.are.normal`
 (Optional, not operationally used currently) Assume the difference of means has a normal distribution. Will be used to build further analyses

Value

`ba.plot` Returns a ggplot data set that can then be plotted

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates a ggplot
# Do note the ggplot function wasn't meant to be used on it's own
# and is generally called via the bland.altman.display.and.draw function

# Passes data to the blandr.statistics function to generate Bland-Altman statistics
statistics.results <- blandr.statistics( measurement1 , measurement2 )

# Generates a ggplot, with no optional arguments
blandr.plot.ggplot( statistics.results )

# Generates a ggplot, with title changed
blandr.plot.ggplot( statistics.results , plotTitle = "Bland-Altman example plot" )

# Generates a ggplot, with title changed, and confidence intervals off
blandr.plot.ggplot( statistics.results , plotTitle = "Bland-Altman example plot" ,
  ciDisplay = FALSE , ciShading = FALSE )
```

`blandr.plot.limits` *Bland-Altman plot limits for R*

Description

Works out plot limits for the Bland-Altman plots. Depends on the `blandr.statistics` function in the package.

Usage

```
blandr.plot.limits(  
  statistics.results,  
  lowest_y_axis = FALSE,  
  highest_y_axis = FALSE  
)
```

Arguments

`statistics.results` A list of statistics generated by the `blandr.statistics` function: see the function's return list to see what variables are passed to this function

`lowest_y_axis` (Optional) Defaults to NULL. If given a continuous variable will use this as the lower boundary of the y axis. Useful if need multiple plots with equivalent y-axes.

`highest_y_axis` (Optional) Defaults to NULL. If given a continuous variable will use this as the upper boundary of the y axis. Useful if need multiple plots with equivalent y-axes.

Value

`x_upper` The upper limit of the X-axis

`x_lower` The lower limit of the X-axis

`y_upper` The upper limit of the Y-axis

`y_lower` The lower limit of the Y-axis

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements  
measurement1 <- rnorm(100)  
measurement2 <- rnorm(100)  
  
# Passes data to the blandr.statistics function to generate Bland-Altman statistics  
statistics.results <- blandr.statistics( measurement1 , measurement2 )  
  
# Calls the function  
blandr.plot.limits( statistics.results )
```

`blandr.plot.normality` *Bland-Altman histogram and density plot*

Description

Generates a combined histogram and density curve for Bland-Altman differences

Usage

```
blandr.plot.normality(statistics.results)
```

Arguments

`statistics.results`

A list of statistics generated by the `blandr.statistics` function: see the function's return list to see what variables are passed to this function

Author(s)

Deepankar Datta deepankar.datta@gmail.com

`blandr.plot.qq` *Bland-Altman differences QQ plot*

Description

Generates a QQ plot for Bland-Altman differences

Usage

```
blandr.plot.qq(statistics.results)
```

Arguments

`statistics.results`

A list of statistics generated by the `blandr.statistics` function: see the function's return list to see what variables are passed to this function

Author(s)

Deepankar Datta deepankar.datta@gmail.com

blandr.plot.rplot	<i>Bland-Altman plotting function, using basic R drawing functions</i>
-------------------	--

Description

Draws a Bland-Altman plot using data calculated using the other functions, using the in-built R graphics

Usage

```
blandr.plot.rplot(
  statistics.results,
  plot.limits,
  method1name = "Method 1",
  method2name = "Method 2",
  plotTitle = "Bland-Altman plot for comparison of 2 methods",
  annotate = FALSE,
  ciDisplay = TRUE,
  ciShading = TRUE,
  normalLow = FALSE,
  normalHigh = FALSE,
  point_size = 0.8
)
```

Arguments

statistics.results	A list of statistics generated by the blandr.statistics function: see the function's return list to see what variables are passed to this function
plot.limits	A list of statistics generated by the blandr.plot.limits function to define the extent of the x- and y- axes: see the function's return list to see what variables are passed to this function
method1name	(Optional) Plotting name for 1st method, default 'Method 1'
method2name	(Optional) Plotting name for 2nd method, default 'Method 2'
plotTitle	(Optional) Title name, default 'Bland-Altman plot for comparison of 2 methods'
annotate	(Optional) TRUE/FALSE switch to provides annotations to plot, default=FALSE
ciDisplay	(Optional) TRUE/FALSE switch to plot confidence intervals for bias and limits of agreement, default=TRUE
ciShading	(Optional) TRUE/FALSE switch to plot confidence interval shading to plot, default=TRUE
normalLow	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its lower boundary
normalHigh	(Optional) If there is a normal range, entering a continuous variable will plot a vertical line on the plot to indicate its higher boundary
point_size	(Optional) Size of marker for each dot. Default is cex=0.8

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates a basic plot
# Do note the blandr.plot.rplot function wasn't meant to be used on it's own
# and is generally called via the bland.altman.display.and.draw function

# Passes data to the blandr.statistics function to generate Bland-Altman statistics
statistics.results <- blandr.statistics( measurement1 , measurement2 )
# Passed data to the blandr.plot.limits function to generate plot limits
plot.limits <- blandr.plot.limits( statistics.results )

# Generates a basic plot, with no optional arguments
blandr.plot.rplot( statistics.results , plot.limits )

# Generates a basic plot, with title changed
blandr.plot.rplot( statistics.results , plot.limits , plotTitle = 'Bland-Altman example plot' )
# Generates a basic plot, with title changed, and confidence intervals off
blandr.plot.rplot( statistics.results , plot.limits , plotTitle = 'Bland-Altman example plot' ,
ciDisplay = FALSE , ciShading = FALSE )
```

blandr.statistics	<i>Bland-Altman statistics for R</i>
-------------------	--------------------------------------

Description

Bland-Altman analysis function for R. Package created as existing functions don't suit my needs, and don't generate 95% for bias and limits of agreement. This base function calculates the basic statistics, and generates return values which can be used in the related `blandr.display` and `bland.altamn.plot` functions. However the return results can be used to generate a custom chart if desired.

Usage

```
blandr.statistics(method1, method2, sig.level = 0.95, LoA.mode = 1)
```

Arguments

method1	Either a formula, or a vector of numbers corresponding to the results from method 1.
method2	A vector of numbers corresponding to the results from method 2. Only needed if X is a vector.

sig.level	(Optional) Two-tailed significance level. Expressed from 0 to 1. Defaults to 0.95.
LoA.mode	(Optional) Switch to change how accurately the limits of agreement (LoA) are calculated from the bias and its standard deviation. The default is LoA.mode=1 which calculates LoA with the more accurate 1.96x multiplier. LoA.mode=2 uses the 2x multiplier which was used in the original papers. This should really be kept at default, except to double check calculations in older papers.

Value

An object of class 'blandr' is returned. This is a list with the following elements:

means	List of arithmetic mean of the two methods
differences	List of differences of the two methods
method1	Returns the 'method1' list in the data frame if further evaluation is needed
method2	Returns the 'method2' list in the data frame if further evaluation is needed
sig.level	Significance level supplied to the function
sig.level.convert.to.z	Significance level convert to Z value
bias	Bias of the two methods
biasUpperCI	Upper confidence interval of the bias (based on significance level)
biasLowerCI	Lower confidence interval of the bias (based on significance level)
biasStdDev	Standard deviation for the bias
biasSEM	Standard error for the bias
LOA_SEM	Standard error for the limits of agreement
upperLOA	Upper limit of agreement
upperLOA_upperCI	Upper confidence interval of the upper limit of agreement
upperLOA_lowerCI	Lower confidence interval of the upper limit of agreement
lowerLOA	Lower limit of agreement
lowerLOA_upperCI	Upper confidence interval of the lower limit of agreement
lowerLOA_lowerCI	Lower confidence interval of the lower limit of agreement
proportion	Differences/means*100
no.of.observations	Number of observations
regression.equation	A regression equation to help determine if there is any proportional bias
regression.fixed.slope	The slope value of the regression equation
regression.fixed.intercept	The intercept value of the regression equation

Note

The function will give similar answers when used on the original Bland-Altman PEFR data sets. They won't be exactly the same as (a) for 95\

The function depends on paired values.

It currently only can currently work out fixed bias.

Improvements for the future: proportional bias charts will need further work

Started 2015-11-14

Last update 2016-02-04

Originally designed for LAVAS and CVLA

Author(s)

Deepankar Datta deepankar.datta@gmail.com

References

Based on: (1) Bland, J. M., & Altman, D. (1986). Statistical methods for assessing agreement between two methods of clinical measurement. *The Lancet*, 327(8476), 307-310. [http://dx.doi.org/10.1016/S0140-6736\(86\)90837-8](http://dx.doi.org/10.1016/S0140-6736(86)90837-8)

Confidence interval work based on follow-up paper: (2) Altman, D. G., & Bland, J. M. (2002). Commentary on quantifying agreement between two methods of measurement. *Clinical chemistry*, 48(5), 801-802. <http://www.clinchem.org/content/48/5/801.full.pdf>

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates Bland-Altman statistics data of the two measurements
blandr.statistics( measurement1 , measurement2 )
```

`blandr.statistics.formula`

Bland-Altman statistics for R - submitting data as a formula

Description

If data is supplied as a formula, the function interprets it and then passes it through to the main `blandr.statistics` function.

Usage

```
blandr.statistics.formula(formula, data = parent.frame(), ...)
```

Arguments

formula	A formula, to be eventually passed to the blandr.statistics function
data	A data frame
...	other arguments.

Note

This was initially packaged in the same file as the blandr.statistics function. Changes to R package checks made it easier to separate it out, but functionally it has not changed.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
# Generates two random measurements
measurement1 <- rnorm(100)
measurement2 <- rnorm(100)

# Generates Bland-Altman statistics data of the two measurements using the formula interface

blandr.statistics.formula( measurement2 ~ measurement1 )

# Example with a real data set
blandr.statistics.formula( Method.B ~ Method.A, data = giavarina.2015 )
```

giavarina.2015	<i>Sample comparison data from Giavarina (2015)</i>
----------------	---

Description

This is sample comparison data, taken from Giavarina's 2015 paper on Bland-Altman analysis. The data is from table 1 of the paper. I do not claim any copyright on the data - this is meant to allow testing of the function. I encourage future package authors to use the .rda file if they so wish.

Usage

```
data("giavarina.2015")
```

Format

A data frame with 30 observations on the following 5 variables.

Method.A a numeric vector

Method.B a numeric vector

Mean a numeric vector
 Difference a numeric vector
 Diff.Mean.Proportion a factor with levels

References

Giavarina D. Understanding Bland Altman analysis. Biochemia Medica. 2015;25(2):141-151. doi:10.11613/BM.2015.015.

Examples

```
data(giavarina.2015)
```

jamoviBAanalysis	<i>Bland-Altman Analysis</i>
------------------	------------------------------

Description

Bland-Altman Analysis

Usage

```
jamoviBAanalysis(  
  data,  
  method1,  
  method2,  
  ciDisplay = TRUE,  
  ciShading = TRUE,  
  plotProportionalBias = FALSE,  
  plotProportionalBias.se = TRUE,  
  overlapping = FALSE  
)
```

Arguments

data	.
method1	.
method2	.
ciDisplay	.
ciShading	.
plotProportionalBias	.
plotProportionalBias.se	.
overlapping	.

Value

A results object containing:

results\$table	a table
results\$plot	an image

Tables can be converted to data frames with asDF or [as.data.frame](#). For example:

```
results$table$asDF
as.data.frame(results$table)
```

jamoviBAplotHistogram	<i>Differences Histogram and Density Curve</i>
-----------------------	--

Description

Differences Histogram and Density Curve

Usage

```
jamoviBAplotHistogram(data, method1, method2)
```

Arguments

data	.
method1	.
method2	.

Value

A results object containing:

results\$plot	an image
---------------	----------

jamoviBAplotQQ	<i>Differences Q-Q Plot</i>
----------------	-----------------------------

Description

Differences Q-Q Plot

Usage

jamoviBAplotQQ(data, method1, method2)

Arguments

data .
method1 .
method2 .

Value

A results object containing:

results\$plot an image

jamoviBASTats	<i>Bland-Altman Raw Statistics</i>
---------------	------------------------------------

Description

Bland-Altman Raw Statistics

Usage

jamoviBASTats(data, method1, method2)

Arguments

data .
method1 .
method2 .

Value

A results object containing:

results\$text a preformatted

print.blandr *Print method for blandr objects*

Description

Compactly print the statistics used to construct a Bland-Altman plot

Usage

```
## S3 method for class 'blandr'
print(x, digits = getOption("digits"), ...)
```

Arguments

x	the results from blandr.statistics.
digits	number of significant digits to be used.
...	other arguments.

Author(s)

Deepankar Datta deepankar.datta@gmail.com

Examples

```
results = blandr.statistics.formula(Method.B ~ Method.A, data = giavarina.2015)
results
```

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