

Package ‘ufs’

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Type Package

Title A Collection of Utilities

Version 0.5.12

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License GPL (>= 3)

Description This is a new version of the 'userfriendlyscience' package, which has grown a bit unwieldy. Therefore, distinct functionalities are being 'consciously uncoupled' into different packages. This package contains the general-purpose tools and utilities (see the 'behaviorchange' package, the 'rosetta' package, and the soon-to-be-released 'scd' package for other functionality), and is the most direct 'successor' of the original 'userfriendlyscience' package. For example, this package contains a number of basic functions to create higher level plots, such as diamond plots, to easily plot sampling distributions, to generate confidence intervals, to plan study sample sizes for confidence intervals, and to do some basic operations such as (dis)attenuate effect size estimates.

URL <https://ufs.opens.science>

BugReports <https://gitlab.com/r-packages/ufs/-/issues>

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Contents

aipedjmv	4
aiperjmv	5
areColors	6
arr	6
associationMatrix	7
associationsDiamondPlot	10
attenuate.d	13
attenuate.r	14
A_VarghaDelaney	15
BAC_plot	15
bfi-data	17
biAxisDiamondPlot	18
biDimColors	20
carelessObject	21
carelessReport	22
cat0	23
checkDataIntegrity	24
checkPkgs	25
CIM	27
cohensdCI	29
computeStatistic_t	32
confintdjmv	34
confIntOmegaSq	35
confIntProp	36
confIntR	37
confintrjmv	38
confIntSD	39
convert	40
convert.cer.to.d	41
convert.d.to.U3	43
convertToNumeric	43
cramersV	44
dataShape	45
descr	49
diamondCoordinates	51
diamondPlot	54
disattenuate.d	57
disattenuate.r	58

duoComparisonDiamondPlot	58
escapeRegex	62
exceptionalScore	63
exceptionalScores	64
exportToHTML	66
extractVarName	66
faConfInt	67
factorLoadingDiamondCIplot	68
factorLoadingHeatmap	70
fa_failsafe	72
findShortestInterval	72
formatCI	73
formatPvalue	74
formatR	75
getData	76
ggBarChart	77
ggBoxplot	78
ggEasyBar	79
ggPie	81
ggProportionPlot	82
ggqq	85
ggSave	87
heading	88
ifelseObj	89
insertFigureCaption	89
invertItem	91
iqrOutlier	92
irpplot	92
is.nr	93
is.odd	94
isTrue	95
kblXtra	95
knitAndSave	96
knitFig	97
makeScales	98
massConvertToNumeric	99
meanConfInt	100
meansDiamondPlot	101
meansDiamondPlotjmv	103
meanSDtoDiamondPlot	104
multiResponse	106
multiResponsejmv	107
multiVarFreq	108
normalHist	109
noZero	111
opts	111
parallelSubscales	112
pomegaSq	113

pwr.bootES	114
pwr.confIntProp	115
pwr.confIntR	116
pwr.omegasq	117
quietRemotesInstall	119
qVec	120
rbind_dfs	121
rbind_df_list	121
regrInfluential	122
repeatStr	123
report	123
safeRequire	124
scaleDiagnosis	124
scaleStructure	127
scatterMatrix	131
setCaptionNumberingKnitrHook	133
sharedSubString	134
simDataSet	135
spearmanBrown	138
strToFilename	138
suspectParticipants	139
testRetestAlpha	140
testRetestCES	141
testRetestReliability	143
testRetestSimData	145
vecTxt	147
viridisPalette	148
wrapVector	148
zotero_construct_export_call	149
zotero_download_and_export_items	150
zotero_get_all_items	151
zotero_nr_of_items	151
%IN%	152

Index 153

aipedjmv	<i>Sample size for accuracy: d</i>
----------	------------------------------------

Description

Sample size for accuracy: d

Usage

aipedjmv(d = 0.5, w = 0.1, conf.level = 95)

Arguments

d .
w .
conf.level .

Value

A results object containing:

results\$text	a html
results\$aipePlot	an image

aiperjmv	<i>Sample size for accuracy: r</i>
----------	------------------------------------

Description

Sample size for accuracy: r

Usage

aiperjmv(r = 0.3, w = 0.1, conf.level = 95)

Arguments

r .
w .
conf.level .

Value

A results object containing:

results\$text	a html
results\$aipePlot	an image

areColors

Check whether elements of a vector are valid colors

Description

This function by Josh O'Brien checks whether elements of a vector are valid colors. It has been copied from a Stack Exchange answer (see <https://stackoverflow.com/questions/13289009/check-if-character-string-is-a-valid-color-representation>).

Usage

```
areColors(x)
```

Arguments

x The vector.

Value

A logical vector.

Author(s)

Josh O'Brien

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
ufs::areColors(c(NA, "black", "blackk", "1", "#00", "#000000"));
```

arr

Absolute Relative Risk and confidence interval

Description

This is a function to conveniently and quickly compute the absolute relative risk (ARR) and its confidence interval.

Usage

```
arr(
  expPos,
  expN,
  conPos,
  conN,
  conf.level = 0.95,
  digits = 2,
  printAsPercentage = TRUE
)

## S3 method for class 'ufsARR'
print(x, digits = x$digits, printAsPercentage = x$printAsPercentage, ...)
```

Arguments

expPos	Number of positive events in the experimental condition.
expN	Total number of cases in the experimental condition.
conPos	Number of positive events in the control condition.
conN	Total number of cases in the control condition.
conf.level	The confidence level for the confidence interval.
digits	The number of digits to round to when printing the results.
printAsPercentage	Whether to multiply with 100 when printing the results.
x	The result of the call to arr.
...	Any additional arguments are neglected.

Value

An object with in estimate, the ARR, and in conf.int, the confidence interval.

Examples

```
ufs::arr(10, 60, 20, 60);
```

associationMatrix	<i>associationMatrix</i>
-------------------	--------------------------

Description

associationMatrix produces a matrix with confidence intervals for effect sizes, point estimates for those effect sizes, and the p-values for the test of the hypothesis that the effect size is zero, corrected for multiple testing.

Usage

```

associationMatrix(
  dat = NULL,
  x = NULL,
  y = NULL,
  conf.level = 0.95,
  correction = "fdr",
  bootstrapV = FALSE,
  info = c("full", "ci", "es"),
  includeSampleSize = "depends",
  bootstrapV.samples = 5000,
  digits = 2,
  pValueDigits = digits + 1,
  colNames = FALSE,
  type = c("R", "html", "latex"),
  file = "",
  statistic = associationMatrixStatDefaults,
  effectSize = associationMatrixESDefaults,
  var.equal = TRUE
)

## S3 method for class 'associationMatrix'
print(x, type = x$input$type, info = x$input$info, file = x$input$file, ...)

## S3 method for class 'associationMatrix'
pander(x, info = x$input$info, file = x$input$file, ...)

```

Arguments

<code>dat</code>	A dataframe with the variables of interest. All variables in this dataframe will be used if both <code>x</code> and <code>y</code> are <code>NULL</code> . If <code>dat</code> is <code>NULL</code> , the user will be presented with a dialog to select a datafile.
<code>x</code>	If not <code>NULL</code> , this should be a character vector with the names of the variables to include in the rows of the association table. If <code>x</code> is <code>NULL</code> , all variables in the dataframe will be used.
<code>y</code>	If not <code>NULL</code> , this should be a character vector with the names of the variables to include in the columns of the association table. If <code>y</code> is <code>NULL</code> , the variables in <code>x</code> will be used for the columns as well (which produces a symmetric matrix, similar to most correlation matrices).
<code>conf.level</code>	Level of confidence of the confidence intervals.
<code>correction</code>	Correction for multiple testing: an element out of the vector <code>c("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none")</code> . NOTE: the p-values are corrected for multiple testing; The confidence intervals are not!
<code>bootstrapV</code>	Whether to use bootstrapping to compute the confidence interval for Cramer's V or whether to use the Fisher's Z conversion.
<code>info</code>	Information to print: either both the confidence interval and the point estimate for the effect size (and the p-value, corrected for multiple testing), or only the

	confidence intervals, or only the point estimate (and the corrected p-value). Must be on element of the vector <code>c("full", "ci", "es")</code> .
<code>includeSampleSize</code>	Whether to include the sample size when the effect size point estimate and p-value are shown. If this is "depends", it will depend on whether all associations have the same sample size (and the sample size will only be printed when they don't). If "always", the sample size will always be added. If anything else, it will never be printed.
<code>bootstrapV.samples</code>	If using bootstrapping for Cramer's V, the number of samples to generate.
<code>digits</code>	Number of digits to round to when printing the results.
<code>pValueDigits</code>	How many digits to use for formatting the p values.
<code>colNames</code>	If true, the column heading will use the variables names instead of numbers.
<code>type</code>	Type of output to generate: must be an element of the vector <code>c("R", "html", "latex")</code> .
<code>file</code>	If a file is specified, the output will be written to that file instead of shown on the screen.
<code>statistic</code>	This is the complicated bit; this is where <code>associationMatrix</code> allows customization of the used statistics to perform null hypothesis significance testing. For everyday use, leaving this at the default value, <code>associationMatrixStatDefaults</code> , works fine. In case you want to customize, read the 'Notes' section below.
<code>effectSize</code>	Like the 'statistics' argument, 'effectSize' also allows customization, in this case of the used effect sizes. Again, the default value, <code>associationMatrixESDefaults</code> , works for everyday use. Again, see the 'Notes' section below if you want to customize.
<code>var.equal</code>	Whether to test for equal variances ('test'), assume equality ('yes'), or assume inequality ('no').
<code>...</code>	Addition arguments are passed on to the <code>print()</code> and <code>pander::pander()</code> functions.

Value

An object with the input and several output variables, one of which is a dataframe with the association matrix in it. When this object is printed, the association matrix is printed to the screen. If the 'file' parameter is specified, a file with this matrix will also be written to disk.

Note

The 'statistic' and 'effectSize' parameter make it possible to use different functions to conduct null hypothesis significance testing and compute effect sizes. In both cases, the parameter needs to be a list containing four lists, named 'dichotomous', 'nominal', 'ordinal', and 'interval'. Each of these lists has to contain four elements, character vectors of length one (i.e. just one string value), again named 'dichotomous', 'nominal', 'ordinal', and 'interval'.

The combination of each of these names (e.g. 'dichotomous' and 'nominal', or 'ordinal' and 'interval', etc) determine which test should be done when computing the p-value to test the association

between two variables of those types, or which effect sizes to compute. When called, `associationMatrix` determines the measurement levels of the relevant variables. It then uses these two levels (their string representation, e.g. 'dichotomous' etc) to find a string in the 'statistic' and 'effectSize' objects. Two functions with these names are then called from two lists, 'computeStatistic' and 'computeEffectSize'. These lists list contain functions that have the same names as the strings in the 'statistic' list.

For example, when the default settings are used, the string (function name) found for two dichotomous variables when searching in `associationMatrixStatDefaults` is 'chisq', and the string found in `associationMatrixESDefaults` is 'v'. `associationMatrix` then calls `computeStatistic[['chisq']]` and `computeEffectSize[['v']]`, providing the two variables as arguments, as well as passing the 'conf.level' argument. These two functions then each return an object that `associationMatrix` extracts the information from. Inspect the source code of these functions (by typing their names without parentheses in the R prompt) to learn how this object should look, if you want to write your own functions.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
### Generate a simple association matrix using all three variables in the
### Orange tree dataframe
associationMatrix(Orange);

### Or four variables from infert:
associationMatrix(infert, c("education", "parity",
                           "induced", "case"), colNames=TRUE);

### Use variable names in the columns and generate html
associationMatrix(Orange, colNames=TRUE, type='html');
```

associationsDiamondPlot

A diamondplot with confidence intervals for associations

Description

This function produces is a diamondplot that plots the confidence intervals for associations between a number of covariates and a criterion. It currently only supports the Pearson's r effect size metric; other effect sizes are converted to Pearson's r.

Usage

```

associationsDiamondPlot(
  dat,
  covariates,
  criteria,
  labels = NULL,
  criteriaLabels = NULL,
  decreasing = NULL,
  sortBy = NULL,
  conf.level = 0.95,
  criteriaColors = viridisPalette(length(criteria)),
  criterionColor = "black",
  returnLayerOnly = FALSE,
  esMetric = "r",
  multiAlpha = 0.33,
  singleAlpha = 1,
  showLegend = TRUE,
  xlab = "Effect size estimates",
  ylab = "",
  theme = ggplot2::theme_bw(),
  lineSize = 1,
  outputFile = NULL,
  outputWidth = 10,
  outputHeight = 10,
  ggsaveParams = ufs::opts$get("ggsaveParams"),
  ...
)

associationsToDiamondPlotDf(
  dat,
  covariates,
  criterion,
  labels = NULL,
  decreasing = NULL,
  conf.level = 0.95,
  esMetric = "r"
)

```

Arguments

<code>dat</code>	The dataframe containing the relevant variables.
<code>covariates</code>	The covariates: the list of variables to associate to the criterion or criteria, usually the predictors.
<code>criteria, criterion</code>	The criteria, usually the dependent variables; one criterion (one dependent variable) can also be specified of course. The helper function <code>associationsToDiamondPlotDf</code> always accepts only one criterion.

labels	The labels for the covariates, for example the questions that were used (as a character vector).
criteriaLabels	The labels for the criteria (in the legend).
decreasing	Whether to sort the covariates by the point estimate of the effect size of their association with the criterion. Use NULL to not sort at all, TRUE to sort in descending order, and FALSE to sort in ascending order.
sortBy	When specifying multiple criteria, this can be used to indicate by which criterion the items should be sorted (if they should be sorted).
conf.level	The confidence of the confidence intervals.
criteriaColors, criterionColor	The colors to use for the different associations can be specified in <code>criteriaColors</code> . This should be a vector of valid colors with at least as many elements as criteria are specified in <code>criteria</code> . If only one criterion is specified, the color in <code>criterionColor</code> is used.
returnLayerOnly	Whether to return the entire object that is generated, or just the resulting <code>ggplot2</code> layer.
esMetric	The effect size metric to plot - currently, only 'r' is supported, and other values will return an error.
multiAlpha, singleAlpha	The transparency (alpha channel) value of the diamonds for each association can be specified in <code>multiAlpha</code> , and if only one criterion is specified, the alpha level of the diamonds can be specified in <code>singleAlpha</code> .
showLegend	Whether to show the legend.
xlab, ylab	The label to use for the x and y axes (for <code>duoComparisonDiamondPlot</code> , must be vectors of two elements). Use NULL to not use a label.
theme	The <code>ggplot()</code> theme to use.
lineSize	The thickness of the lines (the diamonds' strokes).
outputFile	A file to which to save the plot.
outputWidth, outputHeight	Width and height of saved plot (specified in centimeters by default, see <code>ggsaveParams</code>).
ggsaveParams	Parameters to pass to <code>ggsave</code> when saving the plot.
...	Any additional arguments are passed to <code>diamondPlot()</code> and eventually to <code>ggDiamondLayer()</code> .

Details

`associationsToDiamondPlotDf` is a helper function that produces the required dataframe.

This function can be used to quickly plot multiple confidence intervals.

Value

A plot.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com**See Also**[diamondPlot\(\)](#), [ggDiamondLayer\(\)](#)**Examples**

```
### Simple diamond plot with correlations
### and their confidence intervals

associationsDiamondPlot(mtcars,
                        covariates=c('cyl', 'hp', 'drat', 'wt',
                                     'am', 'gear', 'vs', 'carb', 'qsec'),
                        criteria='mpg');

### Same diamond plot, but now with two criteria,
### and colouring the diamonds based on the
### correlation point estimates: a gradient
### is created where red is used for -1,
### green for 1 and blue for 0.

associationsDiamondPlot(mtcars,
                        covariates=c('cyl', 'hp', 'drat', 'wt',
                                     'am', 'gear', 'vs', 'carb', 'qsec'),
                        criteria=c('mpg', 'disp'),
                        generateColors=c("red", "blue", "green"),
                        fullColorRange=c(-1, 1));
```

attenuate.d

*Attenuate a Cohen's d estimate for unreliability in the continuous variable***Description**

Measurement error (i.e. the complement of reliability) results in a downward bias of observed effect sizes. This attenuation can be emulated by this function.

Usage

```
attenuate.d(d, reliability)
```

Arguments

d	The value of Cohen's d (that would be obtained with perfect measurements)
reliability	The reliability of the measurements of the continuous variable

Value

The attenuated value of Cohen's d

Author(s)

Gjalt-Jorn Peters & Stefan Gruijters

References

Bobko, P., Roth, P. L., & Bobko, C. (2001). Correcting the Effect Size of d for Range Restriction and Unreliability. *Organizational Research Methods*, 4(1), 46–61. doi:[10.1177/109442810141003](https://doi.org/10.1177/109442810141003)

Examples

```
attenuate.d(.5, .8);
```

attenuate.r

Attenuate a Pearson's r estimate for unreliability in the measurements

Description

Attenuate a Pearson's r estimate for unreliability in the measurements

Usage

```
attenuate.r(r, reliability1, reliability2)
```

Arguments

r The (disattenuated) value of Pearson's r

reliability1, reliability2

The reliabilities of the two variables

Value

The attenuated value of Pearson's r

Examples

```
attenuate.r(.5, .8, .9);
```

A_VarghaDelaney	<i>Vargha & Delaney's A</i>
-----------------	---------------------------------

Description

Vargha & Delaney's A

Usage

```
A_VarghaDelaney(
  control,
  experimental,
  bootstrap = NULL,
  conf.level = 0.95,
  warn = FALSE
)
```

Arguments

control	A vector with the data for the control condition.
experimental	A vector with the data from the experimental condition.
bootstrap	The number of bootstrap samples to use to compute confidence intervals, or NULL to not compute confidence intervals.
conf.level	The confidence level of the confidence intervals.
warn	Whether to allow the <code>stats::wilcox.test()</code> function to emit warnings, for example if ties are encountered.

Value

A numeric vector of length 1 with the A value, named 'A'.

Examples

```
ufs::A_VarghaDelaney(1:8, 3:12);
```

BAC_plot	<i>Bland-Altman Change plot</i>
----------	---------------------------------

Description

Bland-Altman Change plot

Usage

```

BAC_plot(
  data,
  cols = names(data),
  reliability = NULL,
  pointSize = 2,
  deterioratedColor = "#482576E6",
  unchangedColor = "#25848E80",
  improvedColor = "#7AD151E6",
  zeroLineColor = "black",
  zeroLineType = "dashed",
  ciLineColor = "red",
  ciLineType = "solid",
  conf.level = 0.95,
  theme = ggplot2::theme_minimal(),
  ignoreBias = FALSE,
  iccFromPsych = FALSE,
  iccFromPsychArgs = NULL
)

```

Arguments

<code>data</code>	The data frame; if it only has two columns, the first of which is the pre-change column, <code>cols</code> can be left empty.
<code>cols</code>	The names of the columns with the data; the first is the column with the pre-change data, the second the column after the change.
<code>reliability</code>	The reliability estimate, for example as obtained with the <code>ICC()</code> function in the <code>psych()</code> package; can be omitted, in which case the intraclass correlation is computed.
<code>pointSize</code>	The size of the points in the plot.
<code>deterioratedColor</code> , <code>unchangedColor</code> , <code>improvedColor</code>	The colors to use for cases who deteriorate, stay the same, and improve, respectively.
<code>zeroLineColor</code> , <code>ciLineColor</code>	The colors for the line at 0 (no change) and at the confidence interval bounds (i.e. the point at which a difference becomes indicative of change given the reliability), respectively.
<code>zeroLineType</code> , <code>ciLineType</code>	The line types for the line at 0 (no change) and at the confidence interval bounds (i.e. the point at which a difference becomes indicative of change given the reliability), respectively.
<code>conf.level</code>	The confidence level of the confidence interval.
<code>theme</code>	The <code>ggplot2</code> theme to use.
<code>ignoreBias</code>	Whether to ignore bias (i.e. allow the measurements at the second time to shift upwards or downwards). If <code>FALSE</code> , the variance associated with such a shift is considered error variance (i.e. 'unreliability').

`iccFromPsych` Whether to compute ICC using the `psych::ICC()` function or not.
`iccFromPsychArgs` If using the `psych::ICC()` function, the arguments to pass.

Value

A `ggplot2` plot.

Examples

```
### Create smaller dataset for example
dat <-
  ufs::testRetestSimData[
    1:25,
    c('t0_item1', 't1_item1')
  ];

ufs::BAC_plot(dat, reliability = .5);
ufs::BAC_plot(dat, reliability = .8);
ufs::BAC_plot(dat, reliability = .9);
```

bfi-data

25 Personality items representing 5 factors

Description

This is a dataset lifted from the `psychTools` package (which was originally in the `psych` package). For details, please check that help page (using `"psychTools::bfi"`).

Usage

```
data(bfi)
```

Format

A `data.frame` with 2800 rows and 28 columns.

Examples

```
data(bfi);
```

biAxisDiamondPlot	<i>Diamondplot with two Y axes</i>
-------------------	------------------------------------

Description

This is basically a `meansDiamondPlot()`, but extended to allow specifying subquestions and anchors at the left and right side. This is convenient for psychological questionnaires when the anchors or dimensions were different from item to item. This function is used to function the left panel of the CIBER plot in the behaviorchange package.

Usage

```
biAxisDiamondPlot(
  dat,
  items = NULL,
  leftAnchors = NULL,
  rightAnchors = NULL,
  subQuestions = NULL,
  decreasing = NULL,
  conf.level = 0.95,
  showData = TRUE,
  dataAlpha = 0.1,
  dataColor = "#444444",
  diamondColors = NULL,
  jitterWidth = 0.45,
  jitterHeight = 0.45,
  xbreaks = NULL,
  xLabels = NA,
  xAxisLab = paste0("Scores and ", round(100 * conf.level, 2), "% CIs"),
  drawPlot = TRUE,
  returnPlotOnly = TRUE,
  baseSize = 1,
  dotSize = baseSize,
  baseFontSize = 10 * baseSize,
  theme = ggplot2::theme_bw(base_size = baseFontSize),
  outputFile = NULL,
  outputWidth = 10,
  outputHeight = 10,
  ggsaveParams = ufs::opts$get("ggsaveParams"),
  ...
)
```

Arguments

<code>dat</code>	The dataframe containing the variables.
<code>items</code>	The variables to include.

leftAnchors	The anchors to display on the left side of the left hand panel. If the items were measured with one variable each, this can be used to show the anchors that were used for the respective scales. Must have the same length as items.
rightAnchors	The anchors to display on the left side of the left hand panel. If the items were measured with one variable each, this can be used to show the anchors that were used for the respective scales. Must have the same length as items.
subQuestions	The subquestions used to measure each item. This can also be used to provide pretty names for the variables if the items were not measured by one question each. Must have the same length as items.
decreasing	Whether to sort the items. Specify NULL to not sort at all, TRUE to sort in descending order, and FALSE to sort in ascending order.
conf.level	The confidence levels for the confidence intervals.
showData	Whether to show the individual datapoints.
dataAlpha	The alpha level (transparency) of the individual datapoints. Value between 0 and 1, where 0 signifies complete transparency (i.e. invisibility) and 1 signifies complete 'opaqueness'.
dataColor	The color to use for the individual datapoints.
diamondColors	The colours to use for the diamonds. If NULL, the generateColors argument can be used which will then be passed to <code>diamondPlot()</code> .
jitterWidth	How much to jitter the individual datapoints horizontally.
jitterHeight	How much to jitter the individual datapoints vertically.
xbreaks	Which breaks to use on the X axis (can be useful to override <code>ggplot()</code> 's defaults).
xLabels	Which labels to use for those breaks (can be useful to override <code>ggplot()</code> 's defaults; especially useful in combination with xBreaks of course).
xAxisLab	Axis label for the X axis.
drawPlot	Whether to draw the plot, or only return it.
returnPlotOnly	Whether to return the entire object that is generated (including all intermediate objects) or only the plot.
baseSize	This can be used to efficiently change the size of most plot elements.
dotSize	This is the size of the points used to show the individual data points in the left hand plot.
baseFontSize	This can be used to set the font size separately from the baseSize.
theme	This is the theme that is used for the plots.
outputFile	A file to which to save the plot.
outputWidth, outputHeight	Width and height of saved plot (specified in centimeters by default, see <code>ggsaveParams</code>).
ggsaveParams	Parameters to pass to <code>ggsave</code> when saving the plot.
...	These arguments are passed on to <code>diamondPlot</code> .

Details

This is a diamondplot that can be used for items/questions where the anchors of the response scales could be different for every item. For the rest, it is very similar to [meansDiamondPlot\(\)](#).

Value

Either just a plot (a [gtable::gtable\(\)](#) object) or an object with all produced objects and that plot.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

CIBER() in the behaviorchange package, [associationsDiamondPlot\(\)](#)

Examples

```
biAxisDiamondPlot(dat=mtcars,
  items=c('cyl', 'wt'),
  subQuestions=c('cylinders', 'weight'),
  leftAnchors=c('few', 'light'),
  rightAnchors=c('many', 'heavy'),
  xbreaks=0:8);
```

biDimColors

Create colours for a response scale for an item

Description

Create colours for a response scale for an item

Usage

```
biDimColors(start, mid, end, length, show = TRUE)
```

```
uniDimColors(start, end, length, show = TRUE)
```

Arguments

start	Color to start with
mid	Color in the middle, for bidimensional scales
end	Color to end with
length	The number of response options
show	Whether to show the colours

Value

The colours as hex codes.

Examples

```
uniDimColors("#000000", "#00BB00", length=5, show=FALSE);
```

carelessObject	<i>Compute diagnostics for careless responding</i>
----------------	--

Description

This function is a wrapper for the functions from the `careless` package. Normally, you'd probably call `carelessReport` which calls this function to generate a report of suspect participants.

Usage

```
carelessObject(  
  data,  
  items = names(data),  
  flagUnivar = 0.99,  
  flagMultivar = 0.95,  
  irvSplit = 4,  
  responseTime = NULL  
)
```

Arguments

<code>data</code>	The dataframe.
<code>items</code>	The items to look at.
<code>flagUnivar</code>	How extreme a score has to be for it to be flagged as suspicious univariately.
<code>flagMultivar</code>	This has not been implemented yet.
<code>irvSplit</code>	Whether to split for the IRV, and if so, in how many parts.
<code>responseTime</code>	If not NULL, the name of a column containing the participants' response times.

Value

An object of class `carelessObject`.

Examples

```
carelessObject(mtcars);
```

carelessReport

A report to help diagnosing careless responders

Description

This function wraps functions from the `careless` package to help inspect and diagnose careless participants. It is optimized for using in R Markdown files.

Usage

```
carelessReport(
  data,
  items = names(data),
  nFlags = 1,
  flagUnivar = 0.99,
  flagMultivar = 0.95,
  irvSplit = 4,
  headingLevel = 3,
  datasetName = NULL,
  responseTime = NULL,
  headingSuffix = " {.tabset}",
  digits = 2,
  missingSymbol = "Missing"
)
```

Arguments

<code>data</code>	The dataframe.
<code>items</code>	The items to look at.
<code>nFlags</code>	How many indicators need to be flagged for a participant to be considered suspect.
<code>flagUnivar</code>	How extreme a score has to be for it to be flagged as suspicious univariately.
<code>flagMultivar</code>	This has not been implemented yet.
<code>irvSplit</code>	Whether to split for the IRV, and if so, in how many parts.
<code>headingLevel</code>	The level of the heading in Markdown (the number of #s to include before the heading).
<code>datasetName</code>	The name of the dataset to display (to override, if desired).
<code>responseTime</code>	If not NULL, the name of a column containing the participants' response times.
<code>headingSuffix</code>	The suffix to include; by default, set such that the individual participants IRP plots are placed in separate tabs.
<code>digits</code>	The number of digits to round to.
<code>missingSymbol</code>	How to represent missing values.

Value

NULL, invisibly; and prints the report.

Examples

```
### Get the BFI data taken from the `psych` package
dat <- ufs::bfi;

### Get the variable names for the regular items
bfiVars <-
  setdiff(names(dat),
    c("gender", "education", "age"));

### Inspect suspect participants, very conservatively to
### limit the output (these are 2800 participants).
carelessReport(data = dat,
  items = bfiVars,
  nFlags = 5);
```

cat0

Concatenate to screen without spaces

Description

The cat0 function is to cat what paste0 is to paste; it simply makes concatenating many strings without a separator easier.

Usage

```
cat0(..., sep = "")
```

Arguments

... The character vector(s) to print; passed to [cat](#).

sep The separator to pass to [cat](#), of course, "" by default.

Value

Nothing (invisible NULL, like [cat](#)).

Examples

```
cat0("The first variable is '", names(mtcars)[1], "'.");
```

checkDataIntegrity	<i>Conveniently checking data integrity</i>
--------------------	---

Description

This function is designed to make it easy to perform some data integrity checks, specifically checking for values that are impossible or unrealistic. These values can then be replaced by another value, or the offending cases can be deleted from the dataframe.

Usage

```
checkDataIntegrity(
  x,
  dat,
  newValue = NA,
  removeCases = FALSE,
  validValueSuffix = "_validValue",
  newValueSuffix = "_newValue",
  totalVarName = "numberOfInvalidValues",
  append = TRUE,
  replace = TRUE,
  silent = FALSE,
  rmarkdownOutput = FALSE,
  callingSelf = FALSE
)
```

Arguments

x	This can be either a vector or a list. If it is a vector, it should have two elements, the first one being a regular expression matching one or more variables in the dataframe specified in dat, and second one being the condition the matching variables have to satisfy. If it is a list, it should be a list of such vectors. The conditions should start with a Comparison operator followed by a value (e.g. "<30" or ">=0").
dat	The dataframe containing the variables of which we should check the integrity.
newValue	The new value to be assigned to cases not satisfying the specified conditions.
removeCases	Whether to delete cases that do not satisfy the criterion from the dataframe (if FALSE, they're not deleted, but the offending value is replaced by newValue).
validValueSuffix	Suffix to append to variable names when creating variable names for new variables that contain TRUE and FALSE to specify for each original variable whether its value satisfied the specified criterion.
newValueSuffix	If replace is FALSE, original values are not replaced, but instead new variables are created where the offending values have been replaced. This suffix is appended to each original variable name to create the new variable name.

totalVarName	This is the name of a variable that contains, for each case, the total number of invalid values among all variables checked.
append	Whether to append the columns to the dataframe, or only return the new columns.
replace	Whether to replace the offending values with the value specified in newValue or whether to create new columns (see newValueSuffix).
silent	Whether to display the log, or only set it as attribute of the returned dataframe.
rmarkdownOutput	Whether to format the log so that it's ready to be included in RMarkdown reports.
callingSelf	For internal use; whether the function calls itself.

Value

The dataframe with the corrections, and the log stored in attribute checkDataIntegrity_log.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
### Default behavior: return dataframe with
### offending values replaced by NA

checkDataIntegrity(c('mpg', '<30'),
                  mtcars);

### Check two conditions, and instead of returning the
### dataframe with the results appended, only return the
### columns indicating which cases 'pass', what the new
### values would be, and how many invalid values were
### found for each case (to easily remove cases that
### provided many invalid values)

checkDataIntegrity(list(c('mpg', '<30'),
                       c('gear', '<5')),
                  mtcars,
                  append=FALSE);
```

Description

This function efficiently checks for the presence of a package without loading it (unlike `library()` or `require()`). This is useful to force yourself to use the `package::function` syntax for addressing functions; you can make sure required packages are installed, but their namespace won't attach to the search path.

Usage

```
checkPkgs(
  ...,
  install = FALSE,
  load = FALSE,
  repos = "https://cran.rstudio.com"
)
```

Arguments

<code>...</code>	A series of packages. If the packages are named, the names are the package names, and the values are the minimum required package versions (see the second example).
<code>install</code>	Whether to install missing packages from <code>repos</code> .
<code>load</code>	Whether to load packages (which is exactly <i>not</i> the point of this package, but hey, YMMV).
<code>repos</code>	The repository to use if installing packages; default is the RStudio repository.

Value

Invisibly, a vector of the available packages.

Examples

```
ufs:::checkPkgs('base');

### Require a specific version
ufs:::checkPkgs(ufs = "0.3.1");

### This will show the error message
tryCatch(
  ufs:::checkPkgs(
    base = "99",
    stats = "42.5",
    ufs = 20
  ),
  error = print
);
```

CIM	<i>Conceptual Independence Matrix</i>
-----	---------------------------------------

Description

Conceptual Independence Matrix

Usage

```
CIM(  
  data,  
  scales,  
  conf.level = 0.95,  
  colors = c("#440154FF", "#7AD151FF"),  
  outputFile = NULL,  
  outputWidth = 100,  
  outputHeight = 100,  
  outputUnits = "cm",  
  faMethod = "minres",  
  n.iter = 100,  
  n.repeatOnWarning = 50,  
  warningTolerance = 2,  
  silentRepeatOnWarning = FALSE,  
  showWarnings = FALSE,  
  skipRegex = NULL,  
  headingLevel = 2,  
  printAbbreviations = TRUE,  
  drawPlot = TRUE,  
  returnPlotOnly = TRUE  
)  
  
CIM_partial(  
  x,  
  headingLevel = x$input$headingLevel,  
  quiet = TRUE,  
  echoPartial = FALSE,  
  partialFile = NULL,  
  ...  
)  
  
## S3 method for class 'CIM'  
knit_print(  
  x,  
  headingLevel = x$input$headingLevel,  
  quiet = TRUE,  
  echoPartial = FALSE,  
  partialFile = NULL,
```

```
    ...  
  )
```

Arguments

<code>data</code>	The dataframe containing the variables.
<code>scales</code>	The scales: a named list of character vectors, where the character vectors specify the variable names, and the names of each character vector specifies the relevant scale.
<code>conf.level</code>	The confidence level for the confidence intervals.
<code>colors</code>	The colors used for the factors. The default uses the discrete <code>viridis()</code> palette, which is optimized for perceptual uniformity, maintaining its properties when printed in grayscale, and designed for colourblind readers. A vector can also be supplied; the colors must be valid arguments to <code>colorRamp()</code> (and therefore, to <code>col2rgb()</code>).
<code>outputFile</code>	The file to write the output to.
<code>outputWidth, outputHeight, outputUnits</code>	The width, height, and units for the output file.
<code>faMethod</code>	The method to pass on to <code>psych::fa()</code> .
<code>n.iter</code>	The number of iterations to pass on to <code>psych::fa()</code> .
<code>n.repeatOnWarning</code>	How often to repeat on warnings (in the hopes of getting a run without warnings).
<code>warningTolerance</code>	How many warnings are accepted.
<code>silentRepeatOnWarning</code>	Whether to be chatty or silent when repeating after warnings.
<code>showWarnings</code>	Whether to show the warnings.
<code>skipRegex</code>	A character vector of length 2 containing two regular expressions; if the two scales both match one or both of those regular expressions, that cell is skipped.
<code>headingLevel</code>	The level for the heading; especially useful when knitting an Rmd partial.
<code>printAbbreviations</code>	Whether to print a table with the abbreviations that are used.
<code>drawPlot</code>	Whether to draw the plot or only return it.
<code>returnPlotOnly</code>	Whether to return the plot only, or the entire object.
<code>x</code>	The object to print.
<code>quiet</code>	Whether to be quiet or chatty.
<code>echoPartial</code>	Whether to echo the code in the Rmd partial.
<code>partialFile</code>	Can be used to override the Rmd partial file.
<code>...</code>	Additional arguments are passed on the respective default methods.

Value

A `ggplot2::ggplot()` plot.

Examples

```
### Load dataset `bfi`, originally from psychTools package
data(bfi, package= 'ufs');

### Specify scales
bfiScales <-
  list(Agreeableness = paste0("Agreeableness_item_", 1:5),
        Conscientiousness = paste0("Conscientiousness_item_", 1:5),
        Extraversion = paste0("Extraversion_item_", 1:5),
        Neuroticism = paste0("Neuroticism_item_", 1:5),
        Openness = paste0("Openness_item_", 1:5));

names(bfi) <- c(unlist(bfiScales),
                c('gender', 'education', 'age'));

### Only select first two and the first three items to
### keep it quick; just pass the full 'bfiScales'
### object to run for all five the full scales

CIM(bfi,
    scales=lapply(bfiScales, head, 3)[1:2],
    n.iter=10);
```

cohensdCI

The distribution of Cohen's d

Description

These functions use some conversion to and from the t distribution to provide the Cohen's d distribution. There are four versions that act similar to the standard distribution functions (the `d.`, `p.`, `q.`, and `r.` functions, and their longer aliases `.Cohensd`), three convenience functions (`pdExtreme`, `pdMild`, and `pdInterval`), a function to compute the confidence interval for a Cohen's d estimate `cohensdCI`, and a function to compute the sample size required to obtain a confidence interval around a Cohen's d estimate with a specified accuracy (`pwr.cohensdCI` and its alias `pwr.confIntd`).

Usage

```
cohensdCI(d, n, conf.level = 0.95, plot = FALSE, silent = TRUE)
```

```
dCohensd(
  x,
  df = NULL,
  populationD = 0,
  n = NULL,
  n1 = NULL,
  n2 = NULL,
```

```

    silent = FALSE
  )

pCohensd(q, df, populationD = 0, lower.tail = TRUE)

qCohensd(p, df, populationD = 0, lower.tail = TRUE)

rCohensd(n, df, populationD = 0)

pdInterval(ds, n, populationD = 0)

pdExtreme(d, n, populationD = 0)

pdMild(d, n, populationD = 0)

pwr.cohensdCI(d, w = 0.1, conf.level = 0.95, extensive = FALSE, silent = TRUE)

```

Arguments

<code>n, n1, n2</code>	Desired number of Cohen's d values for <code>rCohensd</code> and <code>rd</code> (n), and the number of participants/datapoints in total (n) or in each group ($n1$ and $n2$) for <code>dd</code> , <code>dCohensd</code> , <code>pdExtreme</code> , <code>pdMild</code> , <code>pdInterval</code> , and <code>cohensdCI</code> .
<code>conf.level</code>	The level of confidence of the confidence interval.
<code>plot</code>	Whether to show a plot of the sampling distribution of Cohen's d and the confidence interval. This can only be used if specifying one value for d , n , and <code>conf.level</code> .
<code>silent</code>	Whether to provide FALSE or suppress (TRUE) warnings. This is useful because function 'qt', which is used under the hood (see qt() for more information), warns that 'full precision may not have been achieved' when the density of the distribution is very close to zero. This is normally no cause for concern, because with sample sizes this big, small deviations have little impact.
<code>x, q, d</code>	Vector of quantiles, or, in other words, the value(s) of Cohen's d .
<code>df</code>	Degrees of freedom.
<code>populationD</code>	The value of Cohen's d in the population; this determines the center of the Cohen's d distribution. I suppose this is the noncentrality parameter.
<code>lower.tail</code>	logical; if TRUE (default), probabilities are the likelihood of finding a Cohen's d smaller than the specified value; otherwise, the likelihood of finding a Cohen's d larger than the specified value.
<code>p</code>	Vector of probabilities (p -values).
<code>ds</code>	A vector with two Cohen's d values.
<code>w</code>	The desired maximum 'half-width' or margin of error of the confidence interval.
<code>extensive</code>	Whether to only return the required sample size, or more extensive results.

Details

The functions use `convert.d.to.t()` and `convert.t.to.d()` to provide the Cohen's d distribution.

The confidence interval functions, `cohensdCI` and `pwr.cohensdCI`, now use the same method as MBESS (a slightly adapted version of the MBESS function `conf.limits.nct` is used).

More details about `cohensdCI` and `pwr.cohensdCI` are provided in Peters & Crutzen (2017).

Value

`dCohensd` (or `dd`) gives the density, `pCohensd` (or `pd`) gives the distribution function, `qCohensd` (or `qd`) gives the quantile function, and `rCohensd` (or `rd`) generates random deviates.

`pdExtreme` returns the probability (or probabilities) of finding a Cohen's d equal to or more extreme than the specified value(s).

`pdMild` returns the probability (or probabilities) of finding a Cohen's d equal to or *less* extreme than the specified value(s).

`pdInterval` returns the probability of finding a Cohen's d that lies in between the two specified values of Cohen's d .

`cohensdCI` provides the confidence interval(s) for a given Cohen's d value.

`pwr.cohensdCI` provides the sample size required to obtain a confidence interval for Cohen's d with a desired width.

Author(s)

Gjalt-Jorn Peters (Open University of the Netherlands), with the exported MBESS function `conf.limits.nct` written by Ken Kelley (University of Notre Dame), and with an error noticed by Guy Prochilo (University of Melbourne).

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Peters, G. J. Y. & Crutzen, R. (2017) Knowing exactly how effective an intervention, treatment, or manipulation is and ensuring that a study replicates: accuracy in parameter estimation as a partial solution to the replication crisis. <https://dx.doi.org/>

Maxwell, S. E., Kelley, K., & Rausch, J. R. (2008). Sample size planning for statistical power and accuracy in parameter estimation. *Annual Review of Psychology*, 59, 537-63. <https://doi.org/10.1146/annurev.psych.59.1030>

Cumming, G. (2013). The New Statistics: Why and How. *Psychological Science*, (November). <https://doi.org/10.1177/0956797613504966>

See Also

`convert.d.to.t()`, `convert.t.to.d()`, `dt()`, `pt()`, `qt()`, `rt()`

Examples

```

### Confidence interval for Cohen's d of .5
### from a sample of 200 participants, also
### showing this visually: this clearly shows
### how wildly our Cohen's d value can vary
### from sample to sample.
cohensdCI(.5, n=200, plot=TRUE);

### How many participants would we need if we
### would want a more accurate estimate, say
### with a maximum confidence interval width
### of .2?
pwr.cohensdCI(.5, w=.1);

### Show that 'sampling distribution':
cohensdCI(.5,
          n=pwr.cohensdCI(.5, w=.1),
          plot=TRUE);

### Generate 10 random Cohen's d values
rCohensd(10, 20, populationD = .5);

### Probability of findings a Cohen's d smaller than
### .5 if it's 0 in the population (i.e. under the
### null hypothesis)
pCohensd(.5, 64);

### Probability of findings a Cohen's d larger than
### .5 if it's 0 in the population (i.e. under the
### null hypothesis)
1 - pCohensd(.5, 64);

### Probability of findings a Cohen's d more extreme
### than .5 if it's 0 in the population (i.e. under
### the null hypothesis)
pdExtreme(.5, 64);

### Probability of findings a Cohen's d more extreme
### than .5 if it's 0.2 in the population.
pdExtreme(.5, 64, populationD = .2);

```

computeStatistic_t *associationMatrix Helper Functions*

Description

These objects contain a number of settings and functions for associationMatrix.

Usage

```

computeStatistic_t(var1, var2, conf.level = 0.95, var.equal = TRUE, ...)

computeStatistic_r(var1, var2, conf.level = 0.95, ...)

computeStatistic_f(var1, var2, conf.level = 0.95, ...)

computeStatistic_chisq(var1, var2, conf.level = 0.95, ...)

computeEffectSize_d(var1, var2, conf.level = 0.95, var.equal = TRUE, ...)

computeEffectSize_r(var1, var2, conf.level = 0.95, ...)

computeEffectSize_etasq(var1, var2, conf.level = 0.95, ...)

computeEffectSize_omegasq(var1, var2, conf.level = 0.95, ...)

computeEffectSize_v(
  var1,
  var2,
  conf.level = 0.95,
  bootstrap = FALSE,
  samples = 5000,
  ...
)

```

Arguments

var1	One of the two variables for which to compute a statistic or effect size
var2	The other variable for which to compute the statistic or effect size
conf.level	The confidence for the confidence interval for the effect size
var.equal	Whether to test for equal variances (test), assume equality (yes), or assume inequality (no).
...	Any additional arguments are sometimes used to specify exactly how statistics and effect sizes should be computed.
bootstrap	Whether to bootstrap to estimate the confidence interval for Cramer's V. If FALSE, the Fisher's Z conversion is used.
samples	If bootstrapping, the number of samples to generate (of course, more samples means more accuracy and longer processing time).

Value

associationMatrixStatDefaults and associationMatrixESDefaults contain the default functions from computeStatistic and computeEffectSize that are called (see the help file for associationMatrix for more details).

The other functions return an object with the relevant statistic or effect size, with a confidence interval for the effect size.

For `computeStatistic`, this object always contains:

<code>statistic</code>	The relevant statistic
<code>statistic.type</code>	The type of statistic
<code>parameter</code>	The degrees of freedom for this statistic
<code>p.raw</code>	The p-value of this statistic for NHST

And in addition, it often contains (among other things, sometimes):

<code>object</code>	The object from which the statistics are extracted
---------------------	--

For `computeEffectSize`, this object always contains:

<code>es</code>	The point estimate for the effect size
<code>esc.type</code>	The type of effect size
<code>ci</code>	The confidence interval for the effect size

And in addition, it often contains (among other things, sometimes):

<code>object</code>	The object from which the effect size is extracted
---------------------	--

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
computeStatistic_f(Orange$Tree, Orange$circumference)
computeEffectSize_etasq(Orange$Tree, Orange$circumference)
```

confintdjm

Effect Size Confidence Interval: Cohens's d

Description

Effect Size Confidence Interval: Cohens's d

Usage

```
confintdjm(d = 0.5, n = 128, conf.level = 95)
```

Arguments

<code>d</code>	.
<code>n</code>	.
<code>conf.level</code>	.

Value

A results object containing:

results\$text	a html
results\$ciPlot	an image

confIntOmegaSq	<i>Confidence intervals for Omega Squared</i>
----------------	---

Description

This function uses the MBESS functions `conf.limits.ncf()` (which has been copied into this package to avoid the dependency on MBESS) and `convert.ncf.to.omegasq()` to compute the point estimate and confidence interval for Omega Squared (which have been lifted out of MBESS to avoid importing the whole package)

Usage

```
confIntOmegaSq(var1, var2, conf.level = 0.95)

## S3 method for class 'confIntOmegaSq'
print(x, ..., digits = 2)
```

Arguments

var1, var2	The two variables: one should be a factor (or will be made a factor), the other should have at least interval level of measurement. If none of the variables is a factor, the function will look for the variable with the least unique values and change it into a factor.
conf.level	Level of confidence for the confidence interval.
x, digits, ...	Respectively the object to print, the number of digits to round to, and any additional arguments to pass on to the print function.

Value

A confIntOmegaSq object is returned, with as elements:

input	The input arguments
intermediate	Objects generated while computing the output
output	The output of the function, consisting of:
output\$es	The point estimate
output\$ci	The confidence interval

Note

Formula 16 in Steiger (2004) is used for the conversion in `convert.ncf.to.omegasq()`.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Steiger, J. H. (2004). Beyond the F test: Effect size confidence intervals and tests of close fit in the analysis of variance and contrast analysis. *Psychological Methods*, 9(2), 164-82. <https://doi.org/10.1037/1082-989X.9.2.164>

Examples

```
confIntOmegaSq(mtcars$mpg, mtcars$cyl);
```

confIntProp

Confidence intervals for proportions, vectorized over all arguments

Description

This function simply computes confidence intervals for proportions.

Usage

```
confIntProp(x, n, conf.level = 0.95, plot = FALSE)
```

Arguments

<code>x</code>	The number of 'successes', i.e. the number of events, observations, or cases that one is interested in.
<code>n</code>	The total number of cases or observations.
<code>conf.level</code>	The confidence level.
<code>plot</code>	Whether to plot the confidence interval in the binomial distribution.

Details

This function is the adapted source code of `binom.test()`. It uses `pbeta()`, with some lines of code taken from the `binom.test()` source. Specifically, the count for the low category is specified as first 'shape argument' to `pbeta()`, and the total count (either the sum of the count for the low category and the count for the high category, or the total number of cases if `compareHiToLo` is `FALSE`) minus the count for the low category as the second 'shape argument'.

Value

The confidence interval bounds in a twodimensional matrix, with the first column containing the lower bound and the second column containing the upper bound.

Author(s)

Unknown (see [binom.test\(\)](#); adapted by Gjalt-Jorn Peters)

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[binom.test\(\)](#) and [ggProportionPlot](#), the function for which this was written.

Examples

```
### Simple case
confIntProp(84, 200);

### Using vectors
confIntProp(c(2,3), c(10, 20), conf.level=c(.90, .95, .99));
```

confIntR

A function to compute a correlation's confidence interval

Description

This function computes the confidence interval for a given correlation and its sample size. This is useful to obtain confidence intervals for correlations reported in papers when informing power analyses.

Usage

```
confIntR(r, N, conf.level = 0.95, plot = FALSE)
```

Arguments

<code>r</code>	The observed correlation coefficient.
<code>N</code>	The sample size of the sample where the correlation was computed.
<code>conf.level</code>	The desired confidence level of the confidence interval.
<code>plot</code>	Whether to show a plot.

Value

The confidence interval(s) in a matrix with two columns. The left column contains the lower bound, the right column the upper bound. The [rownames\(\)](#) are the observed correlations, and the [colnames\(\)](#) are 'lo' and 'hi'. The confidence level and sample size are stored as attributes. The results are returned like this to make it easy to access single correlation coefficients from the resulting object (see the examples).

Author(s)

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References

Bonett, D. G., Wright, T. A. (2000). Sample size requirements for estimating Pearson, Kendall and Spearman correlations. *Psychometrika*, 65, 23-28.

Bonett, D. G. (2014). CIcorr.R and sizeCIcorr.R <https://people.ucsc.edu/~dgbonett/psyc181.html>

Moinester, M., & Gottfried, R. (2014). Sample size estimation for correlations with pre-specified confidence interval. *The Quantitative Methods of Psychology*, 10(2), 124-130. <https://www.tqmp.org/RegularArticles/vol10-2/p124/p124.pdf>

Peters, G. J. Y. & Crutzen, R. (forthcoming) An easy and foolproof method for establishing how effective an intervention or behavior change method is: required sample size for accurate parameter estimation in health psychology.

See Also

[confIntR\(\)](#)

Examples

```
### To request confidence intervals for one correlation
confIntR(.3, 100);

### The lower bound of a single correlation
confIntR(.3, 100)[1];

### To request confidence intervals for multiple correlations:
confIntR(c(.1, .3, .5), 250);

### The upper bound of the correlation of .5:
confIntR(c(.1, .3, .5), 250)['0.5', 'hi'];
```

confintrjmv

Effect Size Confidence Interval: Pearson's r

Description

Effect Size Confidence Interval: Pearson's r

Usage

```
confintrjmv(r = 0.3, N = 400, conf.level = 95)
```

Arguments

r .
N .
conf.level .

Value

A results object containing:

results\$text a html
results\$ciPlot an image

confIntSD	<i>Confidence interval for standard deviation</i>
-----------	---

Description

This function is vectorized.

Usage

```
confIntSD(x, n = NULL, conf.level = 0.95)
```

Arguments

x Either a standard deviation, in which case n must also be provided, or a vector, in which case n must be NULL.
n The sample size is x is a standard deviation.
conf.level The confidence level

Value

A vector or matrix.

Examples

```
ufs::confIntSD(mtcars$mpg);  
ufs::confIntSD(c(6, 7), c(32, 32));
```

convert	<i>conversion functions</i>
---------	-----------------------------

Description

These are a number of functions to convert statistics and effect size measures from/to each other.

Arguments

chisq, cohensf, cohensfsq, d, etasq, f, logodds, means, omegasq, or, p, r, t, z	The value of the relevant statistic or effect size.
ncf	The value of a noncentrality parameter of the F distribution.
n, n1, n2, N, ns	The number of observations that the r or t value is based on, or the number of observations in each of the two groups for an anova, or the total number of participants when specifying a noncentrality parameter.
df, df1, df2	The degrees of freedom for that statistic (for F, the first one is the numerator (i.e. the effect), and the second one the denominator (i.e. the error term)).
proportion	The proportion of participants in each of the two groups in a t-test or anova. This is used to compute the sample size in each group if the group sizes are unknown. Thus, if you only provide df1 and df2 when converting an F value to a Cohen's d value, equal group sizes are assumed.
b	The value of a regression coefficient.
se, sds	The standard error of standard errors of the relevant statistic (e.g. of a regression coefficient) or variables.
minDim	The smallest of the number of columns and the number of rows of the crosstable for which the chisquare is translated to a Cramer's V value.
lower.tail	For the F and chisquare distributions, whether to get the probability of the lower or upper tail.
akfEq8	When converting Cohen's <i>d</i> to <i>r</i> , for small sample sizes, bias is introduced when the commonly suggested formula is used (Aaron, Kromrey & Ferron, 1998). Therefore, by default, this function uses different equations depending on the sample size (for $n < 50$ and for $n > 50$). When akfEq8 is set to TRUE or FALSE, the corresponding action is taken; when akfEq8 is not logical (i.e. TRUE or FALSE), the function depends on the sample size.
var.equal	Whether to compute the value of <i>t</i> or Cohen's <i>d</i> assuming equal variances ('yes'), unequal variances ('no'), or whether to test for the difference ('test').

Details

Note that by default, the behavior of `convert.d.to.r` depends on the sample size (see Bruce, Kromrey & Ferron, 1998).

Value

The converted value as a numeric value.

Author(s)

Gjalt-Jorn Peters and Peter Verboon

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Aaron, B. Kromrey J. D. & Ferron, J. (1998) *Equating "r"-based and "d"-based Effect Size Indices: Problems with a Commonly Recommended Formula*. Paper presented at the Annual Meeting of the Florida Educational Research Association (43rd, Orlando, FL, November 2-4, 1998).

Examples

```
convert.t.to.r(t=-6.46, n=200);
convert.r.to.t(r=-.41, n=200);

### Compute some p-values
convert.t.to.p(4.2, 197);
convert.chisq.to.p(5.2, 3);
convert.f.to.p(8.93, 3, 644);

### Convert d to r using both equations
convert.d.to.r(d=.2, n1=5, n2=5, akfEq8 = FALSE);
convert.d.to.r(d=.2, n1=5, n2=5, akfEq8 = TRUE);
```

convert.cer.to.d	<i>Helper functions for Numbers Needed for Change</i>
------------------	---

Description

These functions are used by `nnc()` in the `behaviorchange` package to compute the Numbers Needed for Change, but are also available for manual use.

Usage

```
convert.cer.to.d(
  cer,
  eer,
  eventDesirable = TRUE,
  eventIfHigher = TRUE,
  dist = "norm",
  distArgs = NULL,
  distNS = "stats"
)

convert.d.to.eer(
  d,
```

```

    cer,
    eventDesirable = TRUE,
    eventIfHigher = TRUE,
    dist = "norm",
    distArgs = list(),
    distNS = "stats"
)

convert.d.to.nnc(d, cer, r = 1, eventDesirable = TRUE, eventIfHigher = TRUE)

convert.eer.to.d(
  eer,
  cer,
  eventDesirable = TRUE,
  eventIfHigher = TRUE,
  dist = "norm",
  distArgs = NULL,
  distNS = "stats"
)

```

Arguments

cer	The Control Event Rate.
eer	The Experimental Event Rate.
eventDesirable	Whether an event is desirable or undesirable.
eventIfHigher	Whether scores above or below the threshold are considered 'an event'.
dist, distArgs, distNS	Used to specify the distribution to use to convert between Cohen's d and the CER and EER. distArgs can be used to specify additional arguments to the corresponding q and p functions, and distNS to specify the namespace (i.e. package) from where to get the distribution functions.
d	The value of Cohen's d .
r	The correlation between the determinant and behavior (for mediated Numbers Needed for Change).

Value

The converted value.

Author(s)

Gjalt-Jorn Peters & Stefan Gruijters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Gruijters, S. L., & Peters, G. Y. (2019). Gauging the impact of behavior change interventions: A tutorial on the Numbers Needed to Treat. *PsyArXiv*. doi:10.31234/osf.io/2bau7

See Also

nnc() in the behaviorchange package.

Examples

```
convert.d.to.eer(d=.5, cer=.25);
convert.d.to.nnc(d=.5, cer=.25);
```

convert.d.to.U3	<i>Convert Cohen's d to U3</i>
-----------------	--------------------------------

Description

This function simply returns the result of `pnorm()` for Cohen's d.

Usage

```
convert.d.to.U3(d)
```

Arguments

d Cohen's d.

Value

An unnames numeric vector with the U3 values.

Examples

```
convert.d.to.U3(.5);
```

convertToNumeric	<i>Conveniently convert vectors to numeric</i>
------------------	--

Description

Tries to 'smartly' convert factor and character vectors to numeric.

Usage

```
convertToNumeric(vector, byFactorLabel = FALSE)
```

Arguments

vector	The vector to convert.
byFactorLabel	When converting factors, whether to do this by their label value (TRUE) or their level value (FALSE).

Value

The converted vector.

Examples

```
ufs::convertToNumeric(as.character(1:8));
```

cramersV	<i>Cramer's V and its confidence interval</i>
----------	---

Description

These functions compute the point estimate and confidence interval for Cramer's V.

Usage

```
cramersV(x, y = NULL, digits = 2)

## S3 method for class 'CramersV'
print(x, digits = x$input$digits, ...)

confIntV(
  x,
  y = NULL,
  conf.level = 0.95,
  samples = 500,
  digits = 2,
  method = c("bootstrap", "fisher"),
  storeBootstrappingData = FALSE
)

## S3 method for class 'confIntV'
print(x, digits = x$input$digits, ...)
```

Arguments

x	Either a crosstable to analyse, or one of two vectors to use to generate that crosstable. The vector should be a factor, i.e. a categorical variable identified as such by the 'factor' class).
y	If x is a crosstable, y can (and should) be empty. If x is a vector, y must also be a vector.

digits	Minimum number of digits after the decimal point to show in the result.
...	Any additional arguments are passed on to the print function.
conf.level	Level of confidence for the confidence interval.
samples	Number of samples to generate when bootstrapping.
method	Whether to use Fisher's Z or bootstrapping to compute the confidence interval.
storeBootstrappingData	Whether to store (or discard) the data generating during the bootstrapping procedure.

Value

A point estimate or a confidence interval for Cramer's V, an effect size to describe the association between two categorical variables.

Examples

```
### Get confidence interval for Cramer's V
### Note that by using 'table', and so removing the raw data, inhibits
### bootstrapping, which could otherwise take a while.
confIntV(table(infert$education, infert$induced));
```

dataShape	<i>normalityAssessment and samplingDistribution</i>
-----------	---

Description

normalityAssessment can be used to assess whether a variable and the sampling distribution of its mean have an approximately normal distribution.

Usage

```
dataShape(
  sampleVector,
  na.rm = TRUE,
  type = 2,
  digits = 2,
  conf.level = 0.95,
  plots = TRUE,
  xLabs = NA,
  yLabs = NA,
  qqCI = TRUE,
  labelOutliers = TRUE,
  sampleSizeOverride = NULL
)
```

```

## S3 method for class 'dataShape'
print(x, digits = x$input$digits, extraNotification = TRUE, ...)

## S3 method for class 'dataShape'
pander(x, digits = x$input$digits, extraNotification = TRUE, ...)

normalityAssessment(
  sampleVector,
  samples = 10000,
  digits = 2,
  samplingDistColor = "#2222CC",
  normalColor = "#00CC00",
  samplingDistLineSize = 2,
  normalLineSize = 1,
  xLabel.sampleDist = NULL,
  yLabel.sampleDist = NULL,
  xLabel.samplingDist = NULL,
  yLabel.samplingDist = NULL,
  sampleSizeOverride = TRUE
)

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

## S3 method for class 'normalityAssessment'
pander(x, headerPrefix = "####", suppressPlot = FALSE, ...)

samplingDistribution(
  popValues = c(0, 1),
  popFrequencies = c(50, 50),
  sampleSize = NULL,
  sampleFromPop = FALSE,
  ...
)

```

Arguments

<code>sampleVector</code>	Numeric vector containing the sample data.
<code>na.rm</code>	Whether to remove missing data first.
<code>type</code>	Type of skewness and kurtosis to compute; either 1 (g1 and g2), 2 (G1 and G2), or 3 (b1 and b2). See Joanes & Gill (1998) for more information.
<code>digits</code>	Number of digits to use when printing results.
<code>conf.level</code>	Confidence of confidence intervals.
<code>plots</code>	Whether to display plots.
<code>xBabs, yLabs</code>	The axis labels for the three plots (should be vectors of three elements; the first specifies the X or Y axis label for the rightmost plot (the histogram), the second for the middle plot (the QQ plot), and the third for the rightmost plot (the box plot).

qqCI	Whether to show the confidence interval for the QQ plot.
labelOutliers	Whether to label outliers with their row number in the box plot.
sampleSizeOverride	Whether to use the sample size of the sample as sample size for the sampling distribution, instead of the sampling distribution size. This makes sense, because otherwise, the sample size and thus sensitivity of the null hypothesis significance tests is a function of the number of samples used to generate the sampling distribution.
x	The object to print/pander.
extraNotification	Whether to be particularly informative.
...	Additional arguments are passed on, usually to the default methods.
samples	Number of samples to use when constructing sampling distribution.
samplingDistColor	Color to use when drawing the sampling distribution.
normalColor	Color to use when drawing the standard normal curve.
samplingDistLineSize	Size of the line used to draw the sampling distribution.
normallineSize	Size of the line used to draw the standard normal distribution.
xLabel.sampleDist	Label of x axis of the distribution of the sample.
yLabel.sampleDist	Label of y axis of the distribution of the sample.
xLabel.samplingDist	Label of x axis of the sampling distribution.
yLabel.samplingDist	Label of y axis of the sampling distribution.
headerPrefix	A prefix to insert before the heading (e.g. to use Markdown headings).
suppressPlot	Whether to suppress (TRUE) or print (FALSE) the plot.
popValues	The possible values (levels) of the relevant variable. For example, for a dichotomous variable, this can be "c(1:2)" (or "c(1, 2)"). Note that samplingDistribution is for manually specifying the frequency distribution (or proportions); if you have a vector with 'raw' data, just call normalityAssessment directly.
popFrequencies	The frequencies corresponding to each value in popValues; must be in the same order! See the examples.
sampleSize	Size of the sample; the sum of the frequencies if not specified.
sampleFromPop	If true, the sample vector is created by sampling from the population information specified; if false, rep() is used to generate the sample vector. Note that if proportions are supplied in popFrequencies, sampling from the population is necessary!

Details

samplingDistribution is a convenient wrapper for normalityAssessment that makes it easy to quickly generate a sample and sampling distribution from frequencies (or proportions).

dataShape computes the skewness and kurtosis.

normalityAssessment provides a number of normality tests and draws histograms of the sample data and the sampling distribution of the mean (most statistical tests assume the latter is normal, rather than the first; normality of the sample data guarantees normality of the sampling distribution of the mean, but if the sample size is sufficiently large, the sampling distribution of the mean is approximately normal even when the sample data are not normally distributed). Note that for the sampling distribution, the degrees of freedom are usually so huge that the normality tests, negligible deviations from normality will already result in very small p-values.

samplingDistribution makes it easy to quickly assess the distribution of a variables based on frequencies or proportions, and dataShape computes skewness and kurtosis.

Value

An object with several results, the most notably of which are:

plot.sampleDist	Histogram of sample distribution
sw.sampleDist	Shapiro-Wilk normality test of sample distribution
ad.sampleDist	Anderson-Darling normality test of sample distribution
ks.sampleDist	Kolmogorov-Smirnof normality test of sample distribution
kurtosis.sampleDist	Kurtosis for sample distribution
skewness.sampleDist	Skewness for sample distribution
plot.samplingDist	Histogram of sampling distribution
sw.samplingDist	Shapiro-Wilk normality test of sampling distribution
ad.samplingDist	Anderson-Darling normality test of sampling distribution
ks.samplingDist	Kolmogorov-Smirnof normality test of sampling distribution
dataShape.samplingDist	Skewness and kurtosis for sampling distribution

Examples

```
### Note: the 'not run' is simply because running takes a lot of time,
###       but these examples are all safe to run!
## Not run:

normalityAssessment(rnorm(35));
```



```

### Create a distribution of three possible values and
### show the sampling distribution for the mean
popValues <- c(1, 2, 3);
popFrequencies <- c(20, 50, 30);
sampleSize <- 100;
samplingDistribution(popValues = popValues,
                    popFrequencies = popFrequencies,
                    sampleSize = sampleSize);

### Create a very skewed distribution of ten possible values
popValues <- c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10);
popFrequencies <- c(2, 4, 8, 6, 10, 15, 12, 200, 350, 400);
samplingDistribution(popValues = popValues,
                    popFrequencies = popFrequencies,
                    sampleSize = sampleSize, digits=5);

## End(Not run)

```

descr

descr (or descriptives)

Description

This function provides a number of descriptives about your data, similar to what SPSS's DESCRIPTIVES (often called with DESCR) does.

Usage

```

descr(
  x,
  digits = 4,
  errorOnFactor = FALSE,
  include = c("central tendency", "spread", "range", "distribution shape", "sample size"),
  maxModes = 1,
  t = FALSE,
  conf.level = 0.95,
  quantileType = 2
)

```

Default S3 method:

```

descr(
  x,
  digits = 4,
  errorOnFactor = FALSE,
  include = c("central tendency", "spread", "range", "distribution shape", "sample size"),
  maxModes = 1,
  t = FALSE,

```

```

    conf.level = 0.95,
    quantileType = 2
)

## S3 method for class 'descr'
print(
  x,
  digits = attr(x, "digits"),
  t = attr(x, "transpose"),
  row.names = FALSE,
  ...
)

## S3 method for class 'descr'
pander(x, headerPrefix = "", headerStyle = "**", ...)

## S3 method for class 'descr'
as.data.frame(x, row.names = NULL, optional = FALSE, ...)

## S3 method for class 'data.frame'
descr(x, ...)

```

Arguments

<code>x</code>	The vector for which to return descriptives.
<code>digits</code>	The number of digits to round the results to when showing them.
<code>errorOnFactor</code>	Whether to show an error when the vector is a factor, or just show the frequencies instead.
<code>include</code>	Which elements to include when showing the results.
<code>maxModes</code>	Maximum number of modes to display: displays "multi" if more than this number of modes if found.
<code>t</code>	Whether to transpose the dataframes when printing them to the screen (this is easier for users relying on screen readers).
<code>conf.level</code>	Confidence of confidence interval around the mean in the central tendency measures.
<code>quantileType</code>	The type of quantiles to be used to compute the interquartile range (IQR). See quantile for more information.
<code>row.names</code>	Whether to show row names (TRUE) or not (FALSE).
<code>...</code>	Additional arguments are passed to the default <code>print</code> and <code>pander</code> methods.
<code>headerPrefix</code>	The prefix for the heading; can be used to insert hashes (#) to create Markdown headings.
<code>headerStyle</code>	A string to insert before and after the heading (to make stuff bold or italic in Markdown).
<code>optional</code>	Provided for compatibility with the default <code>as.data.frame()</code> method - see that help page for details.

Details

Note that R (of course) has many similar functions, such as `summary`, `psych::describe()` in the excellent `psych::psych` package.

The Hartigans' Dip Test may be unfamiliar to users; it is a measure of uni- vs. multidimensionality, computed by `diptest::dip.test()` from the `dip.test` package. Depending on the sample size, values over .025 can be seen as mildly indicative of multimodality, while values over .05 probably warrant closer inspection (the p-value can be obtained using `diptest::dip.test()`; also see Table 1 of Hartigan & Hartigan (1985) for an indication as to critical values).

Value

A list of dataframes with the requested values.

Author(s)

Gjalt-Jorn Peters

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References

Hartigan, J. A.; Hartigan, P. M. The Dip Test of Unimodality. *Ann. Statist.* 13 (1985), no. 1, 70–84. doi:10.1214/aos/1176346577. <https://projecteuclid.org/euclid.aos/1176346577>.

See Also

`summary`, `psych::describe()`

Examples

```
descr(mtcars$mpg);
```

diamondCoordinates	<i>Basic ggplot2 diamond plot layer construction functions</i>
--------------------	--

Description

These functions are used by `diamondPlot()` to construct a diamond plot. It's normally not necessary to call this function directly: instead, use `meansDiamondPlot()`, `meanSDtoDiamondPlot()`, and `factorLoadingDiamondCIplot()`.

Usage

```
diamondCoordinates(  
  values,  
  otherAxisValue = 1,  
  direction = "horizontal",  
  autoSize = NULL,  
  fixedSize = 0.15  
)  
  
ggDiamondLayer(  
  data,  
  ciCols = 1:3,  
  colorCol = NULL,  
  generateColors = NULL,  
  fullColorRange = NULL,  
  color = "black",  
  lineColor = NA,  
  otherAxisCol = 1:nrow(data),  
  autoSize = NULL,  
  fixedSize = 0.15,  
  direction = "horizontal",  
  ...  
)  
  
rawDataDiamondLayer(  
  dat,  
  items = NULL,  
  itemOrder = 1:length(items),  
  dataAlpha = 0.1,  
  dataColor = "#444444",  
  jitterWidth = 0.5,  
  jitterHeight = 0.4,  
  size = 3,  
  ...  
)  
  
varsToDiamondPlotDf(  
  dat,  
  items = NULL,  
  labels = NULL,  
  decreasing = NULL,  
  conf.level = 0.95  
)
```

Arguments

values	A vector of 2 or more values that are used to construct the diamond coordinates. If three values are provided, the middle one becomes the diamond's center. If
--------	--

	two, four, or more values are provided, the median becomes the diamond's center.
otherAxisValue	The value on the other axis to use to compute the coordinates; this will be the Y axis value of the points of the diamond (if direction is 'horizontal') or the X axis value (if direction is 'vertical').
direction	Whether the diamonds should be constructed horizontally or vertically.
autoSize	Whether to make the height of each diamond conditional upon its length (the width of the confidence interval).
fixedSize	If not using relative heights, fixedSize determines the height to use.
data, dat	A dataframe (or matrix) containing lower bounds, centers (e.g. means), and upper bounds of intervals (e.g. confidence intervals) for ggDiamondLayer or items and raw data for varstoDiamondPlotDf and rawDataDiamondLayer.
ciCols	The columns in the dataframe with the lower bounds, centers (e.g. means), and upper bounds (in that order).
colorCol	The column in the dataframe containing the colors for each diamond, or a vector with colors (with as many elements as the dataframe has rows).
generateColors	A vector with colors to use to generate a gradient. These colors must be valid arguments to colorRamp() (and therefore, to col2rgb()).
fullColorRange	When specifying a gradient using generateColors, it is usually desirable to specify the minimum and maximum possible value corresponding to the outer anchors of that gradient. For example, when plotting numbers from 0 to 100 using a gradient from 'red' through 'orange' to 'green', none of the means may actually be 0 or 100; the lowest mean may be, for example, 50. If no fullColorRange is specified, the diamond representing that lowest mean of 50 will be red, not orange. When specifying the fullColorRange, the lowest and highest 'colors' in generateColors are anchored to the minimum and maximum values of fullColorRange.
color	When no colors are automatically generated, all diamonds will have this color.
lineColor	If NA, lines will have the same colors as the diamonds' fill. If not NA, must be a valid color, which is then used as line color. Note that e.g. linetype and color can be used as well, which will be passed on to geom_polygon() .
otherAxisCol	A vector of values, or the index of the column in the dataframe, that specifies the values for the Y axis of the diamonds. This should normally just be a vector of consecutive integers.
...	Any additional arguments are passed to geom_polygon() . This can be used to set, for example, the alpha value of the diamonds. Additional arguments for rawDataDiamondLayer are passed on to geom_jitter() .
items	The items from the dataframe to include in the diamondplot or dataframe.
itemOrder	Order of the items to use (if not sorting).
dataAlpha	This determines the alpha (transparency) of the data points.
dataColor	The color of the data points.
jitterWidth	How much to jitter the individual datapoints horizontally.
jitterHeight	How much to jitter the individual datapoints vertically.

size	The size of the data points.
labels	The item labels to add to the dataframe.
decreasing	Whether to sort the items (rows) in the dataframe decreasing (TRUE), increasing (FALSE), or not at all (NULL).
conf.level	The confidence of the confidence intervals.

Value

ggDiamondLayer returns a `ggplot()` `geom_polygon()` object, which can then be used in `ggplot()` plots (as `diamondPlot()` does).

diamondCoordinates returns a set of four coordinates that together specify a diamond.

varsToDiamondPlotDf returns a dataframe of diamondCoordinates.

rawDataDiamondLayer returns a `geom_jitter()` object.

Author(s)

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See Also

[meansDiamondPlot\(\)](#), [meanSDtoDiamondPlot\(\)](#), [factorLoadingDiamondCIplot\(\)](#), [diamondPlot\(\)](#)

Examples

```
## Not run:
### (Don't run this example as a test, because we
### need the ggplot function which isn't part of
### this package.)

### The coordinates for a simple diamond
diamondCoordinates(values = c(1,2,3));

### Plot this diamond
ggplot() + ggDiamondLayer(data.frame(1,2,3));

## End(Not run)
```

diamondPlot

Basic diamond plot construction function

Description

This function constructs a diamond plot using `ggDiamondLayer()`. It's normally not necessary to call this function directly: instead, use [meansDiamondPlot\(\)](#) [meanSDtoDiamondPlot\(\)](#), and [factorLoadingDiamondCIplot\(\)](#).

Usage

```
diamondPlot(
  data,
  ciCols = 1:3,
  colorCol = NULL,
  otherAxisCol = NULL,
  yValues = NULL,
  yLabels = NULL,
  ylab = NULL,
  autoSize = NULL,
  fixedSize = 0.15,
  xlab = "Effect Size Estimate",
  theme = ggplot2::theme_bw(),
  color = "black",
  returnLayerOnly = FALSE,
  outputFile = NULL,
  outputWidth = 10,
  outputHeight = 10,
  ggsaveParams = ufs::opts$get("ggsaveParams"),
  ...
)
```

Arguments

<code>data</code>	A dataframe (or matrix) containing lower bounds, centers (e.g. means), and upper bounds of intervals (e.g. confidence intervals).
<code>ciCols</code>	The columns in the dataframe with the lower bounds, centers (e.g. means), and upper bounds (in that order).
<code>colorCol</code>	The column in the dataframe containing the colors for each diamond, or a vector with colors (with as many elements as the dataframe has rows).
<code>otherAxisCol</code>	The column in the dataframe containing the values that determine where on the Y axis the diamond should be placed. If this is not available in the dataframe, specify it manually using <code>yValues</code> .
<code>yValues</code>	The values that determine where on the Y axis the diamond should be placed (can also be a column in the dataframe; in that case, use <code>otherAxisCol</code>).
<code>yLabels</code>	The labels to use for for each diamond (placed on the Y axis).
<code>autoSize</code>	Whether to make the height of each diamond conditional upon its length (the width of the confidence interval).
<code>fixedSize</code>	If not using relative heights, <code>fixedSize</code> determines the height to use.
<code>xlab, ylab</code>	The labels of the X and Y axes.
<code>theme</code>	The theme to use.
<code>color</code>	Color to use if colors are specified for each diamond.
<code>returnLayerOnly</code>	Set this to TRUE to only return the <code>ggplot()</code> layer of the diamondplot, which can be useful to include it in other plots.

outputFile A file to which to save the plot.

outputWidth, outputHeight Width and height of saved plot (specified in centimeters by default, see ggsaveParams).

ggsaveParams Parameters to pass to ggsave when saving the plot.

... Additional arguments will be passed to `ggDiamondLayer()`.

Value

A `ggplot2::ggplot()` plot with a `ggDiamondLayer()` is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`meansDiamondPlot()`, `meanSDtoDiamondPlot()`, `ggDiamondLayer()`, `factorLoadingDiamondCIplot()`

Examples

```
tmpDf <- data.frame(lo = c(1, 2, 3),
                    mean = c(1.5, 3, 5),
                    hi = c(2, 4, 10),
                    color = c('green', 'red', 'blue'));

### A simple diamond plot
diamondPlot(tmpDf);

### A diamond plot using the specified colours
diamondPlot(tmpDf, colorCol = 4);

### A diamond plot using automatically generated colours
### using a gradient
diamondPlot(tmpDf, generateColors=c('green', 'red'));

### A diamond plot using automatically generated colours
### using a gradient, specifying the minimum and maximum
### possible values that can be attained
diamondPlot(tmpDf, generateColors=c('green', 'red'),
            fullColorRange=c(1, 10));
```

disattenuate.d	<i>Disattenuate a Cohen's d estimate for unreliability in the continuous variable</i>
----------------	---

Description

Measurement error (i.e. the complement of reliability) results in a downward bias of observed effect sizes. This attenuation can be reversed by disattenuation.

Usage

```
disattenuate.d(d, reliability)
```

Arguments

d	The (attenuated) value of Cohen's d (i.e. the value as observed in the sample, and therefore attenuated (decreased) by measurement error in the continuous variable).
reliability	The reliability of the measurements of the continuous variable

Value

The disattenuated value of Cohen's d

Author(s)

Gjalt-Jorn Peters & Stefan Gruijters

References

Bobko, P., Roth, P. L., & Bobko, C. (2001). Correcting the Effect Size of d for Range Restriction and Unreliability. *Organizational Research Methods*, 4(1), 46–61. doi:10.1177/109442810141003

Examples

```
disattenuate.d(.5, .8);
```

disattenuate.r	<i>Disattenuate a Pearson's r estimate for unreliability</i>
----------------	--

Description

Disattenuate a Pearson's r estimate for unreliability

Usage

```
disattenuate.r(r, reliability1, reliability2)
```

Arguments

r	The (attenuated) value of Pearson's r
reliability1, reliability2	The reliabilities of the two variables

Value

The disattenuated value of Pearson's r

Examples

```
disattenuate.r(.5, .8, .9);
```

duoComparisonDiamondPlot	<i>meansComparisonDiamondPlot and duoComparisonDiamondPlot</i>
--------------------------	--

Description

These are two diamond plot functions to conveniently make diamond plots to compare subgroups or different samples. They are both based on a univariate diamond plot where colors are used to distinguish the data points and diamonds of each subgroup or sample. The means comparison diamond plot produces only this plot, while the duo comparison diamond plot combines it with a diamond plot visualising the effect sizes of the associations. The latter currently only works for two subgroups or samples, while the simple meansComparisonDiamondPlot also works when comparing more than two sets of datapoints. These functions are explained more in detail in Peters (2017).

Usage

```

duoComparisonDiamondPlot(
  dat,
  items = NULL,
  compareBy = NULL,
  labels = NULL,
  compareByLabels = NULL,
  decreasing = NULL,
  conf.level = c(0.95, 0.95),
  showData = TRUE,
  dataAlpha = 0.1,
  dataSize = 3,
  comparisonColors = viridisPalette(length(unique(dat[, compareBy]))),
  associationsColor = "grey",
  alpha = 0.33,
  jitterWidth = 0.5,
  jitterHeight = 0.4,
  xlab = c("Scores and means", "Effect size estimates"),
  ylab = c(NULL, NULL),
  plotTitle = NULL,
  theme = ggplot2::theme_bw(),
  showLegend = TRUE,
  legend.position = "top",
  lineSize = 1,
  drawPlot = TRUE,
  xbreaks = "auto",
  outputFile = NULL,
  outputWidth = 10,
  outputHeight = 10,
  ggsaveParams = ufs::opts$get("ggsaveParams"),
  ...
)

meansComparisonDiamondPlot(
  dat,
  items = NULL,
  compareBy = NULL,
  labels = NULL,
  compareByLabels = NULL,
  decreasing = NULL,
  sortBy = NULL,
  conf.level = 0.95,
  showData = TRUE,
  dataAlpha = 0.1,
  dataSize = 3,
  comparisonColors = viridisPalette(length(unique(dat[, compareBy]))),
  alpha = 0.33,
  jitterWidth = 0.5,

```

```

    jitterHeight = 0.4,
    xlab = "Scores and means",
    ylab = NULL,
    plotTitle = NULL,
    theme = ggplot2::theme_bw(),
    showLegend = TRUE,
    legend.position = "top",
    lineSize = 1,
    xbreaks = "auto",
    outputFile = NULL,
    outputWidth = 10,
    outputHeight = 10,
    ggsaveParams = ufs::opts$get("ggsaveParams"),
    ...
  )

```

Arguments

<code>dat</code>	The dataframe containing the relevant variables.
<code>items</code>	The variables to plot (on the y axis).
<code>compareBy</code>	The variable by which to compare (i.e. the variable indicating to which subgroup or sample a row in the dataframe belongs).
<code>labels</code>	The labels to use on the y axis; these values will replace the variable names in the dataframe (specified in <code>items</code>).
<code>compareByLabels</code>	The labels to use to replace the value labels of the <code>compareBy</code> variable.
<code>decreasing</code>	Whether to sort the variables by their mean values (NULL to not sort, TRUE to sort in descending order (i.e. items with lower means are plotted more to the bottom), and FALSE to sort in ascending order (i.e. items with lower means are plotted more to the top).
<code>conf.level</code>	The confidence level of the confidence intervals specified by the diamonds for the means (for <code>meansComparisonDiamondPlot</code>) and for both the means and effect sizes (for <code>duoComparisonDiamondPlot</code>).
<code>showData</code>	Whether to plot the data points.
<code>dataAlpha</code>	The transparency (alpha channel) value for the data points: a value between 0 and 1, where 0 denotes complete transparency and 1 denotes complete opacity.
<code>dataSize</code>	The size of the data points.
<code>comparisonColors</code>	The colors to use for the different subgroups or samples. This should be a vector of valid colors with at least as many elements as sets of data points that should be plotted.
<code>associationsColor</code>	For <code>duoComparisonDiamondPlot</code> , the color to use to plot the effect sizes in the right-hand plot.
<code>alpha</code>	The alpha channel (transparency) value for the diamonds: a value between 0 and 1, where 0 denotes complete transparency and 1 denotes complete opacity.

jitterWidth, jitterHeight	How much noise to add to the data points (to prevent overplotting) in the horizontal (x axis) and vertical (y axis) directions.
xlab, ylab	The label to use for the x and y axes (for duoComparisonDiamondPlot, must be vectors of two elements). Use NULL to not use a label.
plotTitle	Optionally, for meansComparisonDiamondPlot, a title for the plot (can also be specified for duoComparisonDiamondPlot, in which case it's passed on to meansComparisonDiamondPlot for the left panel - but note that this messes up the alignment of the two panels).
theme	The theme to use for the plots.
showLegend	Whether to show the legend (which color represents which subgroup/sample).
legend.position	Where to place the legend in meansComparisonDiamondPlot (can also be specified for duoComparisonDiamondPlot, in which case it's passed on to meansComparisonDiamondPlot for the left panel - but note that this messes up the alignment of the two panels).
lineSize	The thickness of the lines (the diamonds' strokes).
drawPlot	Whether to draw the plot, or only (invisibly) return it.
xbreaks	Where the breaks (major grid lines, ticks, and labels) on the x axis should be.
outputFile	A file to which to save the plot.
outputWidth, outputHeight	Width and height of saved plot (specified in centimeters by default, see ggsaveParams).
ggsaveParams	Parameters to pass to ggsave when saving the plot.
...	Any additional arguments are passed to diamondPlot() by meansComparisonDiamondPlot and to both meansComparisonDiamondPlot and associationsDiamondPlot() by duoComparisonDiamondPlot.
sortBy	If the variables should be sorted (see decreasing), this variable specified which subgroup should be sorted by. Therefore, the value specified here must be a value label ('level label') of the compareBy variable.

Details

These functions are explained in Peters (2017).

Value

A Diamond plots: a [ggplot2::ggplot\(\)](#) plot meansComparisonDiamondPlot, and a [gtable\(\)](#) by duoComparisonDiamondPlot.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Peters, G.-J. Y. (2017). Diamond Plots: a tutorial to introduce a visualisation tool that facilitates interpretation and comparison of multiple sample estimates while respecting their inaccuracy. *PsyArXiv*. <http://doi.org/10.17605/OSF.IO/9W8YV>

See Also

[diamondPlot\(\)](#), [meansDiamondPlot\(\)](#), the CIBER() function in the behaviorchange package

Examples

```
meansComparisonDiamondPlot(mtcars,
                           items=c('disp', 'hp'),
                           compareBy='vs',
                           xbreaks=c(100,200, 300, 400));
meansComparisonDiamondPlot(chickwts,
                           items='weight',
                           compareBy='feed',
                           xbreaks=c(100,200,300,400),
                           showData=FALSE);
duoComparisonDiamondPlot(mtcars,
                         items=c('disp', 'hp'),
                         compareBy='vs',
                         xbreaks=c(100,200, 300, 400));
```

escapeRegex	<i>Escapes any characters that would have special meaning in a regular expression.</i>
-------------	--

Description

Escapes any characters that would have special meaning in a regular expression.

Usage

```
escapeRegex(string)
```

Arguments

string string being operated on.

Details

escapeRegex will escape any characters that would have special meaning in a regular expression. For any string `grep(regexEscape(string), string)` will always be true.

Value

The value of the string with any characters that would have special meaning in a regular expression escaped.

Note

Note that this function was copied literally from the Hmisc package (to prevent importing the entire package for one line of code).

Author(s)

Charles Dupont
Department of Biostatistics
Vanderbilt University

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[grep](#), [Hmisc](#), <https://hbiostat.org/R/Hmisc/>, <https://github.com/harrelfe/Hmisc>

Examples

```
string <- "this\\(system) {is} [full]."  
escapeRegex(string)
```

exceptionalScore	<i>Find exceptional scores</i>
------------------	--------------------------------

Description

This function can be used to detect exceptionally high or low scores in a vector.

Usage

```
exceptionalScore(  
  x,  
  prob = 0.025,  
  both = TRUE,  
  silent = FALSE,  
  quantileCorrection = 1e-04,  
  quantileType = 8  
)
```

Arguments

x	Vector in which to detect exceptional scores.
prob	Probability that a score is exceptionally positive or negative; i.e. scores with a quartile lower than prob or higher than 1-prob are considered exceptional (if both is TRUE, at least). So, note that a prob of .025 means that if both=TRUE, the most exceptional 5% of the values is marked as such.
both	Whether to consider values exceptional if they're below prob as well as above 1-prob, or whether to only consider values exceptional if they're below prob is prob is < .5, or above prob if prob > .5.
silent	Can be used to suppress messages.
quantileCorrection	By how much to correct the computed quantiles; this is used because when a distribution is very right-skewed, the lowest quantile is the lowest value, which is then also the mode; without subtracting a correction, almost all values would be marked as 'exceptional'.
quantileType	The algorithm used to compute the quantiles; see stats::quantile() .

Details

Note that of course, by definition, prob or $2 * \text{prob}$ percent of the values is exceptional, so it is usually not a wise idea to remove scores based on their 'exceptionalness'. Instead, use [exceptionalScores\(\)](#), which calls this function, to see how often participants answered exceptionally, and remove them based on that.

Value

A logical vector, indicating for each value in the supplied vector whether it is exceptional.

Examples

```
exceptionalScore(
  c(1,1,2,2,2,3,3,3,3,4,4,4,5,5,5,5,6,6,7,8,20),
  prob=.05
);
```

exceptionalScores	<i>Find exceptional scores</i>
-------------------	--------------------------------

Description

A function to detect participants that consistently respond exceptionally.

Usage

```
exceptionalScores(
  dat,
  items = NULL,
  exception = 0.025,
  totalOnly = TRUE,
  append = TRUE,
  both = TRUE,
  silent = FALSE,
  suffix = "_isExceptional",
  totalVarName = "exceptionalScores"
)
```

Arguments

<code>dat</code>	The dataframe containing the variables to inspect, or the vector to inspect (but for vectors, <code>exceptionalScore()</code> might be more useful).
<code>items</code>	The names of the variables to inspect.
<code>exception</code>	When an item will be considered exceptional, passed on as prob to <code>exceptionalScore()</code> .
<code>totalOnly</code>	Whether to return only the number of exceptional scores for each row in the dataframe, or for each inspected item, which values are exceptional.
<code>append</code>	Whether to return the supplied dataframe with the new variable(s) appended (if TRUE), or whether to only return the new variable(s) (if FALSE).
<code>both</code>	Whether to look for both low and high exceptional scores (TRUE) or not (FALSE; see <code>exceptionalScore()</code>).
<code>silent</code>	Can be used to suppress messages.
<code>suffix</code>	If not returning the total number of exceptional values, for each inspected variable, a new variable is returned indicating which values are exceptional. The text string is appended to each original variable name to create the new variable names.
<code>totalVarName</code>	If returning only the total number of exceptional values, and appending these to the provided dataset, this text string is used as variable name.

Value

Either a vector containing the number of exceptional values, a dataset containing, for each inspected variable, which values are exceptional, or the provided dataset where either the total or the exceptional values for each variable are appended.

Examples

```
exceptionalScores(mtcars);
```

exportToHTML	<i>Exporting tables to HTML</i>
--------------	---------------------------------

Description

This function exports data frames or matrices to HTML, sending output to one or more of the console, viewer, and one or more files.

Usage

```
exportToHTML(
  input,
  output = ufs::opts$get("tableOutput"),
  tableOutputCSS = ufs::opts$get("tableOutputCSS")
)
```

Arguments

input	Either a <code>data.frame</code> , <code>table</code> , or <code>matrix</code> , or a list with three elements: <code>pre</code> , <code>input</code> , and <code>post</code> . The <code>pre</code> and <code>post</code> are simply prepended and postpended to the HTML generated based on the <code>input\$input</code> element.
output	The output: a character vector with one or more of "console" (the raw concatenated input, without conversion to HTML), "viewer", which uses the RStudio viewer if available, and one or more filenames in existing directories.
tableOutputCSS	The CSS to use for the HTML table.

Value

Invisibly, the (potentially concatenated) input as character vector.

Examples

```
exportToHTML(mtcars[1:5, 1:5]);
```

extractVarName	<i>Extract variable names</i>
----------------	-------------------------------

Description

Functions often get passed variables from within dataframes or other lists. However, printing these names with all their dollar signs isn't very userfriendly. This function simply uses a regular expression to extract the actual name.

Usage

```
extractVarName(x)
```

Arguments

x A character vector of one or more variable names.

Value

The actual variables name, with all containing objectes stripped off.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
extractVarName('mtcars$mpg');
```

faConfInt

Extract confidence bounds from psych's factor analysis object

Description

This function contains some code from a function in [psych::psych-package](#) that's not exported `print.psych.fa.ci` but useful nonetheless. It basically takes the outcomes of a factor analysis and extracted the confidence intervals.

Usage

```
faConfInt(fa)
```

Arguments

fa The object produced by the [psych::fa\(\)](#) function from the [psych::psych-package](#) package. It is important that the `n.iter` argument of [psych::fa\(\)](#) was set to a realistic number, because otherwise, no confidence intervals will be available.

Details

This function extract confidence interval bounds and combines them with factor loadings using the code from the `print.psych.fa.ci` in [psych::psych-package](#).

Value

A list of dataframes, one for each extracted factor, with in each dataframe three variables:

lo	lower bound of the confidence interval
est	point estimate of the factor loading
hi	upper bound of the confidence interval

Author(s)

William Revelle (extracted by Gjalt-Jorn Peters)

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
## Not run:
### Not run because it takes too long to run to test it,
### and may produce warnings, both because of the bootstrapping
### required to generate the confidence intervals in fa
faConfInt(psych::fa(Thurstone.33, 2, n.iter=100, n.obs=100));

## End(Not run)
```

factorLoadingDiamondCIplot

Two-dimensional visualisation of factor analyses

Description

This function uses the `diamondPlot()` to visualise the results of a factor analyses. Because the factor loadings computed in factor analysis are point estimates, they may vary from sample to sample. The factor loadings for any given sample are usually not relevant; samples are but means to study populations, and so, researchers are usually interested in population values for the factor loadings. However, tables with lots of loadings can quickly become confusing and intimidating. This function aims to facilitate working with and interpreting factor analysis based on confidence intervals by visualising the factor loadings and their confidence intervals.

Usage

```
factorLoadingDiamondCIplot(
  fa,
  xlab = "Factor Loading",
  colors = viridisPalette(max(2, fa$factors)),
  labels = NULL,
  theme = ggplot2::theme_bw(),
  sortAlphabetically = FALSE,
  ...
)
```

Arguments

fa	The object produced by the <code>psych::fa()</code> function from the <code>psych::psych</code> package. It is important that the <code>n.iter</code> argument of <code>psych::fa()</code> was set to a realistic number, because otherwise, no confidence intervals will be available.
xlab	The label for the x axis.

colors	The colors used for the factors. The default uses the discrete viridis palette, which is optimized for perceptual uniformity, maintaining its properties when printed in grayscale, and designed for colourblind readers.
labels	The labels to use for the items (on the Y axis).
theme	The ggplot2 theme to use.
sortAlphabetically	Whether to sort the items alphabetically.
...	Additional arguments will be passed to <code>ggDiamondLayer()</code> . This can be used to set, for example, the transparency (alpha value) of the diamonds to a lower value using e.g. <code>alpha=.5</code> .

Value

A `ggplot2::ggplot()` plot with several `ggDiamondLayer()`s is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`psych::fa()`, `meansDiamondPlot()`, `meansSDtoDiamondPlot()`, `diamondPlot()`, `ggDiamondLayer()`

Examples

```
## Not run:
### (Not run during testing because it takes too long and
### may generate warnings because of the bootstrapping of
### the confidence intervals)

factorLoadingDiamondCIplot(psych::fa(psych::Bechtoldt,
                                     nfactors=2,
                                     n.iter=50,
                                     n.obs=200));

### And using a lower alpha value for the diamonds to
### make them more transparent

factorLoadingDiamondCIplot(psych::fa(psych::Bechtoldt,
                                     nfactors=2,
                                     n.iter=50,
                                     n.obs=200),
                           alpha=.5,
                           size=1);

## End(Not run)
```

factorLoadingHeatmap *Two-dimensional visualisation of factor analyses*

Description

This function uses the `diamondPlot()` to visualise the results of a factor analyses. Because the factor loadings computed in factor analysis are point estimates, they may vary from sample to sample. The factor loadings for any given sample are usually not relevant; samples are but means to study populations, and so, researchers are usually interested in population values for the factor loadings. However, tables with lots of loadings can quickly become confusing and intimidating. This function aims to facilitate working with and interpreting factor analysis based on confidence intervals by visualising the factor loadings and their confidence intervals.

Usage

```
factorLoadingHeatmap(
  fa,
  xlab = "Factor Loading",
  colors = viridisPalette(max(2, fa$factors)),
  labels = NULL,
  showLoadings = FALSE,
  heatmap = FALSE,
  theme = ggplot2::theme_minimal(),
  sortAlphabetically = FALSE,
  digits = 2,
  labs = list(title = NULL, x = NULL, y = NULL),
  themeArgs = list(panel.grid = ggplot2::element_blank(), legend.position = "none",
    axis.text.x = ggplot2::element_blank()),
  ...
)
```

Arguments

<code>fa</code>	The object produced by the <code>psych::fa()</code> function from the <code>psych::psych</code> package. It is important that the <code>n.iter</code> argument of <code>psych::fa()</code> was set to a realistic number, because otherwise, no confidence intervals will be available.
<code>xlab</code>	The label for the x axis.
<code>colors</code>	The colors used for the factors. The default uses the discrete viridis palette, which is optimized for perceptual uniformity, maintaining its properties when printed in grayscale, and designed for colourblind readers.
<code>labels</code>	The labels to use for the items (on the Y axis).
<code>showLoadings</code>	Whether to show the factor loadings or not.
<code>heatmap</code>	Whether to produce a heatmap or use diamond plots.
<code>theme</code>	The ggplot2 theme to use.

sortAlphabetically	Whether to sort the items alphabetically.
digits	Number of digits to round to.
labs	The labels to pass to ggplot2.
themeArgs	Additional theme arguments to pass to ggplot2.
...	Additional arguments will be passed to <code>ggDiamondLayer()</code> . This can be used to set, for example, the transparency (alpha value) of the diamonds to a lower value using e.g. <code>alpha=.5</code> .

Value

A `ggplot2::ggplot()` plot with several `ggDiamondLayer()`s is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`psych::fa()`, `meansDiamondPlot()`, `meanSDtoDiamondPlot()`, `diamondPlot()`, `ggDiamondLayer()`

Examples

```
## Not run:
### (Not run during testing because it takes too long and
### may generate warnings because of the bootstrapping of
### the confidence intervals)

factorLoadingHeatmap(psych::fa(psych::Bechtoldt,
                               nfactors=2,
                               n.iter=50,
                               n.obs=200));

### And using a lower alpha value for the diamonds to
### make them more transparent

factorLoadingHeatmap(psych::fa(psych::Bechtoldt,
                               nfactors=2,
                               n.iter=50,
                               n.obs=200),
                     alpha=.5,
                     size=1);

## End(Not run)
```

fa_failsafe

Do factor-analysis, logging warnings and errors

Description

Do factor-analysis, logging warnings and errors

Usage

```
fa_failsafe(
  ...,
  n.repeatOnWarning = 50,
  warningTolerance = 2,
  silentRepeatOnWarning = FALSE,
  showWarnings = TRUE
)
```

Arguments

... The arguments for fa in psych.

n.repeatOnWarning How often to repeat on warnings (in the hopes of getting a run without warnings).

warningTolerance How many warnings are accepted.

silentRepeatOnWarning Whether to be chatty or silent when repeating after warnings.

showWarnings Whether to show the warnings.

Value

A list with the fa object and a warnings and an errors object.

findShortestInterval

Find the shortest interval

Description

This function takes a numeric vector, sorts it, and then finds the shortest interval and returns its length.

Usage

```
findShortestInterval(x)
```


Arguments

x The numeric vector.

Value

The length of the shortest interval.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
findShortestInterval(c(1, 2, 4, 7, 20, 10, 15));
```

formatCI

Pretty formatting of confidence intervals

Description

Pretty formatting of confidence intervals

Usage

```
formatCI(
  ci,
  sep = "; ",
  prefix = "[",
  suffix = "]",
  digits = 2,
  noZero = FALSE
)
```

Arguments

ci	A confidence interval (a vector of 2 elements; longer vectors work, but I guess that wouldn't make sense).
sep	The separator of the values, usually "; " or ", ".
prefix, suffix	The prefix and suffix, usually a type of opening and closing parenthesis/bracket.
digits	The number of digits to which to round the values.
noZero	Whether to strip the leading zero (before the decimal point), as is typically done when following APA style and displaying correlations, <i>p</i> values, and other numbers that cannot reach 1 or more.

Value

A character vector of one element.

See Also

```
noZero(), formatR(), formatPvalue()
```

Examples

```
### With leading zero ...
formatCI(c(0.55, 0.021));

### ... and without
formatCI(c(0.55, 0.021), noZero=TRUE);
```

formatPvalue	<i>Pretty formatting of p values</i>
--------------	--------------------------------------

Description

Pretty formatting of *p* values

Usage

```
formatPvalue(values, digits = 3, spaces = TRUE, includeP = TRUE)
```

Arguments

values	The p-values to format.
digits	The number of digits to round to. Numbers smaller than this number will be shown as <.001 or <.0001 etc.
spaces	Whether to include spaces between symbols, operators, and digits.
includeP	Whether to include the 'p' and '='-symbol in the results (the '<' symbol is always included).

Value

A formatted P value, roughly according to APA style guidelines. This means that the [noZero](#) function is used to remove the zero preceding the decimal point, and p values that would round to zero given the requested number of digits are shown as e.g. p<.001.

See Also

```
formatCI(), formatR(), noZero()
```

Examples

```
formatPvalue(cor.test(mtcars$mpg,
                      mtcars$disp)$p.value);
formatPvalue(cor.test(mtcars$drat,
                      mtcars$qsec)$p.value);
```

formatR

Pretty formatting of correlation coefficients

Description

Pretty formatting of correlation coefficients

Usage

```
formatR(r, digits = 2)
```

Arguments

r	The Pearson correlation to format.
digits	The number of digits to round to.

Value

The formatted correlation.

See Also

[noZero\(\)](#), [formatCI\(\)](#), [formatPvalue\(\)](#)

Examples

```
formatR(cor(mtcars$mpg, mtcars$disp));
```

 getData

 Use a dialog to load data from an SPSS file

Description

getData() and getDat() provide an easy way to load SPSS datafiles.

Usage

```
getData(
  filename = NULL,
  file = NULL,
  errorMessage = "[defaultErrorMessage]",
  applyRioLabels = TRUE,
  use.value.labels = FALSE,
  to.data.frame = TRUE,
  stringsAsFactors = FALSE,
  silent = FALSE,
  ...
)

getDat(..., dfName = "dat", backup = TRUE)
```

Arguments

filename, file	It is possible to specify a path and filename to load here. If not specified, the default R file selection dialogue is shown. file is still available for backward compatibility but will eventually be phased out.
errorMessage	The error message that is shown if the file does not exist or does not have the right extension; [defaultErrorMessage] is replaced with a default error message (and can be included in longer messages).
applyRioLabels	Whether to apply the labels supplied by Rio. This will make variables that has value labels into factors.
use.value.labels	Only useful when reading from SPSS files: whether to read variables with value labels as factors (TRUE) or numeric vectors (FALSE).
to.data.frame	Only useful when reading from SPSS files: whether to return a dataframe or not.
stringsAsFactors	Whether to read strings as strings (FALSE) or factors (TRUE).
silent	Whether to suppress potentially useful information.
...	Additional options, passed on to the function used to import the data (which depends on the extension of the file).
dfName	The name of the dataframe to create in the parent environment.
backup	Whether to backup an object with name dfName, if one already exists in the parent environment.

Value

getData returns the imported dataframe, with the filename from which it was read stored in the 'filename' attribute.

getDat is a simple wrapper for getData() which creates a dataframe in the parent environment, by default with the name 'dat'. Therefore, calling getDat() in the console will allow the user to select a file, and the data from the file will then be read and be available as 'dat'. If an object with dfName (i.e. 'dat' by default) already exists, it will be backed up with a warning. getDat() also invisibly returns the data.frame.

Note

getData() currently can't read from LibreOffice or OpenOffice files. There doesn't seem to be a platform-independent package that allows this. Non-CRAN package ROpenOffice from Omega-Hat should be able to do the trick, but fails to install (manual download and installation using <https://www.omegahat.org> produces "ERROR: dependency 'Rcompression' is not available for package 'ROpenOffice'" - and manual download and installation of RCompression produces "Please define LIB_ZLIB; ERROR: configuration failed for package 'Rcompression'"). If you have any suggestions, please let me know!

Examples

```
## Not run:
### Open a dialogue to read an SPSS file
getData();

## End(Not run)
```

ggBarChart

Bar chart using ggplot

Description

This function provides a simple interface to create a `ggplot2::ggplot()` bar chart.

Usage

```
ggBarChart(vector, plotTheme = ggplot2::theme_bw(), ...)
```

Arguments

vector	The vector to display in the bar chart.
plotTheme	The theme to apply.
...	And additional arguments are passed to <code>ggplot2::geom_bar()</code> .

Value

A `ggplot2::ggplot()` plot is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`ggplot2::geom_bar()`

Examples

```
ggBarChart(mtcars$cyl);
```

ggBoxplot

Box plot using ggplot

Description

This function provides a simple interface to create a `ggplot` box plot, organising different boxplots by levels of a factor is desired, and showing row numbers of outliers.

Usage

```
ggBoxplot(
  dat,
  y = NULL,
  x = NULL,
  labelOutliers = TRUE,
  outlierColor = "red",
  theme = ggplot2::theme_bw(),
  ...
)
```

Arguments

<code>dat</code>	Either a vector of values (to display in the box plot) or a dataframe containing variables to display in the box plot.
<code>y</code>	If <code>dat</code> is a dataframe, this is the name of the variable to make the box plot of.
<code>x</code>	If <code>dat</code> is a dataframe, this is the name of the variable (normally a factor) to place on the X axis. Separate box plots will be generate for each level of this variable.
<code>labelOutliers</code>	Whether or not to label outliers.
<code>outlierColor</code>	If labeling outliers, this is the color to use.
<code>theme</code>	The theme to use for the box plot.
<code>...</code>	Any additional arguments will be passed to <code>geom_boxplot</code> .

Details

This function is based on JasonAizkalns' answer to a question on Stack Exchange (Cross Validated; see <https://stackoverflow.com/questions/33524669/labeling-outliers-of-boxplots-in-r>).

Value

A `ggplot` plot is returned.

Author(s)

Jason Aizkalns; implemented in this package (and tweaked a bit) by Gjalt-Jorn Peters.

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`geom_boxplot`

Examples

```
### A box plot for miles per gallon in the mtcars dataset:
ggBoxplot(mtcars$mpg);

### And separate for each level of 'cyl' (number of cylinder):
ggBoxplot(mtcars, y='mpg', x='cyl');
```

ggEasyBar

Convenience functions for ggplots based on multiple variables

Description

These are convenience functions to quickly generate plots for multiple variables, with the variables in the y axis.

Usage

```
ggEasyBar(
  data,
  items = NULL,
  labels = NULL,
  sortByMean = TRUE,
  xlab = NULL,
  ylab = NULL,
  scale_fill_function = NULL,
  fontColor = "white",
  fontSize = 2,
  labelMinPercentage = 1,
```

```

    showInLegend = "both",
    legendRows = 2,
    legendValueLabels = NULL,
    biAxisLabels = NULL
  )

  ggEasyRidge(
    data,
    items = NULL,
    labels = NULL,
    sortByMean = TRUE,
    xlab = NULL,
    ylab = NULL
  )

```

Arguments

<code>data</code>	The dataframe containing the variables.
<code>items</code>	The variable names (if not provided, all variables will be used).
<code>labels</code>	Labels can optionally be provided; if they are, these will be used instead of the variable names.
<code>sortByMean</code>	Whether to sort the variables by mean value.
<code>xlab, ylab</code>	The labels for the x and y axes.
<code>scale_fill_function</code>	The function to pass to <code>ggplot()</code> to provide the colors of the bars. If NULL, set to <code>ggplot2::scale_fill_viridis_d(labels = legendValueLabels, guide = ggplot2::guide_legend(title = NULL, nrow=legendRows, byrow=TRUE))</code> .
<code>fontColor, fontSize</code>	The color and size of the font used to display the labels
<code>labelMinPercentage</code>	The minimum percentage that a category must reach before the label is printed (in whole percentages, i.e., on a scale from 0 to 100).
<code>showInLegend</code>	What to show in the legend in addition to the values; nothing ("none"), the frequencies ("freq"), the percentages ("perc"), or both ("both"). This is only used if only one variable is shown in the plot; otherwise, after all, the absolute frequencies and percentages differ for each variable.
<code>legendRows</code>	Number or rows in the legend.
<code>legendValueLabels</code>	Labels to use in the legend; must be a vector of the same length as the number of categories in the variables.
<code>biAxisLabels</code>	This can be used to specify labels to use if you want to use labels on both the left and right side. This is mostly useful when plotting single questions or semantic differentials. This must be a list with two character vectors, <code>leftAnchors</code> and <code>rightAnchors</code> , which must each have the same length as the number of items specified in <code>items</code> . See the examples for, well, examples.

Value

A `ggplot()` plot is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`geom_ridgeline()`, `geom_bar()`

Examples

```
ggEasyBar(mtcars, c('gear', 'carb'));
ggEasyRidge(mtcars, c('disp', 'hp'));

### When plotting single questions, if you want to show the anchors:
ggEasyBar(mtcars, c('gear'),
          biAxisLabels=list(leftAnchors="Fewer",
                           rightAnchors="More"));

### Or for multiple questions (for e.g. semantic differentials):
ggEasyBar(mtcars, c('gear', 'carb'),
          biAxisLabels=list(leftAnchors=c("Fewer", "Lesser"),
                           rightAnchors=c("More", "Greater")));
```

ggPie

A ggplot pie chart

Description

This function creates a pie chart. Note that these are generally quite strongly advised against, as people are not good at interpreting relative frequencies on the basis of pie charts.

Usage

```
ggPie(vector, scale_fill = ggplot2::scale_fill_viridis_d())
```

Arguments

vector	The vector (best to pass a factor).
scale_fill	The ggplot scale fill function to use for the colors.

Value

A ggplot pie chart.

Note

This function is very strongly based on the Mathematical Coffee post at <http://mathematicalcoffee.blogspot.com/2014/06/ggpie-graphs-in-ggplot2.html>.

Examples

```
ggPie(mtcars$cyl);
```

ggProportionPlot	<i>Sample distribution based plotting of proportions</i>
------------------	--

Description

This function visualises percentages, but avoids a clear cut for the sample point estimate, instead using the confidence (as in confidence interval) to create a gradient. This effectively hinders drawing conclusions on the basis of point estimates, thereby urging a level of caution that is consistent with what the data allows.

Usage

```
ggProportionPlot(
  dat,
  items = NULL,
  loCategory = NULL,
  hiCategory = NULL,
  subQuestions = NULL,
  leftAnchors = NULL,
  rightAnchors = NULL,
  compareHiToLo = TRUE,
  showDiamonds = FALSE,
  diamonds.conf.level = 0.95,
  diamonds.alpha = 1,
  na.rm = TRUE,
  barHeight = 0.4,
  conf.steps = seq(from = 0.001, to = 0.999, by = 0.001),
  scale_color = c("#21908CFF", "#FDE725FF"),
  scale_fill = c("#21908CFF", "#FDE725FF"),
  rank.conf = FALSE,
  linetype = 1,
  theme = ggplot2::theme_bw(),
  returnPlotOnly = TRUE
)

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

```
## S3 method for class 'ggProportionPlot'
grid.draw(x, ...)
```

Arguments

<code>dat</code>	The dataframe containing the items (variables), or a vector.
<code>items</code>	The names of the items (variables). If none are specified, all variables in the dataframe are used.
<code>loCategory</code>	The value of the low category (usually 0). If not provided, the minimum value is used.
<code>hiCategory</code>	The value of the high category (usually 1). If not provided, the maximum value is used.
<code>subQuestions</code>	The labels to use for the variables (for example, different questions). The variable names are used if these aren't provided.
<code>leftAnchors</code>	The labels for the low categories. The values are used if these aren't provided.
<code>rightAnchors</code>	The labels for the high categories. The values are used if these aren't provided.
<code>compareHiToLo</code>	Whether to compare the percentage of low category values to the total of the low category values and the high category values, or whether to ignore the high category values and compute the percentage of low category values relative to all cases. This can be useful when a variable has more than two values, and you only want to know/plot the percentage relative to the total number of cases.
<code>showDiamonds</code>	Whether to add diamonds to illustrate the confidence intervals.
<code>diamonds.conf.level</code>	The confidence level of the diamonds' confidence intervals.
<code>diamonds.alpha</code>	The alpha channel (i.e. transparency, or rather 'obliqueness') of the diamonds.
<code>na.rm</code>	Whether to remove missing values.
<code>barHeight</code>	The height of the bars, or rather, half the height. Use .5 to completely fill the space.
<code>conf.steps</code>	The number of steps to use to generate the confidence levels for the proportion.
<code>scale_color, scale_fill</code>	A vector with two values (valid colors), that are used for the colors (stroke) and fill for the gradient; both vectors should normally be the same, but if you feel adventurous, you can play around with the number of <code>conf.steps</code> and this. If you specify only one color, no gradient is used but a single color (i.e. specifying the same single color for both <code>scale_color</code> and <code>scale_fill</code> simply draws bars of that color).
<code>rank.conf</code>	Whether to let the fill and color gradients use the confidence or the ranked confidence.
<code>linetype</code>	The <code>linetype()</code> to use (0 = blank, 1 = solid, 2 = dashed, 3 = dotted, 4 = dotdash, 5 = longdash, 6 = twodash).
<code>theme</code>	The theme to use.
<code>returnPlotOnly</code>	Whether to only return the <code>ggplot2()</code> plot or the full object including intermediate values and objects.
<code>x</code>	The object to print/plot.
<code>...</code>	Any additional arguments are passed on to <code>print</code> and <code>grid.draw</code> .

Details

This function used `confIntProp()` to compute confidence intervals for proportions at different levels of confidence. The confidence interval bounds at those levels of confidence are then used to draw rectangles with colors in a gradient that corresponds to the confidence level.

Note that percentually, the gradient may not look continuous because at the borders between lighter and darker rectangles, the shade of the lighter rectangle is perceived as even lighter than it is, and the shade of the darker rectangle is perceived as even darker. This makes it seem as if each rectangle is coloured with a gradient in the opposite direction.

Value

A `ggplot2()` object (if `returnPlotOnly` is TRUE), or an object containing that `ggplot2()` object and intermediate products.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`confIntProp()` and `binom.test()`

Examples

```
### V/S (no idea what this is: ?mtcars only mentions 'V/S' :-))
### and transmission (automatic vs manual)
ggProportionPlot(mtcars, items=c('vs', 'am'));

### Number of cylinders, by default comparing lowest value
### (4) to highest (8):
ggProportionPlot(mtcars, items=c('cyl'));

## Not run:
### Not running these to save time during package building/checking

### We can also compare 4 to 6:
ggProportionPlot(mtcars, items=c('cyl'),
                  hiCategory=6);

### Now compared to total records, instead of to
### highest value (hiCategory is ignored then)
ggProportionPlot(mtcars, items=c('cyl'),
                  compareHiToLo=FALSE);

### And for 6 cylinders:
ggProportionPlot(mtcars, items=c('cyl'),
                  loCategory=6, compareHiToLo=FALSE);

### And for 8 cylinders:
```

```

ggProportionPlot(mtcars, items=c('cyl'),
                  loCategory=8, compareHiToLo=FALSE);

### And for 8 cylinders with different labels
ggProportionPlot(mtcars, items=c('cyl'),
                  loCategory=8,
                  subQuestions='Cylinders',
                  leftAnchors="Eight",
                  rightAnchors="Four\nor\nsix",
                  compareHiToLo=FALSE);

### ... And showing the diamonds for the confidence intervals
ggProportionPlot(mtcars, items=c('cyl'),
                  loCategory=8,
                  subQuestions='Cylinders',
                  leftAnchors="Eight",
                  rightAnchors="Four\nor\nsix",
                  compareHiToLo=FALSE,
                  showDiamonds=TRUE);

## End(Not run)

### Using less steps for the confidence levels and changing
### the fill colours
ggProportionPlot(mtcars,
                  items=c('vs', 'am'),
                  showDiamonds = TRUE,
                  scale_fill = c("#B63679FF", "#FCFDBFFF"),
                  conf.steps=seq(from=0.0001, to=.9999, by=.2));

```

ggqq

Easy ggplot Q-Q plot

Description

This function creates a qq-plot with a confidence interval.

Usage

```

ggqq(
  x,
  distribution = "norm",
  ...,
  ci = TRUE,
  line.estimate = NULL,
  conf.level = 0.95,
  sampleSizeOverride = NULL,
  observedOnX = TRUE,

```

```

scaleExpected = TRUE,
theoryLab = "Theoretical quantiles",
observeLab = "Observed quantiles",
theme = ggplot2::theme_bw()
)

```

Arguments

<code>x</code>	A vector containing the values to plot.
<code>distribution</code>	The distribution to (a 'd' and 'q' are prepended, and the resulting functions are used, e.g. <code>dnorm</code> and <code>qnorm</code> for the normal curve).
<code>...</code>	Any additional arguments are passed to the quantile function (e.g. <code>qnorm</code>). Because of these dots, any following arguments must be named explicitly.
<code>ci</code>	Whether to show the confidence interval.
<code>line.estimate</code>	Whether to show the line showing the match with the specified distribution (e.g. the normal distribution).
<code>conf.level</code>	The confidence of the confidence level around the estimate for the specified distribution.
<code>sampleSizeOverride</code>	It can be desirable to get the confidence intervals for a different sample size (when the sample size is very large, for example, such as when this plot is generated by the function <code>normalityAssessment</code>). That different sample size can be specified here.
<code>observedOnX</code>	Whether to plot the observed values (if TRUE) or the theoretically expected values (if FALSE) on the X axis. The other is plotted on the Y axis.
<code>scaleExpected</code>	Whether the scale the expected values to match the scale of the variable. This option is provided to be able to mimic SPSS' Q-Q plots.
<code>theoryLab</code>	The label for the theoretically expected values (on the Y axis by default).
<code>observeLab</code>	The label for the observed values (on the Y axis by default).
<code>theme</code>	The theme to use.

Details

This is strongly based on the answer by user Floo0 to a Stack Overflow question at Stack Exchange (see <https://stackoverflow.com/questions/4357031/qqnorm-and-qqline-in-ggplot2/27191036#27191036>), also posted at GitHub (see <https://gist.github.com/rentrop/d39a8406ad8af2a1066c>). That code is in turn based on the `qqPlot()` function from the `car` package.

Value

A `ggplot` plot is returned.

Author(s)

John Fox and Floo0; implemented in this package (and tweaked a bit) by Gjalt-Jorn Peters.

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
ggqq(mtcars$mpg);
```

ggSave

Save a ggplot with specific defaults

Description

This function is vectorized over all argument except 'plot': so if you want to save multiple versions, simply provide vectors. Vectors of length 1 will be recycled using [rep\(\)](#); otherwise vectors have to all be the same length as file.

Usage

```
ggSave(
  file = NULL,
  plot = ggplot2::last_plot(),
  width = ufs::opts$get("ggSaveFigWidth"),
  height = ufs::opts$get("ggSaveFigHeight"),
  units = ufs::opts$get("ggSaveUnits"),
  dpi = ufs::opts$get("ggSaveDPI"),
  device = NULL,
  type = NULL,
  bg = "transparent",
  preventType = ufs::opts$get("ggSavePreventType"),
  ...
)
```

Arguments

file	The file where to save to.
plot	The plot to save; if omitted, the last drawn plot is saved.
height, width	The dimensions of the plot, specified in units.
units	The units, 'cm', 'mm', or 'in'.
dpi	The resolution (dots per inch). This argument is vectorized.
device	The graphic device; is inferred from the file if not specified.
type	An additional arguments for the graphic device.
bg	The background (e.g. 'white').
preventType	Whether to prevent passing a value for the type argument to ggplot2::ggsave() . This is prevented by default since ggplot2::ggplot() switched to using the ragg device by default, resulting in throwing a warning ("Warning: Using ragg device as default. Ignoring type and antialias arguments") if something if passed for 'type'.
...	Any additional arguments are passed on to ggplot2::ggsave() .

Value

The plot, invisibly.

Examples

```
plot <- ufs::ggBoxplot(mtcars, 'mpg');
ggSave(file=tempfile(fileext=".png"), plot=plot);
```

heading	<i>Print a heading</i>
---------	------------------------

Description

This is just a convenience function to print a markdown or HTML heading at a given 'depth'.

Usage

```
heading(
  ...,
  headingLevel = ufs::opts$get("defaultHeadingLevel"),
  output = "markdown",
  cat = TRUE
)
```

Arguments

...	The heading text: pasted together with no separator.
headingLevel	The level of the heading; the default can be set with e.g. <code>ufs::opts\$set(defaultHeadingLevel=1)</code> .
output	Whether to output to HTML ("html") or markdown (anything else).
cat	Whether to cat (print) the heading or just invisibly return it.

Value

The heading, invisibly.

Examples

```
heading("Hello ", "World", headingLevel=5);
### This produces: "\n\n#### Hello World\n\n"
```

ifelseObj	<i>Conditional returning of an object</i>
-----------	---

Description

The ifelseObj function just evaluates a condition, returning one object if it's true, and another if it's false.

Usage

```
ifelseObj(condition, ifTrue, ifFalse)
```

Arguments

condition	Condition to evaluate.
ifTrue	Object to return if the condition is true.
ifFalse	Object to return if the condition is false.

Value

One of the two objects

Examples

```
dat <- ifelseObj(sample(c(TRUE, FALSE), 1), mtcars, Orange);
```

insertFigureCaption	<i>Insert numbered caption</i>
---------------------	--------------------------------

Description

These functions can be used to manually insert a numbered caption. These functions have been designed to work well with [setFigCapNumbering\(\)](#) and [setTabCapNumbering\(\)](#). This is useful when inserting figures or tables in an RMarkdown document when you use automatic caption numbering for knitr chunks, but are inserting a table or figure that isn't produced in a knitr chunk while still retaining the automatic numbering. insertNumberedCaption() is the general-purpose function; you will typically only use insertFigureCaption() and insertTableCaption().

Usage

```

insertFigureCaption(
  captionText = "",
  captionName = "fig.cap",
  prefix = getOption(paste0(optionName, "_prefix"), "Figure %s: "),
  suffix = getOption(paste0(optionName, "_suffix"), ""),
  optionName = paste0("setCaptionNumbering_", captionName),
  resetCounterTo = NULL
)

insertNumberedCaption(
  captionText = "",
  captionName = "fig.cap",
  prefix = getOption(paste0(optionName, "_prefix"), "Figure %s: "),
  suffix = getOption(paste0(optionName, "_suffix"), ""),
  optionName = paste0("setCaptionNumbering_", captionName),
  resetCounterTo = NULL
)

insertTableCaption(
  captionText = "",
  captionName = "tab.cap",
  prefix = getOption(paste0(optionName, "_prefix"), "Table %s: "),
  suffix = getOption(paste0(optionName, "_suffix"), ""),
  optionName = paste0("setCaptionNumbering_", captionName),
  resetCounterTo = NULL
)

```

Arguments

captionText	The text of the caption.
captionName	The name of the caption; by default, for tables, "tab.cap".
prefix, suffix	The prefix and suffix texts; <code>base::sprintf()</code> is used to insert the number in the position taken up by <code>%s</code> .
optionName	The name of the option to use to save the number counter.
resetCounterTo	If a numeric value, the counter is reset to that value.

Value

The caption in a character vector.

Examples

```

insertNumberedCaption("First caption");
insertNumberedCaption("Second caption");
sectionNumber <- 12;
insertNumberedCaption("Third caption",
  prefix = paste0("Table ",

```

```
sectionNumber,
"%s: ");
```

invertItem

invertItems

Description

Inverts items (as in, in a questionnaire), by calling `invertItem` on all relevant items.

Usage

```
invertItem(item, fullRange = NULL, ignorePreviousInversion = FALSE)
```

```
invertItems(dat, items = NULL, ...)
```

Arguments

<code>item</code>	The vector to invert.
<code>fullRange</code>	The full range; will otherwise be derived from the vector.
<code>ignorePreviousInversion</code>	Whether to avoid inverting items that were already inverted.
<code>dat</code>	The dataframe containing the variables to invert.
<code>items</code>	The names or indices of the variables to invert. If not supplied (i.e. <code>NULL</code>), all variables in the dataframe will be inverted.
<code>...</code>	Arguments (parameters) passed on to <code>data.frame</code> when recreating that after having used <code>lapply</code> .

Value

The dataframe with the specified items inverted.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[invertItem](#)

Examples

```
invertItems(mtcars, c('cyl'));
```

`iqrOutlier`*Identify outliers according to the IQR criterion*

Description

The IQR criterion holds that any value lower than one-and-a-half times the interquartile range below the first quartile, or higher than one-and-a-half times the interquartile range above the third quartile, is an outlier. This function returns a logical vector that identifies those outliers.

Usage

```
iqrOutlier(x)
```

Arguments

`x` The vector to scan for outliers.

Value

A logical vector where TRUE identifies outliers.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[IQR](#)

Examples

```
### One outlier in the miles per gallon
iqrOutlier(mtcars$mpg);
```

`irpplot`*Visualising individual response patterns*

Description

Visualising individual response patterns

Usage

```
irpplot(
  data,
  row,
  columns,
  dataName = NULL,
  title = paste("Row", row, "in dataset", dataName)
)
```

Arguments

data	A dataframe with the dataset containing the responses.
row	A vector with indices of the rows for which you want the individual response patterns. These can be either the relevant row numbers, or if character row names are set, the names of the relevant rows.
columns	A vector with the names of the variables you want the individual response patterns for.
dataName, title	Optionally, you can override the dataset name that is used in the title; or, the title (the dataset name is only used in the title).

Value

A `ggplot2::ggplot()`.

Examples

```
### Get a dataset
dat <- ufs::bfi;

### Show the individual responses for
### the tenth participant
irpplot(dat, 10, 1:20);

### Set some missing values
dat[10, c(1, 5, 15)] <- NA;

### Show the individual responses again
irpplot(dat, 10, 1:20);
```

is.nr

NULL and NA 'proof' checking of whether something is a number

Description

Convenience function that returns TRUE if the argument is not null, not NA, and is.numeric.

Usage

```
is.nr(x)
```

Arguments

x The value or vector to check.

Value

TRUE or FALSE.

Examples

```
is.nr(8);     ### Returns TRUE  
is.nr(NULL); ### Returns FALSE  
is.nr(NA);    ### Returns FALSE
```

is.odd

Checking whether numbers are odd or even

Description

Checking whether numbers are odd or even

Usage

```
is.odd(vector)
```

```
is.even(vector)
```

Arguments

vector The vector to process

Value

A logical vector.

Examples

```
is.odd(4);
```

isTrue	<i>More flexible version of isTRUE</i>
--------	--

Description

Returns TRUE for TRUE elements, FALSE for FALSE elements, and whatever is specified in na for NA items.

Usage

```
isTrue(x, na = FALSE)
```

Arguments

x	The vector to check for TRUE, FALSE, and NA values.
na	What to return for NA values.

Value

A logical vector.

Examples

```
isTrue(c(TRUE, FALSE, NA));
isTrue(c(TRUE, FALSE, NA), na=TRUE);
```

kblXtra	<i>Wrapper for kableExtra for consistent ufs table styling</i>
---------	--

Description

Wrapper for kableExtra for consistent ufs table styling

Usage

```
kblXtra(
  x,
  digits = 2,
  format = "html",
  escape = FALSE,
  print = TRUE,
  viewer = FALSE,
  kable_classic = FALSE,
  lighttable_options = "striped",
  html_font = "\"Arial Narrow\"", "\"Source Sans Pro\"", sans-serif,
```

```
    full_width = TRUE,
    table.attr = "style='border:0px solid black !important;'",
    ...
  )
```

Arguments

- x The dataframe to print
- digits, format, escape, table.attr, lightable_options, html_font, full_width Defaults that are passed to `knitr::kable()`
- print Wther to print the table
- viewer Whether to show the table in the viewer
- kable_classic Whether to call `kable_classic`; otherwise, `kable_styling` is called.
- ... Additional arguments are passed to `knitr::kable()`

Value

The table, invisibly.

Examples

```
kblXtra(mtcars);
```

knitAndSave	<i>knitAndSave</i>
-------------	--------------------

Description

knitAndSave

Usage

```
knitAndSave(
  plotToDraw,
  figCaption,
  file = NULL,
  path = NULL,
  figWidth = ufs::opts$get("ggSaveFigWidth"),
  figHeight = ufs::opts$get("ggSaveFigHeight"),
  units = ufs::opts$get("ggSaveUnits"),
  dpi = ufs::opts$get("ggSaveDPI"),
  catPlot = ufs::opts$get("knitAndSave.catPlot"),
  ...
)
```


Arguments

plotToDraw	The plot to knit using <code>knitFig()</code> and save using <code>ggSave()</code> .
figCaption	The caption of the plot (used as filename if no filename is specified).
file, path	The filename to use when saving the plot, or the path where to save the file if no filename is provided (if path is also omitted, <code>getWd()</code> is used).
figWidth, figHeight	The plot dimensions, by default specified in inches (but 'units' can be set which is then passed on to <code>ggSave()</code>).
units, dpi	The units and DPI of the image which are then passed on to <code>ggSave()</code> .
catPlot	Whether to use <code>cat()</code> to print the knitr fragment.
...	Additional arguments are passed on to <code>ggSave()</code> . Note that file (and ...) are vectorized (see the <code>ggSave()</code> manual page).

Value

The `knitFig()` result, visibly.

Examples

```
## Not run: plot <- ggBoxplot(mtcars, 'mpg');
knitAndSave(plot, figCaption="a boxplot", file=tempfile(fileext=".png"));
## End(Not run)
```

knitFig

Easily knit a custom figure fragment

Description

This function was written to make it easy to knit figures with different, or dynamically generated, widths and heights (and captions) in the same chunk when working with R Markdown.

Usage

```
knitFig(
  plotToDraw,
  template = getOption("ufs.knitFig.template", NULL),
  figWidth = ufs::opts$get("ggSaveFigWidth"),
  figHeight = ufs::opts$get("ggSaveFigHeight"),
  figCaption = "A plot.",
  chunkName = NULL,
  returnRaw = FALSE,
  catPlot = ufs::opts$get("knitFig.catPlot"),
  ...
)
```

Arguments

plotToDraw	The plot to draw, e.g. a ggplot plot.
template	A character value with the knit_expand template to use.
figWidth	The width to set for the figure (in inches).
figHeight	The height to set for the figure (in inches).
figCaption	The caption to set for the figure.
chunkName	Optionally, the name for the chunk. To avoid problems because multiple chunks have the name "unnamed-chunk-1", if no chunk name is provided, digest::digest() is used to generate an MD5-hash from Sys.time .
returnRaw	Whether to cat() the result (TRUE) or whether to return it as knitr::asis_output() object (FALSE).
catPlot	Whether to use the base::cat() function to print the code for the plot, and return the result invisibly. If not, the result is returned visible, and so probably printed anyway.
...	Any additional arguments are passed on to knit_expand .

Value

This function returns nothing, but uses [knit_expand](#) and [knit](#) to [cat](#) the result.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[knit_expand](#) and [knit](#)

Examples

```
## Not run: knitFig(ggBoxplot(mtcars, 'mpg'))
```

makeScales	<i>Title</i>
------------	--------------

Description

Title

Usage

```
makeScales(data, scales, append = TRUE)
```

Arguments

data	The dataframe containing the variables (the items).
scales	A list of character vectors with the items in each scale, where each vectors' name is the name of the scale.
append	Whether to return the dataframe including the new variables (TRUE), or a dataframe with only those new variables (FALSE).

Value

Either a dataframe with the newly created variables, or the supplied dataframe with the newly created variables appended.

Examples

```
### First generate a list with the scales
scales <- list(scale1 = c('mpg', 'cyl'), scale2 = c('disp', 'hp'));

### Create the scales and add them to the dataframe
makeScales(mtcars, scales);
```

massConvertToNumeric *Converting many dataframe columns to numeric*

Description

This function makes it easy to convert many dataframe columns to numeric.

Usage

```
massConvertToNumeric(
  dat,
  byFactorLabel = FALSE,
  ignoreCharacter = TRUE,
  stringsAsFactors = FALSE
)
```

Arguments

dat	The dataframe with the columns.
byFactorLabel	When converting factors, whether to do this by their label value (TRUE) or their level value (FALSE).
ignoreCharacter	Whether to convert (FALSE) or ignore (TRUE) character vectors.
stringsAsFactors	In the returned dataframe, whether to return string (character) vectors as factors or not.

Value

A data.frame.

Examples

```
### Create a dataset
a <- data.frame(var1 = factor(1:4),
                var2 = as.character(5:6),
                stringsAsFactors=FALSE);

### Ignores var2
b <- ufs::massConvertToNumeric(a);

### Converts var2
c <- ufs::massConvertToNumeric(a,
                              ignoreCharacter = FALSE);
```

meanConfInt	<i>A confidence interval for the mean</i>
-------------	---

Description

A confidence interval for the mean

Usage

```
meanConfInt(
  vector = NULL,
  mean = NULL,
  sd = NULL,
  n = NULL,
  se = NULL,
  conf.level = 0.95
)

## S3 method for class 'meanConfInt'
print(x, digits = 2, ...)
```

Arguments

- | | |
|------------|---|
| vector | A vector with raw data points - either specify this or a mean and then either an sd and n or an se. |
| mean | A mean. |
| sd, n | A standard deviation and sample size; can be specified to compute the standard error. |
| se | The standard error (cna be specified instead of sd and n). |
| conf.level | The confidence level of the interval. |

`x, digits, ...` Respectively the object to print, the number of digits to round to, and any additional arguments to pass on to the `print` function.

Value

And object with elements `input`, `intermediate`, and `output`, where `output` holds the result in list `ci`.

Examples

```
meanConfInt(mean=5, sd=2, n=20);
```

meansDiamondPlot	<i>Diamond plots</i>
------------------	----------------------

Description

This function generates a so-called diamond plot: a plot based on the forest plots that are commonplace in meta-analyses. The underlying idea is that point estimates are uninformative, and it would be better to focus on confidence intervals. The problem of the points with errorbars that are commonly employed is that the focus the audience's attention on the upper and lower bounds, even though those are the least relevant values. Using diamonds remedies this.

Usage

```
meansDiamondPlot(
  data,
  items = NULL,
  labels = NULL,
  decreasing = NULL,
  conf.level = 0.95,
  showData = TRUE,
  dataAlpha = 0.1,
  dataSize = 3,
  dataColor = "#444444",
  diamondColors = NULL,
  jitterWidth = 0.5,
  jitterHeight = 0.4,
  returnLayerOnly = FALSE,
  xlab = "Scores and means",
  ylab = NULL,
  theme = ggplot2::theme_bw(),
  xbreaks = "auto",
  outputFile = NULL,
  outputWidth = 10,
  outputHeight = 10,
  ggsaveParams = ufs::opts$get("ggsaveParams"),
```

```

    dat = NULL,
    ...
)

```

Arguments

<code>data, dat</code>	The dataframe containing the variables (<code>items</code>) to show in the diamond plot (the name <code>dat</code> for this argument is deprecated but still works for backward compatibility).
<code>items</code>	Optionally, the names (or numeric indices) of the variables (<code>items</code>) to show in the diamond plot. If <code>NULL</code> , all columns (variables, <code>items</code>) will be used.
<code>labels</code>	A character vector of labels to use instead of column names from the dataframe.
<code>decreasing</code>	Whether to sort the variables (rows) in the diamond plot decreasing (<code>TRUE</code>), increasing (<code>FALSE</code>), or not at all (<code>NULL</code>).
<code>conf.level</code>	The confidence of the confidence intervals.
<code>showData</code>	Whether to show the raw data or not.
<code>dataAlpha</code>	This determines the alpha (transparency) of the data points. Note that argument <code>alpha</code> can be used to set the alpha of the diamonds; this is eventually passed on to ggDiamondLayer() .
<code>dataSize</code>	The size of the data points.
<code>dataColor</code>	The color of the data points.
<code>diamondColors</code>	A vector of the same length as there are rows in the dataframe, to manually specify colors for the diamonds.
<code>jitterWidth</code>	How much to jitter the individual datapoints horizontally.
<code>jitterHeight</code>	How much to jitter the individual datapoints vertically.
<code>returnLayerOnly</code>	Set this to <code>TRUE</code> to only return the ggplot() layer of the diamondplot, which can be useful to include it in other plots.
<code>xlab, ylab</code>	The labels of the X and Y axes.
<code>theme</code>	The theme to use.
<code>xbreaks</code>	Where the breaks (major grid lines, ticks, and labels) on the x axis should be.
<code>outputFile</code>	A file to which to save the plot.
<code>outputWidth, outputHeight</code>	Width and height of saved plot (specified in centimeters by default, see ggsaveParams).
<code>ggsaveParams</code>	Parameters to pass to ggsave when saving the plot.
<code>...</code>	Additional arguments are passed to diamondPlot() and eventually to ggDiamondLayer() . This can be used to, for example, specify two or more colors to use to generate a gradient (using generateColors and maybe fullColorRange).

Value

A [ggplot\(\)](#) plot with a [ggDiamondLayer\(\)](#) is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

See [diamondPlot\(\)](#), [meanSDtoDiamondPlot\(\)](#), [ggDiamondLayer\(\)](#), [factorLoadingDiamondCIplot\(\)](#)

Examples

```
tmpDf <- data.frame(item1 = rnorm(50, 1.6, 1),
                    item2 = rnorm(50, 2.6, 2),
                    item3 = rnorm(50, 4.1, 3));

### A simple diamond plot
meansDiamondPlot(tmpDf);

### A diamond plot with manually
### specified labels and colors
meansDiamondPlot(tmpDf,
                 labels=c('First',
                         'Second',
                         'Third'),
                 diamondColors=c('blue', 'magenta', 'yellow'));

### Using a gradient for the colors
meansDiamondPlot(tmpDf,
                 labels=c('First',
                         'Second',
                         'Third'),
                 generateColors = c("magenta", "cyan"),
                 fullColorRange = c(1,5));
```

meansDiamondPlotjmv	<i>Diamond plot: means</i>
---------------------	----------------------------

Description

Diamond plot: means

Usage

```
meansDiamondPlotjmv(data, items, conf.level = 95, showData = TRUE)
```

Arguments

data .
items .
conf.level .
showData .

Value

A results object containing:

results\$text	a html
results\$diamondPlot	an image

meanSDtoDiamondPlot	<i>A diamond plot based on means, standard deviations, and sample sizes</i>
---------------------	---

Description

This function generates a so-called diamond plot: a plot based on the forest plots that are commonplace in meta-analyses. The underlying idea is that point estimates are uninformative, and it would be better to focus on confidence intervals. The problem of the points with errorbars that are commonly employed is that the focus the audience’s attention on the upper and lower bounds, even though those are the least relevant values. Using diamonds remedies this.

Usage

```
meanSDtoDiamondPlot(  
  dat = NULL,  
  means = 1,  
  sds = 2,  
  ns = 3,  
  labels = NULL,  
  colorCol = NULL,  
  conf.level = 0.95,  
  xlab = "Means",  
  outputFile = NULL,  
  outputWidth = 10,  
  outputHeight = 10,  
  ggsaveParams = ufs::opts$get("ggsaveParams"),  
  ...  
)
```


Arguments

<code>dat</code>	The dataset containing the means, standard deviations, sample sizes, and possible labels and manually specified colors.
<code>means</code>	Either the column in the dataframe containing the means, as numeric or as character index, or a vector of means.
<code>sds</code>	Either the column in the dataframe containing the standard deviations, as numeric or as character index, or a vector of standard deviations.
<code>ns</code>	Either the column in the dataframe containing the sample sizes, as numeric or as character index, or a vector of sample sizes.
<code>labels</code>	Optionally, either the column in the dataframe containing labels, as numeric or as character index, or a vector of labels.
<code>colorCol</code>	Optionally, either the column in the dataframe containing manually specified colours, as numeric or as character index, or a vector of manually specified colours.
<code>conf.level</code>	The confidence of the confidence intervals.
<code>xlab</code>	The label for the x axis.
<code>outputFile</code>	A file to which to save the plot.
<code>outputWidth, outputHeight</code>	Width and height of saved plot (specified in centimeters by default, see <code>ggsaveParams</code>).
<code>ggsaveParams</code>	Parameters to pass to <code>ggsave</code> when saving the plot.
<code>...</code>	Additional arguments are passed to <code>diamondPlot()</code> and eventually to <code>ggDiamondLayer()</code> . This can be used to, for example, specify two or more colors to use to generate a gradient (using <code>generateColors</code> and maybe <code>fullColorRange</code>).

Value

A `ggplot()` plot with a `ggDiamondLayer()` is returned.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`meansDiamondPlot()`, `diamondPlot()`, `factorLoadingDiamondCIplot()`, `ggDiamondLayer()`

Examples

```
tmpDf <- data.frame(means = c(1, 2, 3),
                    sds = c(1.5, 3, 5),
                    ns = c(2, 4, 10),
                    labels = c('first', 'second', 'third'),
                    color = c('purple', 'grey', 'orange'));

### A simple diamond plot
```

```

meanSDtoDiamondPlot(tmpDf);

### A simple diamond plot with labels
meanSDtoDiamondPlot(tmpDf, labels=4);

### When specifying column names, specify column
### names for all columns
meanSDtoDiamondPlot(tmpDf, means='means',
                     sds='sds', ns='ns',
                     labels='labels');

### A diamond plot using the specified colours
meanSDtoDiamondPlot(tmpDf, labels=4, colorCol=5);

### A diamond plot using automatically generated colours
### using a gradient
meanSDtoDiamondPlot(tmpDf,
                     generateColors=c('green', 'red'));

### A diamond plot using automatically generated colours
### using a gradient, specifying the minimum and maximum
### possible values that can be attained
meanSDtoDiamondPlot(tmpDf,
                     generateColors=c('red', 'yellow', 'blue'),
                     fullColorRange=c(0, 5));

```

multiResponse

Generate a table for multiple response questions

Description

The multiResponse function mimics the behavior of the table produced by SPSS for multiple response questions.

Usage

```

multiResponse(
  data,
  items = NULL,
  regex = NULL,
  perlRegex = TRUE,
  endorsedOption = 1
)

```

Arguments

data Dataframe containing the variables to display.

items, regex	Arguments items and regex can be used to specify which variables to process. items should contain the variable (column) names (or indices), and regex should contain a regular expression used to match to the column names of the dataframe. If none is provided, all variables in the dataframe are processed.
perlRegex	Whether to use the perl engine to match the regex.
endorsedOption	Which value represents the endorsed option (note that producing this kind of table requires dichotomous items, where each variable is either endorsed or not endorsed, so this is also a way to treat other variables as dichotomous).

Value

A dataframe with columns Option, Frequency, Percentage, and Percentage of (X) cases, where X is the number of cases.

Author(s)

Ananda Mahto; implemented in this package (and tweaked a bit) by Gjalt-Jorn Peters.

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

This function is based on the excellent and extensive Stack Exchange answer by Ananda Mahto at <https://stackoverflow.com/questions/9265003/analysis-of-multiple-response>.

Examples

```
multiResponse(mtcars, c('vs', 'am'));
```

multiResponsejmv	<i>Multi Response</i>
------------------	-----------------------

Description

Multi Response

Usage

```
multiResponsejmv(data, items, endorsedOption = 1)
```

Arguments

```
data          .
items         .
endorsedOption .
```

Value

A results object containing:

`results$table` a table

Tables can be converted to data frames with `asDF` or `as.data.frame`. For example:

```
results$table$asDF
```

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

multiVarFreq

Generate a table collapsing frequencies of multiple variables

Description

This function can be used to efficiently combine the frequencies of variables with the same possible values. The frequencies are collapsed into a table with the variable names as row names and the possible values as column (variable) names.

Usage

```
multiVarFreq(data, items = NULL, labels = NULL, sortByMean = TRUE)
```

Arguments

<code>data</code>	The dataframe containing the variables.
<code>items</code>	The variable names.
<code>labels</code>	Labels can be provided which will be set as row names when provided.
<code>sortByMean</code>	Whether to sort the rows by mean value for each variable (only sensible if the possible values are numeric).

Value

The resulting dataframe, but with class 'multiVarFreq' prepended to allow pretty printing.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[table\(\)](#)

Examples

```
multiVarFreq(mtcars, c('gear', 'carb'));
```

normalHist	<i>normalHist</i>
------------	-------------------

Description

normalHist generates a histogram with a density curve and a normal density curve.

Usage

```
normalHist(
  vector,
  histColor = "#0000CC",
  distributionColor = "#0000CC",
  normalColor = "#00CC00",
  distributionLineSize = 1,
  normalLineSize = 1,
  histAlpha = 0.25,
  xLabel = NULL,
  yLabel = NULL,
  normalCurve = TRUE,
  distCurve = TRUE,
  breaks = 30,
  theme = ggplot2::theme_minimal(),
  rug = NULL,
  jitteredRug = TRUE,
  rugSides = "b",
  rugAlpha = 0.2,
  returnPlotOnly = FALSE
)

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

Arguments

vector	A numeric vector.
histColor	The colour to use for the histogram.
distributionColor	The colour to use for the density curve.
normalColor	The colour to use for the normal curve.
distributionLineSize	The line size to use for the distribution density curve.
normalLineSize	The line size to use for the normal curve.
histAlpha	Alpha value ('opaqueness', as in, versus transparency) of the histogram.
xLabel	Label to use on x axis.

yLabel	Label to use on y axis.
normalCurve	Whether to display the normal curve.
distCurve	Whether to display the curve showing the distribution of the observed data.
breaks	The number of breaks to use (this is equal to the number of bins minus one, or in other words, to the number of bars minus one).
theme	The theme to use.
rug	Whether to add a rug (i.e. lines at the bottom that correspond to individual datapoints).
jitteredRug	Whether to jitter the rug (useful for variables with several datapoints sharing the same value).
rugSides	This is useful when the histogram will be rotated; for example, this can be set to 'r' if the histogram is rotated 270 degrees.
rugAlpha	Alpha value to use for the rug. When there is a lot of overlap, this can help get an idea of the number of datapoints at 'popular' values.
returnPlotOnly	Whether to return the usual normalHist object that also contains all settings and intermediate objects, or whether to only return the <code>ggplot2::ggplot()</code> plot.
x	The object to print.
...	Any additional arguments are passed to the default print method.

Value

An object, with the following elements:

input	The input when the function was called.
intermediate	The intermediate numbers and distributions.
dat	The dataframe used to generate the plot.
plot	The histogram.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
normalHist(mtcars$mpg)
```

noZero	<i>Remove one or more zeroes before the decimal point</i>
--------	---

Description

Remove one or more zeroes before the decimal point

Usage

```
noZero(str)
```

Arguments

str	The character string to process.
-----	----------------------------------

Value

The processed string.

See Also

[formatCI\(\)](#), [formatR\(\)](#), [formatPvalue\(\)](#)

Examples

```
noZero("0.3");
```

opts	<i>Options for the ufs package</i>
------	------------------------------------

Description

The `ufs:::opts` object contains three functions to set, get, and reset options used by the `ufs` package. Use `ufs:::opts$set` to set options, `ufs:::opts$get` to get options, or `ufs:::opts$reset` to reset specific or all options to their default values.

Usage

```
opts
```

Format

An object of class `list` of length 5.

Details

It is normally not necessary to get or set ufs options.

The following arguments can be passed:

... For `ufs::opts$set`, the dots can be used to specify the options to set, in the format `option = value`, for example, `tableOutput = c("console", "viewer")`. For `ufs::opts$reset`, a list of options to be reset can be passed.

option For `ufs::opts$set`, the name of the option to set.

default For `ufs::opts$get`, the default value to return if the option has not been manually specified.

The following options can be set:

tableOutput Where to show some tables.

Examples

```
### Get the default columns in the variable view
ufs::opts$get("tableOutput");

### Set it to a custom version
ufs::opts$set(tableOutput = c("values", "level"));

### Check that it worked
ufs::opts$get("tableOutput");

### Reset this option to its default value
ufs::opts$reset("tableOutput");

### Check that the reset worked, too
ufs::opts$get("tableOutput");
```

parallelSubscales	<i>Split a dataset into two parallel halves</i>
-------------------	---

Description

Split a dataset into two parallel halves

Usage

```
parallelSubscales(dat, convertToNumeric = TRUE)

## S3 method for class 'parallelSubscales'
print(x, digits = 2, ...)
```


Arguments

<code>dat</code>	The dataframe
<code>convertToNumeric</code>	Whether to first convert all columns to numeric
<code>x</code>	The object to print
<code>digits</code>	The number of digits to round to
<code>...</code>	Ignored.

Value

A `parallelSubscales` object that contains the new data frames, and when printed shows the descriptives; or, for the print function, `x`, invisibly.

<code>pomegaSq</code>	<i>The distribution of Omega Squared</i>
-----------------------	--

Description

These functions use some conversion to and from the F distribution to provide the Omega Squared distribution.

Usage

```
pomegaSq(q, df1, df2, populationOmegaSq = 0, lower.tail = TRUE)

qomegaSq(p, df1, df2, populationOmegaSq = 0, lower.tail = TRUE)

romegaSq(n, df1, df2, populationOmegaSq = 0)

domegaSq(x, df1, df2, populationOmegaSq = 0)
```

Arguments

<code>df1, df2</code>	Degrees of freedom for the numerator and the denominator, respectively.
<code>populationOmegaSq</code>	The value of Omega Squared in the population; this determines the center of the Omega Squared distribution. This has not been implemented yet in this version of ufs. If anybody has the inverse of <code>convert.ncf.to.omegasq()</code> for me, I'll happily integrate this.
<code>lower.tail</code>	logical; if TRUE (default), probabilities are the likelihood of finding an Omega Squared smaller than the specified value; otherwise, the likelihood of finding an Omega Squared larger than the specified value.
<code>p</code>	Vector of probabilities (p -values).
<code>n</code>	Desired number of Omega Squared values.
<code>x, q</code>	Vector of quantiles, or, in other words, the value(s) of Omega Squared.

Details

The functions use `convert.omegasq.to.f()` and `convert.f.to.omegasq()` to provide the Omega Squared distribution.

Value

domegasq gives the density, pomegasq gives the distribution function, qomegasq gives the quantile function, and romeegasq generates random deviates.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

`convert.omegasq.to.f()`, `convert.f.to.omegasq()`, `df()`, `pf()`, `qf()`, `rf()`

Examples

```
### Generate 10 random Omega Squared values
romeegasq(10, 66, 3);

### Probability of findings an Omega Squared
### value smaller than .06 if it's 0 in the population
pomegasq(.06, 66, 3);
```

pwr.bootES

Estimate required sample size for accuracy in parameter estimation using bootES

Description

This function uses `bootES::bootES()` to compute

Usage

```
pwr.bootES(data = data, ci.type = "bca", ..., w = 0.1, silent = TRUE)
```

Arguments

data	The dataset, as you would normally supply to <code>bootES::bootES()</code> ; you will probably have to simulate this.
ci.type	The estimation method; by default, the default of <code>bootES::bootES()</code> is used ('bca'), but this is changed to 'basic' if it encounters problems.
...	Other options for <code>bootES::bootES()</code> (see that help page).
w	The desired 'halfwidth' of the confidence interval.
silent	Whether to provide a lot of information about progress ('FALSE') or not ('TRUE').

Value

A single numeric value (the sample size).

References

Kirby, K. N., & Gerlanc, D. (2013). BootES: An R package for bootstrap confidence intervals on effect sizes. *Behavior Research Methods*, 45, 905–927. doi:10.3758/s1342801303305

Examples

```
### This requires the bootES package
if (requireNamespace("bootES", quietly = TRUE)) {

  ### To estimate a mean
  x <- rnorm(500, mean=8, sd=3);
  pwr.bootES(data.frame(x=x),
              R=500,
              w=.5);

  ### To estimate a correlation (the 'effect.type' parameter is
  ### redundant here; with two columns in the data frame, computing
  ### the confidence interval for the Pearson correlation is the default
  ### behavior of bootES)
  y <- x+rnorm(500, mean=0, sd=5);
  cor(x, y);
  requiredN <-
    pwr.bootES(data.frame(x=x,
                          y=y),
                effect.type='r',
                R=500,
                w=.2);
  print(requiredN);
  ### Compare to parametric confidence interval
  ### based on the computed required sample size
  confIntR(r = cor(x, y),
           N = requiredN);
  ### Width of obtained confidence interval
  print(round(diff(as.numeric(confIntR(r = cor(x, y),
                                     N = requiredN))), 2));
}
```

pwr.confIntProp	<i>Estimate required sample size for accuracy in parameter estimation of a proportion</i>
-----------------	---

Description

This function uses `confIntProp()` to compute the required sample size for estimating a proportion with a given accuracy.

Usage

```
pwr.confIntProp(prop, conf.level = 0.95, w = 0.1, silent = TRUE)
```

Arguments

prop	The proportion you expect to find, or a vector of proportions to enable easy sensitivity analyses.
conf.level	The confidence level of the desired confidence interval.
w	The desired 'halfwidth' of the confidence interval.
silent	Whether to provide a lot of information about progress ('FALSE') or not ('TRUE').

Value

A single numeric value (the sample size).

Examples

```
### Required sample size to estimate a prevalence of .03 in the
### population with a confidence interval of a maximum half-width of .01
pwr.confIntProp(.03, w=.01);

### Vectorized over prop, so you can easily see how the required sample
### size varies as a function of the proportion
pwr.confIntProp(c(.03, .05, .10), w=.01);
```

pwr.confIntR	<i>Determine required sample size for a given confidence interval width for Pearson's r</i>
--------------	---

Description

This function computes how many participants you need if you want to achieve a confidence interval of a given width. This is useful when you do a study and you are interested in how strongly two variables are associated.

Usage

```
pwr.confIntR(r, w = 0.1, conf.level = 0.95)
```

Arguments

r	The correlation you expect to find (confidence intervals for a given level of confidence get narrower as the correlation coefficient increases).
w	The required half-width (or margin of error) of the confidence interval.
conf.level	The level of confidence.

Value

The required sample size, or a vector or matrix of sample sizes if multiple correlation coefficients or required (half-)widths were supplied. The row and column names specify the *r* and *w* values to which the sample size in each cell corresponds. The confidence level is set as attribute to the resulting vector or matrix.

Author(s)

Douglas Bonett (UC Santa Cruz, United States), with minor edits by Murray Moinester (Tel Aviv University, Israel) and Gjalt-Jorn Peters (Open University of the Netherlands, the Netherlands).

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

Bonett, D. G., Wright, T. A. (2000). Sample size requirements for estimating Pearson, Kendall and Spearman correlations. *Psychometrika*, 65, 23-28.

Bonett, D. G. (2014). CIcorr.R and sizeCIcorr.R <http://people.ucsc.edu/~dgbonett/psyc181.html>

Moinester, M., & Gottfried, R. (2014). Sample size estimation for correlations with pre-specified confidence interval. *The Quantitative Methods of Psychology*, 10(2), 124-130. <http://www.tqmp.org/RegularArticles/vol10-2/p124/p124.pdf>

Peters, G. J. Y. & Crutzen, R. (forthcoming) An easy and foolproof method for establishing how effective an intervention or behavior change method is: required sample size for accurate parameter estimation in health psychology.

See Also

[pwr.confIntR](#)

Examples

```
pwr.confIntR(c(.4, .6, .8), w=c(.1, .2));
```

pwr.omegasq

Power calculations for Omega Squared.

Description

This function uses [pwr.anova.test](#) from the [pwr](#) package in combination with [convert.cohensf.to.omegasq](#) and [convert.omegasq.to.cohensf](#) to provide power analyses for Omega Squared.

Usage

```
pwr.omegasq(
  k = NULL,
  n = NULL,
  omegasq = NULL,
  sig.level = 0.05,
  power = NULL,
  digits = 4
)

## S3 method for class 'pwr.omegasq'
print(x, digits = x$digits, ...)
```

Arguments

k	The number of groups.
n	The sample size.
omegasq	The Omega Squared value.
sig.level	The significance level (alpha).
power	The power.
digits	The number of digits desired in the output (4, the default, is quite high; but omega squared value tend to be quite low).
x	The object to print.
...	Additional arguments are ignored.

Details

This function was written to work similarly to the power functions in the [pwr](#) package.

Value

An `pwr.htest.ufs` object that contains a number of input and output values, most notably:

power	The (specified or computed) power
n	The (specified or computed) sample size in each group
sig.level	The (specified or computed) significance level (alpha)
sig.level	The (specified or computed) Omega Squared value
cohensf	The computed value for the Cohen's <i>f</i> effect size measure

Author(s)

Gjalt-Jorn Peters & Peter Verboon

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

See Also

[pwr.anova.test](#), [convert.cohensf.to.omegasq](#), [convert.omegasq.to.cohensf](#)

Examples

```
pwr.omegasq(omegasq=.06, k=3, power=.8)
```

quietRemotesInstall	<i>Quietly update a package from a remote repository</i>
---------------------	--

Description

Simple wrapper for remotes functions that fail gracefully (well, don't fail at all, just don't do what they're supposed to do) when there's no internet connection).

Usage

```
quietRemotesInstall(
  x,
  func,
  unloadNamespace = TRUE,
  dependencies = FALSE,
  upgrade = FALSE,
  quiet = TRUE,
  errorInvisible = TRUE,
  ...
)

quietGitLabUpdate(
  x,
  unloadNamespace = TRUE,
  dependencies = FALSE,
  upgrade = FALSE,
  quiet = TRUE,
  errorInvisible = TRUE,
  ...
)
```

Arguments

x	The repository name (e.g. "r-packages/ufs")
func	The remotes function to use
unloadNamespace	Whether to first unload the relevant namespace
dependencies, upgrade	Whether to install dependencies or upgrade

- quiet Whether to suppress messages and warnings
- errorInvisible Whether to suppress errors
- ... Additional arguments are passed on to the remotes function

Value

The result of the call to the remotes function

qVec	<i>Convenience function to quickly copy-paste a vector</i>
------	--

Description

Convenience function to quickly copy-paste a vector

Usage

```
qVec(x, fn = NULL)

qVecSum(x)
```

Arguments

- x A string with numbers, separated by arbitrary whitespace.
- fn An optional function to apply to the vecor before returning it.

Value

The numeric vector or result of calling the function

Examples

```
qVec('23 9 11 14 12 20');
```

`rbind_dfs`*Simple alternative for rbind.fill or bind_rows*

Description

Simple alternative for rbind.fill or bind_rows

Usage

```
rbind_dfs(x, y, clearRowNames = TRUE)
```

Arguments

<code>x</code>	One dataframe
<code>y</code>	Another dataframe
<code>clearRowNames</code>	Whether to clear row names (to avoid duplication)

Value

The merged dataframe

Examples

```
rbind_dfs(Orange, mtcars);
```

`rbind_df_list`*Bind lots of dataframes together rowwise*

Description

Bind lots of dataframes together rowwise

Usage

```
rbind_df_list(x)
```

Arguments

<code>x</code>	A list of dataframes
----------------	----------------------

Value

A dataframe

Examples

```
rbind_df_list(list(Orange, mtcars, ChickWeight));
```

regrInfluential	<i>Detecting influential cases in regression analyses</i>
-----------------	---

Description

This function combines a number of criteria for determining whether a datapoint is an influential case in a regression analysis. It then sum the criteria to compute an index of influentiality. A list of cases with an index of influentiality of 1 or more is then displayed, after which the regression analysis is repeated without those influential cases. A scattermatrix is also displayed, showing the density curves of each variable, and in the scattermatrix, points that are colored depending on how influential each case is.

Usage

```
regrInfluential(formula, data, createPlot = TRUE)

## S3 method for class 'regrInfluential'
print(x, headingLevel = 3, ...)
```

Arguments

formula	The formule of the regression analysis.
data	The data to use for the analysis.
createPlot	Whether to create the scattermatrix (requires the GGally package to be installed).
x	Object to print.
headingLevel	The number of hash symbols to prepend to the heading.
...	Additional arguments are passed on to the regr print function.

Value

A regrInfluential object, which, if printed, shows the influential cases, the regression analyses repeated without those cases, and the scatter matrix.

Author(s)

Gjalt-Jorn Peters & Marwin Snippe
 Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
regrInfluential(mpg ~ hp, mtcars);
```

repeatStr	<i>Repeat a string a number of times</i>
-----------	--

Description

Repeat a string a number of times

Usage

```
repeatStr(n = 1, str = " ")
```

Arguments

n, str	Normally, respectively the frequency with which to repeat the string and the string to repeat; but the order of the inputs can be switched as well.
--------	---

Value

A character vector of length 1.

Examples

```
### 10 spaces:
repStr(10);

### Three euro symbols:
repStr("\u20ac", 3);
```

report	<i>Output report from results</i>
--------	-----------------------------------

Description

This method can be used to format results in a way that can directly be included in a report or manuscript.

Usage

```
report(x, headingLevel = 3, quiet = TRUE, ...)

## Default S3 method:
report(x, headingLevel = 3, quiet = TRUE, ...)
```

Arguments

x	The object to show.
headingLevel	The level of the Markdown heading to provide; basically the number of hashes ('#') to prepend to the headings.
quiet	Passed on to <code>knitr::knit()</code> whether it should be chatty (FALSE) or quiet (TRUE).
...	Passed to the specific method; for the default method, this is passed to the print method.

safeRequire	<i>Load a package, install if not available</i>
-------------	---

Description

Load a package, install if not available

Usage

```
safeRequire(packageName, mirrorIndex = NULL)
```

Arguments

packageName	The package
mirrorIndex	The index of the mirror (1 is used if not specified)

scaleDiagnosis	<i>scaleDiagnosis</i>
----------------	-----------------------

Description

scaleDiagnosis provides a number of diagnostics for a scale (an aggregative measure consisting of several items).

Usage

```
scaleDiagnosis(
  data = NULL,
  items = NULL,
  plotSize = 180,
  sizeMultiplier = 1,
  axisLabels = "none",
  scaleReliability.ci = FALSE,
  conf.level = 0.95,
  normalHist = TRUE,
  poly = TRUE,
```

```

    digits = 3,
    headingLevel = 3,
    scaleName = NULL,
    ...
)

## S3 method for class 'scaleDiagnosis'
print(x, digits = x$digits, ...)

scaleDiagnosis_partial(
  x,
  headingLevel = x$input$headingLevel,
  quiet = TRUE,
  echoPartial = FALSE,
  partialFile = NULL,
  ...
)

## S3 method for class 'scaleDiagnosis'
knit_print(
  x,
  headingLevel = x$headingLevel,
  quiet = TRUE,
  echoPartial = FALSE,
  partialFile = NULL,
  ...
)

```

Arguments

<code>data</code>	A dataframe containing the items in the scale. All variables in this dataframe will be used if <code>items</code> is <code>NULL</code> .
<code>items</code>	If not <code>NULL</code> , this should be a character vector with the names of the variables in the dataframe that represent items in the scale.
<code>plotSize</code>	Size of the final plot in millimeters.
<code>sizeMultiplier</code>	Allows more flexible control over the size of the plot elements
<code>axisLabels</code>	Passed to <code>ggpairs</code> function to set <code>axisLabels</code> .
<code>scaleReliability.ci</code>	TRUE or FALSE: whether to compute confidence intervals for Cronbach's Alpha and Omega (uses bootstrapping function in MBESS, takes a while).
<code>conf.level</code>	Confidence of confidence intervals for reliability estimates (if requested with <code>scaleReliability.ci</code>).
<code>normalHist</code>	Whether to use the default <code>ggpairs</code> histogram on the diagonal of the scattermatrix, or whether to use the <code>normalHist()</code> version.
<code>poly</code>	Whether to also request the estimates based on the polychoric correlation matrix when calling <code>scaleStructure()</code> .

digits	The number of digits to pass to the print method for the descriptives dataframe.
headingLevel	The level of the heading (number of hash characters to insert before the heading, to be rendered as headings of that level in Markdown).
scaleName	Optionally, a name for the scale to print as heading for the results.
...	Additional arguments for scaleDiagnosis() are passed on to <code>scatterMatrix()</code> , and additional arguments for the print method are passed to the default print method.
x	The object to print.
quiet	Whether to be chatty (FALSE) or quiet (TRUE).
echoPartial	Whether to show the code in the partial (TRUE) or hide it (FALSE).
partialFile	The file with the Rmd partial (if you want to overwrite the default).

Details

Function to generate an object with several useful statistics and a plot to assess how the elements (usually items) in a scale relate to each other, such as Cronbach's Alpha, omega, the Greatest Lower Bound, a factor analysis, and a correlation matrix.

Value

An object with the input and several output variables. Most notably:

scaleReliability	The results of scaleReliability.
pca	A Principal Components Analysis
fa	A Factor Analysis
describe	Descriptive statistics about the items
scatterMatrix	A scattermatrix with histograms on the diagonal and correlation coefficients in the upper right half.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

Examples

```
### Note: the 'not run' is simply because running takes a lot of time,
###       but these examples are all safe to run!
## Not run:
### This will prompt the user to select an SPSS file
scaleDiagnosis();

### Generate a datafile to use
exampleData <- data.frame(item1=rnorm(100));
exampleData$item2 <- exampleData$item1+rnorm(100);
exampleData$item3 <- exampleData$item1+rnorm(100);
```

```

exampleData$item4 <- exampleData$item2+rnorm(100);
exampleData$item5 <- exampleData$item2+rnorm(100);

### Use a selection of two variables
scaleDiagnosis(data=exampleData, items=c('item2', 'item4'));

### Use all items
scaleDiagnosis(data=exampleData);

## End(Not run)

```

scaleStructure	<i>scaleStructure</i>
----------------	-----------------------

Description

The `scaleStructure` function (which was originally called `scaleReliability`) computes a number of measures to assess scale reliability and internal consistency. Note that to compute omega, the MBESS and/or the psych packages need to be installed, which are suggested packages and therefore should be installed separately (i.e. won't be installed automatically).

Usage

```

scaleStructure(
  data = NULL,
  items = "all",
  digits = 2,
  ci = TRUE,
  interval.type = "normal-theory",
  conf.level = 0.95,
  silent = FALSE,
  samples = 1000,
  bootstrapSeed = NULL,
  omega.psych = TRUE,
  omega.psych_nfactors = 3,
  omega.psych_flip = TRUE,
  poly = TRUE,
  suppressSuggestedPkgsMsg = FALSE,
  headingLevel = 3
)

## S3 method for class 'scaleStructure'
print(x, digits = x$input$digits, ...)

scaleStructure_partial(
  x,
  headingLevel = x$input$headingLevel,

```

```

    quiet = TRUE,
    echoPartial = FALSE,
    partialFile = NULL,
    ...
)

## S3 method for class 'scaleStructure'
knit_print(
  x,
  headingLevel = x$input$headingLevel,
  quiet = TRUE,
  echoPartial = FALSE,
  partialFile = NULL,
  ...
)

```

Arguments

<code>data</code>	A dataframe containing the items in the scale. All variables in this dataframe will be used if <code>items = 'all'</code> . If <code>dat</code> is <code>NULL</code> , a the getData function will be called to show the user a dialog to open a file.
<code>items</code>	If not <code>'all'</code> , this should be a character vector with the names of the variables in the dataframe that represent items in the scale.
<code>digits</code>	Number of digits to use in the presentation of the results.
<code>ci</code>	Whether to compute confidence intervals as well. This requires the suggested MBESS package, which has to be installed separately. If true, the method specified in <code>interval.type</code> is used. When specifying a bootstrapping method, this can take quite a while!
<code>interval.type</code>	Method to use when computing confidence intervals. The list of methods is explained in the help file for <code>ci.reliability</code> in MBESS. Note that when specifying a bootstrapping method, the method will be set to <code>normal-theory</code> for computing the confidence intervals for the ordinal estimates, because these are based on the polychoric correlation matrix, and raw data is required for bootstrapping.
<code>conf.level</code>	The confidence of the confidence intervals.
<code>silent</code>	If computing confidence intervals, the user is warned that it may take a while, unless <code>silent=TRUE</code> .
<code>samples</code>	The number of samples to compute for the bootstrapping of the confidence intervals.
<code>bootstrapSeed</code>	The seed to use for the bootstrapping - setting this seed makes it possible to replicate the exact same intervals, which is useful for publications.
<code>omega.psych</code>	Whether to also compute the interval estimate for omega using the <code>omega</code> function in the <code>psych</code> package. The default point estimate and confidence interval for omega are based on the procedure suggested by Dunn, Baguley & Brunsden (2013) using the MBESS function <code>ci.reliability</code> (because it has more options for computing confidence intervals, not always requiring bootstrapping),

whereas the psych package point estimate was suggested in Revelle & Zinbarg (2008). The psych estimate usually (perhaps always) results in higher estimates for omega.

<code>omega.psych_nfactors</code>	The number of factor to use in the factor analysis when computing Omega. The default in <code>psych::omega()</code> is 3; to obtain the same results as in jamovi's "Reliability", set this to 1.
<code>omega.psych_flip</code>	Whether to let psych automatically flip items with negative correlations. The default in <code>psych::omega()</code> is TRUE; to obtain the same results as in jamovi's "Reliability", set this to FALSE.
<code>poly</code>	Whether to compute ordinal measures (if the items have sufficiently few categories).
<code>suppressSuggestedPkgsMsg</code>	Whether to suppress the message about the suggested MBESS and psych packages.
<code>headingLevel</code>	The level of the Markdown heading to provide; basically the number of hashes ('#') to prepend to the headings.
<code>x</code>	The object to print
<code>...</code>	Any additional arguments for the default print function.
<code>quiet</code>	Passed on to <code>knitr::knit()</code> whether it should be chatty (FALSE) or quiet (TRUE).
<code>echoPartial</code>	Whether to show the executed code in the R Markdown partial (TRUE) or not (FALSE).
<code>partialFile</code>	This can be used to specify a custom partial file. The file will have object <code>x</code> available, which is the result of a call to <code>scaleStructure()</code> .

Details

If you use this function in an academic paper, please cite Peters (2014), where the function is introduced, and/or Crutzen & Peters (2015), where the function is discussed from a broader perspective.

This function is basically a wrapper for functions from the psych and MBESS packages that compute measures of reliability and internal consistency. For backwards compatibility, in addition to `scaleStructure`, `scaleReliability` can also be used to call this function.

Value

An object with the input and several output variables. Most notably:

<code>input</code>	Input specified when calling the function
<code>intermediate</code>	Intermediate values and objects computed to get to the final results
<code>output</code>	Values of reliability / internal consistency measures, with as most notable elements:
<code>output\$dat</code>	A dataframe with the most important outcomes
<code>output\$omega</code>	Point estimate for omega
<code>output\$glb</code>	Point estimate for the Greatest Lower Bound

```

output$alpha    Point estimate for Cronbach's alpha
output$coefficientH
                Coefficient H
output$omega.ci
                Confidence interval for omega
output$alpha.ci
                Confidence interval for Cronbach's alpha

```

Author(s)

Gjalt-Jorn Peters and Daniel McNeish (University of North Carolina, Chapel Hill, US).

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com

References

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See Also

[psych::omega\(\)](#), [psych::alpha\(\)](#), and [MBESS::ci.reliability\(\)](#).

Examples

```

## Not run:
### (These examples take a lot of time, so they are not run
### during testing.)

```

```

### This will prompt the user to select an SPSS file
scaleStructure();

### Load data from simulated dataset testRetestSimData (which
### satisfies essential tau-equivalence).
data(testRetestSimData);

### Select some items in the first measurement
exampleData <- testRetestSimData[2:6];

### Use all items (don't order confidence intervals to save time
### during automated testing of the example)
ufs::scaleStructure(dat=exampleData, ci=FALSE);

### Use a selection of three variables (without confidence
### intervals to save time
ufs::scaleStructure(
  dat=exampleData,
  items=c('t0_item2', 't0_item3', 't0_item4'),
  ci=FALSE
);

### Make the items resemble an ordered categorical (ordinal) scale
ordinalExampleData <- data.frame(apply(exampleData, 2, cut,
                                       breaks=5, ordered_result=TRUE,
                                       labels=as.character(1:5)));

### Now we also get estimates assuming the ordinal measurement level
ufs::scaleStructure(ordinalExampleData, ci=FALSE);

## End(Not run)

```

scatterMatrix

scatterMatrix

Description

scatterMatrix produces a matrix with jittered scatterplots, histograms, and correlation coefficients.

Usage

```

scatterMatrix(
  dat,
  items = NULL,
  itemLabels = NULL,
  plotSize = 180,
  sizeMultiplier = 1,

```

```

    pointSize = 1,
    axisLabels = "none",
    normalHist = TRUE,
    progress = NULL,
    theme = ggplot2::theme_minimal(),
    hideGrid = TRUE,
    conf.level = 0.95,
    ...
)

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

Arguments

<code>dat</code>	A dataframe containing the items in the scale. All variables in this dataframe will be used if <code>items</code> is <code>NULL</code> .
<code>items</code>	If not <code>NULL</code> , this should be a character vector with the names of the variables in the dataframe that represent items in the scale.
<code>itemLabels</code>	Optionally, labels to use for the items (optionally, named, with the names corresponding to the <code>items</code> ; otherwise, the order of the labels has to match the order of the items)
<code>plotSize</code>	Size of the final plot in millimeters.
<code>sizeMultiplier</code>	Allows more flexible control over the size of the plot elements
<code>pointSize</code>	Size of the points in the scatterplots
<code>axisLabels</code>	Passed to <code>ggpairs</code> function to set <code>axisLabels</code> .
<code>normalHist</code>	Whether to use the default <code>ggpairs</code> histogram on the diagonal of the scattermatrix, or whether to use the <code>normalHist()</code> version.
<code>progress</code>	Whether to show a progress bar; set to <code>FALSE</code> to disable. See <code>GGally::ggpairs()</code> help for more information.
<code>theme</code>	The <code>ggplot2</code> theme to use.
<code>hideGrid</code>	Whether to hide the gridlines in the plot.
<code>conf.level</code>	The confidence level of confidence intervals
<code>...</code>	Additional arguments for <code>scatterMatrix()</code> are passed on to <code>normalHist()</code> , and additional arguments for the <code>print</code> method are passed on to the default <code>print</code> method.
<code>x</code>	The object to print.

Value

An object with the input and several output variables. Most notably:

`output$scatterMatrix`

A scattermatrix with histograms on the diagonal and correlation coefficients in the upper right half.

Author(s)

Gjalt-Jorn Peters

Maintainer: Gjalt-Jorn Peters gjalt-jorn@userfriendlyscience.com**Examples**

```
### Note: the 'not run' is simply because running takes a lot of time,
###       but these examples are all safe to run!
## Not run:

### Generate a datafile to use
exampleData <- data.frame(item1=rnorm(100));
exampleData$item2 <- exampleData$item1+rnorm(100);
exampleData$item3 <- exampleData$item1+rnorm(100);
exampleData$item4 <- exampleData$item2+rnorm(100);
exampleData$item5 <- exampleData$item2+rnorm(100);

### Use all items
scatterMatrix(dat=exampleData);

## End(Not run)
```

setCaptionNumberingKnitrHook

Set a knitr hook for caption numbering

Description

Set a knitr hook to automatically number captions for, e.g., figures and tables. `setCaptionNumberingKnitrHook()` is the general purpose function; you normally use `setFigCapNumbering()` or `setTabCapNumbering()`.

Usage

```
setCaptionNumberingKnitrHook(
  captionName = "fig.cap",
  prefix = "Figure %s: ",
  suffix = "",
  optionName = paste0("setCaptionNumbering_", captionName),
  resetCounterTo = 1
)

setFigCapNumbering(
  captionName = "fig.cap",
  prefix = "Figure %s: ",
  suffix = "",
  optionName = paste0("setCaptionNumbering_", captionName),
  resetCounterTo = 1
)
```

```

)

setTabCapNumbering(
    captionName = "tab.cap",
    prefix = "Table %s: ",
    suffix = "",
    optionName = paste0("setCaptionNumbering_", captionName),
    resetCounterTo = 1
)

```

Arguments

captionName	The name of the caption; for example, fig.cap or tab.cap.
prefix, suffix	The prefix and suffix; any occurrences of %s will be replaced by the number.
optionName	The name to use for the option that keeps track of the numbering.
resetCounterTo	Whether to reset the counter (as stored in the options), and if so, to what value (set to FALSE to prevent resetting).

Value

NULL, invisibly.

Examples

```

### To start automatically numbering figure captions
setFigCapNumbering();

### To start automatically numbering table captions
setTabCapNumbering();

```

sharedSubString	<i>sharedSubString</i>
-----------------	------------------------

Description

A function to find the longest shared substring in a character vector.

Usage

```
sharedSubString(x, y = NULL)
```

Arguments

x	The character vector to process.
y	Optionally, two single values can be specified. This is probably not useful to end users, but it's used by the function when it calls itself.

Value

A vector of length one with either the longest substring that occurs in all values of the character vector, or NA if no overlap can be found.

Author(s)

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Examples

```
sharedSubString(c("t0_responseTime", "t1_responseTime", "t2_responseTime"));  
### Returns "_responseTime"
```

simDataSet

Simulate a dataset

Description

simDataSet can be used to conveniently and quickly simulate a dataset that satisfies certain constraints, such as a specific correlation structure, means, ranges of the items, and measurement levels of the variables. Note that the results are approximate; mvnrm is used to generate the correlation matrix, but the factors are only created after that, so cutting the variable into factors may change the correlations a bit.

Usage

```
simDataSet(  
  n,  
  varNames,  
  correlations = c(0.1, 0.4),  
  specifiedCorrelations = NULL,  
  means = 0,  
  sds = 1,  
  ranges = c(1, 7),  
  factors = NULL,  
  cuts = NULL,  
  labels = NULL,  
  seed = 20160503,  
  empirical = TRUE,  
  silent = FALSE  
)
```

Arguments

<code>n</code>	Number of requires cases (records, entries, participants, rows) in the final dataset.
<code>varNames</code>	Names of the variables in a vector; note that the length of this vector will determine the number of variables simulated.
<code>correlations</code>	The correlations between the variables are randomly sampled from this range using the uniform distribution; this way, it's easy to have a relatively 'messy' correlation matrix without the need to specify every correlation manually.
<code>specifiedCorrelations</code>	The correlations that have to have a specific value can be specified here, as a list of vectors, where each vector's first two elements specify variables names, and the last one the correlation between those two variables. Note that tweaking the correlations may take some time; the <code>MASS::mvrnorm()</code> function will complain that "'Sigma' is not positive definite", or in other words, you supplied a combination of correlations that can't exist simultaneously, if you get it wrong.
<code>means, sds</code>	The means and standard deviations of the variables. Note that is you set ranges for one or more variables (see below), those ranges are used to rescale those variables, overriding any specified means and standard deviations. If only one mean or standard deviation is supplied, it's recycled along the variables.
<code>ranges</code>	The desired ranges of the variables, supplied as a named list where the name of each element corresponds to a variable. The <code>scales::rescale()</code> function will be used to rescale those variables for which a desired scale is specified here. Note that for those variables, the means and standard deviations will be determined by these new ranges.
<code>factors</code>	A vector of variable names that should be converted into factors (using <code>base::cut()</code>). Make sure to specify lists for cuts and labels as well (of the same length).
<code>cuts</code>	A list of vectors that specify, for each factor, where to 'cut' the numeric vector into factor levels.
<code>labels</code>	A list of vectors that specify, for each factor, and for each level, the labels that should be assigned to the factor levels. Each vector in this list has to have one more element than each vector in the cuts list.
<code>seed</code>	The seed to use when generating the dataset (to make sure the exact same dataset can be generated repeatedly).
<code>empirical</code>	Whether to generate the data using the exact <code>empirical = TRUE</code> or approximate (<code>empirical = FALSE</code>) correlation matrix; this is passed on to <code>MASS::mvrnorm()</code> .
<code>silent</code>	Whether to show intermediate and final descriptive information (correlation and covariance matrices as well as summaries).

Details

This function was intended to allow relatively quick generation of datasets that satisfy specific constraints, e.g. including a number of factors, variables with a specified minimum and maximum value or specified means and standard deviations, and of course specific correlations. Because all correlations except those specified are randomly generated from a uniform distribution, it's quite convenient to generate messy kind of real looking datasets quickly. Note that it's mostly a convenience function, and datasets will still require tweaking; for example, factors are simply numeric

vectors that are `cut()` after `MASS::mvrnorm()` generated the data, so the associations will change slightly.

Value

The generated dataframe is returned invisibly.

Examples

```
dat <- simDataSet(
  500,
  varNames=c('age',
             'sex',
             'educationLevel',
             'negativeLifeEventsInPast10Years',
             'problemCoping',
             'emotionCoping',
             'resilience',
             'depression'),
  means = c(40,
            0,
            0,
            5,
            3.5,
            3.5,
            3.5,
            3.5),
  sds = c(10,
         1,
         1,
         1.5,
         1.5,
         1.5,
         1.5,
         1.5),
  specifiedCorrelations =
    list(c('problemCoping', 'emotionCoping', -.5),
         c('problemCoping', 'resilience', .5),
         c('problemCoping', 'depression', -.4),
         c('depression', 'emotionCoping', .6),
         c('depression', 'resilience', -.3)),
  ranges = list(age = c(18, 54),
               negativeLifeEventsInPast10Years = c(0,8),
               problemCoping = c(1, 7),
               emotionCoping = c(1, 7)),
  factors=c("sex", "educationLevel"),
  cuts=list(c(0,
             c(-.5, .5)),
            c('female', 'male'),
            c('lower', 'middle', 'higher')),
  silent=FALSE);
```

spearmanBrown	<i>Spearman-Brown formula</i>
---------------	-------------------------------

Description

Spearman-Brown formula

Usage

```
spearmanBrown(nrOfItems, itemReliability)
spearmanBrown_reversed(nrOfItems, scaleReliability)
spearmanBrown_requiredLength(scaleReliability, itemReliability)
```

Arguments

nrOfItems	Number of items (or 'subtests') in the scale (or 'test').
itemReliability	The reliability of one item (or 'subtest').
scaleReliability	The reliability of the scale (or, desired reliability of the scale).

Value

For `spearmanBrown`, the predicted scale reliability; for `spearmanBrown_requiredLength`, the number of items required to achieve the desired scale reliability; and for `spearmanBrown_reversed`, the reliability of one item.

Examples

```
spearmanBrown(10, .4);
spearmanBrown_reversed(10, .87);
spearmanBrown_requiredLength(.87, .4);
```

strToFilename	<i>Convert a string to a safe filename</i>
---------------	--

Description

Convert a string to a safe filename

Usage

```
strToFilename(str, ext = NULL)
```

Arguments

`str` The string to convert.
`ext` Optionally, an extension to append.

Value

The string, processed to remove potentially problematic characters.

Examples

```
strToFilename("this contains: illegal characters, spaces, et cetera.");
```

`suspectParticipants` *Selects suspect participants from a `carelessObject`*

Description

This function is a wrapper for the `carelessObject()` function, which wraps a number of functions from the `careless` package. Normally, you'd probably call `carelessReport` which calls this function to generate a report of suspect participants.

Usage

```
suspectParticipants(  
  carelessObject,  
  nFlags = 1,  
  digits = 2,  
  missingSymbol = "Missing"  
)
```

Arguments

`carelessObject` The result of the call to `carelessObject()`.
`nFlags` The number of flags required to be considered suspect.
`digits` The number of digits to round to.
`missingSymbol` How to represent missing values.

Value

A logical vector.

Examples

```
suspectParticipants(carelessObject(mtcars),  
  nFlags = 2);
```

testRetestAlpha	<i>Test-Retest Alpha Coefficient</i>
-----------------	--------------------------------------

Description

The testRetestAlpha function computes the test-retest alpha coefficient (Green, 2003).

Usage

```
testRetestAlpha(  
  dat = NULL,  
  moments = NULL,  
  testDat = NULL,  
  retestDat = NULL,  
  sortItems = FALSE,  
  convertToNumeric = TRUE  
)  
  
## S3 method for class 'testRetestAlpha'  
print(x, ...)
```

Arguments

dat	A dataframe containing the items in the scale at both measurement moments. If no dataframe is specified, a dialogue will be launched to allow the user to select an SPSS datafile. If only one dataframe is specified, either the items have to be ordered chronologically (i.e. first all items for the first measurement, then all items for the second measurement), or the vector 'moments' has to be used to indicate, for each item, to which measurement moment it belongs.
moments	Used to indicate to which measurement moment each item in 'dat' belongs; should be a vector with the same length as dat has columns, and with two possible values (e.g. 1 and 2).
testDat, retestDat	Dataframes with the items for each measurement moment: note that the items have to be in the same order (unless sortItems is TRUE).
sortItems	If true, the columns (items) in each dataframe are ordered alphabetically before starting. This can be convenient to ensure that the order of the items at each measurement moment is the same.
convertToNumeric	When TRUE, the function will attempt to convert all vectors in the dataframes to numeric.
x	The object to print
...	Ignored.

Value

An object with the input and several output variables. Most notably:

input	Input specified when calling the function
intermediate	Intermediate values and objects computed to get to the final results
output\$testRetestAlpha	The value of the test-retest alpha coefficient.

References

Green, S. N. (2003). A Coefficient Alpha for Test-Retest Data. *Psychological Methods*, 8(1), 88-101. doi:[10/bxq9r4](https://doi.org/10.1037/1082-989X.8.1.88)

Examples

```
## Not run:
### This will prompt the user to select an SPSS file
testRetestAlpha();

## End(Not run)

### Load data from simulated dataset testRetestSimData (which
### satisfies essential tau-equivalence).
data(testRetestSimData);

### The first column is the true score, so it's excluded in this example.
exampleData <- testRetestSimData[, 2:ncol(testRetestSimData)];

### Compute test-retest alpha coefficient
testRetestAlpha(exampleData);
```

testRetestCES

Test-Retest Coefficient of Equivalence & Stability

Description

The testRetestCES function computes the test-retest Coefficient of Equivalence and Stability (Schmidt, Le & Ilies, 2003).

Usage

```
testRetestCES(
  dat = NULL,
  moments = NULL,
  testDat = NULL,
  retestDat = NULL,
  parallelTests = "means",
```

```

    sortItems = FALSE,
    convertToNumeric = TRUE,
    digits = 4
)

## S3 method for class 'testRetestCES'
print(x, digits = x$input$digits, ...)

```

Arguments

<code>dat</code>	A dataframe. For <code>testRetestCES</code> , this dataframe must contain the items in the scale at both measurement moments. If no dataframe is specified, a dialogue will be launched to allow the user to select an SPSS datafile. If only one dataframe is specified, either the items have to be ordered chronologically (i.e. first all items for the first measurement, then all items for the second measurement), or the vector <code>'moments'</code> has to be used to indicate, for each item, to which measurement moment it belongs. The number of columns in this dataframe MUST be even! Note that instead of providing this dataframe, the items of each measurement moment can be provided separately in <code>testDat</code> and <code>retestDat</code> as well.
<code>moments</code>	Used to indicate to which measurement moment each item in <code>'dat'</code> belongs; should be a vector with the same length as <code>dat</code> has columns, and with two possible values (e.g. 1 and 2).
<code>testDat, retestDat</code>	Dataframes with the items for each measurement moment: note that the items have to be in the same order (unless <code>sortItems</code> is <code>TRUE</code>).
<code>parallelTests</code>	A vector indicating which items belong to which parallel test; like the <code>moments</code> vector, this should have two possible values (e.g. 1 and 2). Alternatively, it can be character value with <code>'means'</code> or <code>'variances'</code> ; in this case, <code>parallelSubscales</code> will be used to create roughly parallel halves.
<code>sortItems</code>	If true, the columns (items) in each dataframe are ordered alphabetically before starting. This can be convenient to ensure that the order of the items at each measurement moment is the same.
<code>convertToNumeric</code>	When <code>TRUE</code> , the function will attempt to convert all vectors in the dataframes to numeric.
<code>digits</code>	Number of digits to print.
<code>x</code>	The object to print
<code>...</code>	Ignored.

Details

This function computes the test-retest Coefficient of Equivalence and Stability (CES) as described in Schmidt, Le & Ilies (2003). Note that this function only computes the test-retest CES for a scale that is administered twice and split into two parallel halves post-hoc (this procedure is explained on page 210, and the equations that are used, 16 and 17a are explained on page 212).

Value

An object with the input and several output variables. Most notably:

input	Input specified when calling the function
intermediate	Intermediate values and objects computed to get to the final results
output\$testRetestCES	The value of the test-retest Coefficient of Equivalence and Stability.

Note

This function uses equations 16 and 17 on page 212 of Schmidt, Le & Ilies (2003): in other words, this function assumes that one scale is administered twice. If you'd like the computation for two different but parallel scales/measures to be implemented, please contact me.

References

Schmidt, F. L., Le, H., & Ilies, R. (2003) Beyond Alpha: An Empirical Examination of the Effects of Different Sources of Measurement Error on Reliability Estimates for Measures of Individual-differences Constructs. *Psychological Methods*, 8(2), 206-224. doi:[10/dzmk7n](https://doi.org/10.1037/1082-989X.8.2.206)

Examples

```
## Not run:
### This will prompt the user to select an SPSS file
testRetestCES();

## End(Not run)

### Load data from simulated dataset testRetestSimData (which
### satisfies essential tau-equivalence).
data(testRetestSimData);

### The first column is the true score, so it's excluded in this example.
exampleData <- testRetestSimData[, 2:ncol(testRetestSimData)];

### Compute test-retest alpha coefficient
testRetestCES(exampleData);
```

testRetestReliability *testRetestReliability*

Description

The testRetestReliability function is a convenient interface to testRetestAlpha and testRetestCES.

Usage

```
testRetestReliability(
  dat = NULL,
  moments = NULL,
  testDat = NULL,
  retestDat = NULL,
  parallelTests = "means",
  sortItems = FALSE,
  convertToNumeric = TRUE,
  digits = 2
)

## S3 method for class 'testRetestReliability'
print(x, digits = x$input$digits, ...)
```

Arguments

<code>dat</code>	A dataframe. This dataframe must contain the items in the scale at both measurement moments. If no dataframe is specified, a dialogue will be launched to allow the user to select an SPSS datafile. If only one dataframe is specified, either the items have to be ordered chronologically (i.e. first all items for the first measurement, then all items for the second measurement), or the vector 'moments' has to be used to indicate, for each item, to which measurement moment it belongs. The number of columns in this dataframe MUST be even! Note that instead of providing this dataframe, the items of each measurement moment can be provided separately in <code>testDat</code> and <code>retestDat</code> as well.
<code>moments</code>	Used to indicate to which measurement moment each item in 'dat' belongs; should be a vector with the same length as <code>dat</code> has columns, and with two possible values (e.g. 1 and 2).
<code>testDat, retestDat</code>	Dataframes with the items for each measurement moment: note that the items have to be in the same order (unless <code>sortItems</code> is TRUE).
<code>parallelTests</code>	A vector indicating which items belong to which parallel test; like the <code>moments</code> vector, this should have two possible values (e.g. 1 and 2). Alternatively, it can be character value with 'means' or 'variances'; in this case, <code>parallelSubscales</code> will be used to create roughly parallel halves.
<code>sortItems</code>	If true, the columns (items) in each dataframe are ordered alphabetically before starting. This can be convenient to ensure that the order of the items at each measurement moment is the same.
<code>convertToNumeric</code>	When TRUE, the function will attempt to convert all vectors in the dataframes to numeric.
<code>digits</code>	Number of digits to show when printing the output
<code>x</code>	The object to print
<code>...</code>	Passed on to the print function

Details

This function calls both testRetestAlpha and testRetestCES to compute and print measures of the test-retest reliability.

Value

An object with the input and several output variables. Most notably:

input	Input specified when calling the function
intermediate	Intermediate values and objects computed to get to the final results
output\$testRetestAlpha	The value of the test-retest alpha coefficient.
output\$testRetestCES	The value of the test-retest Coefficient of Equivalence and Stability.

Examples

```
## Not run:
### This will prompt the user to select an SPSS file
testRetestReliability();

## End(Not run)

### Load data from simulated dataset testRetestSimData (which
### satisfies essential tau-equivalence).
data(testRetestSimData);

### The first column is the true score, so it's excluded in this example.
exampleData <- testRetestSimData[, 2:ncol(testRetestSimData)];

### Compute test-retest alpha coefficient
ufs::testRetestReliability(exampleData);
```

testRetestSimData	<i>testRetestSimData is a simulated dataframe used to demonstrate the testRetestAlpha coefficient function.</i>
-------------------	---

Description

This dataset contains the true scores of 250 participants on some variable, and 10 items of a scale administered twice (at t0 and at t1).

Format

A data frame with 250 observations on the following 21 variables.

trueScore The true scores
t0_item1 Score on item 1 at test
t0_item2 Score on item 2 at test
t0_item3 Score on item 3 at test
t0_item4 Score on item 4 at test
t0_item5 Score on item 5 at test
t0_item6 Score on item 6 at test
t0_item7 Score on item 7 at test
t0_item8 Score on item 8 at test
t0_item9 Score on item 9 at test
t0_item10 Score on item 10 at test
t1_item1 Score on item 1 at retest
t1_item2 Score on item 2 at retest
t1_item3 Score on item 3 at retest
t1_item4 Score on item 4 at retest
t1_item5 Score on item 5 at retest
t1_item6 Score on item 6 at retest
t1_item7 Score on item 7 at retest
t1_item8 Score on item 8 at retest
t1_item9 Score on item 9 at retest
t1_item10 Score on item 10 at retest

Details

This dataset was generated with the code in the reliabilityTest.r test script.

Author(s)

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Examples

```
data(testRetestSimData);  
head(testRetestSimData);  
hist(testRetestSimData$t0_item1);  
cor(testRetestSimData);
```

vecTxt

*Easily parse a vector into a character value***Description**

vecTxtQ, vecTxtB, and vecTxtM are convenience functions with default quotes that can be useful when working in R Markdown documents.

Usage

```
vecTxt(
  vector,
  delimiter = ", ",
  useQuote = "",
  firstDelimiter = NULL,
  lastDelimiter = " & ",
  firstElements = 0,
  lastElements = 1,
  lastHasPrecedence = TRUE
)

vecTxtQ(vector, useQuote = "'", ...)

vecTxtB(vector, useQuote = "`", ...)

vecTxtM(vector, useQuote = "$", ...)
```

Arguments

vector	The vector to process.
delimiter, firstDelimiter, lastDelimiter	The delimiters to use for respectively the middle, first firstElements, and last lastElements elements.
useQuote	This character string is pre- and appended to all elements; so use this to quote all elements (useQuote="'), doublequote all elements (useQuote=''), or anything else (e.g. useQuote=' '). The only difference between vecTxt and vecTxtQ is that the latter by default quotes the elements.
firstElements, lastElements	The number of elements for which to use the first respective last delimiters
lastHasPrecedence	If the vector is very short, it's possible that the sum of firstElements and lastElements is larger than the vector length. In that case, downwardly adjust the number of elements to separate with the first delimiter (TRUE) or the number of elements to separate with the last delimiter (FALSE)?
...	Any addition arguments to vecTxtQ are passed on to vecTxt.

Value

A character vector of length 1.

Examples

```
vecTxtQ(names(mtcars));
```

viridisPalette	<i>Convenience function to get 2-7 color viridis palettes</i>
----------------	---

Description

This function only exists to avoid importing the viridis package.

Usage

```
viridisPalette(x)
```

Arguments

x The number of colors you want (seven at most).

Value

A vector of colours.

wrapVector	<i>Wrap all elements in a vector</i>
------------	--------------------------------------

Description

Wrap all elements in a vector

Usage

```
wrapVector(x, width = 0.9 * getOption("width"), sep = "\n", ...)
```

Arguments

x The character vector
width The number of
sep The glue with which to combine the new lines
... Other arguments are passed to [strwrap\(\)](#).

Value

A character vector

Examples

```
res <- wrapVector(
  c(
    "This is a sentence ready for wrapping",
    "So is this one, although it's a bit longer"
  ),
  width = 10
);

print(res);
cat(res, sep="\n");
```

zotero_construct_export_call

Construct the URL for a Zotero export call

Description

This function is just a convenience function to create a simple URL to download references from a public Zotero group. See https://www.zotero.org/support/dev/web_api/v3/start for details.

Usage

```
zotero_construct_export_call(
  group,
  sort = "dateAdded",
  direction = "asc",
  format = "bibtex",
  start = 0,
  limit = 100
)
```

Arguments

group	The group ID
sort	On which field to sort
direction	The direction to sort in
format	The format to export
start	The index of the first record to return
limit	The number of records to return

Value

The URL in a character vector.

Examples

```
zotero_construct_export_call(2425237);
```

```
zotero_download_and_export_items
```

Download and save all items in a public Zotero group

Description

Download and save all items in a public Zotero group

Usage

```
zotero_download_and_export_items(  
  group,  
  file,  
  format = "bibtex",  
  showKeys = TRUE  
)
```

Arguments

group	The group ID
file	The filename to write to
format	The format to export
showKeys	Whether to show the keys

Value

The bibliography as a character vector

Examples

```
## Not run:  
tmpFile <- tempfile(fileext=".bib");  
zotero_download_and_export_items(  
  2425237,  
  tmpFile  
)  
writtenBibliography <- readLines(tmpFile);  
writtenBibliography[1:7];  
  
## End(Not run)
```

zotero_get_all_items	<i>Get all items in a public Zotero group</i>
----------------------	---

Description

Get all items in a public Zotero group

Usage

```
zotero_get_all_items(group, format = "bibtex")
```

Arguments

group	The group ID
format	The format to export

Value

A character vector

Examples

```
zotero_get_all_items(2425237);
```

zotero_nr_of_items	<i>Get number of items in a public Zotero group</i>
--------------------	---

Description

Get number of items in a public Zotero group

Usage

```
zotero_nr_of_items(group)
```

Arguments

group	The group ID
-------	--------------

Value

The umber of items as a numeric vector.

Examples

```
zotero_nr_of_items(2425237);
```

%IN%	<i>Case insensitive version of %in%</i>
------	---

Description

This is simply 'in', but applies `base::toupper()` to both arguments, first.

Usage

```
find %IN% table
```

Arguments

- | | |
|-------|--|
| find | The element(s) to look up in the vector or matrix. |
| table | The vector or matrix in which to look up the element(s). |

Value

A logical vector.

Examples

```
letters[1:4] %IN% LETTERS
```


Index

- * **bivar**
 - computeStatistic_t, 32
 - confIntOmegaSq, 35
 - cramersV, 44
- * **character**
 - escapeRegex, 62
 - sharedSubString, 134
- * **datasets**
 - opts, 111
 - testRetestSimData, 145
- * **data**
 - bfi-data, 17
- * **file**
 - getData, 76
- * **graphs**
 - ggProportionPlot, 82
- * **hplot**
 - associationsDiamondPlot, 10
 - biAxisDiamondPlot, 18
 - diamondCoordinates, 51
 - diamondPlot, 54
 - duoComparisonDiamondPlot, 58
 - factorLoadingDiamondCIplot, 68
 - factorLoadingHeatmap, 70
 - ggBoxplot, 78
 - ggEasyBar, 79
 - ggProportionPlot, 82
 - ggqq, 85
 - meansDiamondPlot, 101
 - meanSDtoDiamondPlot, 104
- * **htest**
 - confIntProp, 36
 - confIntR, 37
 - pwr.confIntR, 116
 - pwr.omegasq, 117
- * **manip**
 - escapeRegex, 62
- * **programming**
 - escapeRegex, 62
- * **univariate**
 - descr, 49
 - invertItem, 91
 - iqrOutlier, 92
- * **univar**
 - associationMatrix, 7
 - cohensdCI, 29
 - confIntProp, 36
 - getData, 76
 - pomegaSq, 113
 - scaleDiagnosis, 124
 - scaleStructure, 127
 - scatterMatrix, 131
 - testRetestCES, 141
- * **utilities**
 - areColors, 6
 - associationMatrix, 7
 - checkDataIntegrity, 24
 - computeStatistic_t, 32
 - convert, 40
 - convert.cer.to.d, 41
 - dataShape, 45
 - findShortestInterval, 72
 - getData, 76
 - knitFig, 97
 - multiResponse, 106
 - multiVarFreq, 108
 - normalHist, 109
 - scaleDiagnosis, 124
 - scaleStructure, 127
 - scatterMatrix, 131
 - testRetestCES, 141
- * **utils**
 - extractVarName, 66
- %IN%, 152
- A_VarghaDelaney, 15
- aipedjmv, 4
- aiperjmv, 5
- areColors, 6

- arr, 6
- as.data.frame, 108
- as.data.frame(), 50
- as.data.frame.descr(descr), 49
- associationMatrix, 7
- associationMatrixESDefaults
(computeStatistic_t), 32
- associationMatrixStatDefaults
(computeStatistic_t), 32
- associationsDiamondPlot, 10
- associationsDiamondPlot(), 20, 61
- associationsToDiamondPlotDf
(associationsDiamondPlot), 10
- attenuate.d, 13
- attenuate.r, 14

- BAC_plot, 15
- base::cat(), 98
- base::cut(), 136
- base::sprintf(), 90
- base::toupper(), 152
- bfi(bfi-data), 17
- bfi-data, 17
- biAxisDiamondPlot, 18
- biDimColors, 20
- binom.test(), 36, 37, 84
- bootES::bootES(), 114

- carelessObject, 21
- carelessObject(), 139
- carelessReport, 22
- cat, 23, 98
- cat(), 97, 98
- cat0, 23
- checkDataIntegrity, 24
- checkPkgs, 25
- CIM, 27
- CIM_partial(CIM), 27
- cohensdCI, 29
- cohensDdistribution(cohensdCI), 29
- col2rgb(), 28, 53
- colnames(), 37
- colorRamp(), 28, 53
- Comparison, 24
- computeEffectSize_d
(computeStatistic_t), 32
- computeEffectSize_etasq
(computeStatistic_t), 32
- computeEffectSize_omegasq
(computeStatistic_t), 32
- computeEffectSize_r
(computeStatistic_t), 32
- computeEffectSize_v
(computeStatistic_t), 32
- computeStatistic_chisq
(computeStatistic_t), 32
- computeStatistic_f
(computeStatistic_t), 32
- computeStatistic_r
(computeStatistic_t), 32
- computeStatistic_t, 32
- confIntD(cohensdCI), 29
- confIntDjmv, 34
- confIntOmegaSq, 35
- confIntProp, 36
- confIntProp(), 84, 115
- confIntR, 37
- confIntR(), 38
- confintrjmv, 38
- confIntSD, 39
- confIntV(cramersV), 44
- convert, 40
- convert.cer.to.d, 41
- convert.cohensf.to.omegasq, 117, 119
- convert.d.to.eer(convert.cer.to.d), 41
- convert.d.to.nnc(convert.cer.to.d), 41
- convert.d.to.t(), 31
- convert.d.to.U3, 43
- convert.eer.to.d(convert.cer.to.d), 41
- convert.f.to.omegasq(), 114
- convert.ncf.to.omegasq(), 35, 36, 113
- convert.omegasq.to.cohensf, 117, 119
- convert.omegasq.to.f(), 114
- convert.t.to.d(), 31
- convertToNumeric, 43
- cramersV, 44
- cut(), 137

- dataShape, 45
- dCohensd(cohensdCI), 29
- dd(cohensdCI), 29
- descr, 49
- descriptives(descr), 49
- df(), 114
- diamondCoordinates, 51
- diamondPlot, 54

diamondPlot(), [12](#), [13](#), [19](#), [51](#), [54](#), [61](#), [62](#),
 [68–71](#), [102](#), [103](#), [105](#)
 digest::digest(), [98](#)
 diptest::dip.test(), [51](#)
 disattenuate.d, [57](#)
 disattenuate.r, [58](#)
 dnorm, [86](#)
 domegaSq (pomegaSq), [113](#)
 dt(), [31](#)
 duoComparisonDiamondPlot, [58](#)

 escapeBS (escapeRegex), [62](#)
 escapeRegex, [62](#)
 exceptionalScore, [63](#)
 exceptionalScore(), [65](#)
 exceptionalScores, [64](#)
 exceptionalScores(), [64](#)
 exportToHTML, [66](#)
 extractVarName, [66](#)

 fa_failsafe, [72](#)
 faConfInt, [67](#)
 factorLoadingDiamondCIplot, [68](#)
 factorLoadingDiamondCIplot(), [51](#), [54](#), [56](#),
 [103](#), [105](#)
 factorLoadingHeatmap, [70](#)
 findShortestInterval, [72](#)
 formatCI, [73](#)
 formatCI(), [74](#), [75](#), [111](#)
 formatPvalue, [74](#)
 formatPvalue(), [74](#), [75](#), [111](#)
 formatR, [75](#)
 formatR(), [74](#), [111](#)

 geom_bar(), [81](#)
 geom_boxplot, [78](#), [79](#)
 geom_jitter(), [53](#), [54](#)
 geom_polygon(), [53](#), [54](#)
 geom_ridgeline(), [81](#)
 get (opts), [111](#)
 getDat (getData), [76](#)
 getData, [76](#), [128](#)
 GGally::ggpairs(), [132](#)
 ggBarChart, [77](#)
 ggBoxplot, [78](#)
 ggDiamondLayer (diamondCoordinates), [51](#)
 ggDiamondLayer(), [12](#), [13](#), [54](#), [56](#), [69](#), [71](#),
 [102](#), [103](#), [105](#)
 ggEasyBar, [79](#)
 ggEasyPlots (ggEasyBar), [79](#)
 ggEasyRidge (ggEasyBar), [79](#)
 ggPie, [81](#)
 ggplot, [78](#), [79](#), [86](#), [98](#)
 ggplot(), [12](#), [19](#), [54](#), [55](#), [80](#), [81](#), [102](#), [105](#)
 ggplot2(), [83](#), [84](#)
 ggplot2::geom_bar(), [77](#), [78](#)
 ggplot2::ggplot(), [28](#), [56](#), [61](#), [69](#), [71](#), [77](#),
 [78](#), [87](#), [93](#), [110](#)
 ggplot2::ggsave(), [87](#)
 ggProportionPlot, [37](#), [82](#)
 ggqq, [85](#)
 ggSave, [87](#)
 ggSave(), [97](#)
 grep, [63](#)
 grid.draw.ggProportionPlot
 (ggProportionPlot), [82](#)
 gtable(), [61](#)
 gtable::gtable(), [20](#)

 heading, [88](#)

 ifelseObj, [89](#)
 insertFigureCaption, [89](#)
 insertNumberedCaption
 (insertFigureCaption), [89](#)
 insertTableCaption
 (insertFigureCaption), [89](#)
 invertItem, [91](#), [91](#)
 invertItems (invertItem), [91](#)
 IQR, [92](#)
 iqrOutlier, [92](#)
 irpplot, [92](#)
 is.even (is.odd), [94](#)
 is.nr, [93](#)
 is.odd, [94](#)
 isTrue, [95](#)

 kblXtra, [95](#)
 knit, [98](#)
 knit_expand, [98](#)
 knit_print.CIM (CIM), [27](#)
 knit_print.scaleDiagnosis
 (scaleDiagnosis), [124](#)
 knit_print.scaleStructure
 (scaleStructure), [127](#)
 knitAndSave, [96](#)
 knitFig, [97](#)
 knitFig(), [97](#)

knitr::asis_output(), 98
 knitr::kable(), 96
 knitr::knit(), 124, 129

 library(), 26
 linetype(), 83

 makeScales, 98
 MASS::mvrnorm(), 136, 137
 massConvertToNumeric, 99
 MBESS::ci.reliability(), 130
 meanConfInt, 100
 meansComparisonDiamondPlot
 (duoComparisonDiamondPlot), 58
 meansDiamondPlot, 101
 meansDiamondPlot(), 18, 20, 51, 54, 56, 62,
 69, 71, 105
 meansDiamondPlotjmv, 103
 meansSDtoDiamondPlot, 104
 meansSDtoDiamondPlot(), 51, 54, 56, 69, 71,
 103
 multiResponse, 106
 multiResponsejmv, 107
 multiVarFreq, 108

 normalHist, 109
 normalHist(), 125, 132
 normalityAssessment, 86
 normalityAssessment (dataShape), 45
 noZero, 74, 111
 noZero(), 74, 75

 opts, 111

 pandar.associationMatrix
 (associationMatrix), 7
 pandar.dataShape (dataShape), 45
 pandar.descr (descr), 49
 pandar.normalityAssessment (dataShape),
 45
 pandar::pandar(), 9
 parallelSubscales, 112
 pbeta(), 36
 pCohensd (cohensdCI), 29
 pd (cohensdCI), 29
 pdExtreme (cohensdCI), 29
 pdInterval (cohensdCI), 29
 pdMild (cohensdCI), 29
 pf(), 114

 pnorm(), 43
 pomegaSq, 113
 print(), 9
 print.associationMatrix
 (associationMatrix), 7
 print.confIntOmegaSq (confIntOmegaSq),
 35
 print.confIntV (cramersV), 44
 print.CramersV (cramersV), 44
 print.dataShape (dataShape), 45
 print.descr (descr), 49
 print.ggProportionPlot
 (ggProportionPlot), 82
 print.meanConfInt (meanConfInt), 100
 print.normalHist (normalHist), 109
 print.normalityAssessment (dataShape),
 45
 print.parallelSubscales
 (parallelSubscales), 112
 print.pwr.omegasq (pwr.omegasq), 117
 print.regrInfluential
 (regrInfluential), 122
 print.scaleDiagnosis (scaleDiagnosis),
 124
 print.scaleStructure (scaleStructure),
 127
 print.scatterMatrix (scatterMatrix), 131
 print.testRetestAlpha
 (testRetestAlpha), 140
 print.testRetestCES (testRetestCES), 141
 print.testRetestReliability
 (testRetestReliability), 143
 print.ufsARR (arr), 6
 psych::alpha(), 130
 psych::describe(), 51
 psych::fa(), 28, 67–71
 psych::ICC(), 17
 psych::omega(), 129, 130
 psych::psych, 51, 68, 70
 psych::psych-package, 67
 pt(), 31
 pwr, 117, 118
 pwr.anova.test, 117, 119
 pwr.bootES, 114
 pwr.cohensdCI (cohensdCI), 29
 pwr.confIntd (cohensdCI), 29
 pwr.confIntProp, 115
 pwr.confIntR, 116, 117

- pwr.omegasq, [117](#)
- qCohensd (cohendsdCI), [29](#)
- qd (cohendsdCI), [29](#)
- qf(), [114](#)
- qnorm, [86](#)
- qomegaSq (pomegaSq), [113](#)
- qt(), [30](#), [31](#)
- quantile, [50](#)
- quietGitLabUpdate
 - (quietRemotesInstall), [119](#)
- quietRemotesInstall, [119](#)
- qVec, [120](#)
- qVecSum (qVec), [120](#)
- rawDataDiamondLayer
 - (diamondCoordinates), [51](#)
- rbind_df_list, [121](#)
- rbind_dfs, [121](#)
- rCohensd (cohendsdCI), [29](#)
- rd (cohendsdCI), [29](#)
- regrInfluential, [122](#)
- rep(), [87](#)
- repeatStr, [123](#)
- report, [123](#)
- repStr (repeatStr), [123](#)
- require(), [26](#)
- reset (opts), [111](#)
- rf(), [114](#)
- romeqSq (pomeqSq), [113](#)
- rownames(), [37](#)
- rt(), [31](#)
- safeRequire, [124](#)
- samplingDistribution (dataShape), [45](#)
- scaleDiagnosis, [124](#)
- scaleDiagnosis_partial
 - (scaleDiagnosis), [124](#)
- scaleReliability (scaleStructure), [127](#)
- scales::rescale(), [136](#)
- scaleStructure, [127](#)
- scaleStructure(), [125](#)
- scaleStructure_partial
 - (scaleStructure), [127](#)
- scaleStructurePartial (scaleStructure), [127](#)
- scatterMatrix, [131](#)
- scatterMatrix(), [126](#)
- set (opts), [111](#)
- setCaptionNumberingKnitrHook, [133](#)
- setFigCapNumbering
 - (setCaptionNumberingKnitrHook), [133](#)
- setFigCapNumbering(), [89](#)
- setTabCapNumbering
 - (setCaptionNumberingKnitrHook), [133](#)
- setTabCapNumbering(), [89](#)
- sharedSubString, [134](#)
- simDataSet, [135](#)
- spearmanBrown, [138](#)
- spearmanBrown_requiredLength
 - (spearmanBrown), [138](#)
- spearmanBrown_reversed (spearmanBrown), [138](#)
- stats::quantile(), [64](#)
- stats::wilcox.test(), [15](#)
- strToFilename, [138](#)
- strwrap(), [148](#)
- summary, [51](#)
- suspectParticipants, [139](#)
- Sys.time, [98](#)
- table(), [108](#)
- testRetestAlpha, [140](#)
- testRetestCES, [141](#)
- testRetestReliability, [143](#)
- testRetestSimData, [145](#)
- uniDimColors (biDimColors), [20](#)
- varsToDiamondPlotDf
 - (diamondCoordinates), [51](#)
- vecTxt, [147](#)
- vecTxtB (vecTxt), [147](#)
- vecTxtM (vecTxt), [147](#)
- vecTxtQ (vecTxt), [147](#)
- viridisPalette, [148](#)
- wrapVector, [148](#)
- zotero_construct_export_call, [149](#)
- zotero_download_and_export_items, [150](#)
- zotero_get_all_items, [151](#)
- zotero_nr_of_items, [151](#)