

Package ‘scipub’

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Title Summarize Data for Scientific Publication

Version 1.2.1

Description Create and format tables and APA statistics for scientific publication. This includes making a "Table 1" to summarize demographics across groups, correlation tables with significance indicated by stars, and extracting formatted statistical summaries from simple tests for in-text notation. The package also includes functions for Winsorizing data based on a Z-statistic cutoff.

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Encoding UTF-8

LazyData true

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Language en-US

VignetteBuilder knitr

URL <https://github.com/dpagliaccio/scipub>,
<https://dpagliaccio.github.io/scipub/>

BugReports <https://github.com/dpagliaccio/scipub/issues>

Depends R (>= 3.6)

Imports dplyr, forcats, ggplot2, gghalves, purrr, stats, stringr,
tibble, tidyr, tidyselect

Suggests spelling, htmlTable, knitr, rmarkdown

NeedsCompilation no

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apastat	<i>Format simple statistic test results for scientific publication</i>
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Description

The `apastat` function summarizes statistic test results scientific publication. This currently will take `stats::t.test`, `stats::cor.test`, or `stats::lm` results as input. The output is intended to be included as in-text parenthetical statistics in publication.

Usage

```
apastat(test, roundN = 2, es = c(TRUE, FALSE), ci = c(TRUE, FALSE), var = NULL)
```

Arguments

<code>test</code>	The <code>stats::t.test</code> , <code>stats::cor.test</code> , or <code>stats::lm</code> object to be formatted.
<code>roundN</code>	The number of decimal places to round all output to (default=2).
<code>es</code>	Include effect side (Cohen's d for t-test or 2-level factor lm variable), default to TRUE.
<code>ci</code>	Include confidence interval of estimate, default to TRUE.
<code>var</code>	Only for lm object, select name of variable to summarize (default=NULL), if NULL, will summarize overall model fit.

Value

Output formatted statistics

Examples

```
apastat(stats::cor.test(psydat$Age, psydat$Height))
apastat(stats::t.test(Height ~ Sex, data = psydat))
apastat(stats::lm(data = psydat, Height ~ Age + Sex))
apastat(stats::lm(data = psydat, Height ~ Age + Sex), var = "Age")
```

correltable	<i>Create correlation table (with stars for significance) for scientific publication</i>
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Description

The `correltable` function can be used to create correlation table (with stars for significance) for scientific publication. This is intended to summarize correlations between (vars) from an input dataset (data). Correlations are based on `stats::cor`, use and method follow from that function. Stars indicate significance: *p<.05, **p<.01, ***p<.001. For formatting, variables can be renamed, numbers can be rounded, upper or lower triangle only can be selected (or whole matrix), and empty columns/rows can be dropped if using triangles. For more compact columns, variable names can be numbered in the rows and column names will be corresponding numbers. If only cross-correlation between two sets of variables is desired (no correlations within a set of variables), `vars2` and `var_names` can be used. This function will drop any non-numeric variables by default. Requires `tidyverse` and `stats` libraries.

Usage

```
correltable(
  data,
  vars = NULL,
  var_names = vars,
  vars2 = NULL,
  var_names2 = vars2,
  method = c("pearson", "spearman"),
  use = c("pairwise", "complete"),
  round_n = 2,
  tri = c("upper", "lower", "all"),
  cutempty = c(FALSE, TRUE),
  colnum = c(FALSE, TRUE),
  html = c(FALSE, TRUE)
)
```

Arguments

<code>data</code>	The input dataset.
<code>vars</code>	A list of the names of variables to correlate, e.g. <code>c("Age", "height", "WASI")</code> , if <code>NULL</code> , all variables in <code>data</code> will be used.
<code>var_names</code>	An optional list to rename the <code>vars</code> colnames in the output table, e.g. <code>c("Age (years)", "Height (inches)", "IQ")</code> . Must match <code>vars</code> in length. If not supplied, <code>vars</code> will be printed as is.
<code>vars2</code>	If cross-correlation between two sets of variables is desired, add a second list of variables to correlate with <code>vars</code> ; Overrides <code>tri</code> , <code>cutempty</code> , and <code>colnum</code> .
<code>var_names2</code>	An optional list to rename the <code>vars2</code> colnames in the output table. If not supplied, <code>vars2</code> will be printed as is.

method	Type of correlation to calculate c("pearson", "spearman"), based on stats::cor, default = "pearson".
use	Use pairwise.complete.obs or restrict to complete cases c("pairwise", "complete"), based on stats::cor, default = "pairwise".
round_n	The number of decimal places to round all output to (default=2).
tri	Select output formatting c("upper", "lower","all"); KEEP the upper triangle, lower triangle, or all values, default="upper.
cutempty	If keeping only upper/lower triangle with tri, cut empty row/column, default=FALSE.
colnum	For more concise column names, number row names and just use corresponding numbers as column names, default=FALSE, if TRUE overrides cutempty.
html	Format as html in viewer or not (default=F, print in console), needs library(htmlTable) installed.

Value

Output Table 1

Examples

```

correltable(data = psydat)
correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  tri = "lower", html = TRUE
)
correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  var_names = c("Age (months)", "Height (inches)", "IQ"),
  tri = "upper", colnum = TRUE, html = TRUE
)
correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  var_names = c("Age (months)", "Height (inches)", "IQ"),
  vars2 = c("depressT", "anxT"),
  var_names2 = c("Depression T", "Anxiety T"), html = TRUE
)

```

FullTable1

Create Table1 of group summary with stats for scientific publication

Description

The FullTable1 function can be used to create a Table1 for scientific publication. This is intended to summarize demographic and other variables (vars) split by a grouping variable (strata) from an input dataset (data). Continuous variables will be summarized as mean (SD) and tested across groups using t-test or ANOVA (for 3+ level strata). Categorical variables will be summarized as N (%) and tested across groups as chi-squared. Effect sizes for group differences will be calculated as Cohen's d, partial eta-squared, Odds Ratio, Cramer's V depending on the test. Requires tidyverse and stats libraries.

Usage

```
FullTable1(
  data,
  strata = NULL,
  vars = NULL,
  var_names = vars,
  factor_vars = NULL,
  round_n = 2,
  es_col = c(TRUE, FALSE),
  p_col = c(TRUE, FALSE),
  stars = c("col", "name", "stat", "none"),
  html = c(FALSE, TRUE)
)
```

Arguments

<code>data</code>	The input dataset (will be converted to tibble).
<code>strata</code>	The grouping variable of interest (converted to factor), if NULL will make one column table.
<code>vars</code>	A list of variables to summarize, e.g. <code>c("Age", "sex", "WASI")</code> .
<code>var_names</code>	An optional list to rename the variable colnames in the output table, e.g. <code>c("Age (years)", "Sex", "IQ")</code> . Must match <code>vars</code> in length. If not supplied, <code>vars</code> will be printed as is.
<code>factor_vars</code>	An optional list of variables from <code>vars</code> to use as class factor, e.g. <code>c("sex")</code> . Note that any character, factor, or logical class variables will be summarized as categorical by default.
<code>round_n</code>	The number of decimal places to round output to (default=2).
<code>es_col</code>	Include a column for effect size of group difference? (default=T).
<code>p_col</code>	Include a column for p-value of group difference? (default=TRUE).
<code>stars</code>	Where to include stars indicating significance of group differences. Options: "col"=separate column (default), "name"= append to variable name, "stat"= append to group difference statistic, "none" for no stars.
<code>html</code>	Format as html in viewer or not (default=FALSE, print in console), needs <code>library(htmlTable)</code> installed.

Value

Output Table 1

Examples

```
FullTable1(
  data = psydat,
  vars = c("Age", "Height", "depressT"), strata = "Sex"
)
FullTable1(
```

```

data = psydat,
vars = c("Age", "Height", "depressT"), strata = "Sex"
)
FullTable1(
  data = psydat, vars = c("Age", "Sex", "Height", "depressT"),
  var_names = c("Age (months)", "Sex", "Height (inches)", "Depression T"),
  strata = "Income", stars = "name", p_col = FALSE
)
tmp <- FullTable1(data = psydat,
  vars = c("Age", "Height", "depressT"), strata = "Sex")
tmp$caption <- "Write your own caption"
#print(htmlTable(x$table, useViewer=T, rnames=F, caption=x$caption, pos.caption="bottom"))

```

gg_groupplot

Create ggplot to display group differences (box+point+hist)

Description

The `gg_groupplot` function can be used to create group difference plots for scientific publication. This is intended to summarize a continuous outcome (y) based on a factor (x) from an input dataset (`data`). The plot will include standard `ggplot2::geom_boxplot` indicating 25th, median, and 75th percentile for the box and $1.5 * IQR$ for the whiskers. Outliers are not highlighted. Raw data is displayed with standard `ggplot2::geom_point` and lateral but not vertical jittering. Histograms are shown with `gghalves::geom_half_violin` to the right of each boxplot. If `meanline == TRUE` (default), gray dots will indicate the mean for each variable (vs. median in boxplot) connected by a gray line. This function will drop any NA values. Requires `ggplot2` and `gghalves` libraries.

Usage

```
gg_groupplot(data, x = NULL, y = NULL, meanline = c(TRUE, FALSE))
```

Arguments

<code>data</code>	The input dataset.
<code>x</code>	The grouping factor, e.g. "Sex".
<code>y</code>	The numeric outcome variable, e.g. "Age"
<code>meanline</code>	Optional indicator of means

Value

Output group plot

Examples

```
gg_groupplot(data = psydat, x = "Sex", y = "depressT", meanline = TRUE)
```

partial_correltable *Create partial correlation table (with stars for significance) for scientific publication*

Description

The `partial_correltable` function can be used to create partial correlation table (with stars for significance) for scientific publication. This is intended to summarize partial correlations between (vars) from an input dataset (data), residualizing all vars by `partialvars`. This function allows for numeric, binary, and factor variables as `partialvars`. but only numeric vars are used and any non-numeric vars will be dropped. All other flags follow from `scipub::correltable`. Correlations are based on `stats::cor`, use and method follow from that function. Stars indicate significance: *p<.05, **p<.01, ***p<.001. For formatting, variables can be renamed, numbers can be rounded, upper or lower triangle only can be selected (or whole matrix), and empty columns/rows can be dropped if using triangles. For more compact columns, variable names can be numbered in the rows and column names will be corresponding numbers. Requires `tidyverse` and `stats` libraries.

Usage

```
partial_correltable(
  data,
  vars = NULL,
  var_names = vars,
  partialvars = NULL,
  partialvar_names = partialvars,
  method = c("pearson", "spearman"),
  use = c("pairwise", "complete"),
  round_n = 2,
  tri = c("upper", "lower", "all"),
  cutempty = c(FALSE, TRUE),
  colnum = c(FALSE, TRUE),
  html = c(FALSE, TRUE)
)
```

Arguments

<code>data</code>	The input dataset.
<code>vars</code>	A list of the names of 2+ variables to correlate, e.g. <code>c("Age", "height", "WASI")</code> . All variables must be numeric.
<code>var_names</code>	An optional list to rename the vars colnames in the output table, e.g. <code>c("Age (years)", "Height (inches)", "IQ")</code> . Must match <code>vars</code> in length. If not supplied, vars will be printed as is.
<code>partialvars</code>	A list of the names of 1+ variables to partial out, e.g. <code>c("iq", "Sex", "Income")</code> . Can include numeric, binary, factor variables.

<code>partialvar_names</code>	An optional list to rename the <code>partialvars</code> colnames in the output table, e.g. <code>c("IQ (WASI)", "Sex", "Income")</code> . Must match <code>partialvar_names</code> in length. If not supplied, <code>partialvar_names</code> will be printed as is.
<code>method</code>	Type of correlation to calculate <code>c("pearson", "spearman")</code> , based on <code>stats::cor</code> , default = "pearson".
<code>use</code>	Use <code>pairwise.complete.obs</code> or <code>restrict</code> to complete cases <code>c("pairwise", "complete")</code> , based on <code>stats::cor</code> , default = "pairwise".
<code>round_n</code>	The number of decimal places to round all output to (default=2).
<code>tri</code>	Select output formatting <code>c("upper", "lower", "all")</code> ; KEEP the upper triangle, lower triangle, or all values, default = "upper".
<code>cutempty</code>	If keeping only upper/lower triangle with <code>tri</code> , cut empty row/column, default=FALSE.
<code>colnum</code>	For more concise column names, number row names and just use corresponding numbers as column names, default=FALSE, if TRUE overrides <code>cutempty</code> .
<code>html</code>	Format as html in viewer or not (default=F, print in console), needs library(htmlTable) installed.

Value

Output Table 1

Examples

```
partial_correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  partialvars = c("Sex", "Income"),
  tri = "lower", html = TRUE
)

partial_correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  var_names = c("Age (months)", "Height (inches)", "IQ"),
  partialvars = c("Sex", "Income"),
  tri = "upper", colnum = TRUE, html = TRUE
)

partial_correltable(
  data = psydat, vars = c("Age", "Height", "iq"),
  var_names = c("Age (months)", "Height (inches)", "IQ"),
  partialvars = c("anxT"),
  partialvar_names = "Anxiety",
  tri = "all", html = TRUE
)
```

 psydat

Sample demographic and clinical data for 5,000 children

Description

An example dataset containing demographic and clinical data for 5,000 children. The variables are as follows:

Usage

```
data(psydat)
```

Format

A data frame with 5000 rows and 7 variables:

Age age in months (107.2–136.4)

Sex biological sex, 4 missing value (M, F)

Income reported family income, 404 missing values (<50K, >=100K, >=50K&<100K)

Height height in inches, 7 missing values (36.05–84.51)

iq cognition test, 179 missing values (34.86–222.99)

depressT depression symptom severity T-score, 8 missing values (48.53–91.32)

anxT anxiety symptom severity T-score, 8 missing values (48.76–93.67)

 winsorZ

Winsorize outliers based on z-score cutoff to next most extreme non-outlier value

Description

The winsorZ function identifies outliers based on Z-score cutoff and replaces with the next most extreme non-outlier value. This involves z-scoring the variable and identifying/replacing any cases beyond the z-score threshold. The winsorZ_find function is an optional companion to flag any Z-score outliers to tally as needed.

Usage

```
winsorZ(x, zbound = 3)
```

Arguments

x The input variable to Winsorize.

zbound The Z-score cutoff (default=3, i.e. outliers are $Z > 3$ | $Z < -3$).

Value

Output Winsorized variable

Examples

```
winsorZ(psydat$iq)
## Not run:
psydat %>%
  dplyr::select(c(iq, anxT)) %>%
  map(winsorZ)
psydat %>% mutate_at(c("iq", "anxT"), list(~ winsorZ(.)))
psydat %>% mutate_if(is.double, list(~ winsorZ(.)))

## End(Not run)
```

winsorZ_find	<i>Identify outliers based on z-score cutoff that are Winsorized by the winsorZ function</i>
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Description

The winsorZ_find function is an optional companion to the winsorZ function. The winsorZ function identifies Z-score outliers and replaces with the next most extreme non-outlier value. The winsorZ_find function finds/identifies these Z-score outliers (outliers=1, non-outliers=0).

Usage

```
winsorZ_find(x, zbound = 3)
```

Arguments

x	The input variable to check for Z-score outliers.
zbound	The Z-score cutoff (default=3, i.e. outliers are $Z > 3$ $Z < -3$).

Value

Output logical variable of Z-score outliers

Examples

```
summary(winsorZ_find(psydat$iq))
## Not run:
psydat %>% mutate_at(c("iq", "anxT"), list(out = ~ winsorZ_find(.)))

## End(Not run)
```

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