Package 'kim'

October 9, 2023

```
Title A Toolkit for Behavioral Scientists
```

Version 0.5.421

Description A collection of functions for analyzing data typically collected or used by behavioral scientists. Examples of the functions include a function that compares groups in a factorial experimental design, a function that conducts two-way analysis of variance (ANOVA), and a function that cleans a data set generated by Qualtrics surveys. Some of the functions will require installing additional package(s). Such packages and other references are cited within the section describing the relevant functions. Many functions in this package rely heavily on these two popular R packages:

Dowle et al. (2021) https://CRAN.R-project.org/package=ggplot2>. Wickham et al. (2021) https://CRAN.R-project.org/package=ggplot2>.

License GPL-3

```
URL https://github.com/jinkim3/kim, https://jinkim.science
```

BugReports https://github.com/jinkim3/kim/issues

Imports data.table, remotes

Suggests boot, ggplot2, moments, testthat (>= 3.0.0)

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Config/testthat/edition 3

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R topics documented:

akaike_weights	5
barplot_for_counts	5
binomial_test	6
bracket	7
capitalize	9
change_var_names	9
check_modes	0
check_req_pkg	1
chi_squared_test	1
chi_squared_test_pairwise	2
clean_data_from_qualtrics	4
coefficient_of_variation	5
cohen_d	6
cohen_d_borenstein	7
cohen_d_for_one_sample	8
cohen_d_from_cohen_textbook	9
cohen_d_over_n	0
cohen_d_torchiano	1
cohen_d_to_r	2
combine_data_across_cols	2
comma_sep_string_to_numbers	3
compare_datasets	4
compare_dependent_rs	5
compare_effect_sizes	6
compare_groups	7
compare_independent_rs	9
contingency_table	1
convert_cols_to_numeric	2
convert_to_excel_formula	3
correlation_kim	3
correlation_matrix	5
cum_percent_plot	6
desc_stats	6
desc_stats_by_group	7
detach_user_installed_pkgs	
duplicated_values	
excel_formula_convert	
exit_from_parent_function	1
find_duplicates	2
fisher z transform	3
floodlight_2_by_continuous	3
floodlight_2_by_continuous_logistic	7
floodlight_2_by_continuous_mlm_logistic	
floodlight_for_contrasts	
floodlight_multi_by_continuous	
forest_plot	

geomean	
ggsave_quick	
nistogram	
nistogram_by_group	
nistogram_deprecated_1	
nistogram_from_hist	
nistogram_w_outlier_bins	
nolm_adjusted_p	
dentical_all	
d_across_datasets	
nstall_all_dependencies	
curtosis	
enu	
evene_test	. 79
ogistic_regression	
ogistic_regression_table	
ogistic_reg_w_interaction	
oglinear_analysis	
og_odds_ratio	
og_odds_ratio_to_d	. 87
mad_remove_outliers	. 87
mann_whitney	. 89
natrix_prep_dt	. 89
nean_center	. 90
mediation_analysis	
merge_data_tables	. 92
nerge_data_table_list	
nodes_of_objects	. 95
multiple_regression	
noncentrality_parameter	. 98
odds_ratio	. 98
order_rows_specifically_in_dt	. 99
outlier	. 100
overlapping_interval	. 101
00	. 102
	. 103
parallel_analysis	
percentile_rank	. 105
pivot_table	. 105
olot_group_means	. 106
om	. 108
population_variance	. 109
orep	. 109
pretty_round_p_value	. 110
pretty_round_r	. 111
print_loop_progress	. 112
proportion_of_values_in_vector	. 113
stat test homo r	114

163

Index

ead_csv	115
ead_sole_csv	115
egex_match	116
el_pos_of_value_in_vector	117
el_value_of_pos_in_vector	
emove_from_vector	
emove_user_installed_pkgs	119
epeated_measures_anova	
eplace_values_in_dt	121
obust_regression	
ound_flexibly	122
catterplot	
core_scale_items	
etup_r_env	
etwd_to_active_doc	
e_of_mean	
e_of_percentage	
e_of_proportion	
imple_effects_analysis	
kewness	
potlight_2_by_continuous	
tandardize	
tandardized_regression	
tart_kim	140
u	142
abulate_vector	
au_squared	
heme_kim	145
op_median_or_bottom	146
v	
wo_way_anova	148
_test_pairwise	151
ınd	
ınload_user_installed_pkgs	
ppdate_kim	
var_of_log_odds_ratio_to_var_of_d	155
var_of_percentage	
var_of_proportion	156
olookup	156
weighted_mean_effect_size	157
weighted_mean_r	158
weighted_z	159
vilcoxon_rank_sum_test	159
write_csv	160
z_score	161

akaike_weights 5

akaike_weights

Akaike Weights

Description

Compare adequacy of different models by calculating their Akaike weights and the associated evidence ratio.

Usage

```
akaike_weights(aic_values = NULL, print_output_explanation = TRUE)
```

Arguments

```
aic_values a vector of AIC values

print_output_explanation
    logical. Should an explanation about how to read the output be printed? (default = TRUE).
```

Details

Please refer to Wagenmakers & Farrell (2004), doi:10.3758/BF03206482

Value

the output will be a data.table showing AIC weights, their evidence ratio(s), etc.

Examples

```
\# default reference AIC value is the minimum AIC value, e.g., 202 below. akaike_weights(c(204, 202, 206, 206, 214))
```

barplot_for_counts

Barplot for counts

Description

Barplot for counts

Usage

```
barplot_for_counts(data = NULL, x, y)
```

6 binomial_test

Arguments

```
data a data object (a data frame or a data.table)

x name of the variable that will be on the x axis of the barplot

y name of the variable that will be on the y axis of the barplot
```

Examples

```
barplot_for_counts(x = 1:3, y = 7:9)
barplot_for_counts(data = data.frame(
    cyl = names(table(mtcars$cyl)),    count = as.vector(table(mtcars$cyl))),
    x = "cyl", y = "count")
```

binomial_test

Binomial test

Description

Conduct a binomial test. In other words, test whether an observed proportion of "successes" (e.g., proportion of heads in a series of coin tosses) is greater than the expected proportion (e.g., 0.5). This function uses the 'binom.test' function from the 'stats' package.

Usage

```
binomial_test(
  x = NULL,
  success = NULL,
  failure = NULL,
  p = 0.5,
  alternative = "two.sided",
  ci = 0.95,
  round_percentages = 0
)
```

Arguments

X	a vector of values, each of which represents an instance of either a "success" or "failure" (e.g., c("s", "f", "s", "s", "f", "s"))
success	which value(s) indicate "successes"?
failure	(optional) which value(s) indicate "failures"? If no input is provided for this argument, then all the non-NA values that are not declared to be "successes" will be treated as "failures".
р	hypothesized probability of success (default = 0.5)

bracket 7

```
alternative indicates the alternative hypothesis and must be one of "two.sided", "greater", or "less". You can specify just the initial letter. By default, alternative = "two.sided"  

ci width of the confidence interval (default = 0.95)  
round_percentages  
number of decimal places to which to round the percentages in the summary table (default = 0)
```

Examples

```
# sample vector
sample_vector <- c(0, 1, 1, 0, 1, 98, 98, 99, NA)
binomial_test(
x = sample_vector,
success = 1, failure = 0)
binomial_test(
x = sample_vector,
success = 1, failure = 0,
p = 0.1,
alternative = "greater")
binomial_test(
x = sample_vector,
success = c(1, 99), failure = c(0, 98),
p = 0.6,
alternative = "less")</pre>
```

bracket

Draw a bracket on a ggplot

Description

Draw a square bracket with a label on a ggplot

Usage

```
bracket(
   xmin = NULL,
   xmax = NULL,
   ymin = NULL,
   ymax = NULL,
   vertical = NULL,
   horizontal = NULL,
   open = NULL,
   bracket_shape = NULL,
   thickness = 2,
   bracket_color = "black",
   label = NULL,
   label_hjust = NULL,
```

8 bracket

```
label_vjust = NULL,
label_font_size = 5,
label_font_face = "bold",
label_color = "black",
label_parse = FALSE
)
```

Arguments

xmin	xmin
xmax	xmax
ymin	ymin
ymax	ymax
vertical	vertical
horizontal	horizontal
open	open
bracket_shape	bracket_shape
thickness	thickness
bracket_color	bracket_color
label	label
label_hjust	label_hjust
label_vjust	label_vjust
label_font_size	
label_font_face	label_font_size
Tabet_ront_race	label_font_face
label_color	label_font_face
_ label_parse	label_parse
•	

Value

a ggplot object; there will be no meaningful output from this function. Instead, this function should be used with another ggplot object

```
prep(ggplot2)
ggplot(mtcars, aes(x = cyl, y = mpg)) + geom_point() +
bracket(6.1, 6.2, 17, 22, bracket_shape = "]", label = "abc")
```

capitalize 9

capitalize

Capitalize a substring

Description

Capitalizes the first letter (by default) or a substring of a given character string or each element of the character vector

Usage

```
capitalize(x, start = 1, end = 1)
```

Arguments

```
x a character string or a character vector
start starting position of the susbtring (default = 1)
end ending position of the susbtring (default = 1)
```

Value

a character string or a character vector

Examples

```
capitalize("abc")
capitalize(c("abc", "xyx"), start = 2, end = 3)
```

change_var_names

Change variable names in a data set

Description

Change variable names in a data set

Usage

```
change_var_names(
  data = NULL,
  old_var_names = NULL,
  new_var_names = NULL,
  skip_absent = FALSE,
  print_summary = TRUE,
  output_type = "dt"
)
```

10 check_modes

Arguments

data a data object (a data frame or a data.table)

old_var_names a vector of old variable names (i.e., variable names to change)

new_var_names a vector of new variable names

skip_absent If skip_absent = TRUE, old variable names that do not exist in the data set will

be skipped (default = TRUE).

print_summary If print_summary = TRUE, a summary of old and new variable names will printed.

(default = TRUE)

output_type type of the output. If output_type = "dt", the function's output will be a

data.table with changed names. If output_type = "summary", the function's output will be a data.table listing old and new variable names. By default,

output_type = "dt".

Value

a data.table object with changed variable names

Examples

```
change_var_names(
mtcars, old = c("mpg", "cyl"), new = c("mpg_new", "cyl_new"))
```

check_modes

Check modes of objects

Description

Check modes of objects

Usage

```
check_modes(..., mode_to_confirm = NULL)
```

Arguments

```
\begin{tabular}{ll} $\dots$ & R objects. \\ $mode\_to\_confirm \end{tabular}
```

The function will test whether each input is of this mode. For example, check_modes(a, mode_to_confirm = "numeric"), the function will check whether the object a is numeric.

```
check_modes(1L, mode_to_confirm = "numeric")
check_modes(
TRUE, FALSE, 1L, 1:3, 1.1, c(1.2, 1.3), "abc", 1 + 2i, intToBits(1L),
mode_to_confirm = "numeric")
```

check_req_pkg

check_req_pkg

Check for required packages

Description

Check whether required packages are installed.

Usage

```
check_req_pkg(pkg = NULL)
```

Arguments

pkg

a character vector containing names of packages to check

Value

there will be no output from this function. Rather, the function will check whether the packages given as inputs are installed.

Examples

```
check_req_pkg("data.table")
check_req_pkg(c("base", "utils", "ggplot2", "data.table"))
```

chi_squared_test

Chi-squared test

Description

Conduct a chi-squared test and produce a contingency table

Usage

```
chi_squared_test(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  round_chi_sq_test_stat = 2,
  round_p = 3,
  sigfigs_proportion = 2,
  correct = TRUE,
  odds_ratio_ci = 0.95,
```

```
round_odds_ratio_ci_limits = 2,
invert = FALSE
)
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the independent variable

dv_name name of the dependent variable (must be a binary variable)

round_chi_sq_test_stat

number of decimal places to which to round the chi-squared test statistic (default

= 2)

round_p number of decimal places to which to round the p-value from the chi-squared

test (default = 3)

sigfigs_proportion

number of significant digits to round to (for the table of proportions). By default

sigfigs_proportion = 2

correct logical. Should continuity correction be applied? (default = TRUE)

odds_ratio_ci width of the confidence interval for the odds ratio. Input can be any value less

than 1 and greater than or equal to 0. By default, odds_ratio_ci = 0.95. If odds_ratio_ci = TRUE, the default value of 0.95 will be used. If odds_ratio_ci

= FALSE, no confidence interval will be estimated for the odds ratio.

round_odds_ratio_ci_limits

number of decimal places to which to round the limits of the odds ratio's confi-

dence interval (default = 2)

invert logical. Whether the inverse of the odds ratio (i.e., 1 / odds ratio) should be

returned.

Examples

```
chi_squared_test(data = mtcars, iv_name = "cyl", dv_name = "am")
# if the iv has only two levels, odds ratio will also be calculated
chi_squared_test(data = mtcars, iv_name = "vs", dv_name = "am")
```

```
chi_squared_test_pairwise
```

Chi-squared test, pairwise

Description

Conducts a chi-squared test for every possible pairwise comparison with Bonferroni correction

Usage

```
chi_squared_test_pairwise(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  focal_dv_value = NULL,
  contingency_table = TRUE,
  contingency_table_sigfigs = 2,
  percent_and_total = FALSE,
  percentages_only = NULL,
  counts_only = NULL,
  sigfigs = 3,
  chi_sq_test_stats = FALSE,
  correct = TRUE
)
```

Arguments

data a data object (a data frame or a data.table) name of the independent variable (must be a categorical variable) iv_name dv_name name of the dependent variable (must be a binary variable) focal_dv_value focal value of the dependent variable whose frequencies will be calculated (i.e., the value of the dependent variable that will be considered a "success" or a result of interest) contingency_table If contingency_table = TRUE or if contingency_table = "percentages", the percentage of each binary value within each group will be printed. If contingency_table = "counts", a table of frequencies will be printed. If contingency_table = FALSE, no contingency table will be printed. contingency_table_sigfigs number of significant digits that the contingency table's percentage values should be rounded to (default = 2)percent_and_total logical. If percent_and_total = TRUE, tabulate percentages of the focal DV value and a total count of the two values in DV. By default percent_and_total = FALSE percentages_only tabulate percentages of the focal DV value only counts_only tabulate counts of the focal DV value only sigfigs number of significant digits to round to chi_sq_test_stats if chi_sq_test_stats = TRUE, chi-squared test statistic and degrees of freedom will be included in the pairwise comparison data.table. logical. Should continuity correction be applied? (default = TRUE) correct

Examples

```
chi_squared_test_pairwise(data = mtcars, iv_name = "vs", dv_name = "am")
chi_squared_test_pairwise(data = mtcars, iv_name = "vs", dv_name = "am",
percentages_only = TRUE)
# using 3 mtcars data sets combined
chi_squared_test_pairwise(
data = rbind(mtcars, rbind(mtcars, mtcars)),
iv_name = "cyl", dv_name = "am")
# include the total counts
chi_squared_test_pairwise(
data = rbind(mtcars, rbind(mtcars, mtcars)),
iv_name = "cyl", dv_name = "am", percent_and_total = TRUE)
# display counts
chi_squared_test_pairwise(
data = rbind(mtcars, rbind(mtcars, mtcars)),
iv_name = "cyl", dv_name = "am", contingency_table = "counts")
```

clean_data_from_qualtrics

Clean data from Qualtrics

Description

Clean a data set downloaded from Qualtrics

Usage

```
clean_data_from_qualtrics(
  data = NULL,
  remove_survey_preview_data = TRUE,
  remove_test_response_data = TRUE,
  default_cols_by_qualtrics = NULL,
  default_cols_by_qualtrics_new = NULL,
  warn_accuracy_loss = FALSE,
  click_data_cols = "rm",
  page_submit_cols = "move_to_right"
)
```

Arguments

```
data a data object (a data frame or a data.table)

remove_survey_preview_data

logical. Whether to remove data from survey preview (default = TRUE)

remove_test_response_data

logical. Whether to remove data from test response (default = TRUE)
```

coefficent_of_variation 15

default_cols_by_qualtrics

names of columns that Qualtrics includes in the data set by default (e.g., "Start-Date", "Finished"). Accepting the default value default_cols_by_qualtrics = NULL will set the names to be those that Qualtrics uses as of Dec 25, 2020.

default_cols_by_qualtrics_new

new names for columns that Qualtrics includes in the data set by default (e.g., "StartDate", "Finished"). Accepting the default value default_cols_by_qualtrics_new = NULL will set the names to be those that Qualtrics uses as of Dec 25, 2020 converted to snake_case (e.g., "start_date", "finished").

warn_accuracy_loss

logical. whether to warn the user if converting character to numeric leads to loss of accuracy. (default = FALSE)

click_data_cols

if click_data_cols = "rm", columns containing click data (e.g., "_First Click") will be removed. If click_data_cols = "move_to_right", the columns will be moved to the right (end) of the data set.

page_submit_cols

if page_submit_cols = "rm", columns containing page submit data (e.g., "_Page Submit"; "response time" data) will be removed. If page_submit_cols = "move_to_right", the columns will be moved to the right (end) of the data set.

Value

a data.table object

Examples

```
clean_data_from_qualtrics(mtcars)
clean_data_from_qualtrics(mtcars, default_cols_by_qualtrics = "mpg",
default_cols_by_qualtrics_new = "mpg2")
```

coefficent_of_variation

Coefficient of variation

Description

Calculates the (population or sample) coefficient of variation of a given numeric vector

Usage

```
coefficent_of_variation(vector, pop_or_sample = "pop")
```

Arguments

vector a numeric vector

pop_or_sample should coefficient of variation be calculated for a "population" or a "sample"?

16 cohen_d

Value

a numeric value

Examples

```
coefficent_of_variation(1:4, pop_or_sample = "sample")
coefficent_of_variation(1:4, pop_or_sample = "pop")
```

cohen_d

Calculate Cohen's d and its confidence interval using the package 'psych'

Description

To run this function, the following package(s) must be installed: Package 'psych' v2.1.9 (or possibly a higher version) by William Revelle (2021), https://cran.r-project.org/package=psych

Usage

```
cohen_d(
  sample_1 = NULL,
  sample_2 = NULL,
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  ci_range = 0.95,
  output_type = "all"
)
```

Arguments

a vector of values in the first of two samples sample_1 sample_2 a vector of values in the second of two samples a data object (a data frame or a data.table) data name of the independent variable iv_name name of the dependent variable dv_name range of the confidence interval for Cohen's d (default = 0.95) ci_range If output_type == "all" or if output_type == "d_and_ci", the output will be output_type a vector of Cohen's d and its confidence interval. If output_type == "d", the output will be Cohen's d. If output_type == "ci", the output will be a vector of the confidence interval around Cohen's d. By default, output_type == "all".

cohen_d_borenstein 17

Examples

```
## Not run:
cohen_d(sample_1 = 1:10, sample_2 = 3:12)
cohen_d(data = mtcars, iv_name = "vs", dv_name = "mpg", ci_range = 0.99)
sample_dt <- data.table::data.table(iris)[Species != "setosa"]
cohen_d(data = sample_dt, iv_name = "Species", dv_name = "Petal.Width")
## End(Not run)</pre>
```

cohen_d_borenstein

Calculate Cohen's d as illustrated by Borenstein et al. (2009, ISBN: 978-0-470-05724-7)

Description

Calculates Cohen's d, its standard error, and confidence interval, as illustrated in the Borenstein et al. (2009, ISBN: 978-0-470-05724-7).

Usage

```
cohen_d_borenstein(
  sample_1 = NULL,
  sample_2 = NULL,
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  direction = "2_minus_1",
  ci_range = 0.95,
  output_type = "all",
  initial_value = 0
)
```

Arguments

sample_1	a vector of values in the first of two samples
sample_2	a vector of values in the second of two samples
data	a data object (a data frame or a data.table)
iv_name	name of the independent variable
dv_name	name of the dependent variable
direction	If direction == "2_minus_1", Cohen's d will reflect the extent to which the mean of IV level 2 is greater than the mean of IV level 2. If direction == "1_minus_2", Cohen's d will reflect the extent to which the mean of IV level 1 is greater than the mean of IV level 2. By default, direction == "2_minus_1".
ci_range	range of the confidence interval for Cohen's d (default = 0.95)

output_type

If output_type == "all" or if output_type == "d_var_se_and_ci", the output will be a vector of Cohen's d and its variance, SE, and confidence interval. If output_type == "d_se_and_ci", the output will be a vector of Cohen's d and its SE and confidence interval. If output_type == "d_and_ci", the output will be a vector of Cohen's d and its confidence interval. If output_type == "d", the output will be Cohen's d. If output_type == "ci", the output will be a vector of the confidence interval around Cohen's d. If output_type == "se", the output will be the standard error of Cohen's d. By default, output_type == "all".

initial_value

initial value of the noncentrality parameter for optimization (default = 0). Adjust this value if confidence interval results look strange.

Examples

```
cohen_d_borenstein(sample_1 = 1:10, sample_2 = 3:12)
cohen_d_borenstein(
data = mtcars, iv_name = "vs", dv_name = "mpg", ci_range = 0.99)
sample_dt <- data.table::data.table(iris)[Species != "setosa"]
cohen_d_borenstein(
data = sample_dt, iv_name = "Species", dv_name = "Petal.Width",
initial_value = 10)</pre>
```

cohen_d_for_one_sample

Calculate Cohen's d to accompany a one-sample t-test

Description

To run this function, the following package(s) must be installed: Package 'psych' v2.1.9 (or possibly a higher version) by William Revelle (2021), https://cran.r-project.org/package=psych

Usage

```
cohen_d_for_one_sample(x = NULL, mu = NULL)
```

Arguments

x a numeric vector containing values whose mean will be calculated mu the true mean

```
cohen_d_from_cohen_textbook

Cohen's d from Jacob Cohen's textbook (1988)
```

Description

Calculates Cohen's d as described in Jacob Cohen's textbook (1988), Statistical Power Analysis for the Behavioral Sciences, 2nd Edition Cohen, J. (1988) doi:10.4324/9780203771587

Usage

```
cohen_d_from_cohen_textbook(
  sample_1 = NULL,
  sample_2 = NULL,
  data = NULL,
  iv_name = NULL,
  dv_name = NULL
)
```

Arguments

```
sample_1 a vector of values in the first of two samples
sample_2 a vector of values in the second of two samples
data a data object (a data frame or a data.table)
iv_name name of the independent variable
dv_name name of the dependent variable
```

Value

the output will be a Cohen's d value (a numeric vector of length one)

```
cohen_d_from_cohen_textbook(1:10, 3:12)
cohen_d_from_cohen_textbook(
  data = mtcars, iv_name = "vs", dv_name = "mpg"
)
```

cohen_d_over_n

cohen_d_over_n

Cohen's d as a function of sample size

Description

Plot Cohen's d as sample size increases.

Usage

```
cohen_d_over_n(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  save_as_png = FALSE,
  png_name = NULL,
  xlab = NULL,
  ylab = NULL,
  width = 16,
  height = 9
)
```

Arguments

data	a data object (a data frame or a data.table)
iv_name	name of the independent variable (grouping variable)
dv_name	name of the dependent variable (measure variable of interest)
save_as_png	if save = TRUE, the plot will be saved as a PNG file.
png_name	name of the PNG file to be saved. By default, the name will be "cohen_d_over_n_' followed by a timestamp of the current time. The timestamp will be in the format, jan_01_2021_1300_10_000001, where "jan_01_2021" would indicate January 01, 2021; 1300 would indicate 13:00 (i.e., 1 PM); and 10_000001 would indicate 10.000001 seconds after the hour.
xlab	title of the x-axis for the histogram by group. If $xlab = FALSE$, the title will be removed. By default (i.e., if no input is given), dv_name will be used as the title.
ylab	title of the y-axis for the histogram by group. If ylab = FALSE, the title will be removed. By default (i.e., if no input is given), iv_name will be used as the title.
width	width of the plot to be saved. This argument will be directly entered as the width argument for the ggsave function within ggplot2 package (default = 16)
height	height of the plot to be saved. This argument will be directly entered as the height argument for the ggsave function within ggplot2 package (default = 9)

Value

the output will be a list of (1) ggplot object (histogram by group) and (2) a data.table with Cohen's d by sample size

cohen_d_torchiano 21

Examples

```
## Not run:
cohen_d_over_n(data = mtcars, iv_name = "am", dv_name = "mpg")
## End(Not run)
```

cohen_d_torchiano

Calculate Cohen's d and its confidence interval using the package 'effsize'

Description

To run this function, the following package(s) must be installed: Package 'effsize' v0.8.1 (or possibly a higher version) by Marco Torchiano (2020), https://cran.r-project.org/package=effsize

Usage

```
cohen_d_torchiano(
  sample_1 = NULL,
  sample_2 = NULL,
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  ci_range = 0.95
)
```

Arguments

```
sample_1 a vector of values in the first of two samples

a vector of values in the second of two samples

data a data object (a data frame or a data.table)

iv_name name of the independent variable

dv_name range of the confidence interval for Cohen's d (default = 0.95)
```

```
cohen_d_torchiano(1:10, 3:12)
cohen_d_torchiano(
data = mtcars, iv_name = "vs", dv_name = "mpg", ci_range = 0.99)
```

cohen_d_to_r

Convert Cohen's d to r

Description

Convert d (standardized mean difference or Cohen's d) to r (correlation), as illustrated in Borenstein et al. (2009, p. 48, ISBN: 978-0-470-05724-7)

Usage

```
cohen_d_to_r(d = NULL, n1 = NULL, n2 = NULL, d_var = NULL)
```

Arguments

d	Cohen's d (the input can be a vector of values)
n1	sample size in the first of two group (the input can be a vector of values)
n2	sample size in the second of two group (the input can be a vector of values)
d_var	(optional argument) variance of d (the input can be a vector of values). If this argument receives an input, variance of r will be returned as well.

Value

the output will be a vector of correlation values (and variances of r if the argument d_var received an input)

Examples

```
## Not run:
cohen_d_to_r(1)
cohen_d_to_r(d = 1:3)
cohen_d_to_r(d = 1:3, n1 = c(100, 200, 300), n2 = c(50, 250, 900))
cohen_d_to_r(1.1547)
cohen_d_to_r(d = 1.1547, d_var = .0550)
cohen_d_to_r(d = 1:2, d_var = 1:2)
## End(Not run)
```

```
combine_data_across_cols
```

Combine data across columns

Description

Combine data across columns. If NA is the only value across all focal columns for given row(s), NA will be returned for those row(s).

Usage

```
combine_data_across_cols(data = NULL, cols = NULL)
```

Arguments

data a data object (a data frame or a data.table)

cols a character vector containing names of columns, across which to combine data

Value

the output will be a numeric or character vector.

Examples

```
\label{eq:combine_data_across_cols} \begin{array}{l} \text{dt} & \text{cols} = \text{c(NA, 2)} \\ \text{dt} \\ \text{combine\_data\_across\_cols} \\ \text{(data = dt, cols = c("v1", "v2"))} \\ \text{dt} & \text{cols} = \text{c(NA, 4, 3)} \\ \text{dt} \\ \text{combine\_data\_across\_cols} \\ \text{(data = dt, cols = c("v1", "v2"))} \\ \text{dt} & \text{combine\_data\_across\_cols} \\ \text{(data = dt, cols = c(NA, 2, NA))} \\ \text{dt} \\ \text{combine\_data\_across\_cols} \\ \text{(data = dt, cols = c("v1", "v2"))} \\ \end{array}
```

```
comma_sep_string_to_numbers
```

Convert a comma-separated string of numbers

Description

Convert a comma-separated string of numbers

Usage

```
comma_sep_string_to_numbers(string)
```

Arguments

string

a character string consisting of numbers separated by commas

Value

a character string

```
comma_sep_string_to_numbers("1, 2, 3,4, 5 6")
```

24 compare_datasets

compare_datasets

Compare data sets

Description

Compares whether or not data sets are identical

Usage

```
compare_datasets(dataset_1 = NULL, dataset_2 = NULL, dataset_list = NULL)
```

Arguments

```
dataset_1 a data object (a data frame or a data.table)
dataset_2 another data object (a data frame or a data.table)
dataset_list list of data objects (data.frame or data.table)
```

Value

the output will be a data.table showing differences in data sets

```
# catch differences in class attributes of the data sets
compare_datasets(
dataset_1 = data.frame(a = 1:2, b = 3:4),
dataset_2 = data.table::data.table(a = 1:2, b = 3:4))
# catch differences in number of columns
compare_datasets(
dataset_1 = data.frame(a = 1:2, b = 3:4, c = 5:6),
dataset_2 = data.frame(a = 1:2, b = 3:4))
# catch differences in number of rows
compare_datasets(
dataset_1 = data.frame(a = 1:2, b = 3:4),
dataset_2 = data.frame(a = 1:10, b = 11:20))
# catch differences in column names
compare_datasets(
dataset_1 = data.frame(A = 1:2, B = 3:4),
dataset_2 = data.frame(a = 1:2, b = 3:4))
# catch differences in values within corresponding columns
compare_datasets(
dataset_1 = data.frame(a = 1:2, b = c(3, 400)),
dataset_2 = data.frame(a = 1:2, b = 3:4))
compare_datasets(
dataset_1 = data.frame(a = 1:2, b = 3:4, c = 5:6),
dataset_2 = data.frame(a = 1:2, b = c(3, 4), c = c(5, 6)))
# check if data sets in a list are identical
compare_datasets(
dataset_list = list(
```

compare_dependent_rs 25

```
dt1 = data.frame(a = 1:2, b = 3:4, c = 5:6),
dt2 = data.frame(a = 1:2, b = 3:4),
dt3 = data.frame(a = 1:2, b = 3:4, c = 5:6)))
```

compare_dependent_rs Compare dependent correlations

Description

Compares whether two dependent correlations from the same sample are significantly different each other.

Usage

```
compare_dependent_rs(
  data = NULL,
  var_1_name = NULL,
  var_2_name = NULL,
  var_3_name = NULL,
  one_tailed = FALSE,
  round_r = 3,
  round_p = 3,
  round_t = 2,
  print_summary = TRUE,
  return_dt = FALSE
)
```

Arguments

data	a data object (a data frame or a data.table)
var_1_name	name of the variable whose correlations with two other variables will be compared.
var_2_name	name of the first of the two variables whose correlations with var_1_name will be compared.
var_3_name	name of the second of the two variables whose correlations with var_1_name will be compared.
one_tailed	logical. Should the p value based on a one-tailed t-test? (default = FALSE)
round_r	number of decimal places to which to round correlation coefficients (default = 2)
round_p	number of decimal places to which to round p-values (default = 3)
round_t	number of decimal places to which to round the t-statistic (default = 2)
print_summary	logical. Should the summary be printed? (default = TRUE)
return_dt	logical. Should the function return a summary table as an output, as opposed to returning the output through the "invisible" function? (default = FALSE)

Details

Suppose that Variables A, B, and C are measured from a group of subjects. This function tests whether A is related to B differently than to C. Put differently, this function tests H0: r(A, B) = r(A, C)

For more information on formulas used in this function, please refer to Steiger (1980) doi:10.1037/00332909.87.2.245 and Chen & Popovich (2002) doi:10.4135/9781412983808

Value

the output will be a summary of the test comparing two dependent correlations

Examples

Description

Compares effect sizes See p. 156 of Borenstein et al. (2009, ISBN: 978-0-470-05724-7).

Usage

```
compare_effect_sizes(
  effect_sizes = NULL,
  effect_size_variances = NULL,
  round_stats = TRUE,
  round_p = 3,
  round_se = 2,
  round_z = 2,
  pretty_round_p_value = TRUE
)
```

Arguments

```
a vector of estimated effect sizes
effect_sizes
effect_size_variances
                  a vector of variances of the effect sizes
                  logical. Should the statistics be rounded? (default = TRUE)
round_stats
                  number of decimal places to which to round p-values (default = 3)
round_p
round_se
                  number of decimal places to which to round the standard errors of the difference
                  (default = 2)
round_z
                  number of decimal places to which to round the z-statistic (default = 2)
pretty_round_p_value
                  logical. Should the p-values be rounded in a pretty format (i.e., lower threshold:
                  "<.001"). By default, pretty_round_p_value = TRUE.
```

compare_groups 27

Examples

```
compare_effect_sizes(
effect_sizes = c(0.6111, 0.3241, 0.5),
effect_size_variances = c(.0029, 0.0033, 0.01))
```

compare_groups

Compare groups

Description

Compares groups by (1) creating histogram by group; (2) summarizing descriptive statistics by group; and (3) conducting pairwise comparisons (t-tests and Mann-Whitney tests).

Usage

```
compare_groups(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  sigfigs = 3,
  stats = "basic",
  cohen_d = TRUE,
  cohen_d_w_ci = TRUE,
  adjust_p = "holm",
  bonferroni = NULL,
 mann_whitney = TRUE,
  t_test_stats = TRUE,
  t_test_df_decimals = 1,
  round_p = 3,
  save_as_png = FALSE,
  png_name = NULL,
  xlab = NULL,
  ylab = NULL,
  x_limits = NULL,
  x_breaks = NULL,
  x_{labels} = NULL,
 width = 5000,
  height = 3600,
  units = "px",
  res = 300,
  layout_matrix = NULL,
  col_names_nicer = TRUE,
  convert_dv_to_numeric = TRUE
)
```

28 compare_groups

Arguments

data a data object (a data frame or a data.table)

iv_name name of the independent variable (grouping variable)

dv_name name of the dependent variable (measure variable of interest)

sigfigs number of significant digits to round to

stats statistics to calculate for each group. If stats = "basic", group size, mean,

standard deviation, median, minimum, and maximum will be calculated. If ${\tt stats} = {\tt "all"}$, in addition to the aforementioned statistics, standard error, 95% confidence and prediction intervals, skewness, and kurtosis will also be calculated. The ${\tt stats}$ argument can also be a character vector with types of statistics to calculate. For example, entering ${\tt stats} = {\tt c("mean", "median")}$ will calculate.

late mean and median. By default, stats = "basic"

cohen_d if cohen_d = TRUE, Cohen's d statistics will be included in the pairwise compar-

ison data.table.

cohen_d_w_ci if cohen_d_w_ci = TRUE, Cohen's d with 95% CI will be included in the output

data.table.

adjust_p the name of the method to use to adjust p-values. If adjust_p = "holm", the

Holm method will be used; if adjust_p = "bonferroni", the Bonferroni method

will be used. By default, adjust_p = "holm"

bonferroni The use of this argument is deprecated. Use the 'adjust_p' argument instead.

If bonferroni = TRUE, Bonferroni tests will be conducted for t-tests or Mann-

Whitney tests.

mann_whitney if TRUE, Mann-Whitney test results will be included in the pairwise comparison

data.table. If FALSE, Mann-Whitney tests will not be performed.

t_test_stats if t_test_stats = FALSE, t-test statistic and degrees of freedom will be ex-

cluded in the pairwise comparison data.table.

t_test_df_decimals

number of decimals for the degrees of freedom in t-tests (default = 1)

round_p number of decimal places to which to round p-values (default = 3)

save_as_png if save = "all" or if save = TRUE, the histogram by group, descriptive statistics

by group, and pairwise comparison results will be saved as a PNG file.

png_name name of the PNG file to be saved. By default, the name will be "compare_groups_results_"

followed by a timestamp of the current time. The timestamp will be in the format, jan_01_2021_1300_10_000001, where "jan_01_2021" would indicate January 01, 2021; 1300 would indicate 13:00 (i.e., 1 PM); and 10_000001 would

indicate 10.000001 seconds after the hour.

xlab title of the x-axis for the histogram by group. If xlab = FALSE, the title will be

removed. By default (i.e., if no input is given), dv_name will be used as the title.

ylab title of the y-axis for the histogram by group. If ylab = FALSE, the title will be

removed. By default (i.e., if no input is given), iv_name will be used as the title.

x_limits a numeric vector with values of the endpoints of the x axis.

x_breaks a numeric vector indicating the points at which to place tick marks on the x axis.

x_labels

width	width of the PNG file (default = 4000)	
height	height of the PNG file (default = 3000)	
units	the units for the width and height arguments. Can be "px" (pixels), "in" (inches), "cm", or "mm". By default, units = "px".	
res	The nominal resolution in ppi which will be recorded in the png file, if a positive integer. Used for units other than the default. If not specified, taken as 300 ppi to set the size of text and line widths.	
layout_matrix	The layout argument for arranging plots and tables using the grid.arrange function.	
col_names_nicer		
layout_matrix	The nominal resolution in ppi which will be recorded in the png file, if a positive integer. Used for units other than the default. If not specified, taken as 300 ppi to set the size of text and line widths. The layout argument for arranging plots and tables using the grid.arrange function.	

a vector containing labels for the place tick marks on the x axis.

if col_names_nicer = TRUE, column names will be converted from snake_case to an easier-to-eye format.

convert_dv_to_numeric

logical. Should the values in the dependent variable be converted to numeric for

plotting the histograms? (default = TRUE)

holm if holm = TRUE, the relevant p values will be adjusted using Holm method (also

known as the Holm-Bonferroni or Bonferroni-Holm method)

Value

the output will be a list of (1) ggplot object (histogram by group) (2) a data.table with descriptive statistics by group; and (3) a data.table with pairwise comparison results. If save_as_png = TRUE, the plot and tables will be also saved on local drive as a PNG file.

Examples

```
## Not run:
compare_groups(data = iris, iv_name = "Species", dv_name = "Sepal.Length")
compare_groups(data = iris, iv_name = "Species", dv_name = "Sepal.Length",
x_breaks = 4:8)
## End(Not run)
```

compare_independent_rs

Compare independent correlations

Description

Compares whether two correlations from two independent samples are significantly different each other. See Field et al. (2012, ISBN: 978-1-4462-0045-2).

Usage

```
compare_independent_rs(
  r1 = NULL,
  n1 = NULL,
  r2 = NULL,
  n2 = NULL,
  one_tailed = FALSE,
  round_p = 3,
  round_z_diff = 2,
  round_r = 2,
  print_summary = TRUE,
  output_type = NULL
)
```

Arguments

r1	correlation in the first sample
n1	size of the first sample
r2	correlation in the second sample
n2	size of the first sample
one_tailed	logical. Should the p value based on a one-tailed t-test? (default = FALSE)
round_p	(only for displaying purposes) number of decimal places to which to round the p -value (default = 3)
round_z_diff	(only for displaying purposes) number of decimal places to which to round the z-score (default = 2)
round_r	(only for displaying purposes) number of decimal places to which to round correlation coefficients (default $= 2$)
print_summary	logical. Should the summary be printed? (default = TRUE)
output_type	type of the output. If output_type = "z", the function's output will be the z-score of the difference between the two correlations. If output_type = "p", the function's output will be the p-value associated with the z-score of the difference between the two correlations. By default, output_type = NULL, and the function will not return any value other than the printed summary.

Value

the output will be the results of a test comparing two independent correlations.

```
compare_independent_rs(r1 = .1, n1 = 100, r2 = .2, n2 = 200) compare_independent_rs( r1 = .1, n1 = 100, r2 = .2, n2 = 200, one_tailed = TRUE) compare_independent_rs(r1 = .506, n1 = 52, r2 = .381, n2 = 51)
```

contingency_table 31

contingency_table

Contingency table

Description

Create a contingency table that takes two variables as inputs

Usage

```
contingency_table(
  data = NULL,
  row_var_name = NULL,
  col_var_name = NULL,
  row = NULL,
  col = NULL,
  output_type = "table"
)
```

Arguments

a data object (a data frame or a data.table)

row_var_name

name of the variable whose values will fill the rows of the contingency table

row

a vector whose values will fill the rows of the contingency table

a vector whose values will fill the rows of the contingency table

a vector whose values will fill the columns of the contingency table

output_type

If output_type == "dt" the output will be a contingency table as a data.table object. If output_type == "table" the output will be a contingency table as a table object. If output_type == "df" the output will be a contingency table as a data.frame object. By default, output_type == "table".

```
contingency_table(
data = mtcars,
row_var_name = "am",
col_var_name = "cyl")
contingency_table(row = mtcars$cyl, col = mtcars$am)
contingency_table(mtcars, "am", "cyl", output_type = "dt")
```

```
convert_cols_to_numeric
```

Convert columns to numeric

Description

Check whether each column in a data.table can be converted to numeric, and if so, convert every such column.

Usage

```
convert_cols_to_numeric(
  data = NULL,
  classes = "character",
  warn_accuracy_loss = TRUE,
  print_summary = TRUE,
  silent = FALSE
)
```

Arguments

data a data object (a data frame or a data.table)

classes a character vector specifying classes of columns that will be converted. For

example, if classes = "character", all columns of the class "character" will be converted—if they can be converted. The current version of the function only

supports converting character columns to numeric.

warn_accuracy_loss

logical. whether to warn the user if converting character to numeric leads to loss

of accuracy. (default = TRUE)

print_summary = TRUE, a summary of converted columns will printed. (de-

fault = TRUE)

silent If silent = FALSE, a message regarding conversion for a data.frame will be

printed. If silent = TRUE, this message will be suppressed. By default, silent

= FALSE.

```
data_frame_1 <- data.frame(a = c("1", "2"), b = c("1", "b"), c = 1:2)
convert_cols_to_numeric(data = data_frame_1)
data_table_1 <- data.table::data.table(
a = c("1", "2"), b = c("1", "b"), c = 1:2)
convert_cols_to_numeric(data = data_table_1)</pre>
```

```
convert_to_excel_formula
```

Convert character to Excel formula

Description

Convert elements of a character vector to Excel formulas to preserve the character (string) format when opened in an Excel file.

Usage

```
convert_to_excel_formula(vector = NULL)
```

Arguments

vector

a character vector

Value

the output will be a character vector formatted as an Excel formula. For example, if an element in the input vector was ".500", this element will be converted to =".500", which will show up as ".500" in Excel, rather than as "0.5"

Examples

```
## Not run:
# compare the two csv files below
# example 1
dt <- data.table::data.table(a = ".500")
data.table::fwrite(dt, "example1.csv") # the csv will show "0.5"
# example 2
dt <- data.table::data.table(a = convert_to_excel_formula(".500"))
data.table::fwrite(dt, "example2.csv") # the csv will show ".500"
## End(Not run)</pre>
```

correlation_kim

Estimate the correlation between two variables

Description

Estimate the correlation between two variables

34 correlation_kim

Usage

```
correlation_kim(
  x = NULL,
  y = NULL,
  data = NULL,
  x_var_name = NULL,
  y_var_name = NULL,
  ci_range = 0.95,
  round_r = 2,
  round_p = 3,
  output_type = "summary"
)
```

Arguments

X	a numeric vector of data values
у	a numeric vector of data values
data	(optional) a data object (a data frame or a data.table)
x_var_name	(optional) name of the first variable (if using a data set as an input)
y_var_name	(optional) name of the second variable (if using a data set as an input)
ci_range	range of the confidence interval for the correlation coefficient. If ci_range = FALSE, no confidence interval will be estimated. By default, ci_range = 0.95.
round_r	number of decimal places to which to round correlation coefficients (default = 2)
round_p	number of decimal places to which to round p-values (default = 3)
output_type	type of the output. If output_type = "dt", the function's output will be a data.table with the results from the correlation analysis. If output_type = "summary", the function's output will be a statement (a string) summarizing the results from the correlation analysis. By default, output_type = "summary"

```
## Not run:
correlation_kim(x = 1:4, y = c(1, 3, 2, 4))
correlation_kim(x = 1:4, y = c(1, 3, 2, 4), ci_range = FALSE)
# output as a data table
correlation_kim(x = 1:4, y = c(1, 3, 2, 4), output_type = "dt")
## End(Not run)
```

correlation_matrix 35

correlation_matrix

correlation matrix

Description

Creates a correlation matrix

Usage

```
correlation_matrix(
  data = NULL,
  var_names = NULL,
  row_var_names = NULL,
  col_var_names = NULL,
  round_r = 2,
  round_p = 3,
  output_type = "r"
)
```

Arguments

data	a data object (a data frame or a data.table)
var_names	names of the variables for which to calculate all pairwise correlations
row_var_names	names of the variables that will go on the rows of the correlation matrix
col_var_names	names of the variables that will go on the columns of the correlation matrix
round_r	number of decimal places to which to round correlation coefficients (default $=$ 2)
round_p	number of decimal places to which to round p-values (default = 3)
output_type	which value should be filled in cells of the correlation matrix? If output_type = "r", correlation coefficients; if output_type = "p", p-values; if output_type = "rp", correlation coefficients with significance symbols based on p-values; if output_type = "n", sizes of the samples used to calculate the correlation coefficients

Value

the output will be a correlation matrix in a data.table format

```
correlation_matrix(data = mtcars, var_names = c("mpg", "cyl", "wt"))
correlation_matrix(data = mtcars,
row_var_names = c("mpg", "cyl", "hp"), col_var_names = c("wt", "am"))
```

36 desc_stats

cum_percent_plot

Cumulative percentage plot

Description

Plots or tabulates cumulative percentages associated with elements in a vector

Usage

```
cum_percent_plot(vector, output_type = "plot")
```

Arguments

vector a numeric vector

output_type if output_type = "plot", return a cumulative percentage plot; if output_type

= "dt", return a data.table with cumulative percentages. By default, output_type

= "plot"

Examples

```
cum_percent_plot(c(1:100, NA, NA))
cum_percent_plot(mtcars$mpg)
cum_percent_plot(vector= mtcars$mpg, output_type = "dt")
```

desc_stats

Descriptive statistics

Description

Returns descriptive statistics for a numeric vector.

Usage

```
desc_stats(
  vector = NULL,
  output_type = "vector",
  sigfigs = 3,
  se_of_mean = FALSE,
  ci = FALSE,
  pi = FALSE,
  skewness = FALSE,
  kurtosis = FALSE,
  notify_na_count = NULL,
  print_dt = FALSE
)
```

desc_stats_by_group 37

Arguments

	vector	a numeric vector
	output_type	<pre>if output_type = "vector", return a vector of descriptive statistics; if output_type = "dt", return a data.table of descriptive statistics (default = "vector")</pre>
	sigfigs	number of significant digits to round to (default = 3)
	se_of_mean	logical. Should the standard errors around the mean be included in the descriptive stats? (default = FALSE)
	ci	logical. Should 95% CI be included in the descriptive stats? (default = FALSE)
	pi	logical. Should 95% PI be included in the descriptive stats? (default = FALSE)
	skewness	logical. Should the skewness statistic be included in the descriptive stats? (default = FALSE)
	kurtosis	logical. Should the kurtosis statistic be included in the descriptive stats? (default = FALSE)
notify_na_count		
		if TRUE, notify how many observations were removed due to missing values. By default, NA count will be printed only if there are any NA values.
	print_dt	if TRUE, print the descriptive stats data.table

Value

if output_type = "vector", the output will be a named numeric vector of descriptive statistics; if output_type = "dt", the output will be data.table of descriptive statistics.

Examples

```
desc_stats(1:100)
desc_stats(1:100, ci = TRUE, pi = TRUE, sigfigs = 2)
desc_stats(1:100, se_of_mean = TRUE,
ci = TRUE, pi = TRUE, sigfigs = 2,
skewness = TRUE, kurtosis = TRUE)
desc_stats(c(1:100, NA))
example_dt <- desc_stats(vector = c(1:100, NA), output_type = "dt")
example_dt</pre>
```

desc_stats_by_group

Descriptive statistics by group

Description

Returns descriptive statistics by group

Usage

```
desc_stats_by_group(
  data = NULL,
  var_for_stats = NULL,
  grouping_vars = NULL,
  stats = "all",
  sigfigs = NULL,
  cols_to_round = NULL
)
```

Arguments

data a data object (a data frame or a data.table) var_for_stats name of the variable for which descriptive statistics will be calculated grouping_vars name(s) of grouping variables stats statistics to calculate. If stats = "basic", group size, mean, standard deviation, median, minimum, and maximum will be calculated. If stats = "all", in addition to the aforementioned statistics, standard error, 95% confidence and prediction intervals, skewness, and kurtosis will also be calculated. The stats argument can also be a character vector with types of statistics to calculate. For example, entering stats = c("mean", "median") will calculate mean and median. By default, stats = "all" sigfigs number of significant digits to round to names of columns whose values will be rounded cols_to_round

Value

the output will be a data.table showing descriptive statistics of the variable for each of the groups formed by the grouping variables.

Examples

```
desc_stats_by_group(data = mtcars, var_for_stats = "mpg",
grouping_vars = c("vs", "am"))
desc_stats_by_group(data = mtcars, var_for_stats = "mpg",
grouping_vars = c("vs", "am"), sigfigs = 3)
desc_stats_by_group(data = mtcars, var_for_stats = "mpg",
grouping_vars = c("vs", "am"), stats = "basic", sigfigs = 2)
desc_stats_by_group(data = mtcars, var_for_stats = "mpg",
grouping_vars = c("vs", "am"), stats = "basic", sigfigs = 2,
cols_to_round = "all")
desc_stats_by_group(data = mtcars, var_for_stats = "mpg",
grouping_vars = c("vs", "am"), stats = c("mean", "median"), sigfigs = 2,
cols_to_round = "all")
```

```
detach_user_installed_pkgs
```

Detach all user-installed packages

Description

Detach all user-installed packages

Usage

```
detach_user_installed_pkgs(exceptions = NULL, force = FALSE, keep_kim = TRUE)
```

Arguments

exceptions a character vector of names of packages to keep attached

force logical. Should a package be detached even though other attached packages

depend on it? By default, force = FALSE

keep_kim logical. If keep_kim = FALSE, Package 'kim' will be detached along with all

other user-installed packages. If keep_kim = TRUE, Package 'kim' will not be

detached. By default, keep_kim = FALSE

Examples

```
## Not run:
detach_user_installed_pkgs()
## End(Not run)
```

 $duplicated_values$

Duplicated values in a vector

Description

Return all duplicated values in a vector. This function is a copy of the earlier function, find_duplicates, in Package 'kim'

```
duplicated_values(vector = NULL, na.rm = TRUE, sigfigs = 2, output = "summary")
```

Arguments

vector	a vector whose elements will be checked for duplicates
na.rm	logical. If na.rm = TRUE, NA values in the vector will be removed before searching for duplicates. If na.rm = FALSE, NA values will be included in the search as potentially duplicated values. By default, na.rm = TRUE.
sigfigs	number of significant digits to round to in the percent column of the summary $(default = 2)$
output	type of output. If output = "summary", the function's output will be a data.table summarizing duplicated values and their counts. If output = "duplicated_values", the function's output will be a vector of duplicated values. If output = "non_duplicated_values", the function's output will be a vector of non-duplicated values (default = "summary")

Value

the output will be a data.table object (summary), a vector of duplicated values, or a vector non-duplicated values.

Examples

```
duplicated_values(mtcars$cyl)
duplicated_values(mtcars$cyl, output = "duplicated_values")
duplicated_values(vector = c(mtcars$cyl, 11:20, NA, NA))
duplicated_values(vector = c(mtcars$cyl, 11:20, NA, NA), na.rm = FALSE)
duplicated_values(vector = c(mtcars$cyl, 11:20, NA, NA),
na.rm = FALSE, sigfigs = 4, output = "duplicated_values")

excel_formula_convert Excel formula, convert (to)
```

Description

Alias for the 'convert_to_excel_formula' function. Convert elements of a character vector to Excel formulas to preserve the character (string) format when opened in an Excel file.

Usage

```
excel_formula_convert(vector = NULL)
```

Arguments

vector a character vector

Value

the output will be a character vector formatted as an Excel formula. For example, if an element in the input vector was ".500", this element will be converted to =".500", which will show up as ".500" in Excel, rather than as "0.5"

Examples

```
## Not run:
# compare the two csv files below
# example 1
dt <- data.table::data.table(a = ".500")
data.table::fwrite(dt, "example1.csv") # the csv will show "0.5"
# example 2
dt <- data.table::data.table(a = excel_formula_convert(".500"))
data.table::fwrite(dt, "example2.csv") # the csv will show ".500"
## End(Not run)</pre>
```

```
exit_from_parent_function
```

Exit from a Parent Function

Description

Exit from a Parent Function

Usage

```
exit_from_parent_function(
  n = 1,
  silent = FALSE,
  message = "Exiting from a parent function"
)
```

Arguments

```
n the number of generations to go back (default = 1)
silent logical. If silent = TRUE, a message will be printed.
message message to print
```

Examples

```
fn1 <- function() {
  print(1)
  print(2)
}
fn1()
fn2 <- function() {
  print(1)
  exit_from_parent_function()
  print(2)
}
fn2()</pre>
```

find_duplicates

find_duplicates	Find duplicated values in a vector	

Description

Find duplicated values in a vector

Usage

```
find_duplicates(vector = NULL, na.rm = TRUE, sigfigs = 2, output = "summary")
```

Arguments

vector	a vector whose elements will be checked for duplicates
na.rm	logical. If na.rm = TRUE, NA values in the vector will be removed before searching for duplicates. If na.rm = FALSE, NA values will be included in the search as potentially duplicated values. By default, na.rm = TRUE.
sigfigs	number of significant digits to round to in the percent column of the summary $(default = 2)$
output	type of output. If output = "summary", the function's output will be a data.table summarizing duplicated values and their counts. If output = "duplicated_values", the function's output will be a vector of duplicated values. If output = "non_duplicated_values", the function's output will be a vector of non-duplicated values (default = "summary")

Value

the output will be a data.table object (summary), a vector of duplicated values, or a vector non-duplicated values.

Examples

```
find_duplicates(mtcars$cyl)
find_duplicates(mtcars$cyl, output = "duplicated_values")
find_duplicates(vector = c(mtcars$cyl, 11:20, NA, NA))
find_duplicates(vector = c(mtcars$cyl, 11:20, NA, NA), na.rm = FALSE)
find_duplicates(vector = c(mtcars$cyl, 11:20, NA, NA),
na.rm = FALSE, sigfigs = 4, output = "duplicated_values")
```

fisher_z_transform 43

```
fisher_z_transform Fisher's Z transformation
```

Description

Perform Fisher's r-to-Z transformation for given correlation coefficient(s).

Usage

```
fisher_z_transform(r = NULL)
```

Arguments

r a (vector of) correlation coefficient(s)

Value

the output will be a vector of Z values which were transformed from the given r values.

Examples

```
fisher_z_transform(0.99)
fisher_z_transform(r = seq(0.1, 0.5, 0.1))
```

```
floodlight_2_by_continuous
```

Floodlight 2 by Continuous

Description

Conduct a floodlight analysis for 2 x Continuous design.

```
floodlight_2_by_continuous(
   data = NULL,
   iv_name = NULL,
   dv_name = NULL,
   mod_name = NULL,
   covariate_name = NULL,
   interaction_p_include = TRUE,
   iv_level_order = NULL,
   output = "reg_lines_plot",
   jitter_x_percent = 0,
   jitter_y_percent = 0,
   dot_alpha = 0.5,
```

```
dot_size = 4,
  interaction_p_value_font_size = 8,
  jn_point_label_add = TRUE,
  jn_point_font_size = 8,
  jn_point_label_hjust = NULL,
  lines_at_mod_extremes = FALSE,
  interaction_p_vjust = -3,
  plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt"),
  legend_position = "right",
  reg_line_types = c("solid", "dashed"),
  jn_line_types = c("solid", "solid"),
  jn_line_thickness = 1.5,
  colors_for_iv = c("red", "blue"),
  sig_region_color = "green",
  sig_region_alpha = 0.08,
  nonsig_region_color = "gray",
  nonsig_region_alpha = 0.08,
  x_axis_title = NULL,
  y_axis_title = NULL,
  legend_title = NULL,
  round_decimals_int_p_value = 3,
  line_of_fit_thickness = 1,
  round_jn_point_labels = 2
)
```

Arguments

```
data
                  a data object (a data frame or a data.table)
iv_name
                  name of the binary independent variable
dv_name
                   name of the dependent variable
mod_name
                   name of the continuous moderator variable
covariate_name name of the variables to control for
interaction_p_include
                  logical. Should the plot include a p-value for the interaction term?
iv_level_order order of levels in the independent variable for legend. By default, it will be set
                   as levels of the independent variable ordered using R's base function sort.
output
                   type of output (default = "reg_lines_plot"). Possible inputs: "interactions_pkg_results",
                   "simple_effects_plot", "jn_points", "regions", "reg_lines_plot"
jitter_x_percent
                   horizontally jitter dots by a percentage of the range of x values
jitter_y_percent
                   vertically jitter dots by a percentage of the range of y values
dot_alpha
                   opacity of the dots (0 = completely transparent, 1 = completely opaque). By
                   default, dot_alpha = 0.5
dot_size
                   size of the dots (default = 4)
```

interaction_p_value_font_size

font size for the interaction p value (default = 8)

jn_point_label_add

logical. Should the labels for Johnson-Neyman point labels be added to the plot? (default = TRUE)

jn_point_font_size

font size for Johnson-Neyman point labels (default = 8)

jn_point_label_hjust

a vector of hjust values for Johnson-Neyman point labels. By default, the hjust value will be 0.5 for all the points.

lines_at_mod_extremes

logical. Should vertical lines be drawn at the observed extreme values of the moderator if those values lie in significant region(s)? (default = FALSE)

interaction_p_vjust

By how much should the label for the interaction p-value be adjusted vertically? By default, interaction_p_vjust = -3)

plot_margin margin for the plot By default plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt")

legend_position

position of the legend (default = "right"). If legend_position = "none", the legend will be removed.

reg_line_types types of the regression lines for the two levels of the independent variable. By default, reg_line_types = c("solid", "dashed")

jn_line_thickness

thickness of the lines at Johnson-Neyman points (default = 1.5)

colors_for_iv colors for the two values of the independent variable (default = c("red", "blue")) sig_region_color

color of the significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is statistically significant.

sig_region_alpha

opacity for sig_region_color. (0 = completely transparent, 1 = completely opaque). By default, sig_region_alpha = 0.08

nonsig_region_color

color of the non-significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is not statistically significant.

nonsig_region_alpha

opacity for nonsig_region_color. (0 = completely transparent, 1 = completely opaque). By default, nonsig_region_alpha = 0.08

x_axis_title title of the x axis. By default, it will be set as input for mod_name. If x_axis_title = FALSE, it will be removed.

Details

The following package(s) must be installed prior to running this function: Package 'interactions' v1.1.1 (or possibly a higher version) by Jacob A. Long (2020), https://cran.r-project.org/package=interactions See the following references: Spiller et al. (2013) doi:10.1509/jmr.12.0420 Kim (2021) doi:10.5281/zenodo.4445388

Examples

```
# typical example
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "gsec")
# add covariates
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg"
mod_name = "qsec",
covariate_name = c("cyl", "hp"))
# adjust the jn point label positions
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am"
dv_name = "mpg",
mod_name = "qsec",
jn_point_label_hjust = c(1, 0)
# return regions of significance and nonsignificance
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am"
dv_name = "mpg",
mod_name = "qsec"
output = "regions")
# draw lines at the extreme values of the moderator
```

```
# if they are included in the significant region
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "qsec",
lines_at_mod_extremes = TRUE)

# remove the labels for jn points
floodlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "qsec",
jn_point_label_add = FALSE)
```

floodlight_2_by_continuous_logistic

Floodlight 2 by Continuous for a Logistic Regression

Description

Conduct a floodlight analysis for a logistic regression with a 2 x Continuous design involving a binary dependent variable.

```
floodlight_2_by_continuous_logistic(
  data = NULL,
  iv_name = NULL,
 dv_name = NULL,
 mod_name = NULL,
  interaction_p_include = TRUE,
  iv_level_order = NULL,
  dv_level_order = NULL,
  jn_points_disregard_threshold = NULL,
  output = "reg_lines_plot",
  num_of_spotlights = 20,
  jitter_x_percent = 0,
  jitter_y_percent = 5,
  dot_alpha = 0.3,
  dot_size = 6,
  interaction_p_value_font_size = 8,
  jn_point_font_size = 8,
  jn_point_label_hjust = NULL,
  interaction_p_vjust = -3,
  plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt"),
  legend_position = "right",
```

```
line_types_for_pred_values = c("solid", "dashed"),
line_thickness_for_pred_values = 2.5,
jn_line_types = c("solid", "solid"),
jn_line_thickness = 1.5,
sig_region_color = "green",
sig_region_alpha = 0.08,
nonsig_region_color = "gray",
nonsig_region_alpha = 0.08,
x_axis_title = NULL,
y_axis_title = NULL,
legend_title = NULL,
round_decimals_int_p_value = 3,
round_jn_point_labels = 2
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the binary independent variable

dv_name name of the binary dependent variable

mod_name name of the continuous moderator variable

interaction_p_include

logical. Should the plot include a p-value for the interaction term?

iv_level_order order of levels in the independent variable for legend. By default, it will be set as levels of the independent variable ordered using R's base function sort.

dv_level_order order of levels in the dependent variable. By default, it will be set as levels of the dependent variable ordered using R's base function sort.

jn_points_disregard_threshold

the Minimum Distance in the unit of the moderator variable that will be used for various purposes, such as (1) to disregard the second Johnson-Neyman point that is different from the first Johnson-Neyman (JN) point by less than the Minimum Distance; (2) to determine regions of significance, which will calculate the p-value of the IV's effect (the focal dummy variable's effect) on DV at a candidate JN point + / - the Minimum Distance. This input is hard to explain, but a user can enter a really low value for this argument (e.g., jn_points_disregard_threshold = 0.1 for a moderator measured on a 100-point scale) or use the default. By default, jn_points_disregard_threshold = range of the moderator / 10000 For example, if the observed moderator values range from 1 to 7 (because it is a 7-point scale), then jn_points_disregard_threshold = (7 - 1) / 10000 = 0.0006

output

type of output (default = "reg_lines_plot"). Possible inputs: "interactions_pkg_results", "simple_effects_plot", "jn_points", "regions", "reg_lines_plot"

num_of_spotlights

How many spotlight analyses should be conducted to plot the predicted values at various values of the moderator? (default = 20)

jitter_x_percent

horizontally jitter dots by a percentage of the range of x values (default = 0)

jitter_y_percent vertically jitter dots by a percentage of the range of y values (default = 5) dot_alpha opacity of the dots (0 = completely transparent, 1 = completely opaque). By default, dot_alpha = 0.3 dot_size size of the dots (default = 6) interaction_p_value_font_size font size for the interaction p value (default = 8) jn_point_font_size font size for Johnson-Neyman point labels (default = 8) jn_point_label_hjust a vector of hjust values for Johnson-Neyman point labels. By default, the hjust value will be 0.5 for all the points. interaction_p_vjust By how much should the label for the interaction p-value be adjusted vertically? By default, interaction_p_vjust = -3) margin for the plot By default plot_margin = ggplot2::unit(c(75, 7, 7, plot_margin 7), "pt") legend_position position of the legend (default = "right"). If legend_position = "none", the legend will be removed. line_types_for_pred_values types of the lines for plotting the predicted values By default, line_types_for_pred_values = c("solid", "dashed") line_thickness_for_pred_values thickness of the lines for plotting the predicted values (default = 2.5) types of the lines for Johnson-Neyman points. By default, jn_line_types = jn_line_types c("solid", "solid") jn_line_thickness thickness of the lines at Johnson-Neyman points (default = 1.5) sig_region_color color of the significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is statistically significant. sig_region_alpha opacity for sig_region_color . (0 = completely transparent, 1 = completely opaque). By default, sig_region_alpha = 0.08 nonsig_region_color color of the non-significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is not statistically significant. nonsig_region_alpha opacity for nonsig_region_color. (0 = completely transparent, 1 = completelyopaque). By default, nonsig_region_alpha = 0.08 x_axis_title title of the x axis. By default, it will be set as input for mod_name. If x_axis_title = FALSE, it will be removed.

Details

```
See the following reference(s): Spiller et al. (2013) doi:10.1509/jmr.12.0420 Kim (2023) https://jinkim.science/docs/floodlight.pdf
```

Examples

```
floodlight_2_by_continuous_logistic(
data = mtcars,
iv_name = "am",
dv_name = "vs",
mod_name = "mpg")
# adjust the number of spotlights
# (i.e., predict values at only 4 values of the moderator)
floodlight_2_by_continuous_logistic(
data = mtcars,
iv_name = "am",
dv_name = "vs",
mod_name = "mpg",
num_of_spotlights = 4)
```

```
floodlight_2_by_continuous_mlm_logistic
```

Floodlight 2 by Continuous for a Multilevel Logistic Regression

Description

Conduct a floodlight analysis for a multilevellogistic regression with a 2 x Continuous design involving a binary dependent variable.

Usage

```
floodlight_2_by_continuous_mlm_logistic(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
 mod_name = NULL,
  interaction_p_include = TRUE,
  iv_level_order = NULL,
  dv_level_order = NULL,
  jn_points_disregard_threshold = NULL,
  output = "reg_lines_plot",
  num_of_spotlights = 20,
  jitter_x_percent = 0,
  jitter_y_percent = 5,
  dot_alpha = 0.3,
  dot_size = 6,
  interaction_p_value_font_size = 8,
  jn_point_font_size = 8,
  jn_point_label_hjust = NULL,
  interaction_p_vjust = -3,
  plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt"),
  legend_position = "right",
  line_types_for_pred_values = c("solid", "dashed"),
  line_thickness_for_pred_values = 2.5,
  jn_line_types = c("solid", "solid"),
  jn_line_thickness = 1.5,
  sig_region_color = "green",
  sig_region_alpha = 0.08,
  nonsig_region_color = "gray",
  nonsig_region_alpha = 0.08,
  x_axis_title = NULL,
  y_axis_title = NULL,
  legend_title = NULL,
  round_decimals_int_p_value = 3,
  round_jn_point_labels = 2
)
```

Arguments

```
data a data object (a data frame or a data.table)

iv_name name of the binary independent variable

dv_name name of the binary dependent variable

mod_name name of the continuous moderator variable

interaction_p_include

logical. Should the plot include a p-value for the interaction term?

iv_level_order order of levels in the independent variable for legend. By default, it will be set as levels of the independent variable ordered using R's base function sort.
```

dv_level_order order of levels in the dependent variable. By default, it will be set as levels of the dependent variable ordered using R's base function sort.

jn_points_disregard_threshold

the Minimum Distance in the unit of the moderator variable that will be used for various purposes, such as (1) to disregard the second Johnson-Neyman point that is different from the first Johnson-Neyman (JN) point by less than the Minimum Distance; (2) to determine regions of significance, which will calculate the p-value of the IV's effect (the focal dummy variable's effect) on DV at a candidate JN point + / - the Minimum Distance. This input is hard to explain, but a user can enter a really low value for this argument (e.g., jn_points_disregard_threshold = 0.1 for a moderator measured on a 100-point scale) or use the default. By default, jn_points_disregard_threshold = range of the moderator / 10000 For example, if the observed moderator values range from 1 to 7 (because it is a 7-point scale), then jn_points_disregard_threshold = (7 - 1) / 10000 = 0.0006

output

type of output (default = "reg_lines_plot"). Possible inputs: "interactions_pkg_results", "simple_effects_plot", "jn_points", "reg_lines_plot"

num_of_spotlights

How many spotlight analyses should be conducted to plot the predicted values at various values of the moderator? (default = 20)

jitter_x_percent

horizontally jitter dots by a percentage of the range of x values (default = 0)

jitter_y_percent

vertically jitter dots by a percentage of the range of y values (default = 5)

dot_alpha

opacity of the dots (0 = completely transparent, 1 = completely opaque). By default, dot_alpha = 0.3

dot_size

size of the dots (default = 6)

interaction_p_value_font_size

font size for the interaction p value (default = 8)

jn_point_font_size

font size for Johnson-Neyman point labels (default = 8)

jn_point_label_hjust

a vector of hjust values for Johnson-Neyman point labels. By default, the hjust value will be 0.5 for all the points.

interaction_p_vjust

By how much should the label for the interaction p-value be adjusted vertically? By default, interaction_p_vjust = -3)

plot_margin

margin for the plot By default plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt")

legend_position

position of the legend (default = "right"). If legend_position = "none", the legend will be removed.

line_types_for_pred_values

types of the lines for plotting the predicted values By default, line_types_for_pred_values = c("solid", "dashed")

```
line_thickness_for_pred_values
                  thickness of the lines for plotting the predicted values (default = 2.5)
                  types of the lines for Johnson-Neyman points. By default, jn_line_types =
jn_line_types
                  c("solid", "solid")
jn_line_thickness
                  thickness of the lines at Johnson-Neyman points (default = 1.5)
sig_region_color
                  color of the significant region, i.e., range(s) of the moderator variable for which
                  simple effect of the independent variable on the dependent variable is statisti-
                  cally significant.
sig_region_alpha
                  opacity for sig_region_color. (0 = completely transparent, 1 = completely
                  opaque). By default, sig_region_alpha = 0.08
nonsig_region_color
                  color of the non-significant region, i.e., range(s) of the moderator variable for
                  which simple effect of the independent variable on the dependent variable is not
                  statistically significant.
nonsig_region_alpha
                  opacity for nonsig_region_color. (0 = completely transparent, 1 = completely
                  opaque). By default, nonsig_region_alpha = 0.08
x_axis_title
                  title of the x axis. By default, it will be set as input for mod_name. If x_axis_title
                  = FALSE, it will be removed.
y_axis_title
                  title of the y axis. By default, it will be set as input for dv_name. If y_axis_title
                  = FALSE, it will be removed.
legend_title
                  title of the legend. By default, it will be set as input for iv_name. If legend_title
                  = FALSE, it will be removed.
round_decimals_int_p_value
                  To how many digits after the decimal point should the p value for the interaction
                  term be rounded? (default = 3)
round_jn_point_labels
                  To how many digits after the decimal point should the jn point labels be rounded?
```

Details

See the following reference(s): Spiller et al. (2013) doi:10.1509/jmr.12.0420 Kim (2023) https://jinkim.science/docs/floodlight.pdf

Examples

```
floodlight_2_by_continuous_logistic(
data = mtcars,
iv_name = "am",
dv_name = "vs",
mod_name = "mpg")
# adjust the number of spotlights
```

(default = 2)

```
# (i.e., predict values at only 4 values of the moderator)
floodlight_2_by_continuous_logistic(
data = mtcars,
iv_name = "am",
dv_name = "vs",
mod_name = "mpg",
num_of_spotlights = 4)
```

floodlight_for_contrasts

Floodlight Analyses for a Set of Contrasts

Description

Conduct a floodlight analysis for a set of contrasts with a continuous moderator variable.

```
floodlight_for_contrasts(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
 mod_name = NULL,
  contrasts = NULL,
  contrasts_for_floodlight = NULL,
  covariate_name = NULL,
  interaction_p_include = TRUE,
  iv_category_order = NULL,
  heteroskedasticity_consistent_se = "HC4",
  round_r_squared = 3,
  round_f = 2,
  sigfigs = 2,
  jn_points_disregard_threshold = NULL,
  print_floodlight_plots = TRUE,
  output = "reg_lines_plot",
  jitter_x_percent = 0,
  jitter_y_percent = 0,
  dot_alpha = 0.5,
  dot_size = 4,
  interaction_p_value_font_size = 6,
  jn_point_font_size = 6,
  jn_point_label_hjust = NULL,
  interaction_p_vjust = -3,
  plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt"),
  legend_position = "right",
  line_of_fit_types = c("solid", "dashed"),
```

```
line_of_fit_thickness = 1.5,
  jn_line_types = c("solid", "solid"),
  jn_line_thickness = 1.5,
  sig_region_color = "green",
  sig_region_alpha = 0.08,
  nonsig_region_color = "gray",
  nonsig_region_alpha = 0.08,
  x_axis_title = NULL,
  y_axis_title = NULL,
  legend_title = NULL,
  round_decimals_int_p_value = 3,
  round_jn_point_labels = 2
)
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the multicategorical independent variable; this variable must have three

or more categories.

dv_name name of the dependent variable

mod_name name of the continuous moderator variable

contrasts names of the contrast variables

contrasts_for_floodlight

names of the contrast variables for which floodlight analyses will be conducted

covariate_name name of the variables to control for

interaction_p_include

logical. Should the plot include a p-value for the interaction term?

iv_category_order

order of levels in the independent variable for legend. By default, it will be set as levels of the independent variable ordered using R's base function sort.

heteroskedasticity_consistent_se

which kind of heteroskedasticity-consistent (robust) standard errors should be calculated? (default = "HC4")

round_r_squared

number of decimal places to which to round r-squared values (default = 3)

round_f number of decimal places to which to round the f statistic for model comparison

(default = 2)

sigfigs number of significant digits to round to (for values in the regression tables, ex-

cept for p values). By default sigfigs = 2

jn_points_disregard_threshold

the Minimum Distance in the unit of the moderator variable that will be used for various purposes, such as (1) to disregard the second Johnson-Neyman point that is different from the first Johnson-Neyman (JN) point by less than the Minimum Distance; (2) to determine regions of significance, which will calculate the p-value of the IV's effect (the focal dummy variable's effect) on DV at a candidate JN point + / - the Minimum Distance. This input is hard to explain, but a user can

enter a really low value for this argument (e.g., jn_points_disregard_threshold = 0.1 for a moderator measured on a 100-point scale) or use the default. By default, jn_points_disregard_threshold = range of the moderator / 10000 For example, if the observed moderator values range from 1 to 7 (because it is a 7-point scale), then jn_points_disregard_threshold = (7 - 1) / 10000 = 0.0006

print_floodlight_plots

If print_floodlight_plots = TRUE, a floodlight plot for each dummy variable will be printed. By default, print_floodlight_plots = TRUE

output of the function (default = "all"). Possible inputs: "reg_models", "reg_tables", "reg_tables rounded", "all"

jitter_x_percent

horizontally jitter dots by a percentage of the range of x values

jitter_y_percent

vertically jitter dots by a percentage of the range of y values

dot_alpha opacity of the dots (0 = completely transparent, 1 = completely opaque). By default, dot_alpha = 0.5

 dot_size size of the dots (default = 4)

interaction_p_value_font_size

font size for the interaction p value (default = 8)

jn_point_font_size

font size for Johnson-Neyman point labels (default = 6)

jn_point_label_hjust

a vector of hjust values for Johnson-Neyman point labels. By default, the hjust value will be 0.5 for all the points.

interaction_p_vjust

By how much should the label for the interaction p-value be adjusted vertically? By default, interaction_p_vjust = -3)

plot_margin margin for the plot By default plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt")

legend_position

position of the legend (default = "right"). If legend_position = "none", the legend will be removed.

line_of_fit_types

types of the lines of fit for the two levels of the independent variable. By default, line_of_fit_types = c("solid", "dashed")

line_of_fit_thickness

thickness of the lines of fit (default = 1.5)

jn_line_thickness

thickness of the lines at Johnson-Neyman points (default = 1.5)

sig_region_color

color of the significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is statistically significant.

```
sig_region_alpha
                  opacity for sig_region_color. (0 = completely transparent, 1 = completely
                  opaque). By default, sig_region_alpha = 0.08
nonsig_region_color
                  color of the non-significant region, i.e., range(s) of the moderator variable for
                  which simple effect of the independent variable on the dependent variable is not
                  statistically significant.
nonsig_region_alpha
                  opacity for nonsig_region_color. (0 = completely transparent, 1 = completely
                  opaque). By default, nonsig_region_alpha = 0.08
                  title of the x axis. By default, it will be set as input for mod_name. If x_axis_title
x_axis_title
                  = FALSE, it will be removed.
                  title of the y axis. By default, it will be set as input for dv_name. If y_axis_title
y_axis_title
                  = FALSE, it will be removed.
                  title of the legend. By default, it will be set as input for iv_name. If legend_title
legend_title
                  = FALSE, it will be removed.
round_decimals_int_p_value
                  To how many digits after the decimal point should the p value for the interaction
                  term be rounded? (default = 3)
round_jn_point_labels
                  To how many digits after the decimal point should the jn point labels be rounded?
                  (default = 2)
```

Details

See the following reference, which covers a related topic: Hayes & Montoya (2017) doi:10.1080/19312458.2016.1271116

Examples

```
## Not run:
# typical example
# copy and modify the 'mtcars' data
mtcars2 <- setDT(data.table::copy(mtcars))
# make sure the data table package is attached
mtcars2[, contrast_1 := fcase(cyl == 4, -2, cyl %in% c(6, 8), 1)]
mtcars2[, contrast_2 := fcase(cyl == 4, 0, cyl == 6, 1, cyl == 8, -1)]
floodlight_for_contrasts(
data = mtcars2,
iv_name = "cyl",
dv_name = "mpg",
mod_name = "qsec",
contrasts = paste0("contrast_", 1:2),
contrasts_for_floodlight = "contrast_2")
## End(Not run)</pre>
```

floodlight_multi_by_continuous

Floodlight Multicategorical by Continuous

Description

Conduct a floodlight analysis for a Multicategorical IV x Continuous Moderator design.

```
floodlight_multi_by_continuous(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
 mod_name = NULL,
  coding = "indicator",
 baseline_category = NULL,
  covariate_name = NULL,
  interaction_p_include = TRUE,
  iv_category_order = NULL,
  heteroskedasticity_consistent_se = "HC4",
  round_r_squared = 3,
  round_f = 2,
  sigfigs = 2,
  jn_points_disregard_threshold = NULL,
  print_floodlight_plots = TRUE,
  output = "all",
  jitter_x_percent = 0,
  jitter_y_percent = 0,
  dot_alpha = 0.5,
  dot_size = 4,
  interaction_p_value_font_size = 8,
  jn_point_font_size = 8,
  jn_point_label_hjust = NULL,
  interaction_p_vjust = -3,
  plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt"),
  legend_position = "right",
  line_of_fit_types = c("solid", "dashed"),
  line_of_fit_thickness = 1.5,
  jn_line_types = c("solid", "solid"),
  jn_line_thickness = 1.5,
  colors_for_iv = c("red", "blue").
  sig_region_color = "green",
  sig_region_alpha = 0.08,
  nonsig_region_color = "gray",
  nonsig_region_alpha = 0.08,
  x_axis_title = NULL,
```

```
y_axis_title = NULL,
  legend_title = NULL,
  round_decimals_int_p_value = 3,
  round_jn_point_labels = 2
)
```

Arguments

data a data object (a data frame or a data.table)

name of the multicategorical independent variable; this variable must have three iv_name

or more categories.

dv_name name of the dependent variable

name of the continuous moderator variable mod_name

coding name of the coding scheme to use; the current version of the function allows

only the "indicator" coding scheme. By default, coding = "indicator"

baseline_category

value of the independent variable that will be the reference value against which

other values of the independent variable will be compared

covariate_name name of the variables to control for

interaction_p_include

logical. Should the plot include a p-value for the interaction term?

iv_category_order

order of levels in the independent variable for legend. By default, it will be set as levels of the independent variable ordered using R's base function sort.

heteroskedasticity_consistent_se

which kind of heteroskedasticity-consistent (robust) standard errors should be

calculated? (default = "HC4")

round_r_squared

number of decimal places to which to round r-squared values (default = 3)

round_f number of decimal places to which to round the f statistic for model comparison

(default = 2)

sigfigs number of significant digits to round to (for values in the regression tables, ex-

cept for p values). By default sigfigs = 2

jn_points_disregard_threshold

the Minimum Distance in the unit of the moderator variable that will be used for various purposes, such as (1) to disregard the second Johnson-Neyman point that is different from the first Johnson-Neyman (JN) point by less than the Minimum Distance; (2) to determine regions of significance, which will calculate the pvalue of the IV's effect (the focal dummy variable's effect) on DV at a candidate JN point + / - the Minimum Distance. This input is hard to explain, but a user can enter a really low value for this argument (e.g., jn_points_disregard_threshold = 0.1 for a moderator measured on a 100-point scale) or use the default. By default, in_points_disregard_threshold = range of the moderator / 10000 For example, if the observed moderator values range from 1 to 7 (because it is a 7-point scale), then jn_points_disregard_threshold = (7 - 1) / 10000 = 0.0006

print_floodlight_plots If print_floodlight_plots = TRUE, a floodlight plot for each dummy variable will be printed. By default, print_floodlight_plots = TRUE output of the function (default = "all"). Possible inputs: "reg_models", "reg_tables", output "reg tables rounded", "all" jitter_x_percent horizontally jitter dots by a percentage of the range of x values jitter_y_percent vertically jitter dots by a percentage of the range of y values dot_alpha opacity of the dots (0 = completely transparent, 1 = completely opaque). By default, dot_alpha = 0.5 dot_size size of the dots (default = 4) interaction_p_value_font_size font size for the interaction p value (default = 8) jn_point_font_size font size for Johnson-Neyman point labels (default = 8) jn_point_label_hjust a vector of hjust values for Johnson-Neyman point labels. By default, the hjust value will be 0.5 for all the points. interaction_p_vjust By how much should the label for the interaction p-value be adjusted vertically? By default, interaction_p_vjust = -3) plot_margin margin for the plot By default plot_margin = ggplot2::unit(c(75, 7, 7, 7), "pt") legend_position position of the legend (default = "right"). If legend_position = "none", the legend will be removed. line_of_fit_types types of the lines of fit for the two levels of the independent variable. By default, line_of_fit_types = c("solid", "dashed") line_of_fit_thickness thickness of the lines of fit (default = 1.5) jn_line_types types of the lines for Johnson-Neyman points. By default, jn_line_types = c("solid", "solid") jn_line_thickness thickness of the lines at Johnson-Neyman points (default = 1.5) colors for the two values of the independent variable (default = c("red", "blue")) colors_for_iv sig_region_color color of the significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is statistically significant. sig_region_alpha opacity for sig_region_color . (0 = completely transparent, 1 = completely

opaque). By default, sig_region_alpha = 0.08

forest_plot 61

```
nonsig_region_color
```

color of the non-significant region, i.e., range(s) of the moderator variable for which simple effect of the independent variable on the dependent variable is not statistically significant.

nonsig_region_alpha

opacity for nonsig_region_color. (0 = completely transparent, 1 = completely opaque). By default, nonsig_region_alpha = 0.08

x_axis_title title of the x axis. By default, it will be set as input for mod_name. If x_axis_title = FALSE, it will be removed.

y_axis_title title of the y axis. By default, it will be set as input for dv_name. If y_axis_title = FALSE, it will be removed.

legend_title title of the legend. By default, it will be set as input for iv_name. If legend_title = FALSE, it will be removed.

round_decimals_int_p_value

To how many digits after the decimal point should the p value for the interaction term be rounded? (default = 3)

round_jn_point_labels

To how many digits after the decimal point should the jn point labels be rounded? (default = 2)

Details

See the following reference: Hayes & Montoya (2017) doi:10.1080/19312458.2016.1271116 Williams (2004) on r-squared values when calculating robust standard errors https://web.archive.org/web/20230627025457/https://www.stata.com/statalist/archive/2004-05/msg00107.html

Examples

```
## Not run:
# typical example
floodlight_multi_by_continuous(
data = mtcars,
iv_name = "cyl",
dv_name = "mpg",
mod_name = "qsec")
## End(Not run)
```

forest_plot

Forest plot

Description

Create a forest plot using outputs from 'metafor' package

62 forest_plot

Usage

```
forest_plot(
  estimates = NULL,
  estimate_ci_ll = NULL,
  estimate_ci_ul = NULL,
  point_size_range = c(2, 10),
  error_bar_size = 1,
  error_bar_tip_height = 0.3,
  weights = NULL,
  diamond_x = NULL,
  diamond_ci_11 = NULL,
  diamond_ci_ul = NULL,
  diamond_height = 1.2,
  diamond_gap_height = 0.3,
  diamond_1_tip_at_top_y = -0.5,
  diamond_colors = "black",
  study_labels = NULL,
  diamond_labels = NULL,
  diamond_label_size = 6,
  diamond_label_hjust = 0,
  diamond_label_fontface = "bold",
  diamond_estimate_label_hjust = 0,
  diamond_estimate_label_size = 6,
  diamond_estimate_label_fontface = "bold",
  round_estimates = 2,
  x_axis_title = "Observed Outcome",
  vline_size = 1,
  vline_intercept = 0,
  vline_type = "dotted",
  study_label_hjust = 0,
  study_label_begin_x = NULL,
  study_label_begin_x_perc = 60,
  study_label_size = 6,
  study_label_fontface = "plain",
  estimate_label_begin_x = NULL,
  estimate_label_begin_x_perc = 25,
  estimate_label_hjust = 0,
  estimate_label_size = 6,
  estimate_label_fontface = "plain",
  x_axis_tick_marks = NULL,
  x_axis_tick_mark_label_size = 6,
  legend_position = "none",
  plot_margin = NULL
)
```

Arguments

estimates default = NULL

forest_plot 63

```
estimate_ci_ll default = NULL
estimate_ci_ul default = NULL
point_size_range
                 default = c(2, 10)
error_bar_size default = 1
error_bar_tip_height
                 default = 0.3
                 default = NULL
weights
diamond_x
                default = NULL
diamond_ci_ll
                default = NULL
diamond_ci_ul
                default = NULL
diamond_height default = 1.2
diamond_gap_height
                 default = 0.3
diamond_1_tip_at_top_y
                default = -0.5
diamond_colors default = "black"
study_labels
                 default = NULL
diamond_labels default = NULL
diamond_label_size
                 default = 6
diamond_label_hjust
                 default = 0
diamond_label_fontface
                default = "bold"
diamond_estimate_label_hjust
                 default = 0
diamond_estimate_label_size
                 default = 6
{\tt diamond\_estimate\_label\_fontface}
                 default = "bold"
round_estimates
                 default = 2
x_axis_title
                 default = "Observed Outcome"
vline_size
                 default = 1
vline_intercept
                 default = 0
                 default = "dotted"
vline_type
study_label_hjust
                 default = 0
study_label_begin_x
                 default = NULL
study_label_begin_x_perc
                 default = 60
```

64 geomean

```
study_label_size
                 default = 6
study_label_fontface
                 default = "plain"
estimate_label_begin_x
                 default = NULL
estimate_label_begin_x_perc
                 default = 25
estimate_label_hjust
                 default = 0
estimate_label_size
                 default = 6
estimate_label_fontface
                 default = "plain"
x_axis_tick_marks
                 default = NULL
x_axis_tick_mark_label_size
                 default = 6
{\tt legend\_position}
                 default = "none"
                 default = NULL
plot_margin
```

Examples

```
forest_plot(
  estimates = c(2, 3, 4),
  estimate_ci_ll = c(1, 2, 3),
  estimate_ci_ul = c(3, 4, 6),
  weights = 1:3,
  diamond_x = 2,
  diamond_labels = "RE",
  diamond_ci_ll = 1.8,
  diamond_ci_ul = 2.2,
  estimate_label_begin_x_perc = 40,
  x_axis_tick_marks = seq(-2, 6, 2))
```

geomean

Geometric mean

Description

Calculate the geometric mean of a numeric vector

```
geomean(x = NULL, zero_or_neg_convert_to = NA)
```

ggsave_quick 65

Arguments

```
x a numeric vector
zero_or_neg_convert_to
the value to which zero or negative values will be converted to. If zero_or_neg_convert_to
== NA, zero or negative values will be converted to NA values and thus be excluded when calculating the geometric mean. (default = NA)
```

Examples

```
## Not run:
geomean(c(1, 4))
geomean(c(1, 100))
geomean(c(1, 100, NA))
geomean(c(1, 100, NA, 0, -1, -2))
geomean(
x = c(1, 100, NA, 0, -1, -2),
zero_or_neg_convert_to = 1)
geomean(c(1, 100, NA, 1, 1, 1))
## End(Not run)
```

ggsave_quick

ggsave quick

Description

quickly save the current plot with a timestamp

Usage

```
ggsave_quick(
  name = NULL,
  file_name_extension = "png",
  timestamp = NULL,
  width = 16,
  height = 9
)
```

Arguments

name

a character string of the png file name. By default, if no input is given (name = NULL), the file name will begin with "ggplot". If the desired output file name is "myplot.png", enter name = "myplot", timestamp = FALSE

```
file_name_extension
```

file name extension (default = "png"). If file_name_extension = "svg", Package syglite needs to be installed.

66 histogram

if timestamp = TRUE, a timestamp of the current time will be appended to the file name. The timestamp will be in the format, jan_01_2021_1300_10_000001, where "jan_01_2021" would indicate January 01, 2021; 1300 would indicate 13:00 (i.e., 1 PM); and 10_000001 would indicate 10.000001 seconds after the hour. By default, timestamp will be set as TRUE, if no input is given for the name argument, and as FALSE, if an input is given for the name argument.

width width of the plot to be saved. This argument will be directly entered as the width argument for the ggsave function within ggplot2 package (default = 16)

height height argument for the ggsave function within ggplot2 package (default = 9)

Value

the output will be a .png image file in the working directory.

Examples

```
## Not run:
kim::histogram(rep(1:30, 3))
ggsave_quick()
## End(Not run)
```

histogram

Histogram

Description

Create a histogram based on the output of the hist function in the graphics package.

```
histogram(
  vector = NULL,
  breaks = NULL,
  counts = NULL,
  percent = FALSE,
  bin_fill_color = "green4",
  bin_border_color = "black",
  bin_border_thickness = 1,
  notify_na_count = NULL,
  x_axis_tick_marks = NULL,
  y_axis_tick_marks = NULL,
  cap_axis_lines = TRUE,
  x_axis_title = "Value",
  y_axis_title = NULL,
  y_axis_title_vjust = 0.85
)
```

histogram 67

Arguments

```
vector
                  a numeric vector
breaks
                  a numeric vector indicating breaks for the bins. By default, no input is required
                  for this argument.
                  a numeric vector containing counts for the bins (i.e., heights of the bins). By
counts
                  default, no input is required for this argument.
                  logical. If percent = TRUE, percentages will be plotted rather than frequencies
percent
                  (default = FALSE).
bin_fill_color color of the area inside each bin (default = "green4")
bin_border_color
                  color of the border around each bin (default = "black")
bin_border_thickness
                  thickness of the border around each bin (default = 1)
notify_na_count
                  if TRUE, notify how many observations were removed due to missing values. By
                  default, NA count will be printed only if there are any NA values.
x_axis_tick_marks
                  a vector of values at which to place tick marks on the x axis (e.g., setting
                  x_axis_tick_marks = seq(0, 10, 5) will put tick marks at 0, 5, and 10.)
y_axis_tick_marks
                  a vector of values at which to place tick marks on the y axis (e.g., setting
                  y_axis_tick_marks = seq(0, 10, 5) will put tick marks at 0, 5, and 10.)
cap_axis_lines logical. Should the axis lines be capped at the outer tick marks? (default =
                  FALSE)
x_axis_title
                  title for x axis (default = "Value")
y_axis_title
                  title for y axis (default = "Count" or "Percentage", depending on the value of
                  percent)
y_axis_title_vjust
                  position of the y axis title (default = 0.85).
```

Value

the output will be a histogram, a ggplot object.

Examples

```
histogram(1:100)
histogram(c(1:100, NA))
histogram(vector = mtcars[["mpg"]])
histogram(vector = mtcars[["mpg"]], percent = TRUE)
histogram(vector = mtcars[["mpg"]],
x_axis_tick_marks = c(10, 25, 35), y_axis_title_vjust = 0.5,
y_axis_title = "Freq", x_axis_title = "Values of mpg")
```

68 histogram_by_group

histogram_by_group Histogram by group

Description

Creates histograms by group to compare distributions.

Usage

```
histogram_by_group(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  order_of_groups_top_to_bot = NULL,
  number_of_bins = 40,
  space_between_histograms = 0.15,
  draw_baseline = FALSE,
  xlab = NULL,
  ylab = NULL,
  x_limits = NULL,
  x_breaks = NULL,
  x_{labels} = NULL,
  sigfigs = 3,
  convert_dv_to_numeric = TRUE
)
```

Arguments

data	a data object (a data frame or a data.table)		
iv_name	name of the independent variable		
dv_name	name of the dependent variable		
order_of_groups	der_of_groups_top_to_bot		
	a character vector indicating the desired presentation order of levels in the in- dependent variable (from the top to bottom). Omitting a group in this argument will remove the group in the set of histograms.		
number_of_bins	number of bins for the histograms (default = 40)		
space_between_histograms			
	space between histograms (minimum = 0 , maximum = 1 , default = 0.15)		
draw_baseline	logical. Should the baseline and the trailing lines to either side of the histogram be drawn? (default = FALSE)		
xlab	title of the x-axis for the histogram by group. If $xlab = FALSE$, the title will be removed. By default (i.e., if no input is given), dv_n will be used as the title.		
ylab	title of the y-axis for the histogram by group. If ylab = FALSE, the title will be removed. By default (i.e., if no input is given), iv_name will be used as the title.		

logical. Should the values in the dependent variable be converted to numeric for plotting the histograms? (default = TRUE)

Details

The following package(s) must be installed prior to running this function: Package 'ggridges' v0.5.3 (or possibly a higher version) by Claus O. Wilke (2021), https://cran.r-project.org/package=ggridges

Value

the output will be a set of vertically arranged histograms (a ggplot object), i.e., one histogram for each level of the independent variable.

Examples

```
histogram_by_group(data = mtcars, iv_name = "cyl", dv_name = "mpg")
histogram_by_group(
  data = mtcars, iv_name = "cyl", dv_name = "mpg",
  order_of_groups_top_to_bot = c("8", "4"), number_of_bins = 10,
  space_between_histograms = 0.5
)
histogram_by_group(
data = iris, iv_name = "Species", dv_name = "Sepal.Length", x_breaks = 4:8,
  x_limits = c(4, 8))
```

histogram_deprecated_1

Histogram

Description

Create a histogram

```
histogram_deprecated_1(
  vector = NULL,
  number_of_bins = 30,
  x_tick_marks = NULL,
```

```
y_tick_marks = NULL,
fill_color = "cyan4",
border_color = "black",
y_axis_title_vjust = 0.85,
x_axis_title = NULL,
y_axis_title = NULL,
cap_axis_lines = FALSE,
notify_na_count = NULL)
```

Arguments

```
vector
                  a numeric vector
number_of_bins number of bins for the histogram (default = 30)
x_tick_marks
                  a vector of values at which to place tick marks on the x axis (e.g., setting
                  x_{tick_marks} = seq(0, 10, 5) will put tick marks at 0, 5, and 10.)
                  a vector of values at which to place tick marks on the y axis (e.g., setting
y_tick_marks
                  y_{tick_marks} = seq(0, 10, 5) will put tick marks at 0, 5, and 10.)
fill_color
                  color for inside of the bins (default = "cyan4")
border_color
                  color for borders of the bins (default = "black")
y_axis_title_vjust
                  position of the y axis title (default = 0.85).
                  title for x axis (default = "Value")
x_axis_title
y_axis_title
                  title for y axis (default = "Count")
cap_axis_lines logical. Should the axis lines be capped at the outer tick marks? (default =
                  FALSE)
notify_na_count
                  if TRUE, notify how many observations were removed due to missing values. By
                  default, NA count will be printed only if there are any NA values.
```

Value

the output will be a histogram, a ggplot object.

Examples

```
histogram_deprecated_1(1:100)
histogram_deprecated_1(c(1:100, NA))
histogram_deprecated_1(vector = mtcars[["mpg"]])
histogram_deprecated_1(
vector = mtcars[["mpg"]], x_tick_marks = seq(10, 36, 2))
histogram_deprecated_1(
vector = mtcars[["mpg"]], x_tick_marks = seq(10, 36, 2),
y_tick_marks = seq(0, 8, 2), y_axis_title_vjust = 0.5,
y_axis_title = "Freq", x_axis_title = "Values of mpg")
```

histogram_from_hist 71

histogram_from_hist Histogram from hist function

Description

Create a histogram based on the output of the hist function in the graphics package.

Usage

```
histogram_from_hist(
  vector = NULL,
  breaks = NULL,
  counts = NULL,
  percent = FALSE,
  bin_fill_color = "green4",
  bin_border_color = "black",
  bin_border_thickness = 1,
  notify_na_count = NULL,
  x_axis_tick_marks = NULL,
  y_axis_tick_marks = NULL,
  cap_axis_lines = TRUE,
  x_axis_title = "Value",
 y_axis_title = NULL,
 y_axis_title_vjust = 0.85
)
```

Arguments

```
vector
                  a numeric vector
                  a numeric vector indicating breaks for the bins. By default, no input is required
breaks
                  for this argument.
                  a numeric vector containing counts for the bins (i.e., heights of the bins). By
counts
                  default, no input is required for this argument.
                  logical. If percent = TRUE, percentages will be plotted rather than frequencies
percent
                  (default = FALSE).
bin_fill_color color of the area inside each bin (default = "green4")
bin_border_color
                  color of the border around each bin (default = "black")
bin_border_thickness
                  thickness of the border around each bin (default = 1)
notify_na_count
                  if TRUE, notify how many observations were removed due to missing values. By
                  default, NA count will be printed only if there are any NA values.
x_axis_tick_marks
                  a vector of values at which to place tick marks on the x axis (e.g., setting
                  x_axis_tick_marks = seq(0, 10, 5) will put tick marks at 0, 5, and 10.)
```

Value

the output will be a histogram, a ggplot object.

Examples

```
histogram_from_hist(1:100)
histogram_from_hist(c(1:100, NA))
histogram_from_hist(vector = mtcars[["mpg"]])
histogram_from_hist(vector = mtcars[["mpg"]], percent = TRUE)
histogram_from_hist(vector = mtcars[["mpg"]],
x_axis_tick_marks = c(10, 25, 35), y_axis_title_vjust = 0.5,
y_axis_title = "Freq", x_axis_title = "Values of mpg")
```

histogram_w_outlier_bins

Histogram with outlier bins

Description

Create a histogram with outlier bins

```
histogram_w_outlier_bins(
  vector = NULL,
  bin_cutoffs = NULL,
  outlier_bin_left = TRUE,
  outlier_bin_right = TRUE,
  x_tick_marks = NULL,
  x_tick_mark_labels = NULL,
  y_tick_marks = NULL,
  outlier_bin_fill_color = "coral",
  non_outlier_bin_fill_color = "cyan4",
```

```
border_color = "black",
      y_axis_title_vjust = 0.85,
      x_axis_title = NULL,
      y_axis_title = NULL,
      notify_na_count = NULL,
      plot_proportion = TRUE,
      plot_frequency = FALSE,
      mean = TRUE,
      ci = TRUE,
      median = TRUE,
      median_position = 15,
      error_bar_size = 3
    )
Arguments
    vector
                       a numeric vector
    bin_cutoffs
                       cutoff points for bins
    outlier_bin_left
                       logical. Should the leftmost bin treated as an outlier bin? (default = TRUE)
    outlier_bin_right
                      logical. Should the rightmost bin treated as an outlier bin? (default = TRUE)
                       a vector of values at which to place tick marks on the x axis. Note that the first
    x_tick_marks
                       bar spans from 0.5 to 1.5, second bar from 1.5 to 2.5, ... nth bar from n - 0.5 to
                       n + 0.5. See the example. By default, tick marks will be placed at every cutoff
                       point for bins
    x_tick_mark_labels
                       a character vector to label tick marks. By default, the vector of cutoff points for
                       bins will also be used as labels.
                       a vector of values at which to place tick marks on the y axis (e.g., setting
    y_tick_marks
                       y_{\text{tick\_marks}} = \text{seq}(0, 10, 5) will put tick marks at 0, 5, and 10.)
    outlier_bin_fill_color
                      color to fill inside of the outlier bins (default = "coral")
    non_outlier_bin_fill_color
                       color to fill inside of the non-outlier bins (default = "cyan4")
    border_color
                       color for borders of the bins (default = "black")
    y_axis_title_vjust
                       position of the y axis title (default = 0.85).
                       title for x axis (default = "Value"). If x_axis_title = FALSE, x axis title will be
    x_axis_title
                       removed from the plot.
    y_axis_title
                       title for y axis. By default, it will be either "Proportion" or "Count".
    notify_na_count
                       if TRUE, notify how many observations were removed due to missing values. By
```

default, NA count will be printed only if there are any NA values.

74 holm_adjusted_p

plot_proportion

logical. Should proportions be plotted, as opposed to frequencies? (default =

TRUE)

plot_frequency logical. Should frequencies be plotted, as opposed to proportions? (default

= FALSE). If plot_frequency = TRUE, plot_proportion will switch to be

FALSE.

mean logical. Should mean marked on the histogram? (default = TRUE)

ci logical. Should 95% confidence interval marked on the histogram? (default =

TRUE)

median logical. Should median marked on the histogram? (default = TRUE)

median_position

position of the median label as a percentage of height of the tallest bin (default

= 15)

error_bar_size size of the error bars (default = 3)

Value

a ggplot object

Examples

```
histogram_w_outlier_bins(vector = 1:100, bin_cutoffs = seq(0, 100, 10))
histogram_w_outlier_bins(vector = 0:89, bin_cutoffs = seq(0, 90, 10),
x_tick_marks = seq(0.5, 9.5, 3), x_tick_mark_labels = seq(0, 90, 30))
histogram_w_outlier_bins(vector = 1:10, bin_cutoffs = seq(0, 10, 2.5))
histogram_w_outlier_bins(vector = 1:5, bin_cutoffs = seq(0, 10, 2.5))
histogram_w_outlier_bins(vector = 1:15, bin_cutoffs = c(5.52, 10.5))
```

holm_adjusted_p

Holm-adjusted p-values

Description

Adjust a vector of p-values using the method proposed by Holm

Usage

```
holm_adjusted_p(p = NULL)
```

Arguments

р

a numeric vector of p-values

identical_all 75

Details

See the following reference: Holm 1979 https://www.jstor.org/stable/4615733 Manual for the 'p.adjust' function in the 'stats' package https://stat.ethz.ch/R-manual/R-devel/library/stats/html/p.adjust.html

Examples

```
holm_adjusted_p(c(.05, .01))
holm_adjusted_p(c(.05, .05, .05))
```

identical_all

Check whether all inputs are identical

Description

Check whether all inputs are identical

Usage

```
identical_all(...)
```

Arguments

... two or more R objects. If a vector or list is entered as an input, the function will test whether the vector's or list's elements are identical.

Value

the output will be TRUE if all inputs are identical or FALSE if not

```
identical_all(1:3, 1:3) # should return TRUE
identical_all(1:3, 1:3, 1:3, 1:3, 1:3) # should return TRUE
identical_all(1:3, 1:3, 1:3, 1:3, 1:4) # should return FALSE
identical_all(1:10) # should return FALSE
identical_all(rep(1, 100)) # should return TRUE
identical_all(list(1, 1, 1)) # should return TRUE
identical_all(TRUE, FALSE) # should return FALSE
identical_all(FALSE, TRUE) # should return FALSE
```

76 id_across_datasets

id_across_datasets ID across datasets

Description

Create an ID column in each of the data sets. The ID values will span across the data sets.

Usage

```
id_across_datasets(
  dt_list = NULL,
  id_col_name = "id",
  id_col_position = "first",
  silent = FALSE
)
```

Arguments

dt_list a list of data.table objects id col name name of the column that will contain ID values. By default, id_col_name = "id".

id_col_position

position of the newly created ID column. If id_col_position = "first", the new ID column will be placed as the first column in respective data sets. If id_col_position = "last", the new ID column will be placed as the last column in respective data sets.

silent If silent = TRUE, a summary of starting and ending ID values in each data set

will not be printed. If silent = FALSE, a summary of starting and ending ID

values in each data set will be printed. (default = FALSE)

Value

the output will be a list of data.table objects.

```
# running the examples below requires importing the data.table package.
prep(data.table)
id_across_datasets(
dt_list = list(setDT(copy(mtcars)), setDT(copy(iris))))
id_across_datasets(
dt_list = list(setDT(copy(mtcars)), setDT(copy(iris)), setDT(copy(women))),
id_col_name = "newly_created_id_col",
id_col_position = "last")
```

```
install_all_dependencies
```

Install all dependencies for all functions

Description

Install all dependencies for all functions in Package 'kim'.

Usage

```
install_all_dependencies()
```

Value

there will be no output from this function. Rather, dependencies of all functions in Package 'kim' will be installed.

Examples

```
## Not run:
install_all_dependencies()
## End(Not run)
```

kurtosis

Kurtosis

Description

Calculate kurtosis of the sample using a formula for either the (1) biased estimator or (2) an unbiased estimator of the population kurtosis. Formulas were taken from DeCarlo (1997), doi:10.1037/1082-989X.2.3.292

Usage

```
kurtosis(vector = NULL, unbiased = TRUE)
```

Arguments

vector a numeric vector

unbiased logical. If unbiased = TRUE, the unbiased estimate of the population kurtosis

will be calculated. If unbiased = FALSE, the biased estimate of the population

kurtosis will be calculated. By default, unbiase = TRUE.

Value

a numeric value, i.e., kurtosis of the given vector

78 lenu

Examples

```
# calculate the unbiased estimator (e.g., kurtosis value that
# Excel 2016 will produce)
kim::kurtosis(c(1, 2, 3, 4, 5, 10))
# calculate the biased estimator (e.g., kurtosis value that
# R Package 'moments' will produce)
kim::kurtosis(c(1, 2, 3, 4, 5, 10), unbiased = FALSE)
# compare with kurtosis from 'moments' package
moments::kurtosis(c(1, 2, 3, 4, 5, 10))
```

lenu

lenu: Length of unique values

Description

Extract unique elements and get the length of those elements

Usage

```
lenu(x = NULL)
```

Arguments

Х

a vector or a data frame or an array or NULL.

Value

a vector, data frame, or array-like 'x' but with duplicate elements/rows removed.

```
unique(c(10, 3, 7, 10))
lenu(c(10, 3, 7, 10))
unique(c(10, 3, 7, 10, NA))
lenu(c(10, 3, 7, 10, NA))
lenu(c("b", "z", "b", "a", NA, NA, NA))
```

levene_test 79

levene_test

Levene's test

Description

Conduct Levene's test (i.e., test the null hypothesis that the variances in different gorups are equal)

Usage

```
levene_test(
  data = NULL,
  dv_name = NULL,
  iv_1_name = NULL,
  iv_2_name = NULL,
  round_f = 2,
  round_p = 3,
  output_type = "text"
)
```

Arguments

data a data object (a data frame or a data.table) name of the dependent variable dv_name name of the first independent variable iv_1_name iv_2_name name of the second independent variable number of decimal places to which to round the F-statistic from Levene's test round_f (default = 2)number of decimal places to which to round the p-value from Levene's test (deround_p fault = 3If output_type = "text", the output will be the results of Levene's test in a text output_type format (i.e., character). If output_type = "list", the output will be the results of Levene's test in a list format (e.g., p value, F stat, etc. as a list). By default, output_type = "text"

Value

the output of the function depends on the input for output_type. By default, the output will be the results of Levene's test in a text format (i.e., character).

```
## Not run:
levene_test(
data = mtcars, dv_name = "mpg",
iv_1_name = "vs", iv_2_name = "am")
## End(Not run)
```

80 logistic_regression

Logistic regression logistic_regression

Description

Conduct a logistic regression analysis

Usage

```
logistic_regression(
  data = NULL,
  formula = NULL,
  formula_1 = NULL,
  formula_2 = NULL,
  z_values_keep = FALSE,
  constant_row_clean = TRUE,
  odds_ratio_cols_combine = TRUE,
  round_b_and_se = 3,
  round_z = 3,
  round_p = 3,
  round_odds_ratio = 3,
  round_r_sq = 3,
  round_model_chi_sq = 3,
  pretty_round_p_value = TRUE,
  print_glm_default_summary = FALSE,
 print_summary_dt_list = TRUE,
  print_model_comparison = TRUE,
 output_type = "summary_dt_list"
)
```

Arguments

data

```
a data object (a data frame or a data.table)
                  formula for estimating a single logistic regression model
formula
formula_1
                  formula for estimating logistic regression model 1 of 2
formula_2
                  formula for estimating logistic regression model 2 of 2
z_values_keep
                  logical. Should the z values be kept in the table? (default = FALSE)
constant_row_clean
                  logical. Should the row for the constant be cleared except for b and standard
                  error of b? (default = TRUE)
odds_ratio_cols_combine
                  logical. Should the odds ratio columns be combined? (default = TRUE)
round_b_and_se number of decimal places to which to round b and standard error of b (default =
                  3)
round_z
                  number of decimal places to which to round z values (default = 3)
```

logistic_regression 81

```
round_p
                  number of decimal places to which to round p-values (default = 3)
round_odds_ratio
                  number of decimal places to which to round odds ratios (default = 3)
round_r_sq
                  number of decimal places to which to round R-squared values (default = 3)
round_model_chi_sq
                  number of decimal places to which to round model chi-squared values (default
                  = 3)
pretty_round_p_value
                  logical. Should the p-values be rounded in a pretty format (i.e., lower threshold:
                  "<.001"). By default, pretty_round_p_value = TRUE.
print_glm_default_summary
                  logical. Should the default summary output of the glm objects be printed? (de-
                  fault = FALSE)
print_summary_dt_list
                  logical. Should the summaries of logistic regressions in a data table format be
                  printed? (default = TRUE)
print_model_comparison
                  logical. Should the comparison of two logistic regression models be printed?
                  (default = TRUE)
                  If output_type = "summary_dt_list" (default), the output of the function will
output_type
                  be summaries of the two logistic regressions in a data.table format. If output_type
                  = "glm_object_list", the output of the function will be the two glm objects es-
                  timating logistic regression models. If output_type = "glm_default_summary_list",
                  the output of the function will be the R's default summary output for the two glm
                  objects estimating logistic regression models. If output_type = "model_comparison_stats",
                  the output of the function will be statistics from comparison of the two logistic
                  regression models. If output_type = "all", the output of the function will be
                  a list of the aforementioned outputs.
```

Value

the output will be a summary of logistic regression results, unless set otherwise by the output_type argument to the function.

```
logistic_regression(data = mtcars, formula = am ~ mpg)
logistic_regression(
data = mtcars,
formula_1 = am ~ mpg,
formula_2 = am ~ mpg + wt)
```

```
logistic_regression_table
```

Logistic regression table

Description

Construct a table of logistic regression results from the given glm object estimating a logistic regression model.

Usage

```
logistic_regression_table(
  logistic_reg_glm_object = NULL,
  z_values_keep = FALSE,
  constant_row_clean = TRUE,
  odds_ratio_cols_combine = TRUE,
  round_b_and_se = 3,
  round_z = 3,
  round_p = 3,
  round_odds_ratio = 3,
  round_r_sq = 3,
  round_model_chi_sq = 3,
  pretty_round_p_value = TRUE
)
```

Arguments

```
logistic_reg_glm_object
                  a glm object estimating a logistic regression model
                 logical. Should the z values be kept in the table? (default = FALSE)
z_values_keep
constant_row_clean
                  logical. Should the row for the constant be cleared except for b and standard
                  error of b? (default = TRUE)
odds_ratio_cols_combine
                  logical. Should the odds ratio columns be combined? (default = TRUE)
round_b_and_se number of decimal places to which to round b and standard error of b (default =
round_z
                  number of decimal places to which to round z values (default = 3)
round_p
                  number of decimal places to which to round p-values (default = 3)
round_odds_ratio
                  number of decimal places to which to round odds ratios (default = 3)
round_r_sq
                  number of decimal places to which to round R-squared values (default = 3)
round_model_chi_sq
                  number of decimal places to which to round model chi-squared values (default
                  = 3)
```

```
\label{logical} \begin{tabular}{l} pretty\_round\_p\_value \\ logical. Should the p-values be rounded in a pretty format (i.e., lower threshold: "<.001"). By default, pretty\_round\_p\_value = TRUE. \\ \end{tabular}
```

Value

the output will be a summary of logistic regression results.

Examples

```
logistic_regression_table(logistic_reg_glm_object =
glm(formula = am ~ mpg, family = binomial(), data = mtcars))
logistic_regression_table(logistic_reg_glm_object =
glm(formula = am ~ mpg, family = binomial(), data = mtcars),
z_values_keep = TRUE, constant_row_clean = FALSE,
odds_ratio_cols_combine = FALSE)
```

logistic_reg_w_interaction

Logistic regression with an interaction term

Description

Conduct logistic regression for a model with an interaction between two predictor variables

Usage

```
logistic_reg_w_interaction(
  data = NULL,
  dv_name = NULL,
  iv_1_name = NULL,
  iv_2_name = NULL,
  round_p = 3,
  round_chi_sq = 2,
  dv_ordered_levels = NULL,
  iv_1_ordered_levels = NULL,
  iv_2_ordered_levels = NULL,
  one_line_summary_only = FALSE,
  p_value_interaction_only = FALSE,
  return_dt_w_binary = FALSE
```

Arguments

```
data a data object (a data frame or a data.table)

dv_name name of the dependent variable (must be a binary variable)

iv_1_name name of the first independent variable
```

84 loglinear_analysis

iv_2_name name of the second independent variable

round_p number of decimal places to which to round p-values (default = 3)

round_chi_sq number of decimal places to which to round chi square statistics (default = 2) dv_ordered_levels

a vector with the ordered levels of the dependent variable, the first and second elements of which will be coded as 0 and 1, respectively, to run logistic regression. E.g., dv_ordered_levels = c("fail", "pass")

iv_1_ordered_levels

(only if the first independent variable is a binary variable) a vector with the ordered levels of the first independent variable, the first and second elements of which will be coded as 0 and 1, respectively, to run logistic regression. E.g., iv_1_ordered_levels = c("control", "treatment")

iv_2_ordered_levels

(only if the second independent variable is a binary variable) a vector with the ordered levels of the first independent variable, the first and second elements of which will be coded as 0 and 1, respectively, to run logistic regression. E.g., iv_2_ordered_levels = c("male", "female")

one_line_summary_only

logical. Should the output simply be a printout of a one-line summary on the interaction term? (default = FALSE)

p_value_interaction_only

logical. Should the output simply be a p-value of the interaction term in the logistic regression model? (default = FALSE)

return_dt_w_binary

logical. If return_dt_w_binary = TRUE, the function will return a data.table with binary variables coded as 0 or 1 (default = FALSE)

Value

the output will be a summary of logistic regression results, unless set otherwise by arguments to the function.

Examples

```
logistic_reg_w_interaction(data = mtcars, dv_name = "vs",
iv_1_name = "mpg", iv_2_name = "am")
```

loglinear_analysis

Loglinear analysis

Description

Conduct a loglinear analysis

loglinear_analysis 85

Usage

```
loglinear_analysis(
  data = NULL,
  dv_name = NULL,
  iv_1_name = NULL,
  iv_2_name = NULL,
  iv_1_values = NULL,
  iv_2_values = NULL,
  output = "all",
  round_p = 3,
  round_chi_sq = 2,
  mosaic_plot = TRUE,
  report_as_field = FALSE
)
```

Arguments

data	a data object (a data frame or a data.table)
dv_name	name of the dependent variable
iv_1_name	name of the first independent variable
iv_2_name	name of the second independent variable
iv_1_values	restrict all analyses to observations having these values for the first independent variable
iv_2_values	restrict all analyses to observations having these values for the second independent variable
output	type of the output. If output_type = "all", the function will return a results summary and print a mosaic plot. (default = "all")
round_p	number of decimal places to which to round p-values (default = 3)
round_chi_sq	number of decimal places to which to round chi-squared test statistics (default = 2)
mosaic_plot	If mosaic_plot = TRUE, a mosaic plot will be printed (default = TRUE)
report_as_field	
	If report_as_field = TRUE, reports summary will follow the format suggested by Andy Field (2012) (ISBN: 978-1-4462-0045-2, p. 851)

```
loglinear_analysis(data = data.frame(Titanic), "Survived", "Sex", "Age")
```

86 log_odds_ratio

log_odds_ratio

Log odds ratio

Description

Calculate log odds ratio (i.e., ln of odds ratio), as illustrated in Borenstein et al. (2009, p. 36, ISBN: 978-0-470-05724-7)

Usage

```
log_odds_ratio(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  contingency_table = NULL,
  ci = 0.95,
  var_include = FALSE,
  invert = FALSE
)
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the independent variable (grouping variable)

dv_name name of the dependent variable (binary outcome)

contingency_table

a contingency table, which can be directly entered as an input for calculating the

odds ratio

ci width of the confidence interval. Input can be any value less than 1 and greater

than or equal to 0. By default, ci = 0.95. If ci = TRUE, the default value of 0.95

will be used. If ci = FALSE, no confidence interval will be estimated.

var_include logical. Should the output include variance of the log of odds ratio? (default =

FALSE)

invert logical. Whether the inverse of the odds ratio (i.e., 1 / odds ratio) should be

returned.

```
## Not run:
log_odds_ratio(data = mtcars, iv_name = "vs", dv_name = "am")
log_odds_ratio(contingency_table = matrix(c(5, 10, 95, 90), nrow = 2))
log_odds_ratio(contingency_table = matrix(c(5, 10, 95, 90), nrow = 2),
invert = TRUE)
log_odds_ratio(contingency_table = matrix(c(34, 39, 16, 11), nrow = 2))
log_odds_ratio(contingency_table = matrix(c(34, 39, 16, 11), nrow = 2),
var_include = TRUE)
```

log_odds_ratio_to_d 87

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

```
log_odds_ratio_to_d
Convert log odds ratio to Cohen's d
```

Description

Convert log odds ratio to Cohen'd (standardized mean difference), as illustrated in Borenstein et al. (2009, p. 47, ISBN: 978-0-470-05724-7)

Usage

```
log_odds_ratio_to_d(log_odds_ratio = NULL, unname = TRUE)
```

Arguments

```
log_odds_ratio log odds ratio (the input can be a vector of values), which will be converted to Cohen's d
unname logical. Should the names from the input be removed? (default = TRUE)
```

Examples

```
## Not run:
log_odds_ratio_to_d(log(1))
log_odds_ratio_to_d(log(2))
## End(Not run)
```

mad_remove_outliers

Remove outliers using the MAD method

Description

Detect outliers in a numeric vector using the Median Absolute Deviation (MAD) method and remove or convert them. For more information on MAD, see Leys et al. (2013) doi:10.1016/j.jesp.2013.03.013

```
mad_remove_outliers(
  x = NULL,
  threshold = 2.5,
  constant = 1.4826,
  convert_outliers_to = NA,
  output_type = "converted_vector"
)
```

Arguments

x a numeric vector

threshold the threshold value for determining outliers. If threshold == 2.5, the median

plus or minus 2.5 times the MAD will be the cutoff values for determining outliers. In other words, values less than the median minus 2.5 times the MAD and values greater than the median plus 2.5 times the MAD will be considered

outliers. By default, threshold == 2.5

constant scale factor for the 'mad' function in the 'stats' package. It is the constant linked

to the assumed distribution. In case of normality, constant = 1.4826. By default,

constant == 1.4826.

convert_outliers_to

the value to which outliers will be converted. For example, if convert_outliers_to = NA, the outlier values will be converted to NA values. If convert_outliers_to = 1000, the outlier values will be converted to 1000. By default, convert_outliers_to

== NA.

output_type

type of the output. If output_type = "converted_vector", the function's output will be a vector with outliers converted to the value set by the argument convert_outliers_to. If output_type = "outliers", the function's output will be outliers in the original vector as determined by the MAD method. If output_type = "cutoff_values", the function's output will be the cutoff values for determining outliers. For example, if outliers will be values less than 0 and greater than 10, the cutoff values will be 0 and 10. If output_type = "non_outlier_values", the function's output will be a vector consisting only of the values that are not outliers; here, the outliers will be removed from the vector, rather than being converted to NA values. By default, output_type = "converted_vector".

```
## Not run:
mad\_remove\_outliers(x = c(1, 3, 3, 6, 8, 10, 10, 1000))
mad_remove_outliers(x = c(1, 3, 3, 6, 8, 10, 10, 1000, -10000))
# return the vector with the outlier converted to NA values
mad_remove_outliers(
x = c(1, 3, 3, 6, 8, 10, 10, 1000, -10000),
output_type = "converted_vector")
# return the cutoff values for determining outliers
mad_remove_outliers(
x = c(1, 3, 3, 6, 8, 10, 10, 1000, -10000),
output_type = "cutoff_values")
# return the outliers
mad_remove_outliers(
x = c(1, 3, 3, 6, 8, 10, 10, 1000, -10000),
output_type = "outliers")
mad_remove_outliers(
x = c(1, 3, 3, 6, 8, 10, 10, 1000, -10000),
output_type = "non_outlier_values")
## End(Not run)
```

mann_whitney 89

mann_whitney

Mann-Whitney U Test (Also called Wilcoxon Rank-Sum Test)

Description

A nonparametric equivalent of the independent t-test

Usage

```
mann_whitney(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  iv_level_order = NULL,
  sigfigs = 3
)
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the independent variable (grouping variable)

dv_name name of the dependent variable (measure variable of interest)

iv_level_order order of levels in the independent variable. By default, it will be set as levels of

the independent variable ordered using R's base function sort.

sigfigs number of significant digits to round to

Value

the output will be a data.table object with all pairwise Mann-Whitney test results

Examples

```
mann_whitney(data = iris, iv_name = "Species", dv_name = "Sepal.Length")
```

matrix_prep_dt

Prepare a two-column data.table that will be used to fill values in a matrix

Description

Prepare a two-column data.table that will be used to fill values in a matrix

```
matrix_prep_dt(row_var_names = NULL, col_var_names = NULL)
```

90 mean_center

Arguments

row_var_names a vector of variable names, each of which will be header of a row in the eventual matrix

col_var_names a vector of variable names, each of which will be header of a column in the

eventual matrix

Examples

```
matrix_prep_dt(
  row_var_names = c("mpg", "cyl"),
  col_var_names = c("hp", "gear")
)
```

mean_center

Mean center

Description

Mean-center a variable, i.e., subtract the mean of a numeric vector from each value in the numeric vector

Usage

```
mean_center(x)
```

Arguments

Х

a numeric vector; though not thoroughly tested, the function can accept a matrix as an input.

```
mean_center(1:5)
mean_center(1:6)
# if the input is a matrix
matrix(1:9, nrow = 3)
mean_center(matrix(1:9, nrow = 3))
```

mediation_analysis 91

mediation_analysis

Mediation analysis

Description

Conducts a mediation analysis to estimate an independent variable's indirect effect on dependent variable through a mediator variable. The current version of the package only supports a simple mediation model consisting of one independent variable, one mediator variable, and one dependent variable.

Usage

```
mediation_analysis(
  data = NULL,
  iv_name = NULL,
  mediator_name = NULL,
  dv_name = NULL,
  covariates_names = NULL,
  robust_se = TRUE,
  iterations = 1000,
  sigfigs = 3,
  output_type = "summary_dt",
  silent = FALSE
)
```

Arguments

data a data object (a data frame or a data.table)

iv_namename of the independent variablemediator_namename of the mediator variabledv_namename of the dependent variable

covariates_names

names of covariates to control for

robust_se if TRUE, heteroskedasticity-consistent standard errors will be used in quasi-Bayesian

simulations. By default, it will be set as FALSE if nonparametric bootstrap is used

and as TRUE if quasi-Bayesian approximation is used.

iterations number of bootstrap samples. The default is set at 1000, but consider increasing

the number of samples to 5000, 10000, or an even larger number, if slower

handling time is not an issue.

sigfigs number of significant digits to round to

output_type if output_type = "summary_dt", return the summary data.table; if output_type

= "mediate_output", return the output from the mediate function in the 'mediate' package; if output_type = "indirect_effect_p", return the p value associated with the indirect effect estimated in the mediation model (default =

"summary_dt")

92 merge_data_tables

silent

if silent = FALSE, mediation analysis summary, estimation method, sample size, and number of simulations will be printed; if silent = TRUE, nothing will be printed. (default = FALSE)

Details

This function requires installing Package 'mediation' v4.5.0 (or possibly a higher version) by Tingley et al. (2019), and uses the source code from a function in the package. https://cran.r-project.org/package=mediation

Value

if output_type = "summary_dt", which is the default, the output will be a data.table showing a summary of mediation analysis results; if output_type = "mediate_output", the output will be the output from the mediate function in the 'mediate' package; if output_type = "indirect_effect_p", the output will be the p-value associated with the indirect effect estimated in the mediation model (a numeric vector of length one).

Examples

```
mediation_analysis(
  data = mtcars, iv_name = "cyl",
  mediator_name = "disp", dv_name = "mpg", iterations = 100
)
mediation_analysis(
  data = iris, iv_name = "Sepal.Length",
  mediator_name = "Sepal.Width", dv_name = "Petal.Length",
  iterations = 100
)
```

merge_data_tables

Merge data tables

Description

Merge two data.table objects. If there are any duplicated ID values and column names across the two data tables, the cell values in the first data.table will remain intact and the cell values in the second data.table will be discarded for the resulting merged data table.

```
merge_data_tables(dt1 = NULL, dt2 = NULL, id = NULL, silent = TRUE)
```

merge_data_tables 93

Arguments

dt1	the first data.table which will remain intact
dt2	the second data.table which will be joined outside of (around) the first data.table. If there are any duplicated ID values and column names across the two data tables, the cell values in the first data.table will remain intact and the cell values in the second data.table will be discarded for the resulting merged data table.
id	$name(s) \ of \ the \ column(s) \ that \ will \ contain \ the \ ID \ values \ in \ the \ two \ data \ tables.$ The $name(s) \ of \ the \ ID \ column(s) \ must \ be \ identical \ in \ the \ two \ data \ tables.$
silent	If silent = TRUE, no message will be printed regarding how many ID values and column names were duplicated. If silent = FALSE, messages will be printed regarding how many column names were duplicated. In cases where only one column was used as the 'id' column (which is the most common case), silent = FALSE will also print messages regarding how many input ID values were duplicated. By default, silent = FALSE.

Value

a data.table object, which merges (joins) the second data.table around the first data.table.

```
## Example 1: Typical Usage
data_1 <- data.table::data.table(</pre>
id_{col} = c(4, 2, 1, 3),
a = 3:6,
b = 5:8,
c = c("w", "x", "y", "z"))
data_2 <- data.table::data.table(</pre>
id_{col} = c(1, 99, 4),
e = 6:8,
b = c("p", "q", "r"),
d = c(TRUE, FALSE, FALSE))
# check the two example data tables
data_1
data_2
# check the result of merging the two data tables above and
# note how data_1 (the upper left portion) is intact in the resulting
merge_data_tables(dt1 = data_1, dt2 = data_2, id = "id_col")
# compare the result with above with the result from the `merge` function
merge(data_1, data_2, by = "id_col", all = TRUE)
## Example 2: Some values can be converted
data_3 <- data.table::data.table(</pre>
id_{col} = 99,
a = "abc",
b = TRUE,
c = TRUE)
data_1
data_3
merge_data_tables(data_1, data_3, id = "id_col")
```

```
# In the example above, note how the value of TRUE gets
# converted to 1 in the last row of Column 'b' in the resulting data table
## Example 3: A simpler case
data_4 <- data.table::data.table(</pre>
id_{col} = c(5, 3),
a = c("a", NA))
data_5 <- data.table::data.table(</pre>
id_{col} = 1,
a = 2
# check the two example data tables
data_4
data_5
merge_data_tables(data_4, data_5, id = "id_col")
## Example 4: Merging data tables using multiple ID columns
data_6 <- data.table::data.table(</pre>
id_{col_1} = 3:1,
id_col_2 = c("a", "b", "c"),
id_{col_3} = 4:6,
a = 7:9,
b = 10:12
data_7 <- data.table::data.table(</pre>
id_{col_1} = c(3, 2),
id_{col_3} = c(3, 5),
id_{col_2} = c("a", "b"),
c = 13:14,
a = 15:16
# check the example data sets
data_6
data_7
# merge data sets using the three id columns
suppressWarnings(merge_data_tables(
dt1 = data_6,
dt2 = data_7,
id = c("id_col_1", "id_col_2", "id_col_3")))
```

merge_data_table_list Merge a list of data tables

Description

Successively merge a list of data.table objects in a recursive fashion. That is, merge the (second data table in the list) around the first data table in the list; then, around this resulting data table, merge the third data table in the list; and so on.

```
merge_data_table_list(dt_list = NULL, id = NULL, silent = TRUE)
```

modes_of_objects 95

Arguments

id name(s) of the column(s) that will contain the ID values in the two data tables.

The name(s) of the ID column(s) must be identical in the two data tables.

silent If silent = TRUE, no message will be printed regarding how many ID values and column names were duplicated. If silent = FALSE, messages will be printed regarding how many column names were duplicated. In cases where only one column was used as the 'id' column (which is the most common case), silent = FALSE will also print messages regarding how many input ID values were du-

plicated. By default, silent = FALSE.

Details

If there are any duplicated ID values and column names across the data tables, the cell values in the earlier data table will remain intact and the cell values in the later data table will be discarded for the resulting merged data table in each recursion.

Value

a data.table object, which successively merges (joins) a data table around (i.e., outside) the previous data table in the list of data tables.

Examples

```
data_1 <- data.table::data.table(</pre>
id_{col} = c(4, 2, 1, 3),
a = 3:6,
b = 5:8,
c = c("w", "x", "y", "z"))
data_2 <- data.table::data.table(</pre>
id_{col} = c(1, 4, 99),
d = 6:8,
b = c("p", "q", "r"),
e = c(TRUE, FALSE, FALSE))
data_3 <- data.table::data.table(</pre>
id_{col} = c(200, 3),
f = 11:12,
b = c(300, "abc"))
merge_data_table_list(
dt_list = list(data_1, data_2, data_3), id = "id_col")
```

modes_of_objects

Find modes of objects

Description

Find modes of objects

96 multiple_regression

Usage

```
modes_of_objects(...)
```

Arguments

```
... R objects.
```

Value

the output will be a data.table listing objects and their mods.

Examples

```
modes_of_objects( TRUE, FALSE, 1L, 1:3, 1.1, c(1.2, 1.3), "abc", 1 + 2i, intToBits(1L))
```

multiple_regression

Multiple regression

Description

Conduct multiple regression analysis and summarize the results in a data.table.

Usage

```
multiple_regression(
  data = NULL,
  formula = NULL,
  vars_to_mean_center = NULL,
  mean_center_vars = NULL,
  sigfigs = NULL,
  round_digits_after_decimal = NULL,
  round_p = NULL,
  pretty_round_p_value = TRUE,
  return_table_upper_half = FALSE,
  round_r_squared = 3,
  round_f_stat = 2,
  prettify_reg_table_col_names = TRUE,
  silent = FALSE
)
```

Arguments

```
data a data object (a data frame or a data.table)
formula a formula object for the regression equation
```

multiple_regression 97

```
vars_to_mean_center
                  (deprecated) a character vector specifying names of variables that will be mean-
                  centered before the regression model is estimated
mean_center_vars
                  a character vector specifying names of variables that will be mean-centered be-
                  fore the regression model is estimated
sigfigs
                  number of significant digits to round to
round_digits_after_decimal
                  round to nth digit after decimal (alternative to sigfigs)
                  number of decimal places to round p values (overrides all other rounding argu-
round_p
                  ments)
pretty_round_p_value
                  logical. Should the p-values be rounded in a pretty format (i.e., lower threshold:
                  "<.001"). By default, pretty_round_p_value = TRUE.
return_table_upper_half
                  logical. Should only the upper part of the table be returned? By default, return_table_upper_half
                  = FALSE.
round_r_squared
                  number of digits after the decimal both r-squared and adjusted r-squared values
                  should be rounded to (default 3)
                  number of digits after the decimal the f statistic of the regression model should
round_f_stat
                  be rounded to (default 2)
prettify_reg_table_col_names
                  logical. Should the column names of the regression table be made pretty (e.g.,
                  change "std_beta" to "Std. Beta")? (Default = TRUE)
                  If silent = FALSE, a message regarding mean-centered variables will be printed.
silent
                  If silent = TRUE, this message will be suppressed. By default, silent = FALSE.
```

Details

To include standardized beta(s) in the regression results table, the following package(s) must be installed prior to running the function: Package 'lm.beta' v1.5-1 (or possibly a higher version) by Stefan Behrendt (2014), https://cran.r-project.org/package=lm.beta

Value

the output will be a data.table showing multiple regression results.

```
multiple_regression(data = mtcars, formula = mpg ~ gear * cyl)
multiple_regression(
data = mtcars, formula = mpg ~ gear * cyl,
mean_center_vars = "gear",
round_digits_after_decimal = 2)
```

98 odds_ratio

```
noncentrality_parameter
```

Find noncentrality parameter

Description

Find noncentrality parameter

Usage

```
noncentrality_parameter(t_stat, df, initial_value = 0, ci = 0.95)
```

Arguments

t_stat the t-statistic associated with the noncentrality parameters

df degrees of freedom associated with the noncentrality parameters

initial_value initial value of the noncentrality parameter for optimization (default = 0). Adjust

this value if results look strange.

ci width of the confidence interval associated with the noncentrality parameters

(default = 0.95)

Examples

```
noncentrality_parameter(4.29, 9)
```

odds_ratio

Odds ratio

Description

Calculate odds ratio, as illustrated in Borenstein et al. (2009, pp. 33-36, ISBN: 978-0-470-05724-7)

```
odds_ratio(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  contingency_table = NULL,
  ci = 0.95,
  round_ci_limits = 2,
  invert = FALSE
)
```

Arguments

data a data object (a data frame or a data.table)

iv_namename of the independent variable (grouping variable)dv_namename of the dependent variable (binary outcome)

contingency_table

a contingency table, which can be directly entered as an input for calculating the

odds ratio

ci width of the confidence interval. Input can be any value less than 1 and greater

than or equal to 0. By default, ci = 0.95. If ci = TRUE, the default value of 0.95

will be used. If ci = FALSE, no confidence interval will be estimated.

round_ci_limits

number of decimal places to which to round the limits of the confidence interval

(default = 2)

invert logical. Whether the inverse of the odds ratio (i.e., 1 / odds ratio) should be

returned.

Examples

```
## Not run:
odds_ratio(data = mtcars, iv_name = "vs", dv_name = "am")
odds_ratio(data = mtcars, iv_name = "vs", dv_name = "am", ci = 0.9)
odds_ratio(contingency_table = matrix(c(5, 10, 95, 90), nrow = 2))
odds_ratio(contingency_table = matrix(c(5, 10, 95, 90), nrow = 2),
invert = TRUE)
odds_ratio(contingency_table = matrix(c(34, 39, 16, 11), nrow = 2))
## End(Not run)
```

```
order_rows_specifically_in_dt
```

Order rows specifically in a data table

Description

Order rows in a data.table in a specific order

```
order_rows_specifically_in_dt(
  dt = NULL,
  col_to_order_by = NULL,
  specific_order = NULL
)
```

100 outlier

Arguments

dt a data.table object

col_to_order_by

a character value indicating the name of the column by which to order the

data.table

specific_order a vector indicating a specific order of the values in the column by which to order

the data.table.

Value

the output will be a data.table object whose rows will be ordered as specified.

Examples

```
order_rows_specifically_in_dt(mtcars, "carb", c(3, 2, 1, 4, 8, 6))
```

outlier

Outlier

Description

Return outliers in a vector

Usage

```
outlier(x = NULL, iqr = 1.5, na.rm = TRUE, type = 7, unique_outliers = FALSE)
```

Arguments

type

Х	a numeric vector
iqr	a nonnegative constant by which interquartile range (IQR) will be multiplied to build a "fence," outside which observations will be considered outliers. For example, if $iqr = 1.5$, $IQR * 1.5$ will be the "fence" outside which observations will be considered to be outliers. By default, $iqr = 1.5$.
na.rm	logical. na.rm argument to be passed onto the 'quantile' function in the 'stats' package. If true, any NA and NaN's are removed from x before the quantiles are computed.

type argument to be passed onto the 'quantile' function in the 'stats' package. An integer between 1 and 9 selecting one of the nine quantile algorithms detailed

below to be used. Type '?stats::quantile' for details. By default, type = 7

unique_outliers

logical. If unique_outliers = TRUE, the function will return the unique outlier values. If unique_outliers = FALSE, the function will return all the outlier values in the vector x. By default, unique_outliers = FALSE.

overlapping_interval 101

Value

the output will be a numeric vector with outliers removed.

Examples

```
# Example 1
outlier(c(1:10, 100))
# The steps below show how the outlier, 100, was obtained
# v1 is the vector of interest
v1 <- c(1:10, 100)
# quantile
stats::quantile(v1)
# first and third quartiles
q1 <- stats::quantile(v1, 0.25)
q3 <- stats::quantile(v1, 0.75)
# interquartile range
interquartile_range <- unname(q3 - q1)</pre>
# fence, using the default 1.5 as the factor to multiply the IQR
cutoff_low <- unname(q1 - 1.5 * interquartile_range)</pre>
cutoff_high <- unname(q3 + 1.5 * interquartile_range)</pre>
v1[v1 < cutoff_low | v1 > cutoff_high]
```

overlapping_interval Find the overlapping interval of two ranges.

Description

This function should be applied to cases where the two ranges are inclusive of both endpoints. For example, the function can work for a pair of ranges like [0, 1] and [3, 4] but not for pairs like $[0, 1\setminus)$ and $(3, 5\setminus)$

Usage

```
overlapping_interval(
  interval_1_begin = NULL,
  interval_1_end = NULL,
  interval_2_begin = NULL,
  interval_2_end = NULL
)
```

Arguments

```
interval_1_begin

a number at which the first interval begins (the left INCLUSIVE endpoint of interval 1)

interval_1_end

a number at which the first interval ends (the right INCLUSIVE endpoint of interval 1)
```

102 p0

```
interval_2_begin
```

a number at which the second interval begins (the left INCLUSIVE endpoint of interval 2)

interval_2_end a number at which the second interval ends (the right INCLUSIVE endpoint of interval 2)

Value

the output will be NULL if there is no overlapping region or a vector of the endpoints of the overlapping interval.

Examples

```
overlapping_interval(1, 3, 2, 4)
overlapping_interval(1, 2.22, 2.22, 3)
```

p0

Paste0

Description

A shorthand for the function paste0 Concatenate vectors after converting to character.

Usage

```
p0(..., collapse = NULL, recycle0 = FALSE)
```

Arguments

... one or more R objects, to be converted to character vectors. This is the same

argument that would be used in the paste0 function.

collapse an optional character string to separate the results. Not NA_character_. This is

the same argument that would be used in the paste0 function.

recycle0 logical indicating if zero-length character arguments should lead to the zero-

length character(0) after the sep-phase (which turns into "" in the collapse-phase, i.e., when collapse is not NULL). This is the same argument that would be used

in the paste0 function.

```
paste0("a", "b")
p0("a", "b")
```

package_list_default 103

```
package_list_default Packages - List the default packages
```

Description

List the default packages in R

Usage

```
package_list_default(package_type = c("base", "recommended"))
```

Arguments

```
package_type a vector of package types. By default, package_type = c("base", "recommended")
```

Examples

```
package_list_default()
package_list_default(package_type = "base")
```

parallel_analysis

Parallel analysis

Description

Conducts a parallel analysis to determine how many factors to retain in a factor analysis.

```
parallel_analysis(
  data = NULL,
  names_of_vars = NULL,
  iterations = NULL,
  percentile_for_eigenvalue = 95,
  line_types = c("dashed", "solid"),
  colors = c("red", "blue"),
  eigenvalue_random_label_x_pos = NULL,
  eigenvalue_random_label_y_pos = NULL,
  unadj_eigenvalue_label_x_pos = NULL,
  unadj_eigenvalue_label_y_pos = NULL,
  label_offset_percent = 2,
  label_size = 6,
  dot_size = 5,
  line_thickness = 1.5,
  y_axis_title_vjust = 0.8,
  title_text_size = 26,
  axis_text_size = 22
)
```

104 parallel_analysis

Arguments

```
data
                  a data object (a data frame or a data.table)
names_of_vars
                  names of the variables
iterations
                  number of random data sets. If no input is entered, this value will be set as 30 *
                  number of variables.
percentile_for_eigenvalue
                  percentile used in estimating bias (default = 95).
line_types
                  types of the lines connecting eigenvalues. By default, line_types = c("dashed",
                   "solid")
                  size of the dots denoting eigenvalues (default = 5).
colors
eigenvalue_random_label_x_pos
                  (optional) x coordinate of the label for eigenvalues from randomly generated
eigenvalue_random_label_y_pos
                  (optional) y coordinate of the label for eigenvalues from randomly generated
                  data.
unadj_eigenvalue_label_x_pos
                  (optional) x coordinate of the label for unadjusted eigenvalues
unadj_eigenvalue_label_y_pos
                  (optional) y coordinate of the label for unadjusted eigenvalues
label_offset_percent
                  How much should labels for the eigenvalue curves be offset, as a percentage of
                  the plot's x and y range? (default = 2)
label_size
                  size of the labels for the eigenvalue curves (default = 6).
dot_size
                  size of the dots denoting eigenvalues (default = 5).
line_thickness thickness of the eigenvalue curves (default = 1.5).
y_axis_title_vjust
                  position of the y axis title as a proportion of the range (default = 0.8).
title_text_size
                  size of the plot title (default = 26).
axis_text_size size of the text on the axes (default = 22).
```

Details

The following package(s) must be installed prior to running the function: Package 'paran' v1.5.2 (or possibly a higher version) by Alexis Dinno (2018), https://cran.r-project.org/package=paran

```
parallel_analysis(
  data = mtcars, names_of_vars = c("disp", "hp", "drat"))
# parallel_analysis(
# data = mtcars, names_of_vars = c("carb", "vs", "gear", "am"))
```

percentile_rank 105

percentile_rank

Percentile rank

Description

Calculate percentile rank of each value in a vector

Usage

```
percentile_rank(vector)
```

Arguments

vector

a numeric vector

Examples

```
percentile_rank(1:5)
percentile_rank(1:10)
percentile_rank(1:100)
```

pivot_table

Pivot Table

Description

Create a pivot table.

```
pivot_table(
  data = NULL,
  row_names = NULL,
  col_names = NULL,
  function_as_character = NULL,
  sigfigs = 3,
  output = "dt",
  remove_col_names = TRUE
)
```

plot_group_means

Arguments

data a data object (a data frame or a data.table)
row_names names of variables for constructing rows

function_as_character

function to perform for each cell in the pivot table

sigfigs number of significant digits to which to round values in the pivot table (default

= 3)

output type of output. If output = "dt", the function's output will be a pivot table in a

data.table format. If output = "subsets", the function's output will be a list of data tables that are subsets representing each cell in the pivot table. By default,

output = "dt"

remove_col_names

logical. Should the column names (i.e., v1, v2, ...) be removed in the data table

output?

Value

the output will be a contingency table in a data.table format

Examples

```
pivot_table(
  data = mtcars, col_names = "am", row_names = c("cyl", "vs"),
  function_as_character = "mean(mpg)")
pivot_table(
  data = mtcars, col_names = "am", row_names = c("cyl", "vs"),
  function_as_character = "sum(mpg < 17)")
pivot_table(
  data = mtcars, col_names = "am", row_names = c("cyl", "vs"),
  function_as_character =
  "round(sum(mpg < 17) / sum(!is.na(mpg)) * 100, 0)")</pre>
```

plot_group_means

Plot group means

Description

Creates a plot of sample means and error bars by group.

```
plot_group_means(
  data = NULL,
  dv_name = NULL,
  iv_name = NULL,
```

107 plot_group_means

```
na.rm = TRUE,
  error_bar = "ci",
  error_bar_range = 0.95,
  error_bar_tip_width = 0.13,
  error_bar_thickness = 1,
  error_bar_caption = TRUE,
  lines_connecting_means = TRUE,
  line_types = NULL,
  line_thickness = 1,
  line_size = NULL,
  dot_size = 3,
  position_dodge = 0.13,
  legend_position = "right",
 y_axis_title_vjust = 0.85
)
data
dv name
iv name
```

Arguments

line_size

dot_size

a data object (a data frame or a data.table) name of the dependent variable name(s) of the independent variable(s). Up to two independent variables can be supplied. logical. If na.rm = TRUE, NA values in independent and dependent variables will na.rm be removed before calculating group means. if error_bar = "se"; error bars will be +/-1 standard error, if error_bar = "ci" error_bar error bars will be a confidence interval; if error_bar = "pi", error bars will be a prediction interval error_bar_range width of the confidence or prediction interval (default = 0.95 for 95 percent confidence or prediction interval). This argument will not apply when error_bar = error_bar_tip_width graphically, width of the segments at the end of error bars (default = 0.13) error_bar_thickness thickness of the error bars (default = 1) error_bar_caption should a caption be included to indicate the width of the error bars? (default = TRUE). lines_connecting_means logical. Should lines connecting means within each group be drawn? (default = line_types types of the lines connecting means (default = NULL) If the second IV has two levels, then by default, line_types = c("solid", "dashed") line_thickness thickness of the lines connecting group means (default = 1)

Deprecated. Use the 'linewidth' argument instead. (default = 1)

size of the dots indicating group means (default = 3)

108 pm

= 0.85, the y axis title will be positioned at 85% of the way up from the bottom of the plot.

Value

by default, the output will be a ggplot object. If output = "table", the output will be a data.table object.

Examples

```
plot_group_means(data = mtcars, dv_name = "mpg", iv_name = c("vs", "am"))
plot_group_means(
   data = mtcars, dv_name = "mpg", iv_name = c("vs", "am"),
   error_bar = "se"
)
plot_group_means(
   data = mtcars, dv_name = "mpg", iv_name = c("vs", "am"),
   error_bar = "pi", error_bar_range = 0.99
)
```

pm

Paste for message

Description

Combines the base functions paste0 and message

Usage

```
pm(..., collapse = NULL)
```

Arguments

one or more R objects, to be converted to character vectors. Input(s) to this argument will be passed onto the paste0 function.

collapse an optional character string to separate the resu

an optional character string to separate the results. Not NA_character_. Input(s) to this argument will be passed onto the paste0 function.

population_variance 109

Value

there will be no output from this function. Rather, a message will be generated from the arguments.

Examples

```
pm("hello", 123)
pm(c("hello", 123), collapse = ", ")
```

population_variance

Population variance of a vector

Description

Calculates the population variance, rather than the sample variance, of a vector

Usage

```
population_variance(vector, na.rm = TRUE)
```

Arguments

vector a numeric vector

na.rm if TRUE, NA values will be removed before calculation

Examples

```
population_variance(1:4)
var(1:4)
```

prep

Prepare package(s) for use

Description

Installs, loads, and attaches package(s). If package(s) are not installed, installs them prior to loading and attaching.

```
prep(
    ...,
    pkg_names_as_object = FALSE,
    silent_if_successful = FALSE,
    silent_load_pkgs = NULL
)
```

Arguments

names of packages to load and attach, separated by commas, e.g., "ggplot2", data.table. The input can be any number of packages, whose names may or may not be wrapped in quotes.

pkg_names_as_object

logical. If pkg_names_as_object = TRUE, the input will be evaluated as one object containing package names. If pkg_names_as_object = FALSE, the input will be considered as literal packages names (default = FALSE).

silent_if_successful

logical. If silent_if_successful = TRUE, no message will be printed if preparation of package(s) is successful. If silent_if_successful = FALSE, a message indicating which package(s) were successfully loaded and attached will be printed (default = FALSE).

silent_load_pkgs

a character vector indicating names of packages to load silently (i.e., suppress messages that get printed when loading the packaged). By default, silent_load_pkgs = NULL

Value

there will be no output from this function. Rather, packages given as inputs to the function will be installed, loaded, and attached.

Examples

```
prep(data.table)
prep("data.table", silent_if_successful = TRUE)
prep("base", utils, ggplot2, "data.table")
pkgs <- c("ggplot2", "data.table")
prep(pkgs, pkg_names_as_object = TRUE)
prep("data.table", silent_load_pkgs = "data.table")</pre>
```

Description

Round p-values to the desired number of decimals and remove leading 0s before the decimal.

```
pretty_round_p_value(
   p_value_vector = NULL,
   round_digits_after_decimal = 3,
   include_p_equals = FALSE
)
```

pretty_round_r 111

Arguments

Value

the output will be a character vector with p values, e.g., a vector of strings like "< .001" (or "p < .001").

Examples

```
pretty_round_p_value(0.0495, 3)
pretty_round_p_value(
    p_value_vector = 0.049,
    round_digits_after_decimal = 2, include_p_equals = FALSE)
pretty_round_p_value(c(0.0015, 0.0014, 0.0009), include_p_equals = TRUE)
```

pretty_round_r

Pretty round r

Description

Round correlation coefficients in APA style (7th Ed.)

Usage

```
pretty_round_r(r = NULL, round_digits_after_decimal = 2)
```

Arguments

```
r a (vector of) correlation coefficient(s)
round_digits_after_decimal
how many digits after the decimal point should the p-value be rounded to? (default = 2)
```

Value

the output will be a character vector of correlation coefficient(s).

```
pretty_round_r(r = -0.123) pretty_round_r(c(-0.12345, 0.45678), round_digits_after_decimal = 3) pretty_round_r(c(-0.12, 0.45), round_digits_after_decimal = 4)
```

print_loop_progress

Description

Print current progress inside a loop (e.g., for loop or lapply)

Usage

```
print_loop_progress(
  iteration_number = NULL,
  iteration_start = 1,
  iteration_end = NULL,
  text_before = "",
  percent = 1,
  output_method = "cat"
)
```

Arguments

```
for (i in seq_len(250)) {
   Sys.sleep(0.001)
   print_loop_progress(
      iteration_number = i,
      iteration_end = 250)
}
unlist(lapply(seq_len(7), function (i) {
   Sys.sleep(0.1)
   print_loop_progress(
      iteration_number = i,
      iteration_end = 7)
   return(i)
}))
```

```
proportion_of_values_in_vector
```

Proportion of given values in a vector

Description

Proportion of given values in a vector

Usage

```
proportion_of_values_in_vector(
  values = NULL,
  vector = NULL,
  na.exclude = TRUE,
  output_type = "proportion",
  silent = FALSE,
  conf.level = 0.95,
  correct_yates = TRUE
)
```

Arguments

values a set of values that will count as successes (hits) a numeric or character vector containing successes (hits) and failures (misses) vector na.exclude if TRUE, NA values will be removed both from vector and values before calculation (default = TRUE). output_type By default, output_type = "proportion". If output_type = "proportion", the function will return the calculated proportion; if output_type = "se", the function will return the standard error of the sample proportion; if output_type = "dt", the function will return the the data table of proportion and confidence intervals. silent If silent = TRUE, no message will be printed regarding number of NA values or confidence interval. (default = FALSE) conf.level confidence level of the returned confidence interval. Input to this argument will be passed onto the conf.level argument in the prop. test function from the default stats package. a logical indicating whether Yates' continuity correction should be applied where correct_yates possible (default = TRUE). Input to this argument will be passed onto the correct argument in the prop. test function from the default stats package.

```
proportion_of_values_in_vector(
  values = 2:3, vector = c(rep(1:3, each = 10), rep(NA, 10))
)
```

114 q_stat_test_homo_r

```
proportion_of_values_in_vector(
  values = 2:3, vector = c(rep(1:3, each = 10), rep(NA, 10)),
  output_type = "se"
)
proportion_of_values_in_vector(
  values = 2:3, vector = c(rep(1:3, each = 10), rep(NA, 10)),
  conf.level = 0.99
)
proportion_of_values_in_vector(
  values = c(2:3, NA), vector = c(rep(1:3, each = 10), rep(NA, 10)),
  na.exclude = FALSE
)
```

q_stat_test_homo_r

Q statistic for testing homogeneity of correlations

Description

Calculate the Q statistic to test for homogeneity of correlation coefficients. See p. 235 of the book Hedges & Olkin (1985), Statistical Methods for Meta-Analysis (ISBN: 0123363802).

Usage

```
q_stat_test_homo_r(z = NULL, n = NULL)
```

Arguments

z a vector of z values

n a vector of sample sizes which will be used to calculate the weights, which in turn will be used to calculate the weighted z.

Value

the output will be a weighted z value.

```
q_stat_test_homo_r(1:3, c(100, 200, 300))
q_stat_test_homo_r(z = c(1:3, NA), n = c(100, 200, 300, NA))
```

read_csv 115

read_csv

Read a csv file

Description

Read a csv file

Usage

```
read_csv(name = NULL, head = FALSE, dirname = NULL, ...)
```

Arguments

name a character string of the csv file name without the ".csv" extension. For example,

if the csv file to read is "myfile.csv", enter name = "myfile"

head logical. if head = TRUE, prints the first five rows of the data set.

dirname a character string of the directory containing the csv file, e.g., dirname = "c:/Users/Documents"

optional arguments for the fread function from the data.table package. Any

arguments for data.table's fread function can be used, e.g., fill = TRUE, nrows

= 100

Value

the output will be a data.table object, that is, an output from the data.table function, fread

Examples

```
## Not run:
mydata <- read_csv("myfile")
## End(Not run)</pre>
```

read_sole_csv

Read the sole csv file in the working directory

Description

Read the sole csv file in the working directory

```
read_sole_csv(head = FALSE, ...)
```

116 regex_match

Arguments

head logical. if head = TRUE, prints the first five rows of the data set.

optional arguments for the fread function from the data.table package. Any arguments for data.table's fread function can be used, e.g., fill = TRUE, nrows

1aa

= 100

Value

the output will be a data.table object, that is, an output from the data.table function, fread

Examples

```
mydata <- read_sole_csv()
mydata <- read_sole_csv(head = TRUE)
mydata <- read_sole_csv(fill = TRUE, nrows = 5)</pre>
```

regex_match

Regular expression matches

Description

Returns elements of a character vector that match the given regular expression

Usage

```
regex_match(regex = NULL, vector = NULL, silent = FALSE, perl = FALSE)
```

Arguments

regex a regular expression provided, a default theme will be used.

vector a character vector in which to search for regular expression matches, or a data

table whose column names will be searched

silent logical. If silent = FALSE, a report on regular expression matches will be

printed. If silent = TRUE, the report on regular expression matches will not

be printed. By default, silent = FALSE

perl logical. Should Perl-compatible regexps be used?

```
regex_match("p$", names(mtcars))
colnames_ending_with_p <- regex_match("p$", names(mtcars))</pre>
```

```
rel_pos_of_value_in_vector
```

Find relative position of a value in a vector

Description

Find relative position of a value in a vector that may or may not contain the value

Usage

```
rel_pos_of_value_in_vector(value = NULL, vector = NULL)
```

Arguments

value a value whose relative position is to be searched in a vector

vector a numeric vector

Value

a number indicating the relative position of the value in the vector

Examples

```
rel_pos_of_value_in_vector(value = 3, vector = c(2, 4))
rel_pos_of_value_in_vector(value = 3, vector = c(2, 6))
rel_pos_of_value_in_vector(value = 3, vector = 1:3)
```

```
rel_value_of_pos_in_vector
```

Find relative value of a position in a vector

Description

Find relative value of a position in a vector

Usage

```
rel_value_of_pos_in_vector(vector = NULL, position = NULL)
```

Arguments

vector a numeric vector
position position of a vector

118 remove_from_vector

Value

a number indicating the relative value of the position in the vector

Examples

```
rel_value_of_pos_in_vector(vector = c(0, 100), position = 1.5)
rel_value_of_pos_in_vector(vector = 2:4, position = 2)
rel_value_of_pos_in_vector(vector = c(2, 4, 6), position = 2.5)
```

remove_from_vector

Remove from a vector

Description

Remove certain values from a vector

Usage

```
remove_from_vector(values = NULL, vector = NULL, silent = FALSE)
```

Arguments

values	a single value or a vector of values which will be removed from the target vector
vector	a character or numeric vector
silent	if silent = FALSE, a summary of values removed will be printed; if silent = TRUE, such summary will not be printed. By default, silent = FALSE

Value

the output will be a vector with the given values removed.

```
remove_from_vector(values = 1, vector = 1:3)
remove_from_vector(values = NA, vector = c(1:3, NA))
remove_from_vector(values = c(1, NA), vector = c(1:3, NA))
remove_from_vector(values = 1:5, vector = 1:10)
```

```
remove_user_installed_pkgs
```

Remove all user installed packages

Description

Remove all user installed packages

Usage

```
remove_user_installed_pkgs(
  exceptions = NULL,
  type_of_pkg_to_keep = c("base", "recommended"),
  keep_kim = FALSE
)
```

Arguments

```
exceptions a character vector of names of packages to keep
```

type_of_pkg_to_keep

a character vector indicating types of packages to keep. The default, $type_of_pkg_to_keep$

= c("base", "recommended"), keeps all base and recommended packages that

come with R when R is installed.

keep_kim

logical. If keep_kim = FALSE, Package 'kim' will be removed along with all other user-installed packages. If keep_kim = TRUE, Package 'kim' will not be

removed. By default, keep_kim = FALSE

Examples

```
## Not run:
remove_user_installed_pkgs()
## End(Not run)
```

repeated_measures_anova

Repeated-Measures ANVOA

Description

Conduct a repeated-measures analysis of variance (ANOVA). This analysis will be appropriate for within-subjects experimental design.

Usage

```
repeated_measures_anova(
  data = NULL,
  p_col_name = NULL,
  measure_vars = NULL,
  histograms = TRUE,
  round_w = 2,
  round_epsilon = 2,
  round_df_model = 2,
  round_df_error = 2,
  round_f = 2,
  round_ges = 2
)
```

Arguments

data	a data object (a data frame or a data.table)
p_col_name	name of the column identifying participants
measure_vars	names of the columns containing repeated measures (within-subjects variables)
histograms	logical. If histograms = TRUE, histograms of the repeated measures will be plotted. If histograms = FALSE, no histograms will be plotted.
round_w	number of decimal places to which to round W statistic from Mauchly's test $(default = 2)$
round_epsilon	number of decimal places to which to round the epsilon statistic from Greenhouse-Geisser or Huynh-Feldt correction (default $= 2$)
round_df_model	number of decimal places to which to round the corrected degrees of freedom for model (default = 2)
round_df_error	number of decimal places to which to round the corrected degrees of freedom for error (default = 2)
round_f	number of decimal places to which to round the F statistic (default = 2)
round_ges	number of decimal places to which to round generalized eta-squared (default $=$ 2)

Details

The following package(s) must be installed prior to running the function: Package 'ez' v4.4-0 (or possibly a higher version) by Michael A Lawrence (2016), https://cran.r-project.org/package=ez

```
## Not run:
repeated_measures_anova(
  data = mtcars, p_col_name = "cyl", measure_vars = c("wt", "qsec"))
## End(Not run)
```

replace_values_in_dt 121

```
replace_values_in_dt Replace values in a data table
```

Description

Replace values in a data.table

Usage

```
replace_values_in_dt(
  data = NULL,
  old_values = NULL,
  new_values = NULL,
  silent = FALSE
)
```

Arguments

a data object (a data frame or a data.table)

old_values a vector of old values that need to be replaced

new_values a new value or a vector of new values that will replace the old values

silent If silent = FALSE, a message will be printed regarding how many values were replaced. If silent = TRUE, no message will be printed regarding how many values were replaced. (default = FALSE)

Examples

```
replace_values_in_dt(data = mtcars, old_values = 21.0, new_values = 888)
replace_values_in_dt(data = mtcars, old_values = c(0, 1), new_values = 999)
replace_values_in_dt(
data = mtcars, old_values = c(0, 1), new_values = 990:991)
replace_values_in_dt(
data = data.table::data.table(a = NA_character_, b = NA_character_),
old_values = NA, new_values = "")
```

robust_regression Robust regression (bootstrapped regression)

Description

Estimate coefficients in a multiple regression model by bootstrapping.

round_flexibly

Usage

```
robust_regression(
  data = NULL,
  formula = NULL,
  sigfigs = NULL,
  round_digits_after_decimal = NULL,
  iterations = 1000
)
```

Arguments

data a data object (a data frame or a data.table)

formula a formula object for the regression equation

sigfigs number of significant digits to round to

round_digits_after_decimal

round to nth digit after decimal (alternative to sigfigs)

iterations number of bootstrap samples. The default is set at 1000, but consider increasing

the number of samples to 5000, 10000, or an even larger number, if slower

handling time is not an issue.

Details

The following package(s) must be installed prior to running this function: Package 'boot' v1.3-26 (or possibly a higher version) by Canty & Ripley (2021), https://cran.r-project.org/package=boot

Examples

```
## Not run:
robust_regression(
  data = mtcars, formula = mpg ~ cyl * hp,
  iterations = 100
)
## End(Not run)
```

round_flexibly

Round flexibly

Description

Round numbers to a flexible number of significant digits. "Flexible" rounding refers to rounding all numbers to the highest level of precision seen among numbers that would have resulted from the 'signif()' function in base R. The usage examples of this function demonstrate flexible rounding (see below).

round_flexibly 123

Usage

```
round_flexibly(x = NULL, sigfigs = 3)
```

Arguments

```
x a numeric vector
sigfigs number of significant digits to flexibly round to. By default, sigfigs = 3.
```

Value

the output will be a numeric vector with values rounded to the highest level of precision seen among numbers that result from the 'signif()' function in base R.

```
# Example 1
# First, observe results from the 'signif' function:
c(0.00012345, pi)
signif(c(0.00012345, pi), 3)
# In the result above, notice how info is lost on some digits
# (e.g., 3.14159265 becomes 3.140000).
# In contrast, flexible rounding retains the lost info in the digits
round_flexibly(x = c(0.00012345, pi), sigfigs = 3)
# Example 2
# Again, first observe results from the 'signif' function:
c(0.12345, 1234, 0.12, 1.23, .01)
signif(c(0.12345, 1234, 0.12, 1.23, .01), 3)
# In the result above, notice how info is lost on some digits
# (e.g., 1234 becomes 1230.000).
# In contrast, flexible rounding retains the lost info in the digits.
# Specifically, in the example below, 0.12345 rounded to 3 significant
# digits (default) is signif(0.12345, 3) = 0.123 (3 decimal places).
# Because this 3 decimal places is the highest precision seen among
# all numbers, all other numbers will also be rounded to 3 decimal places.
round_flexibly(
c(0.12345, 1234, 0.12, 1.23, .01))
# If the input is a character vector, the original input will be returned.
round_flexibly(c("a", "b", "c"))
# Example 4
# If the input is a list (e.g., a data.frame) that contains at least
# one numeric vector, the numeric vector element(s) will be rounded
round_flexibly(data.frame(a = c(1.2345, 123.45), b = c("a", "b")))
# Example 5
# If the input is a matrix, all numbers will be rounded flexibly
round_flexibly(matrix(
```

124 scatterplot

```
c(1.23, 2.345, 3.4567, 4.56789), ncol = 2), sigfigs = 3)
```

scatterplot

Scatterplot

Description

Creates a scatter plot and calculates a correlation between two variables.

```
scatterplot(
 data = NULL,
 x_var_name = NULL,
  y_var_name = NULL,
 dot_label_var_name = NULL,
 weight_var_name = NULL,
  alpha = 1,
  annotate_stats = TRUE,
  annotate_y_pos = 5,
  annotated_stats_color = "green4",
  annotated_stats_font_size = 6,
  annotated_stats_font_face = "bold",
  line_of_fit_type = "lm",
  ci_for_line_of_fit = FALSE,
  line_of_fit_color = "blue",
  line_of_fit_thickness = 1,
  dot_color = "black",
  x_axis_label = NULL,
  y_axis_label = NULL,
  dot_size = 2,
  dot_label_size = NULL,
  dot_size_range = c(3, 12),
  jitter_x_percent = 0,
  jitter_y_percent = 0,
  jitter_x_y_percent = 0,
  cap_axis_lines = TRUE,
  color_dots_by = NULL,
  png_name = NULL,
  save_as_png = FALSE,
 width = 16,
  height = 9
)
```

scatterplot 125

Arguments

data a data object (a data frame or a data.table) name of the variable that will go on the x axis x_var_name name of the variable that will go on the y axis y_var_name dot_label_var_name name of the variable that will be used to label individual observations weight_var_name name of the variable by which to weight the individual observations for calculating correlation and plotting the line of fit opacity of the dots (0 = completely transparent, 1 = completely opaque)alpha annotate_stats if TRUE, the correlation and p-value will be annotated at the top of the plot (default = TRUE) annotate_y_pos position of the annotated stats, expressed as a percentage of the range of y values by which the annotated stats will be placed above the maximum value of y in the data set (default = 5). If annotate_y_pos = 5, and the minimum and maximum y values in the data set are 0 and 100, respectively, the annotated stats will be placed at 5% of the y range (100 - 0) above the maximum y value, y = 0.05 *(100 - 0) + 100 = 105.annotated_stats_color color of the annotated stats (default = "green4"). annotated_stats_font_size font size of the annotated stats (default = 6). annotated_stats_font_face font face of the annotated stats (default = "bold"). line_of_fit_type ifline_of_fit_type = "lm", a regression line will be fit; if line_of_fit_type = "loess", a local regression line will be fit; if line_of_fit_type = "none", no line will be fit ci_for_line_of_fit if ci_for_line_of_fit = TRUE, confidence interval for the line of fit will be shaded line_of_fit_color color of the line of fit (default = "blue") line_of_fit_thickness thickness of the line of fit (default = 1) dot_color color of the dots (default = "black") alternative label for the x axis x_axis_label y_axis_label alternative label for the y axis dot_size size of the dots on the plot (default = 2) dot_label_size size for dots' labels on the plot. If no input is entered for this argument, it will be set as dot_label_size = 5 by default. If the plot is to be weighted by some variable, this argument will be ignored, and dot sizes will be determined by the argument dot_size_range

126 scatterplot

dot_size_range minimum and maximum size for dots on the plot when they are weighted

jitter_x_percent horizontally jitter dots by a percentage of the range of x values. jitter_y_percent vertically jitter dots by a percentage of the range of y values jitter_x_y_percent horizontally and vertically jitter dots by a percentage of the range of x and y cap_axis_lines logical. Should the axis lines be capped at the outer tick marks? (default = TRUE) name of the variable that will determine colors of the dots color_dots_by png_name name of the PNG file to be saved. By default, the name will be "scatterplot_" followed by a timestamp of the current time. The timestamp will be in the format, jan 01 2021 1300 10 000001, where "jan 01 2021" would indicate January 01, 2021; 1300 would indicate 13:00 (i.e., 1 PM); and 10_000001 would indicate 10.000001 seconds after the hour. if save = TRUE, the plot will be saved as a PNG file. save_as_png width of the plot to be saved. This argument will be directly entered as the width width

Details

height

If a weighted correlation is to be calculated, the following package(s) must be installed prior to running the function: Package 'weights' v1.0 (or possibly a higher version) by John Pasek (2018), https://cran.r-project.org/package=weights

argument for the ggsave function within ggplot2 package (default = 16)

height of the plot to be saved. This argument will be directly entered as the height argument for the ggsave function within ggplot2 package (default = 9)

Value

the output will be a scatter plot, a ggplot object.

```
## Not run:
scatterplot(data = mtcars, x_var_name = "wt", y_var_name = "mpg")
scatterplot(
   data = mtcars, x_var_name = "wt", y_var_name = "mpg",
   dot_label_var_name = "hp", weight_var_name = "drat",
   annotate_stats = TRUE)
scatterplot(
   data = mtcars, x_var_name = "wt", y_var_name = "mpg",
   dot_label_var_name = "hp", weight_var_name = "cyl",
   dot_label_size = 7, annotate_stats = TRUE)
## End(Not run)
```

score_scale_items 127

score_scale_items

Score scale items

Description

Score items in a scale (e.g., Likert scale items) by computing the sum or mean of the items.

Usage

```
score_scale_items(
  item_list = NULL,
  reverse_item_list = NULL,
  operation = "mean",
  na.rm = FALSE,
  na_summary = TRUE,
  reverse_code_minuend = NULL
)
```

Arguments

item_list a list of scale items (i.e., list of vectors of ratings) to code normally (as opposed to reverse coding).

reverse_item_list

a list of scale items to reverse code.

operation if operation = "mean", mean of the scale items will be calculated; if operation

= "sum", sum of the scale items will be calculated (default = "mean").

na.rm logical. The na.rm argument that will be passed onto the base R's rowMeans or

rowSums function (default = FALSE).

na_summary logical. If na_summary = TRUE a summary of NA values will be printed; if

na_summary = FALSE the summary will not be printed (default = TRUE).

reverse_code_minuend

required for reverse coding; the number from which to subtract item ratings when reverse-coding. For example, if the items to reverse code are measured on a 7-point scale, enter reverse_code_minuend = 8.

```
score_scale_items(item_list = list(1:5, rep(3, 5)),
reverse_item_list = list(rep(5, 5)), reverse_code_minuend = 6)
score_scale_items(item_list = list(c(1, 1), c(1, 5)),
reverse_item_list = list(c(5, 3)),
reverse_code_minuend = 6, na_summary = FALSE)
score_scale_items(item_list = list(c(1, 1), c(1, 5)),
reverse_item_list = list(c(5, 1)),
reverse_code_minuend = 6, operation = "sum")
score_scale_items(item_list = list(1:5, rep(3, 5)))
score_scale_items(item_list = list(c(1, NA, 3), c(NA, 2, 3)))
score_scale_items(item_list = list(c(1, NA, 3), c(NA, 2, 3))), na.rm = TRUE)
```

128 setwd_to_active_doc

setup_r_env

Set up R environment

Description

Set up R environment by (1) clearing the console; (2) removing all objects in the global environment; (3) setting the working directory to the active document (in RStudio only); (4) unloading and loading the kim package.

Usage

```
setup_r_env(
  clear_console = TRUE,
  clear_global_env = TRUE,
  setwd_to_active_doc = TRUE,
  prep_kim = TRUE
)
```

Arguments

Examples

```
## Not run:
setup_r_env()
## End(Not run)
```

setwd_to_active_doc

Set working directory to active document in RStudio

Description

Set working directory to location of the active document in RStudio

```
setwd_to_active_doc()
```

se_of_mean 129

Value

there will be no output from this function. Rather, the working directory will be set as location of the active document.

Examples

```
## Not run:
setwd_to_active_doc()
## End(Not run)
```

se_of_mean

Standard error of the mean

Description

Standard error of the mean

Usage

```
se_of_mean(vector, na.rm = TRUE, notify_na_count = NULL)
```

Arguments

vector a numeric vector

na.rm if TRUE, NA values will be removed before calculation

notify_na_count

if TRUE, notify how many observations were removed due to missing values. By

default, NA count will be printed only if there are any NA values.

Value

the output will be a numeric vector of length one, which will be the standard error of the mean for the given numeric vector.

```
se_of_mean(c(1:10, NA))
```

se_of_proportion

se_of_percentage

Standard Error (SE) of a percentage

Description

Calculate the standard error of a percentage. See Fowler, Jr. (2014, p. 34, ISBN: 978-1-4833-1240-8)

Usage

```
se_of_percentage(percent = NULL, n = NULL)
```

Arguments

percent a vector of percentages; each of the percentage values must be between 0 and

100

n a vector of sample sizes; number of observations used to calculate each of the

percentage values

Examples

```
se_of_percentage(percent = 40, n = 50)
se_of_percentage(percent = 50, n = 10)
```

se_of_proportion

Standard Error (SE) of a proportion

Description

Calculate the standard error of a proportion. See Anderson and Finn (1996, p. 364, ISBN: 978-1-4612-8466-6)

Usage

```
se_of_proportion(p = NULL, n = NULL)
```

Arguments

p a vector of proportions; each of the proportion values must be between 0 and 1 n a vector of sample sizes; number of observations used to calculate each of the

percentage values

```
se_of_proportion(p = 0.56, n = 400)

se_of_proportion(p = 0.5, n = 10)
```

```
simple_effects_analysis

Simple Effects Analysis
```

Description

Conduct a simple effects analysis to probe a two-way interaction effect. See Field et al. (2012, ISBN: 978-1-4462-0045-2).

Usage

```
simple_effects_analysis(
  data = NULL,
  dv_name = NULL,
  iv_1_name = NULL,
  iv_2_name = NULL,
  iv_1_levels = NULL,
  iv_2_levels = NULL,
  print_contrast_table = "weights_sums_and_products",
  output = NULL
)
```

Arguments

data a data object (a data frame or a data.table) dv_name name of the dependent variable (DV) name of the first independent variable (IV1), whose main effects will be examiv_1_name ined in the first set of contrasts iv_2_name name of the second independent variable (IV2), whose simple effects at each level of IV1 will be examined in the second set of contrasts iv_1_levels ordered levels of IV1 iv_2_levels ordered levels of IV2 print_contrast_table If print_contrast_table = "weights_sums_and_products", contrasts' weights, sums of the weights and products will be printed. If print_contrast_table = "weights_only", only the contrasts will be printed. output can be one of the following: "lm_object", "table", "weights_only", output "weights_sums_and_products", "all" By default, output = NULL, and there will be no output from the function other than the tables of simple effects and constrasts which will be printed on the console by default.

Value

By default, the function will print a table of contrasts and a table of simple effects.

132 skewness

Examples

```
two_way_anova(
  data = mtcars, dv_name = "mpg", iv_1_name = "vs",
  iv_2_name = "am", iterations = 100, plot = TRUE)
simple_effects_analysis(
  data = mtcars, dv_name = "mpg", iv_1_name = "vs",
  iv_2_name = "am")
```

skewness

Skewness

Description

Calculate skewness using one of three formulas: (1) the traditional Fisher-Pearson coefficient of skewness; (2) the adjusted Fisher-Pearson standardized moment coefficient; (3) the Pearson 2 skewness coefficient. Formulas were taken from Doane & Seward (2011), doi:10.1080/10691898.2011.11889611

Usage

```
skewness(vector = NULL, type = "adjusted")
```

Arguments

vector a numeric vector

type a character string indicating the type of skewness to calculate. If type = "adjusted",

the adjusted Fisher-Pearson standardized moment coefficient will be calculated. If type = "traditional", the traditional Fisher-Pearson coefficient of skewness will be calculated. If type = "pearson_2", the Pearson 2 skewness coeffi-

cient will be calculated. By default, type = "adjusted".

Value

a numeric value, i.e., skewness of the given vector

```
# calculate the adjusted Fisher-Pearson standardized moment coefficient
kim::skewness(c(1, 2, 3, 4, 5, 10))
# calculate the traditional Fisher-Pearson coefficient of skewness
kim::skewness(c(1, 2, 3, 4, 5, 10), type = "traditional")
# compare with skewness from 'moments' package
moments::skewness(c(1, 2, 3, 4, 5, 10))
# calculate the Pearson 2 skewness coefficient
kim::skewness(c(1, 2, 3, 4, 5, 10), type = "pearson_2")
```

```
spotlight_2_by_continuous
Spotlight 2 by Continuous
```

Description

Conduct a spotlight analysis for a 2 x Continuous design. See Spiller et al. (2013) doi:10.1509/jmr.12.0420

```
spotlight_2_by_continuous(
  data = NULL,
  iv_name = NULL,
 dv_name = NULL,
 mod_name = NULL,
 logistic = NULL,
  covariate_name = NULL,
  focal_values = NULL,
  interaction_p_include = TRUE,
  iv_level_order = NULL,
  output_type = "plot",
  colors = c("red", "blue"),
  dot_size = 3,
  observed_dots = FALSE,
  reg_lines = FALSE,
  reg_line_size = 1,
  lines_connecting_est_dv = TRUE,
  lines_connecting_est_dv_size = 1,
  estimated_dv_dot_shape = 15,
  estimated_dv_dot_size = 6,
 error_bar = "ci",
  error_bar_range = 0.95,
  error_bar_tip_width = NULL,
  error_bar_tip_width_percent = 8,
  error_bar_thickness = 1,
  error_bar_offset = NULL,
  error_bar_offset_percent = 8,
  simp_eff_bracket_leg_ht = NULL,
  simp_eff_bracket_leg_ht_perc = 2,
  simp_eff_bracket_offset = NULL,
  simp_eff_bracket_offset_perc = 1,
 simp_eff_bracket_color = "black",
  simp_eff_bracket_line_size = 1,
  simp_eff_text_offset = NULL,
  simp_eff_text_offset_percent = 7,
  simp_eff_text_hjust = 0.5,
```

```
simp_eff_text_part_1 = "Simple Effect\n",
  simp_eff_text_color = "black",
  simp_eff_font_size = 5,
  interaction_p_value_x = NULL,
  interaction_p_value_y = NULL,
  interaction_p_value_font_size = 6,
  interaction_p_value_vjust = -1,
  interaction_p_value_hjust = 0.5,
  x_axis_breaks = NULL,
  x_axis_limits = NULL,
  x_axis_tick_mark_labels = NULL,
  y_axis_breaks = NULL,
 y_axis_limits = NULL,
  x_axis_space_left_perc = 10,
  x_axis_space_right_perc = 30,
  y_axis_tick_mark_labels = NULL,
  x_axis_title = NULL,
  y_axis_title = NULL,
  legend_title = NULL,
  legend_position = "right",
  y_axis_title_vjust = 0.85,
  round_decimals_int_p_value = 3,
  jitter_x_percent = 0,
  jitter_y_percent = 0,
  dot_alpha = 0.2,
  reg_line_alpha = 0.5,
  jn_point_font_size = 6,
  reg_line_types = c("solid", "dashed"),
  caption = NULL,
  plot_margin = ggplot2::unit(c(60, 30, 7, 7), "pt"),
  silent = FALSE
)
```

Arguments

data a data object (a data frame or a data.table) name of the binary independent variable (IV) iv_name dv_name name of the dependent variable (DV) mod name name of the continuous moderator variable (MOD) logistic logical. Should logistic regressions be conducted, rather than ordinary least squares regressions? By default, ordinary least squares regressions will be conducted. covariate_name name(s) of the variable(s) to control for in estimating conditional values of the focal_values focal values of the moderator variable at which to estimate IV's effect on DV. interaction_p_include

logical. Should the plot include a p-value for the interaction term?

iv_level_order order of levels in the independent variable for legend. By default, it will be set as levels of the independent variable ordered using R's base function sort. type of output (default = "plot"). Other possible values include "spotlight_results", output_type "dt_for_plotting", "modified_dt" colors set colors for the two levels of the independent variable By default, colors = c("red", "blue"). size of the observed_dots (default = 3) dot_size observed_dots logical. If observed_dots = TRUE, the observed values of all IV, DV, and MOD combinations will be plotted as dots. On top of these dots the spotlight analysis plot will be laid. If observed_dots = FALSE, these dots will not be plotted. By default, observed_dots = FALSE. logical. If reg_lines = TRUE, the regression lines from regressing DV on MOD reg_lines at each value of IV will be plotted. If reg_lines = FALSE, these regression lines will not be plotted. By default, observed_dots = FALSE. reg_line_size thickness of the regression lines (default = 1). lines_connecting_est_dv logical. Should lines connecting the estimated values of DV be drawn? (default = TRUE)lines_connecting_est_dv_size thickness of the lines connecting the estimated values of DV (default = 1). estimated_dv_dot_shape ggplot value for shape of the dots at estimated values of DV (default = 15, a square shape). estimated_dv_dot_size size of the dots at estimated values of DV (default = 6). if error_bar = "se"; error bars will be +/-1 standard error, if error_bar = "ci" error_bar error bars will be a confidence interval. By default, error_bar = "ci". error_bar_range width of the confidence interval (default = 0.95 for a 95 percent confidence interval). This argument will not apply when error_bar = "se" error_bar_tip_width graphically, width of the segments at the end of error bars (default = 0.13) error_bar_tip_width_percent (default) error_bar_thickness thickness of the error bars (default = 1) error_bar_offset (default) error_bar_offset_percent (default) simp_eff_bracket_leg_ht (default) simp_eff_bracket_leg_ht_perc (default)

```
simp_eff_bracket_offset
                 (default)
simp_eff_bracket_offset_perc
                  (default)
simp_eff_bracket_color
                 (default)
simp_eff_bracket_line_size
                 (default)
simp_eff_text_offset
                 (default)
simp_eff_text_offset_percent
                 (default)
simp_eff_text_hjust
                 (default)
simp\_eff\_text\_part\_1
                 The first part of the text for labeling simple effects. By default, simp_eff_text_part_1
                  = "Simple Effect\n"
simp_eff_text_color
                  color for the text indicating p-values of simple effects (default = "black").
simp_eff_font_size
                 font size of the text indicating p-values of simple effects (default = 5).
interaction_p_value_x
                 (default)
interaction_p_value_y
                 (default)
interaction_p_value_font_size
                 font size for the interaction p value (default = 6)
interaction_p_value_vjust
                 (default)
interaction_p_value_hjust
                 (default)
x_axis_breaks
                 (default)
x_axis_limits
                 (default)
x_axis_tick_mark_labels
                 (default)
y_axis_breaks
                 (default)
y_axis_limits
                 (default)
x_axis_space_left_perc
                 (default)
x_axis_space_right_perc
                 (default)
y_axis_tick_mark_labels
                 (default)
```

```
title of the x axis. By default, it will be set as input for mod_name. If x_axis_title
x_axis_title
                   = FALSE, it will be removed.
                  title of the y axis. By default, it will be set as input for dv_name. If y_axis_title
y_axis_title
                   = FALSE, it will be removed.
legend_title
                  title of the legend. By default, it will be set as input for iv_name. If legend_title
                   = FALSE, it will be removed.
legend_position
                   position of the legend (default = "right"). If legend_position = "none", the
                   legend will be removed.
y_axis_title_vjust
                   position of the y axis title (default = 0.85). If default is used, y_axis_title_vjust
                   = 0.85, the y axis title will be positioned at 85% of the way up from the bottom
                  of the plot.
round_decimals_int_p_value
                  To how many digits after the decimal point should the p value for the interaction
                  term be rounded? (default = 3)
jitter_x_percent
                   horizontally jitter dots by a percentage of the range of x values
jitter_y_percent
                   vertically jitter dots by a percentage of the range of y values
                   opacity of the dots (0 = \text{completely transparent}, 1 = \text{completely opaque}). By
dot_alpha
                  default, dot_alpha = 0.2
reg_line_alpha (default)
jn_point_font_size
                  (default)
reg_line_types types of the regression lines for the two levels of the independent variable. By
                  default, reg_line_types = c("solid", "dashed")
caption
                  (default)
plot_margin
                  margin for the plot By default plot_margin = ggplot2::unit(c(60, 30, 7,
                  7), "pt")
                  If silent = FALSE, (various) messages will be printed. If silent = TRUE, the
silent
                  messages will be suppressed. By default, silent = FALSE.
```

```
spotlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "qsec")
# control for variables
spotlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
```

138 standardize

```
mod_name = "qsec",
covariate_name = c("cyl", "hp"))
# control for variables and adjust simple effect labels
spotlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "qsec",
covariate_name = c("cyl", "hp"),
reg_lines = TRUE,
observed_dots = TRUE,
error_bar_offset_percent = 3,
error_bar_tip_width_percent = 3,
simp_eff_text_offset_percent = 3,
simp_eff_bracket_leg_ht_perc = 2,
dot_alpha = 0.2,
simp_eff_text_part_1 = "")
# spotlight at specific values
spotlight_2_by_continuous(
data = mtcars,
iv_name = "am",
dv_name = "mpg",
mod_name = "qsec",
covariate_name = c("cyl", "hp"),
focal\_values = seq(15, 22, 1),
reg_lines = TRUE,
observed_dots = TRUE,
dot_alpha = 0.2,
simp_eff_text_part_1 = "",
simp_eff_font_size = 4,
error_bar_offset_percent = 3,
error_bar_tip_width_percent = 3,
simp_eff_text_offset_percent = 3,
simp_eff_bracket_leg_ht_perc = 1,
x_axis_breaks = seq(15, 22, 1))
# spotlight for logistic regression
spotlight_2_by_continuous(
data = mtcars,
iv_name = "am";
dv_name = "vs",
mod_name = "drat",
logistic = TRUE)
```

standardize

Standardize

Description

Standardize (i.e., normalize, obtain z-scores, or obtain the standard scores)

standardized_regression 139

Usage

```
standardize(x = NULL)
```

Arguments

Χ

a numeric vector

Value

the output will be a vector of the standard scores of the input.

Examples

```
standardize(1:10)
```

standardized_regression

Standardized Regression

Description

This function standardizes all variables for a regression analysis (i.e., dependent variable and all independent variables) and then conducts a regression with the standardized variables.

Usage

```
standardized_regression(
  data = NULL,
  formula = NULL,
  reverse_code_vars = NULL,
  sigfigs = NULL,
  round_digits_after_decimal = NULL,
  round_p = 3,
  pretty_round_p_value = TRUE,
  return_table_upper_half = FALSE,
  round_r_squared = 3,
  round_f_stat = 2,
  prettify_reg_table_col_names = TRUE)
```

Arguments

```
data a data object (a data frame or a data.table)

formula a formula object for the regression equation
reverse_code_vars
names of binary variables to reverse code
```

140 start_kim

```
number of significant digits to round to
sigfigs
round_digits_after_decimal
                  round to nth digit after decimal (alternative to sigfigs)
round_p
                  number of decimal places to which to round p-values (default = 3)
pretty_round_p_value
                  logical. Should the p-values be rounded in a pretty format (i.e., lower threshold:
                  "<.001"). By default, pretty_round_p_value = TRUE.
return_table_upper_half
                  logical. Should only the upper part of the table be returned? By default, return_table_upper_half
                  = FALSE.
round_r_squared
                  number of digits after the decimal both r-squared and adjusted r-squared values
                  should be rounded to (default 3)
                  number of digits after the decimal the f statistic of the regression model should
round_f_stat
                  be rounded to (default 2)
prettify_reg_table_col_names
                  logical. Should the column names of the regression table be made pretty (e.g.,
                  change "std_beta" to "Std. Beta")? (Default = TRUE)
```

Value

the output will be a data.table showing multiple regression results.

Examples

```
standardized_regression(data = mtcars, formula = mpg ~ gear * cyl)
standardized_regression(
data = mtcars, formula = mpg ~ gear + gear:am + disp * cyl,
round_digits_after_decimal = 3)
```

start_kim

Start kim

Description

Start kim (update kim; attach default packages; set working directory, etc.) This function requires installing Package 'remotes' v2.4.2 (or possibly a higher version) by Csardi et al. (2021), https://cran.r-project.org/package=remotes

start_kim 141

Usage

```
start_kim(
  update = TRUE,
  upgrade_other_pkg = FALSE,
  setup_r_env = TRUE,
  default_packages = c("data.table", "ggplot2"),
  silent_load_pkgs = c("data.table", "ggplot2")
)
```

Arguments

update

If update = "force", force updating the package 'kim'. If update = TRUE, compares the currently installed package 'kim' with the most recent version on GitHub and, if the version on GitHub is more recent, ask the user to confirm the update. If confirmed, then update the package. If update = FALSE, skip updating the package. By default, update = "force"

upgrade_other_pkg

input for the upgrade argument to be passed on to remotes::install_github. One of "default", "ask", "always", "never", TRUE, or FALSE. "default" respects the value of the R_REMOTES_UPGRADE environment variable if set, and falls back to "ask" if unset. "ask" prompts the user for which out of date packages to upgrade. For non-interactive sessions "ask" is equivalent to "always". TRUE and FALSE correspond to "always" and "never" respectively. By default, upgrade_other_pkg = FALSE.

setup_r_env

logical. If update = TRUE, runs the function setup_r_env in the package "kim". Type "?kim::setup_r_env" to learn more. By default, setup_r_env = TRUE

default_packages

a vector of names of packages to load and attach. By default, default_packages = c("data.table", "ggplot2")

silent_load_pkgs

a character vector indicating names of packages to load silently (i.e., suppress messages that get printed when loading the packages). By default, silent_load_pkgs = c("data.table", "ggplot2")

```
## Not run:
start_kim()
start_kim(default_packages = c("dplyr", "ggplot2"))
start_kim(update = TRUE, setup_r_env = FALSE)
## End(Not run)
```

142 tabulate_vector

su

su: Sorted unique values

Description

Extract unique elements and sort them

Usage

```
su(x = NULL, na.last = TRUE, decreasing = FALSE)
```

Arguments

x a vector or a data frame or an array or NULL.

na.last an argument to be passed onto the 'sort' function (in base R) for controlling the

treatment of NA values. If na.last = TRUE, NA values in the data are put last; if na.last = FALSE, NA values are put first; if na.last = NA, NA values are

removed. By default, na.last = TRUE

decreasing logical. Should the sort be increasing or decreasing? An argument to be passed

onto the 'sort' function (in base R). By default, decreasing = FALSE

Value

a vector, data frame, or array-like 'x' but with duplicate elements/rows removed.

Examples

```
su(c(10, 3, 7, 10, NA))
su(c("b", "z", "b", "a", NA, NA, NA))
```

tabulate_vector

Tabulate vector

Description

Shows frequency and proportion of unique values in a table format

```
tabulate_vector(
  vector = NULL,
  na.rm = TRUE,
  sort_by_decreasing_count = NULL,
  sort_by_increasing_count = NULL,
  sort_by_decreasing_value = NULL,
```

tabulate_vector 143

```
sort_by_increasing_value = NULL,
total_included = TRUE,
sigfigs = NULL,
round_digits_after_decimal = NULL,
output_type = "dt"
)
```

Arguments

```
vector
                  a character or numeric vector
                  if TRUE, NA values will be removed before calculating frequencies and propor-
na.rm
                  tions.
sort_by_decreasing_count
                  if TRUE, the output table will be sorted in the order of decreasing frequency.
sort_by_increasing_count
                  if TRUE, the output table will be sorted in the order of increasing frequency.
sort_by_decreasing_value
                  if TRUE, the output table will be sorted in the order of decreasing value.
sort_by_increasing_value
                  if TRUE, the output table will be sorted in the order of increasing value.
total_included if TRUE, the output table will include a row for total counts.
                  number of significant digits to round to
sigfigs
round_digits_after_decimal
                  round to nth digit after decimal (alternative to sigfigs)
                  if output_type = "df", return a data.frame. By default, output_type = "dt",
output_type
                  which will return a data.table.
```

Value

if output_type = "dt", which is the default, the output will be a data.table showing the count and proportion (percent) of each element in the given vector; if output_type = "df", the output will be a data.frame showing the count and proportion (percent) of each value in the given vector.

```
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA))
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_increasing_count = TRUE
)
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_decreasing_value = TRUE
)
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_increasing_value = TRUE
)
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
    sigfigs = 4
)
```

144 tau_squared

```
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
  round_digits_after_decimal = 1
)
tabulate_vector(c("a", "b", "b", "c", "c", "c", NA),
  output_type = "df"
)
```

tau_squared

Tau-squared (between-studies variance for meta analysis)

Description

Calculate tau-squared, the between-studies variance (the variance of the effect size parameters across the population of studies), as illustrated in Borenstein et al. (2009, pp. 72-73, ISBN: 978-0-470-05724-7).

Usage

```
tau_squared(effect_sizes = NULL, effect_size_variances = NULL)
```

Arguments

```
effect_sizes effect sizes (e.g., standardized mean differences)
effect_size_variances
    within-study variances
```

Details

Negative values of tau-squared are converted to 0 in the output (see Cheung, 2013; https://web.archive.org/web/20230512225

```
## Not run:
tau_squared(effect_sizes = c(1, 2), effect_size_variances = c(3, 4))
# a negative tau squared value is converted to 0:
tau_squared(effect_sizes = c(1.1, 1.4), effect_size_variances = c(1, 4))
## End(Not run)
```

theme_kim 145

theme_kim

Theme Kim

Description

A custom ggplot theme

Usage

```
theme_kim(
  legend_position = "none",
  legend_spacing_y = 1,
  legend_key_size = 3,
  base_size = 20,
  axis_tick_font_size = 20,
  axis_title_font_size = 24,
  y_axis_title_vjust = 0.85,
  axis_title_margin_size = 24,
  cap_axis_lines = FALSE
)
```

Arguments

```
legend_position
                  position of the legend (default = "none")
legend_spacing_y
                  vertical spacing of the legend keys in the unit of "cm" (default = 1)
legend_key_size
                  size of the legend keys in the unit of "lines" (default = 3)
base_size
                  base font size
axis_tick_font_size
                  font size for axis tick marks
axis_title_font_size
                  font size for axis title
y_axis_title_vjust
                  position of the y axis title (default = 0.85). If default is used, y_axis_title_vjust
                  = 0.85, the y axis title will be positioned at 85% of the way up from the bottom
                  of the plot.
axis_title_margin_size
                  size of the margin between axis title and the axis line
cap_axis_lines logical. Should the axis lines be capped at the outer tick marks? (default =
                  FALSE)
```

Details

If a axis lines are to be capped at the ends, the following package(s) must be installed prior to running the function: Package 'lemon' v0.4.4 (or possibly a higher version) by Edwards et al. (2020), https://cran.r-project.org/package=lemon

Value

```
a ggplot object; there will be no meaningful output from this function. Instead, this function should be used with another ggplot object, e.g., ggplot(mtcars, aes(x = disp, y = mpg)) + theme_kim()
```

Examples

```
prep(ggplot2)
ggplot2::ggplot(mtcars, aes(x = cyl, y = mpg)) +
geom_point() + theme_kim()
```

Description

Indicates whether each value in a vector belongs to top, median, or bottom

Usage

```
top_median_or_bottom(vector)
```

Arguments

vector a numeric vector

Value

a character vector indicating whether each element in a vector belongs to "top", "median", or "bottom"

```
top_median_or_bottom(c(1, 2, 3, NA)) top_median_or_bottom(c(1, 2, 2, NA)) top_median_or_bottom(c(1, 1, 2, NA))
```

tv 147

tv Tabulate vector

Description

Shows frequency and proportion of unique values in a table format. This function is a copy of the earlier function, tabulate vector, in Package 'kim'

Usage

```
tv(
  vector = NULL,
  na.rm = FALSE,
  sort_by_decreasing_count = NULL,
  sort_by_increasing_count = NULL,
  sort_by_decreasing_value = NULL,
  sort_by_increasing_value = NULL,
  total_included = TRUE,
  sigfigs = NULL,
  round_digits_after_decimal = NULL,
  output_type = "dt"
)
```

Arguments

vector a character or numeric vector na.rm if TRUE, NA values will be removed before calculating frequencies and proportions. By default, FALSE. sort_by_decreasing_count if TRUE, the output table will be sorted in the order of decreasing frequency. sort_by_increasing_count if TRUE, the output table will be sorted in the order of increasing frequency. sort_by_decreasing_value if TRUE, the output table will be sorted in the order of decreasing value. sort_by_increasing_value if TRUE, the output table will be sorted in the order of increasing value. total_included if TRUE, the output table will include a row for total counts. sigfigs number of significant digits to round to round_digits_after_decimal round to nth digit after decimal (alternative to sigfigs) if output_type = "df", return a data.frame. By default, output_type = "dt", output_type which will return a data.table.

148 two_way_anova

Value

if output_type = "dt", which is the default, the output will be a data.table showing the count and proportion (percent) of each element in the given vector; if output_type = "df", the output will be a data.frame showing the count and proportion (percent) of each value in the given vector.

Examples

```
tv(c("a", "b", "b", "c", "c", "c", NA))
tv(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_increasing_count = TRUE
)
tv(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_decreasing_value = TRUE
)
tv(c("a", "b", "b", "c", "c", "c", NA),
    sort_by_increasing_value = TRUE
)
tv(c("a", "b", "b", "c", "c", "c", NA),
    sigfigs = 4
)
tv(c("a", "b", "b", "c", "c", "c", NA),
    round_digits_after_decimal = 1
)
tv(c("a", "b", "b", "c", "c", "c", NA),
    output_type = "df"
)
```

two_way_anova

Two-way ANOVA

Description

Conduct a two-way analysis of variance (ANOVA).

Usage

```
two_way_anova(
  data = NULL,
  dv_name = NULL,
  iv_1_name = NULL,
  iv_2_name = NULL,
  iv_1_values = NULL,
  iv_2_values = NULL,
  sigfigs = 3,
  robust = FALSE,
  iterations = 2000,
  plot = FALSE,
  error_bar = "ci",
```

two_way_anova 149

```
error_bar_range = 0.95,
error_bar_tip_width = 0.13,
error_bar_thickness = 1,
error_bar_caption = TRUE,
line_thickness = 1,
dot_size = 3,
position_dodge = 0.13,
legend_position = "right",
output = "anova_table",
png_name = NULL,
width = 7000,
height = 4000,
units = "px",
res = 300,
layout_matrix = NULL
```

Arguments

data a data object (a data frame or a data.table) dv_name name of the dependent variable iv_1_name name of the first independent variable iv_2_name name of the second independent variable iv_1_values restrict all analyses to observations having these values for the first independent variable iv_2_values restrict all analyses to observations having these values for the second independent variable sigfigs number of significant digits to which to round values in anova table (default = if TRUE, conduct a robust ANOVA in addition. robust number of bootstrap samples for robust ANOVA. The default is set at 2000, but iterations consider increasing the number of samples to 5000, 10000, or an even larger number, if slower handling time is not an issue. plot if TRUE, print a plot and enable returning an output. if error_bar = "se"; error bars will be +/-1 standard error; if error_bar = error_bar "ci" error bars will be a confidence interval error_bar_range width of the confidence interval (default = 0.95 for 95 percent confidence interval). This argument will not apply when error_bar = "se" error_bar_tip_width graphically, width of the segments at the end of error bars (default = 0.13) error_bar_thickness thickness of the error bars (default = 1) error_bar_caption should a caption be included to indicate the width of the error bars? (default = TRUE).

150 two_way_anova

line_thickness thickness of the lines connecting group means, (default = 1)
dot_size size of the dots indicating group means (default = 3)

position_dodge by how much should the group means and error bars be horizontally offset from

each other so as not to overlap? (default = 0.13)

 $legend_position$

position of the legend: "none", "top", "right", "bottom", "left", "none"

(default = "right")

output type can be one of the following: "anova_table", "group_stats",

"plot", "robust_anova_results", "robust_anova_post_hoc_results", "robust_anova_post_hoc

"all"

png_name name of the PNG file to be saved. If png_name = TRUE, the name will be "two_way_anova_"

followed by a timestamp of the current time. The timestamp will be in the format, jan_01_2021_1300_10_000001, where "jan_01_2021" would indicate January 01, 2021; 1300 would indicate 13:00 (i.e., 1 PM); and 10_000001 would

indicate 10.000001 seconds after the hour.

width width of the PNG file (default = 7000) height height of the PNG file (default = 4000)

units the units for the width and height arguments. Can be "px" (pixels), "in"

(inches), "cm", or "mm". By default, units = "px".

res The nominal resolution in ppi which will be recorded in the png file, if a positive

integer. Used for units other than the default. If not specified, taken as 300 ppi

to set the size of text and line widths.

layout_matrix The layout argument for arranging plots and tables using the grid.arrange

function.

Details

The following package(s) must be installed prior to running this function: Package 'car' v3.0.9 (or possibly a higher version) by Fox et al. (2020), https://cran.r-project.org/package=car

If robust ANOVA is to be conducted, the following package(s) must be installed prior to running the function: Package 'WRS2' v1.1-1 (or possibly a higher version) by Mair & Wilcox (2021), https://cran.r-project.org/package=WRS2

Value

by default, the output will be "anova_table"

```
two_way_anova(
  data = mtcars, dv_name = "mpg", iv_1_name = "vs",
  iv_2_name = "am", iterations = 100)
anova_results <- two_way_anova(
  data = mtcars, dv_name = "mpg", iv_1_name = "vs",
  iv_2_name = "am", output = "all")
anova_results</pre>
```

t_test_pairwise 151

t_test_pairwise t-tests, pairwise

Description

Conducts a t-test for every possible pairwise comparison with Holm or Bonferroni correction

Usage

```
t_test_pairwise(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  sigfigs = 3,
  cohen_d = TRUE,
  cohen_d_w_ci = TRUE,
  adjust_p = "holm",
  bonferroni = NULL,
 mann_whitney = TRUE,
  t_test_stats = FALSE,
  t_test_df_decimals = 1,
  sd = FALSE,
  round_p = 3,
  anova = TRUE,
  round_f = 2
)
```

Arguments data

data	a data object (a data frame or a data.table)
iv_name	name of the independent variable
dv_name	name of the dependent variable
sigfigs	number of significant digits to round to
cohen_d	if cohen_d = TRUE, Cohen's d statistics will be included in the output data.table.
cohen_d_w_ci	if cohen_d_w_ci = TRUE, Cohen's d with 95% CI will be included in the output data.table.
adjust_p	the name of the method to use to adjust p-values. If $adjust_p = "holm"$, the Holm method will be used; if $adjust_p = "bonferroni"$, the Bonferroni method will be used. By default, $adjust_p = "holm"$
bonferroni	The use of this argument is deprecated. Use the 'adjust_p' argument instead. If bonferroni = TRUE, Bonferroni tests will be conducted for t-tests or Mann-Whitney tests.
mann_whitney	if TRUE, Mann-Whitney test results will be included in the output data.table. If FALSE, Mann-Whitney tests will not be performed.

152 und

t_test_stats if t_test_stats = TRUE, t-test statistic and degrees of freedom will be included in the output data.table.

t_test_df_decimals

number of decimals for the degrees of freedom in t-tests (default = 1)

sd if sd = TRUE, standard deviations will be included in the output data.table.

round_p number of decimal places to which to round p-values (default = 3)

anova Should a one-way ANOVA be conducted and reported? (default = TRUE) round_f number of decimal places to which to round the f statistic (default = 2)

Value

the output will be a data.table showing results of all pairwise comparisons between levels of the independent variable.

Examples

```
## Not run:
t_test_pairwise(data = iris, iv_name = "Species", dv_name = "Sepal.Length")
t_test_pairwise(data = iris, iv_name = "Species",
dv_name = "Sepal.Length", t_test_stats = TRUE, sd = TRUE)
t_test_pairwise(data = iris, iv_name = "Species", dv_name = "Sepal.Length",
mann_whitney = FALSE)
## End(Not run)
```

und

Undocumented functions

Description

A collection of miscellaneous functions lacking documentations

Usage

```
und(fn, ...)
```

Arguments

fn name of the function
... arguments for the function

Value

the output will vary by function

Examples

```
# correlation
und(corr_text, x = 1:5, y = c(1, 2, 2, 2, 3))
# mean center
und(mean_center, 1:10)
# compare results with base function
scale(1:10, scale = TRUE)
# find the modes
und(mode, c(3, 3, 3, 1, 2, 2))
# return values that are not outliers
und(outlier_rm, c(12:18, 100))
kim::outlier(c(1:10, 100))
```

```
unload_user_installed_pkgs
```

Unload all user-installed packages

Description

Unload all user-installed packages

Usage

```
unload_user_installed_pkgs(exceptions = NULL, force = FALSE, keep_kim = TRUE)
```

Arguments

exceptions a character vector of names of packages to keep loaded

force logical. Should a package be unloaded even though other attached packages

depend on it? By default, force = FALSE

keep_kim logical. If keep_kim = FALSE, Package 'kim' will be detached along with all

other user-installed packages. If keep_kim = TRUE, Package 'kim' will not be

detached. By default, keep_kim = FALSE

```
## Not run:
unload_user_installed_pkgs()
## End(Not run)
```

154 update_kim

update_kim

Update the package 'kim'

Description

Updates the current package 'kim' by installing the most recent version of the package from GitHub This function requires installing Package 'remotes' v2.4.2 (or possibly a higher version) by Csardi et al. (2021), https://cran.r-project.org/package=remotes

Usage

```
update_kim(force = TRUE, upgrade_other_pkg = FALSE, confirm = TRUE)
```

Arguments

force

logical. If force = TRUE, force installing the update. If force = FALSE, do not force installing the update. By default, force = TRUE.

upgrade_other_pkg

input for the upgrade argument to be passed on to remotes::install_github. One of "default", "ask", "always", "never", TRUE, or FALSE. "default" respects the value of the R_REMOTES_UPGRADE environment variable if set, and falls back to "ask" if unset. "ask" prompts the user for which out of date packages to upgrade. For non-interactive sessions "ask" is equivalent to "always". TRUE and FALSE correspond to "always" and "never" respectively. By default, upgrade_other_pkg = FALSE.

confirm

logical. If confirm = TRUE, the user will need to confirm the update. If confirm = FALSE, the confirmation step will be skipped. By default, confirm = TRUE.

Value

there will be no output from this function. Rather, executing this function will update the current 'kim' package by installing the most recent version of the package from GitHub.

```
## Not run:
if (interactive()) {update_kim()}
## End(Not run)
```

```
var_of_log_odds_ratio_to_var_of_d
```

Convert variance of log odds ratio to variance of d

Description

Convert the variance of a log odds ratio to the variance of a Cohen'd (standardized mean difference), as illustrated in Borenstein et al. (2009, p. 47, ISBN: 978-0-470-05724-7)

Usage

```
var_of_log_odds_ratio_to_var_of_d(var_of_log_odds_ratio = NULL)
```

Arguments

```
var_of_log_odds_ratio
```

the variance of a log odds ratio (the input can be a vector of values)

Examples

```
## Not run:
var_of_log_odds_ratio_to_var_of_d(1)
## End(Not run)
```

var_of_percentage

Variance of a percentage

Description

Calculate the variance of a percentage. See Fowler, Jr. (2014, p. 34, ISBN: 978-1-4833-1240-8)

Usage

```
var_of_percentage(percent = NULL, n = NULL)
```

Arguments

percent a vector of percentages; each of the percentage values must be between 0 and

100

n a vector of sample sizes; number of observations used to calculate each of the

percentage values

```
var_of_percentage(percent = 40, n = 50)
var_of_percentage(percent = 50, n = 10)
```

156 vlookup

var_of_proportion

Variance of a proportion

Description

Calculate the variance of a proportion. See Anderson and Finn (1996, p. 364, ISBN: 978-1-4612-8466-6)

Usage

```
var_of_proportion(p = NULL, n = NULL)
```

Arguments

p a vector of proportions; each of the proportion values must be between 0 and 1

a vector of sample sizes; number of observations used to calculate each of the percentage values

Examples

n

vlookup

Vlookup

Description

Look up values in a reference data.table and return values associated with the looked-up values contained in the reference data.table

Usage

```
vlookup(
  lookup_values = NULL,
  reference_dt = NULL,
  col_name_for_lookup_values = NULL,
  col_name_for_output_values = NULL
)
```

Arguments

Examples

```
\label{eq:vlookup} $$ vlookup(lookup\_values = c(2.620, 2.875), reference\_dt = mtcars[1:9, ], $$ col\_name\_for\_lookup\_values = "wt", col\_name\_for\_output\_values = "qsec") $$
```

```
weighted_mean_effect_size
```

Estimate the mean effect size in a meta analysis

Description

Estimate the mean effect size in a meta analysis, as illustrated in Borenstein et al. (2009, pp. 73-74, ISBN: 978-0-470-05724-7)

Usage

```
weighted_mean_effect_size(
  effect_sizes = NULL,
  effect_size_variances = NULL,
  ci = 0.95,
  one_tailed = FALSE,
  random_vs_fixed = "random"
)
```

Arguments

weighted_mean_r

Examples

```
## Not run:
weighted_mean_effect_size(
effect_sizes = c(1, 2), effect_size_variances = c(3, 4))
weighted_mean_effect_size(
effect_sizes = c(0.095, 0.277, 0.367, 0.664, 0.462, 0.185),
effect_size_variances = c(0.033, 0.031, 0.050, 0.011, 0.043, 0.023))
# if effect sizes have a variance of 0, they will be excluded from
# the analysis
weighted_mean_effect_size(
effect_sizes = c(1.1, 1.2, 1.3, 1.4),
effect_size_variances = c(1, 0, 0, 4))
## End(Not run)
```

weighted_mean_r

Weighted mean correlation

Description

Calculate the weighted mean correlation coefficient for a given correlations and sample sizes. This function uses the Hedges-Olkin Method with random effects. See Field (2001) doi:10.1037/1082-989X.6.2.161

Usage

```
weighted_mean_r(r = NULL, n = NULL, ci = 0.95, sigfigs = 3, silent = FALSE)
```

Arguments

r	a (vector of) correlation coefficient(s)
n	a (vector of) sample size(s)
ci	width of the confidence interval. Input can be any value less than 1 and greater than or equal to 0. By default, $ci = 0.95$. If $ci = TRUE$, the default value of 0.95 will be used. If $ci = FALSE$, no confidence interval will be estimated.
sigfigs	number of significant digits to round to (default = 3)
silent	logical. If silent = FALSE, a message regarding the weighted mean correlation and its p-value and CI will be printed. If silent = TRUE, this message will be suppressed. By default, silent = FALSE.

Value

the output will be a list of vector of correlation coefficient(s).

weighted_z

Examples

```
weighted_mean_r(r = c(0.2, 0.4), n = c(100, 100)) weighted_mean_r(r = c(0.2, 0.4), n = c(100, 20000)) # example consistent with using MedCalc weighted_mean_r( r = c(0.51, 0.48, 0.3, 0.21, 0.6, 0.46, 0.22, 0.25), n = c(131, 129, 155, 121, 111, 119, 112, 145))
```

weighted_z

Weighted z

Description

Calculate the weighted z (for calculating weighted mean correlation). See p. 231 of the book Hedges & Olkin (1985), Statistical Methods for Meta-Analysis (ISBN: 0123363802).

Usage

```
weighted_z(z = NULL, n = NULL)
```

Arguments

z a vector of z values

a vector of sample sizes which will be used to calculate the weights, which in turn will be used to calculate the weighted z.

Value

the output will be a weighted z value.

Examples

```
weighted_z(1:3, c(100, 200, 300)) weighted_z(z = c(1:3, NA), n = c(100, 200, 300, NA))
```

wilcoxon_rank_sum_test

Wilcoxon Rank-Sum Test (Also called the Mann-Whitney U Test)

Description

A nonparametric equivalent of the independent t-test

160 write_csv

Usage

```
wilcoxon_rank_sum_test(
  data = NULL,
  iv_name = NULL,
  dv_name = NULL,
  sigfigs = 3
)
```

Arguments

data a data object (a data frame or a data.table)

iv_name name of the independent variable (grouping variable)

dv_name name of the dependent variable (measure variable of interest)

sigfigs number of significant digits to round to

Value

the output will be a data.table object with all pairwise Wilcoxon rank-sum test results

Examples

```
wilcoxon_rank_sum_test(
data = iris, iv_name = "Species", dv_name = "Sepal.Length")
```

write_csv

Write to a csv file

Description

Write to a csv file

Usage

```
write_csv(data = NULL, name = NULL, timestamp = NULL)
```

Arguments

data a data object (a data frame or a data.table)

name a character string of the csv file name without the ".csv" extension. For example,

if the csv file to write to is "myfile.csv", enter name = "myfile"

timestamp logical. Should the timestamp be appended to the file name?

Value

the output will be a .csv file in the working directory, that is, an output from the data.table function, fwrite

z_score 161

Examples

```
## Not run:
write_csv(mtcars, "mtcars_from_write_csv")
write_csv(mtcars)
## End(Not run)
```

z_score

z score

Description

Calculate z-scores (i.e., standardize or obtain the standard scores)

Usage

```
z_score(x = NULL, na.rm = TRUE)
```

Arguments

x a numeric vector

na.rm

logical. If na.rm = TRUE, NA values in the vector will be removed before calculating z-scores (default = TRUE).

Value

the output will be a vector of z-scores.

Examples

```
z_score(1:10)
```

 $z_{to}_r_{transform}$

Z to r transformation (Inverse of Fisher's Z transformation)

Description

Perform the Z-to-r transformation (i.e., the inverse of Fisher's r-to-Z transformation) for given Z value(s).

Usage

```
z_{to} = NULL
```

Arguments

z

a (vector of) Z values

z_to_r_transform

Value

the output will be a vector of correlation coefficient(s) that are the result(s) of the Z-to-r transformation.

```
z_to_r_transform(2.646652)
z_to_r_transform(z = -3:3)
```

Index

akaike_weights,5	exit_from_parent_function, 41
barplot_for_counts, 5	find_duplicates, 42
binomial_test, 6	fisher_z_transform, 43
bracket, 7	floodlight_2_by_continuous, 43
<pre>capitalize, 9 change_var_names, 9</pre>	<pre>floodlight_2_by_continuous_logistic,</pre>
check_modes, 10	50
check_req_pkg, 11	floodlight_for_contrasts, 54
<pre>chi_squared_test, 11</pre>	floodlight_multi_by_continuous, 58
<pre>chi_squared_test_pairwise, 12</pre>	forest_plot, 61
clean_data_from_qualtrics, 14	101 C3t_p10t, 01
<pre>coefficent_of_variation, 15</pre>	geomean, 64
cohen_d, 16	ggsave_quick, 65
cohen_d_borenstein, 17	36
<pre>cohen_d_for_one_sample, 18</pre>	histogram, 66
cohen_d_from_cohen_textbook, 19	histogram_by_group,68
cohen_d_over_n, 20	histogram_deprecated_1,69
cohen_d_to_r, 22	histogram_from_hist,71
cohen_d_torchiano, 21	histogram_w_outlier_bins,72
combine_data_across_cols, 22	holm_adjusted_p, 74
comma_sep_string_to_numbers, 23	
compare_datasets, 24	id_across_datasets, 76
compare_dependent_rs, 25	identical_all, 75
compare_effect_sizes, 26	<pre>install_all_dependencies, 77</pre>
compare_groups, 27	kurtosis,77
compare_independent_rs, 29	Kui 10313, 77
contingency_table, 31	lenu, 78
convert_cols_to_numeric, 32	levene_test, 79
<pre>convert_to_excel_formula, 33 correlation_kim, 33</pre>	log_odds_ratio,86
correlation_matrix, 35	log_odds_ratio_to_d,87
cum_percent_plot, 36	<pre>logistic_reg_w_interaction, 83</pre>
cum_per cert_prot, 30	logistic_regression, 80
desc_stats, 36	logistic_regression_table,82
desc_stats_by_group, 37	loglinear_analysis,84
detach_user_installed_pkgs, 39	
duplicated_values, 39	mad_remove_outliers, 87
	mann_whitney, 89
excel_formula_convert,40	matrix_prep_dt,89

INDEX

mean_center, 90	skewness, 132
mediation_analysis, 91	spotlight_2_by_continuous, 133
merge_data_table_list, 94	standardize, 138
merge_data_tables, 92	standardized_regression, 139
modes_of_objects, 95	start_kim, 140
multiple_regression, 96	su, 142
	33, 112
noncentrality_parameter, 98	t_test_pairwise, 151
	tabulate_vector, 142
odds_ratio, 98	tau_squared, 144
order_rows_specifically_in_dt,99	theme_kim, 145
outlier, 100	top_median_or_bottom, 146
overlapping_interval, 101	tv, 147
	two_way_anova, 148
p0, 102	
package_list_default, 103	und, 152
parallel_analysis, 103	unload_user_installed_pkgs, 153
percentile_rank, 105	update_kim, 154
pivot_table, 105	· [· · · · · · - · · · · · · · · · · ·
plot_group_means, 106	<pre>var_of_log_odds_ratio_to_var_of_d, 155</pre>
pm, 108	var_of_percentage, 155
population_variance, 109	var_of_proportion, 156
prep, 109	vlookup, 156
pretty_round_p_value, 110	• •
pretty_round_r, 111	<pre>weighted_mean_effect_size, 157</pre>
print_loop_progress, 112	weighted_mean_r, 158
proportion_of_values_in_vector, 113	weighted_z, 159
	wilcoxon_rank_sum_test, 159
q_stat_test_homo_r, 114	write_csv, 160
mand and 115	464
read_csv, 115	z_score, 161
read_sole_csv, 115	z_to_r_transform, 161
regex_match, 116	
rel_pos_of_value_in_vector, 117	
rel_value_of_pos_in_vector, 117	
remove_from_vector, 118	
remove_user_installed_pkgs, 119	
repeated_measures_anova, 119	
replace_values_in_dt, 121	
robust_regression, 121	
round_flexibly, 122	
scatterplot, 124	
score_scale_items, 127	
se_of_mean, 129	
se_of_percentage, 130	
se_of_proportion, 130	
setup_r_env, 128	
setwd_to_active_doc, 128	
simple_effects_analysis, 131	
Simpic_errects_anarysis, 131	