Package ‘statsExpressions’

August 11, 2022

Type Package

Title Tidy Dataframes and Expressions with Statistical Details

Version 1.3.3

Maintainer Indrajeet Patil <patilindrajeet.science@gmail.com>

Description Utilities for producing dataframes with rich details for the most common types of statistical approaches and tests: parametric, nonparametric, robust, and Bayesian t-test, one-way ANOVA, correlation analyses, contingency table analyses, and meta-analyses. The functions are pipe-friendly and provide a consistent syntax to work with tidy data. These dataframes additionally contain expressions with statistical details, and can be used in graphing packages. This package also forms the statistical processing backend for 'ggstatsplot'.

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BugReports https://github.com/IndrajeetPatil/statsExpressions/issues

Depends R (>= 4.0.0)

Imports BayesFactor (>= 0.9.12-4.4), correlation (>= 0.8.2), datawizard (>= 0.5.0), dplyr, effectsize (>= 0.7.0.5), glue, insight (>= 0.18.2), magrittr, parameters (>= 0.18.2), performance (>= 0.9.2), rlang, stats, tibble, tidyr, WRS2 (>= 1.1-3), zeallot

Suggests afex, ggplot2, knitr, metaBMA, metafor, metaplus, PMCMRplus, purrr, rmarkdown, spelling, survival, testthat (>= 3.1.4), utils, withr

VignetteBuilder knitr

Encoding UTF-8

Language en-US

LazyData true
Description

The {statsExpressions} package has two key aims:

- to provide a consistent syntax to do statistical analysis with tidy data (in pipe-friendly manner),
- to provide statistical expressions (pre-formatted in-text statistical results) for plotting functions.

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statsExpressions-package

statsExpressions: Tidy Dataframes and Expressions with Statistical Details

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Statistical packages exhibit substantial diversity in terms of their syntax and expected input type. This can make it difficult to switch from one statistical approach to another. For example, some functions expect vectors as inputs, while others expect dataframes. Depending on whether it is a repeated measures design or not, different functions might expect data to be in wide or long format. Some functions can internally omit missing values, while other functions error in their presence. Furthermore, if someone wishes to utilize the objects returned by these packages downstream in their workflow, this is not straightforward either because even functions from the same package can return a list, a matrix, an array, a dataframe, etc., depending on the function.

This is where \{statsExpressions\} comes in: It can be thought of as a unified portal through which most of the functionality in these underlying packages can be accessed, with a simpler interface and no requirement to change data format.

This package forms the statistical processing backend for \texttt{ggstatsplot} package.

For more documentation, see the dedicated \texttt{website}.

**Details**

statsExpressions

**Author(s)**

Maintainer: Indrajeet Patil \texttt{<patilindrajeet.science@gmail.com>} (ORCID) (@patilindrajeets) [copyright holder]

**See Also**

Useful links:

- \url{https://indrajeetpatil.github.io/statsExpressions/}
- \url{https://github.com/IndrajeetPatil/statsExpressions}
- Report bugs at \url{https://github.com/IndrajeetPatil/statsExpressions/issues}

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**add_expression_col**

*Template for expressions with statistical details*

**Description**

Creates an expression from a dataframe containing statistical details. Ideally, this dataframe would come from having run \texttt{tidy_model_parameters} function on your model object.

This function is currently \textbf{not} stable and should not be used outside of this package context.
add_expression_col

Usage

add_expression_col(
  data,
  paired = FALSE,
  statistic.text = NULL,
  effsize.text = NULL,
  prior.type = NULL,
  n = NULL,
  n.text = ifelse(paired, list(quote(italic("n")["pairs"])),
                  list(quote(italic("n")["obs"])),
                  k = 2L,
                  k.df = 0L,
                  k.df.error = k.df,
                  ...
)

Arguments

  data               A dataframe containing details from the statistical analysis and should contain
                     some or all of the the following columns:
                     • statistic: the numeric value of a statistic.
                     • df.error: the numeric value of a parameter being modeled (often degrees of
                       freedom for the test); note that if there are no degrees of freedom (e.g., for
                       non-parametric tests), this column will be irrelevant.
                     • df: relevant only if the statistic in question has two degrees of freedom.
                     • p.value: the two-sided p-value associated with the observed statistic.
                     • method: method describing the test carried out.
                     • effectsize: name of the effect size (if not present, same as method).
                     • estimate: estimated value of the effect size.
                     • conf.level: width for the confidence intervals.
                     • conf.low: lower bound for effect size estimate.
                     • conf.high: upper bound for effect size estimate.
                     • bf10: Bayes Factor value (if bayesian = TRUE).

  paired             Logical that decides whether the experimental design is repeated measures/within-
                     subjects or between-subjects. The default is FALSE.

  statistic.text     A character that specifies the relevant test statistic. For example, for tests with
                     t-statistic, statistic.text = "t".

  effsize.text       A character that specifies the relevant effect size.

  prior.type         The type of prior.

  n                  An integer specifying the sample size used for the test.

  n.text             A character that specifies the design, which will determine what the n stands
                     for. It defaults to quote(italic("n")["pairs"]) if paired = TRUE, and to
                     quote(italic("n")["obs"]) if paired = FALSE. If you wish to customize this
                     further, you will need to provide object of language type.
**Description**

Tidy version of the "Bugs" dataset.

**Usage**

`bugs_long`

**Format**

A data frame with 372 rows and 6 variables

- `subject`: Dummy identity number for each participant.
- gender. Participant’s gender (Female, Male).
- region. Region of the world the participant was from.
- education. Level of education.
- condition. Condition of the experiment the participant gave rating for (LDLF: low freighteningness and low disgustingness; LFHD: low freighteningness and high disgustingness; HFHD: high freighteningness and low disgustingness; HFHD: high freighteningness and high disgustingness).
- desire. The desire to kill an arthropod was indicated on a scale from 0 to 10.

Details

This data set, "Bugs", provides the extent to which men and women want to kill arthropods that vary in freighteningness (low, high) and disgustingness (low, high). Each participant rates their attitudes towards all anthropods. Subset of the data reported by Ryan et al. (2013).

Source


Examples

dim(bugs_long)
head(bugs_long)
dplyr::glimpse(bugs_long)

Description

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

<table>
<thead>
<tr>
<th>Type</th>
<th>Measure</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>mean</td>
<td>parameters::describe_distribution()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>median</td>
<td>parameters::describe_distribution()</td>
</tr>
<tr>
<td>Robust</td>
<td>trimmed mean</td>
<td>parameters::describe_distribution()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>MAP (maximum a posteriori probability) estimate</td>
<td>parameters::describe_distribution()</td>
</tr>
</tbody>
</table>
Usage

centrality_description(data, x, y, type = "parametric", tr = 0.2, k = 2L, ...)

Arguments

data A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from {dplyr} should be ungrouped before they are entered as data.
x The grouping (or independent) variable in data.
y The response (or outcome or dependent) variable from data.
type A character specifying the type of statistical approach:
  • "parametric"
  • "nonparametric"
  • "robust"
  • "bayes"
   You can specify just the initial letter.
tr Trim level for the mean when carrying out robust tests. In case of an error, try reducing the value of tr, which is by default set to 0.2. Lowering the value might help.
k Number of digits after decimal point (should be an integer) (Default: k = 2L).
... Currently ignored.

Details

This function describes a distribution for y variable for each level of the grouping variable in x by a set of indices (e.g., measures of centrality, dispersion, range, skewness, kurtosis, etc.). It additionally returns an expression containing a specified centrality measure. The function internally relies on datawizard::describe_distribution() function.

Examples

set.seed(123)

# parametric -----------------------
centrality_description(iris, Species, Sepal.Length)

# non-parametric -------------------
centrality_description(mtcars, am, wt, type = "n")

# robust ---------------------------
centrality_description(ToothGrowth, supp, len, type = "r")

# Bayesian -------------------------
centrality_description(sleep, group, extra, type = "b")
### Contingency Table Analyses

**Description**

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

**Two-way Table:**

#### Hypothesis Testing

<table>
<thead>
<tr>
<th>Type</th>
<th>Design</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric/Non-parametric</td>
<td>Unpaired</td>
<td>Pearson’s chi-squared test</td>
<td>stats::chisq.test()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Unpaired</td>
<td>Bayesian Pearson’s chi-squared test</td>
<td>BayesFactor::contingencyTableBF()</td>
</tr>
<tr>
<td>Parametric/Non-parametric</td>
<td>Paired</td>
<td>McNemar’s chi-squared test</td>
<td>stats::mcnemar.test()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Paired</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

#### Effect Size Estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>Design</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric/Non-parametric</td>
<td>Unpaired</td>
<td>Cramer’s V</td>
<td>Yes</td>
<td>effectsize::cramers_v()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Unpaired</td>
<td>Cramer’s V</td>
<td>Yes</td>
<td>effectsize::cramers_v()</td>
</tr>
<tr>
<td>Parametric/Non-parametric</td>
<td>Paired</td>
<td>Cohen’s g</td>
<td>Yes</td>
<td>effectsize::cohens_g()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Paired</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

**One-way Table:**

#### Hypothesis Testing

<table>
<thead>
<tr>
<th>Type</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric/Non-parametric</td>
<td>Goodness of fit chi-squared test</td>
<td>stats::chisq.test()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Bayesian Goodness of fit chi-squared test (custom)</td>
<td></td>
</tr>
</tbody>
</table>

#### Effect Size Estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric/Non-parametric</td>
<td>Pearson’s C</td>
<td>Yes</td>
<td>effectsize::pearsons_c()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
Usage

contingency_table(
  data,
  x,
  y = NULL,
  paired = FALSE,
  type = "parametric",
  counts = NULL,
  ratio = NULL,
  k = 2L,
  conf.level = 0.95,
  sampling.plan = "indepMulti",
  fixed.margin = "rows",
  prior.concentration = 1,
  ...)

Arguments

data A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from {dplyr} should be ungrouped before they are entered as data.

x The variable to use as the rows in the contingency table.

y The variable to use as the columns in the contingency table. Default is NULL. If NULL, one-sample proportion test (a goodness of fit test) will be run for the x variable. Otherwise association test will be carried out.

paired Logical indicating whether data came from a within-subjects or repeated measures design study (Default: FALSE). If TRUE, McNemar’s test expression will be returned. If FALSE, Pearson’s chi-square test will be returned.

type A character specifying the type of statistical approach:
  • "parametric"
  • "nonparametric"
  • "robust"
  • "bayes"

You can specify just the initial letter.

counts The variable in data containing counts, or NULL if each row represents a single observation.

ratio A vector of proportions: the expected proportions for the proportion test (should sum to 1). Default is NULL, which means the null is equal theoretical proportions across the levels of the nominal variable. This means if there are two levels this will be ratio = c(0.5, 0.5) or if there are four levels this will be ratio = c(0.25, 0.25, 0.25, 0.25), etc.

k Number of digits after decimal point (should be an integer) (Default: k = 2L).
contingency_table

conf.level  Scalar between 0 and 1. If unspecified, the defaults return 95% confidence/credible intervals (0.95).

sampling.plan  Character describing the sampling plan. Possible options are "indepMulti" (independent multinomial; default), "poisson", "jointMulti" (joint multinomial), "hypergeom" (hypergeometric). For more, see ?BayesFactor::contingencyTableBF().

fixed.margin  For the independent multinomial sampling plan, which margin is fixed ("rows" or "cols"). Defaults to "rows".

prior.concentration  Specifies the prior concentration parameter, set to 1 by default. It indexes the expected deviation from the null hypothesis under the alternative, and corresponds to Gunel and Dickey’s (1974) "a" parameter.

...  Additional arguments (currently ignored).

Value

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- statistic: the numeric value of a statistic
- df: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- df.error and df: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- p.value: the two-sided p-value associated with the observed statistic
- method: the name of the inferential statistical test
- estimate: estimated value of the effect size
- conf.low: lower bound for the effect size estimate
- conf.high: upper bound for the effect size estimate
- conf.level: width of the confidence interval
- conf.method: method used to compute confidence interval
- conf.distribution: statistical distribution for the effect
- effectsize: the name of the effect size
- n.obs: number of observations
- expression: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by set.seed() calls for reproducibility.

Examples

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)
```
# # ------------------------ non-Bayesian -----------------------------

# association test
contingency_table(
    data = mtcars,
    x = am,
    y = cyl,
    paired = FALSE
)

# goodness-of-fit test
contingency_table(
    data = as.data.frame(HairEyeColor),
    x = Eye,
    counts = Freq,
    ratio = c(0.2, 0.2, 0.3, 0.3)
)

# ------------------------ Bayesian -----------------------------

# association test
contingency_table(
    data = mtcars,
    x = am,
    y = cyl,
    paired = FALSE,
    type = "bayes"
)

# goodness-of-fit test
contingency_table(
    data = as.data.frame(HairEyeColor),
    x = Eye,
    counts = Freq,
    ratio = c(0.2, 0.2, 0.3, 0.3),
    type = "bayes"
)

---

corr_test

**Correlation analyses**

**Description**

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

**Hypothesis testing** and **Effect size estimation**
<table>
<thead>
<tr>
<th>Type</th>
<th>Test</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>Pearson's correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>Spearman's rank correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Robust</td>
<td>Winsorized Pearson correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Bayesian Pearson's correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
</tbody>
</table>

Usage

```r
corr_test(
  data, x, y,
type = "parametric",
k = 2L, conf.level = 0.95,
tr = 0.2, bf.prior = 0.707,
...)
```

Arguments

data: A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from {dplyr} should be ungrouped before they are entered as data.

x: The column in data containing the explanatory variable to be plotted on the x-axis.

y: The column in data containing the response (outcome) variable to be plotted on the y-axis.

type: A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
  - "bayes"

You can specify just the initial letter.

k: Number of digits after decimal point (should be an integer) (Default: k = 2L).

conf.level: Scalar between 0 and 1. If unspecified, the defaults return 95% confidence/credible intervals (0.95).

tr: Trim level for the mean when carrying out robust tests. In case of an error, try reducing the value of tr, which is by default set to 0.2. Lowering the value might help.

bf.prior: A number between 0.5 and 2 (default 0.707), the prior width to use in calculating Bayes factors and posterior estimates. In addition to numeric arguments, several named values are also recognized: "medium", "wide", and "ultrawide".
corresponding to \( r \) scale values of 1/2, \( \sqrt{2}/2 \), and 1, respectively. In case of an ANOVA, this value corresponds to scale for fixed effects.

Additional arguments (currently ignored).

**Value**

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- **statistic**: the numeric value of a statistic
- **df**: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- **df.error** and **df**: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- **p.value**: the two-sided \( p \)-value associated with the observed statistic
- **method**: the name of the inferential statistical test
- **estimate**: estimated value of the effect size
- **conf.low**: lower bound for the effect size estimate
- **conf.high**: upper bound for the effect size estimate
- **conf.level**: width of the confidence interval
- **conf.method**: method used to compute confidence interval
- **conf.distribution**: statistical distribution for the effect
- **effectsize**: the name of the effect size
- **n.obs**: number of observations
- **expression**: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by `set.seed()` calls for reproducibility.

**Examples**

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)

# without changing defaults
corr_test(
  data = ggplot2::midwest,
  x = area,
  y = percblack
)

# changing defaults
corr_test(
  data = ggplot2::midwest,
```

```
iris_long

Edgar Anderson’s Iris Data in long format.

Description
Edgar Anderson’s Iris Data in long format.

Usage
iris_long

Format
A data frame with 600 rows and 5 variables

- id. Dummy identity number for each flower (150 flowers in total).
- Species. The species are Iris setosa, versicolor, and virginica.
- attribute. What attribute is being measured ("Sepal" or "Pepal").
- measure. What aspect of the attribute is being measured ("Length" or "Width").
- value. Value of the measurement.

Details
This famous (Fisher’s or Anderson’s) iris data set gives the measurements in centimeters of the variables sepal length and width and petal length and width, respectively, for 50 flowers from each of 3 species of iris. The species are Iris setosa, versicolor, and virginica.
This is a modified dataset from datasets package.

Examples
dim(iris_long)
head(iris_long)
dplyr::glimpse(iris_long)
long_to_wide_converter

Convert long/tidy data frame to wide format with NAs removed

Description

This conversion is helpful mostly for repeated measures design, where removing NAs by participant can be a bit tedious.

It does not make sense to spread the data frame to wide format when the measure is not repeated, so if paired = TRUE, spread argument will be ignored.

Usage

long_to_wide_converter(
  data,
  x,
  y,
  subject.id = NULL,
  paired = TRUE,
  spread = TRUE,
  ...
)

Arguments

data A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from \{dplyr\} should be ungrouped before they are entered as data.

x The grouping (or independent) variable from data. In case of a repeated measures or within-subjects design, if subject.id argument is not available or not explicitly specified, the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted, the results can be inaccurate when there are more than two levels in x and there are NAs present. The data is expected to be sorted by user in subject-1, subject-2, ..., pattern.

y The response (or outcome or dependent) variable from data.

subject.id Relevant in case of a repeated measures or within-subjects design (paired = TRUE, i.e.), it specifies the subject or repeated measures identifier. **Important:** Note that if this argument is NULL (which is the default), the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted and you leave this argument unspecified, the results can be inaccurate when there are more than two levels in x and there are NAs present.

paired Logical that decides whether the experimental design is repeated measures/within-subjects or between-subjects. The default is FALSE.
spread

Logical that decides whether the data frame needs to be converted from long/tidy to wide (default: TRUE). Relevant only if paired = TRUE.

... Currently ignored.

Value

A dataframe with NAs removed while respecting the between-or-within-subjects nature of the dataset.

Examples

# for reproducibility
library(statsExpressions)
set.seed(123)

# repeated measures design
long_to_wide_converter(
data = bugs_long,
x = condition,
y = desire,
subject.id = subject,
paired = TRUE
)

# independent measures design
long_to_wide_converter(
data = ggplot2::msleep,
x = vore,
y = brainwt,
paired = FALSE
)

---

meta_analysis

Random-effects meta-analyses

Description

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

Hypothesis testing and Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>Test</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>Pearson’s correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>Spearman’s rank correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Robust</td>
<td>Winsorized Pearson correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Bayesian Pearson’s correlation coefficient</td>
<td>Yes</td>
<td>correlation::correlation()</td>
</tr>
</tbody>
</table>
Usage

```r
meta_analysis(
  data,
  type = "parametric",
  random = "mixture",
  k = 2L,
  conf.level = 0.95,
  ...
)
```

Arguments

- **data**: A dataframe. It **must** contain columns named `estimate` (effect sizes or outcomes) and `std.error` (corresponding standard errors). These two columns will be used:
  - as `yi` and `sei` arguments in `metafor::rma` (for **parametric** test) or `metaplus::metaplus` (for **robust** test)
  - as `y` and `SE` arguments in `metaBMA::meta_random` (for **Bayesian** test).

- **type**: A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
  - "bayes"

You can specify just the initial letter.

- **random**: The type of random effects distribution. One of "normal", "t-dist", "mixture", for standard normal, t-distribution or mixture of normals respectively.

- **k**: Number of digits after decimal point (should be an integer) (Default: `k = 2L`).

- **conf.level**: Scalar between 0 and 1. If unspecified, the defaults return 95% confidence/credible intervals (0.95).

- **...**: Additional arguments passed to the respective meta-analysis function.

Value

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- **statistic**: the numeric value of a statistic
- **df**: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- **df.error** and **df**: relevant only if the statistic in question has two degrees of freedom (e.g. `anova`)
- **p.value**: the two-sided p-value associated with the observed statistic
- **method**: the name of the inferential statistical test
- **estimate**: estimated value of the effect size
- **conf.low**: lower bound for the effect size estimate
• `conf.high`: upper bound for the effect size estimate
• `conf.level`: width of the confidence interval
• `conf.method`: method used to compute confidence interval
• `conf.distribution`: statistical distribution for the effect
• `effectsize`: the name of the effect size
• `n.obs`: number of observations
• `expression`: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by `set.seed()` calls for reproducibility.

Note

Important: The function assumes that you have already downloaded the needed package (`metafor`, `{metaplus}`, or `{metaBMA}`) for meta-analysis. If they are not available, you will be asked to install them.

Examples

```r
# a dataframe with estimates and standard errors (`mag` dataset from `{metaplus}`)
df <- structure(list(
  estimate = c(-0.8303483, -1.056053, -1.27834, -0.0434851, 0.2231435, -2.40752, -1.280934, -1.191703, -0.695748, -2.208274, -2.03816, -0.8501509, -0.7932307, -0.2993399, -1.570789, 0.0575873),
  std.error = c(1.24701799987009, 0.41407060026039, 0.808139200261935, 1.42950999996502, 0.489168400451215, 1.07220799987689, 1.280934, -1.191703, -0.695748, -2.208274, -2.03816, -0.8501509, -0.7932307, -0.2993399, -1.570789, 0.0575873)),
  row.names = c(NA, -16L), class = "data.frame")

# setup
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)

meta_analysis(df) # parametric
# meta_analysis(df, type = "random", random = "normal") # robust
```
movies_long

# meta_analysis(df, type = "bayes") # Bayesian

movies_long  

<table>
<thead>
<tr>
<th>Description</th>
<th>Movie information and user ratings from IMDB.com (long format).</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Usage</th>
<th>movies_long</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Format</th>
<th>A data frame with 1,579 rows and 8 variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>• title</td>
<td>Title of the movie.</td>
</tr>
<tr>
<td>• year</td>
<td>Year of release.</td>
</tr>
<tr>
<td>• budget</td>
<td>Total budget (if known) in US dollars</td>
</tr>
<tr>
<td>• length</td>
<td>Length in minutes.</td>
</tr>
<tr>
<td>• rating</td>
<td>Average IMDB user rating.</td>
</tr>
<tr>
<td>• votes</td>
<td>Number of IMDB users who rated this movie.</td>
</tr>
<tr>
<td>• mpaa</td>
<td>MPAA rating.</td>
</tr>
<tr>
<td>• genre</td>
<td>Different genres of movies (action, animation, comedy, drama, documentary, romance, short).</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Details</th>
<th>Modified dataset from ggplot2movies package.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>The internet movie database, <a href="https://imdb.com/">https://imdb.com/</a>, is a website devoted to collecting movie data supplied by studios and fans. It claims to be the biggest movie database on the web and is run by amazon.</td>
</tr>
<tr>
<td></td>
<td>Movies were are identical to those selected for inclusion in movies_wide but this dataset has been constructed such that every movie appears in one and only one genre category.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Source</th>
<th><a href="https://CRAN.R-project.org/package=ggplot2movies">https://CRAN.R-project.org/package=ggplot2movies</a></th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Examples</th>
<th>dim(movies_long)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>head(movies_long)</td>
</tr>
<tr>
<td></td>
<td>dplyr::glimpse(movies_long)</td>
</tr>
</tbody>
</table>
movies_wide

Movie information and user ratings from IMDB.com (wide format).

Description

Movie information and user ratings from IMDB.com (wide format).

Usage

movies_wide

Format

A data frame with 1,579 rows and 13 variables

- title. Title of the movie.
- year. Year of release.
- budget. Total budget in millions of US dollars
- length. Length in minutes.
- rating. Average IMDB user rating.
- votes. Number of IMDB users who rated this movie.
- mpaa. MPAA rating.
- action, animation, comedy, drama, documentary, romance, short. Binary variables representing if movie was classified as belonging to that genre.
- NumGenre. The number of different genres a film was classified in an integer between one and four.

Details

Modified dataset from ggplot2movies package.

The internet movie database, https://imdb.com/, is a website devoted to collecting movie data supplied by studios and fans. It claims to be the biggest movie database on the web and is run by amazon.

Movies were selected for inclusion if they had a known length and had been rated by at least one imdb user. Small categories such as documentaries and NC-17 movies were removed.

Source

https://CRAN.R-project.org/package=ggplot2movies

Examples

dim(movies_wide)
head(movies_wide)
dplyr::glimpse(movies_wide)
## One-way analysis of variance (ANOVA)

### Description

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

### between-subjects:
#### Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>&gt; 2</td>
<td>Fisher’s or Welch’s one-way ANOVA</td>
<td>stats::oneway.test()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>&gt; 2</td>
<td>Kruskal-Wallis one-way ANOVA</td>
<td>stats::kruskal.test()</td>
</tr>
<tr>
<td>Robust</td>
<td>&gt; 2</td>
<td>Heteroscedastic one-way ANOVA for trimmed means</td>
<td>WRS2::t1way()</td>
</tr>
<tr>
<td>Bayes Factor</td>
<td>&gt; 2</td>
<td>Fisher’s ANOVA</td>
<td>BayesFactor::anovaBF()</td>
</tr>
</tbody>
</table>

#### Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>&gt; 2</td>
<td>partial eta-squared, partial omega-squared</td>
<td>Yes</td>
<td>effectsize::omega_squared()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>&gt; 2</td>
<td>rank epsilon squared</td>
<td>Yes</td>
<td>effectsize::rank_epsilon_squared()</td>
</tr>
<tr>
<td>Robust</td>
<td>&gt; 2</td>
<td>Explanatory measure of effect size</td>
<td>Yes</td>
<td>WRS2::t1way()</td>
</tr>
<tr>
<td>Bayes Factor</td>
<td>&gt; 2</td>
<td>Bayesian R-squared</td>
<td>Yes</td>
<td>performance::r2_bayes()</td>
</tr>
</tbody>
</table>

### within-subjects:
#### Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>&gt; 2</td>
<td>One-way repeated measures ANOVA</td>
<td>afex::aov_ez()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>&gt; 2</td>
<td>Friedman rank sum test</td>
<td>stats::friedman.test()</td>
</tr>
<tr>
<td>Robust</td>
<td>&gt; 2</td>
<td>Heteroscedastic one-way repeated measures ANOVA for trimmed means</td>
<td>WRS2::rmanova()</td>
</tr>
<tr>
<td>Bayes Factor</td>
<td>&gt; 2</td>
<td>One-way repeated measures ANOVA</td>
<td>BayesFactor::anovaBF()</td>
</tr>
</tbody>
</table>

#### Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>&gt; 2</td>
<td>partial eta-squared, partial omega-squared</td>
<td>Yes</td>
<td>effectsize::omega_squared()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>&gt; 2</td>
<td>Kendall’s coefficient of concordance</td>
<td>Yes</td>
<td>effectsize::kendalls_w()</td>
</tr>
<tr>
<td>Robust</td>
<td>&gt; 2</td>
<td>Algina-Keselman-Penfield robust standardized difference average</td>
<td>Yes</td>
<td>WRS2::wmcpAKP()</td>
</tr>
</tbody>
</table>
Usage

```r
oneway_anova(
  data,
  x,
  y,
  subject.id = NULL,
  type = "parametric",
  paired = FALSE,
  k = 2L,
  conf.level = 0.95,
  effsize.type = "omega",
  var.equal = FALSE,
  bf.prior = 0.707,
  tr = 0.2,
  nboot = 100L,
  ...
)
```

Arguments

data  A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from `{dplyr}` should be ungrouped before they are entered as data.

x     The grouping (or independent) variable from data. In case of a repeated measures or within-subjects design, if `subject.id` argument is not available or not explicitly specified, the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted, the results can be inaccurate when there are more than two levels in x and there are NAs present. The data is expected to be sorted by user in subject-1, subject-2, ..., pattern.

y     The response (or outcome or dependent) variable from data.

subject.id Relevant in case of a repeated measures or within-subjects design (paired = TRUE, i.e.), it specifies the subject or repeated measures identifier. Important: Note that if this argument is NULL (which is the default), the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted and you leave this argument unspecified, the results can be inaccurate when there are more than two levels in x and there are NAs present.

type A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
Value

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- **statistic**: the numeric value of a statistic
- **df**: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- **df.error** and **df**: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- **p.value**: the two-sided $p$-value associated with the observed statistic
- **method**: the name of the inferential statistical test
- **estimate**: estimated value of the effect size
- **conf.low**: lower bound for the effect size estimate
- **conf.high**: upper bound for the effect size estimate
- **conf.level**: width of the confidence interval
- **conf.method**: method used to compute confidence interval
- **conf.distribution**: statistical distribution for the effect
- **effectsize**: the name of the effect size
- `n.obs`: number of observations
- `expression`: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by `set.seed()` calls for reproducibility.

**Examples**

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)

# ----------------------- parametric -------------------------------------

# between-subjects
oneway_anova(
  data = ggplot2::msleep,
  x = vore,
  y = sleep_rem
)

if (require("afex", quietly = TRUE)) {
  # within-subjects design
  oneway_anova(
    data = iris_long,
    x = condition,
    y = value,
    subject.id = id,
    paired = TRUE
  )
}

# ----------------------- non-parametric ----------------------------------

# between-subjects
oneway_anova(
  data = ggplot2::msleep,
  x = vore,
  y = sleep_rem,
  type = "np"
)

# within-subjects design
oneway_anova(
  data = iris_long,
  x = condition,
  y = value,
  subject.id = id,
  paired = TRUE,
  type = "np"
)
```
one_sample_test

# ----------------------- robust -------------------------------------

# between-subjects
oneway_anova(
    data = ggplot2::msleep,
    x = vore,
    y = sleep_rem,
    type = "r"
)

# within-subjects design
oneway_anova(
    data = iris_long,
    x = condition,
    y = value,
    subject.id = id,
    paired = TRUE,
    type = "r"
)

# ----------------------- Bayesian -------------------------------------

# between-subjects
oneway_anova(
    data = ggplot2::msleep,
    x = vore,
    y = sleep_rem,
    type = "bayes"
)

# within-subjects design
oneway_anova(
    data = iris_long,
    x = condition,
    y = value,
    subject.id = id,
    paired = TRUE,
    type = "bayes"
)

---

one_sample_test

Description

The table below provides summary about:

- statistical test carried out for inferential statistics
• type of effect size estimate and a measure of uncertainty for this estimate
• functions used internally to compute these details

Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>One-sample Student's ( t )-test</td>
<td>stats::t.test</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>One-sample Wilcoxon test</td>
<td>stats::wilcox.test</td>
</tr>
<tr>
<td>Robust</td>
<td>Bootstrap-( t ) method for one-sample test</td>
<td>WRS2::trimcibt</td>
</tr>
<tr>
<td>Bayesian</td>
<td>One-sample Student's ( t )-test</td>
<td>BayesFactor::ttestBF</td>
</tr>
</tbody>
</table>

Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>Cohen's ( d ), Hedge's ( g )</td>
<td>Yes</td>
<td>effectsize::cohens.d(), effectsize::hedges.g()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>( r ) (rank-biserial correlation)</td>
<td>Yes</td>
<td>effectsize::rank_biserial()</td>
</tr>
<tr>
<td>Robust</td>
<td>trimmed mean</td>
<td>Yes</td>
<td>WRS2::trimcibt()</td>
</tr>
<tr>
<td>Bayes Factor</td>
<td>difference</td>
<td>Yes</td>
<td>bayestestR::describe_posterior()</td>
</tr>
</tbody>
</table>

Usage

```r
one_sample_test(
  data,    
  x,       
  type = "parametric",  
  test.value = 0,       
  alternative = "two.sided",  
  k = 2L,             
  conf.level = 0.95,   
  tr = 0.2,           
  bf.prior = 0.707,   
  effsize.type = "g",  
  ...                 
)
```

Arguments

data          A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from \{dplyr\} should be ungrouped before they are entered as data.

\( x \)          A numeric variable from the dataframe \( \text{data} \).

\( \text{type} \)  A character specifying the type of statistical approach:
• "parametric"
• "nonparametric"
one_sample_test

- "robust"
- "bayes"

You can specify just the initial letter.

test.value
A number indicating the true value of the mean (Default: \(0\)).
alternative
a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.
k
Number of digits after decimal point (should be an integer) (Default: \(k = 2\L\)).
conf.level
Scalar between \(0\) and \(1\). If unspecified, the defaults return 95\% confidence/credible intervals (0.95).
tr
Trim level for the mean when carrying out robust tests. In case of an error, try reducing the value of \(tr\), which is by default set to 0.2. Lowering the value might help.
bf.prior
A number between 0.5 and 2 (default 0.707), the prior width to use in calculating Bayes factors and posterior estimates. In addition to numeric arguments, several named values are also recognized: "medium", "wide", and "ultrawide", corresponding to \(r\) scale values of 1/2, \(\sqrt{2}/2\), and 1, respectively. In case of an ANOVA, this value corresponds to scale for fixed effects.
effsize.type
Type of effect size needed for parametric tests. The argument can be "d" (for Cohen’s \(d\)) or "g" (for Hedge’s \(g\)).

... Currently ignored.

**Value**

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- **statistic**: the numeric value of a statistic
- **df**: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- **df.error**: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- **p.value**: the two-sided \(p\)-value associated with the observed statistic
- **method**: the name of the inferential statistical test
- **estimate**: estimated value of the effect size
- **conf.low**: lower bound for the effect size estimate
- **conf.high**: upper bound for the effect size estimate
- **conf.level**: width of the confidence interval
- **conf.method**: method used to compute confidence interval
- **conf.distribution**: statistical distribution for the effect
- **effectsize**: the name of the effect size
- **n.obs**: number of observations
- **expression**: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by set.seed() calls for reproducibility.
Examples

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)

# ----------------------- parametric ---------------------------------------
one_sample_test(
  data = ggplot2::msleep,
  x = brainwt,
  test.value = 0.275,
  type = "parametric"
)

# ----------------------- non-parametric -----------------------------------
one_sample_test(
  data = ggplot2::msleep,
  x = brainwt,
  test.value = 0.275,
  type = "nonparametric"
)

# ----------------------- robust --------------------------------------------
one_sample_test(
  data = ggplot2::msleep,
  x = brainwt,
  test.value = 0.275,
  type = "robust"
)

# ---------------------------- Bayesian -----------------------------------
one_sample_test(
  data = ggplot2::msleep,
  x = brainwt,
  test.value = 0.275,
  type = "bayes",
  bf.prior = 0.8
)
```

pairwise_comparisons  Multiple pairwise comparison for one-way design
Description
Calculate parametric, non-parametric, robust, and Bayes Factor pairwise comparisons between group levels with corrections for multiple testing.

The table below provides summary about:

• statistical test carried out for inferential statistics
• type of effect size estimate and a measure of uncertainty for this estimate
• functions used internally to compute these details

between-subjects:

Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>Equal variance?</th>
<th>Test</th>
<th>p-value adjustment?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>No</td>
<td>Games-Howell test</td>
<td>Yes</td>
<td>PMCMRplus::gamesHowellTest()</td>
</tr>
<tr>
<td>Parametric</td>
<td>Yes</td>
<td>Student’s t-test</td>
<td>Yes</td>
<td>stats::pairwise.t.test()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>No</td>
<td>Dunn test</td>
<td>Yes</td>
<td>PMCMRplus::kwAllPairsDunnTest()</td>
</tr>
<tr>
<td>Robust</td>
<td>No</td>
<td>Yuen’s trimmed means test</td>
<td>Yes</td>
<td>WRS2::lincon()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>NA</td>
<td>Student’s t-test</td>
<td>NA</td>
<td>BayesFactor::ttestBF()</td>
</tr>
</tbody>
</table>

Effect size estimation
Not supported.

within-subjects:

Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>Test</th>
<th>p-value adjustment?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>Student’s t-test</td>
<td>Yes</td>
<td>stats::pairwise.t.test()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>Durbin-Conover test</td>
<td>Yes</td>
<td>PMCMRplus::durbinAllPairsTest()</td>
</tr>
<tr>
<td>Robust</td>
<td>Yuen’s trimmed means test</td>
<td>Yes</td>
<td>WRS2::rmmcp()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>Student’s t-test</td>
<td>NA</td>
<td>BayesFactor::ttestBF()</td>
</tr>
</tbody>
</table>

Effect size estimation
Not supported.

Usage

```r
pairwise_comparisons(
data, x, y, subject.id = NULL, type = "parametric", paired = FALSE, var.equal = FALSE, tr = 0.2,
```
pairwise_comparisons

```r
bf.prior = 0.707,
p.adjust.method = "holm",
k = 2L,
...
```

**Arguments**

- **data**
  A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will not be accepted. Additionally, grouped data frames from `{dplyr}` should be ungrouped before they are entered as `data`.

- **x**
  The grouping (or independent) variable from `data`. In case of a repeated measures or within-subjects design, if `subject_id` argument is not available or not explicitly specified, the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted, the results can be inaccurate when there are more than two levels in `x` and there are `NA`S present. The data is expected to be sorted by user in subject-1, subject-2, ..., pattern.

- **y**
  The response (or outcome or dependent) variable from `data`.

- **subject_id**
  Relevant in case of a repeated measures or within-subjects design (paired = TRUE, i.e.), it specifies the subject or repeated measures identifier. **Important**: Note that if this argument is NULL (which is the default), the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is not sorted and you leave this argument unspecified, the results can be inaccurate when there are more than two levels in `x` and there are `NA`S present.

- **type**
  A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
  - "bayes"
  You can specify just the initial letter.

- **paired**
  Logical that decides whether the experimental design is repeated measures/within-subjects or between-subjects. The default is FALSE.

- **var.equal**
  A logical variable indicating whether to treat the two variances as being equal. If TRUE then the pooled variance is used to estimate the variance otherwise the Welch (or Satterthwaite) approximation to the degrees of freedom is used.

- **tr**
  Trim level for the mean when carrying out robust tests. In case of an error, try reducing the value of `tr`, which is by default set to 0.2. Lowering the value might help.

- **bf.prior**
  A number between 0.5 and 2 (default 0.707), the prior width to use in calculating Bayes factors and posterior estimates. In addition to numeric arguments, several named values are also recognized: "medium", "wide", and "ultrawide", corresponding to r scale values of 1/2, sqrt(2)/2, and 1, respectively. In case of an ANOVA, this value corresponds to scale for fixed effects.
pairwise comparisons

p.adjust.method
Adjustment method for p-values for multiple comparisons. Possible methods are: "holm" (default), "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".

k
Number of digits after decimal point (should be an integer) (Default: k = 2L).

Additional arguments passed to other methods.

Value
The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- statistic: the numeric value of a statistic
- df: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- df.error and df: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- p.value: the two-sided p-value associated with the observed statistic
- method: the name of the inferential statistical test
- estimate: estimated value of the effect size
- conf.low: lower bound for the effect size estimate
- conf.high: upper bound for the effect size estimate
- conf.level: width of the confidence interval
- conf.method: method used to compute confidence interval
- conf.distribution: statistical distribution for the effect
- effectsize: the name of the effect size
- n.obs: number of observations
- expression: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.
Note that all examples are preceded by set.seed() calls for reproducibility.

References
For more, see: https://indrajeetpatil.github.io/ggstatsplot/articles/web_only/pairwise.html

Examples

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
library(PMCMRplus)

# show all columns and make the column titles bold
# as a user, you don't need to do this; this is just for the package website
```
pairwise_comparisons

```r
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE, pillar.subtle_num = TRUE)

#------------------- between-subjects design ----------------------------

# parametric
# if `var.equal = TRUE`, then Student's t-test will be run
pairwise_comparisons(
  data = mtcars,
  x = cyl,
  y = wt,
  type = "parametric",
  var.equal = TRUE,
  paired = FALSE,
  p.adjust.method = "none"
)

# if `var.equal = FALSE`, then Games-Howell test will be run
pairwise_comparisons(
  data = mtcars,
  x = cyl,
  y = wt,
  type = "parametric",
  var.equal = FALSE,
  paired = FALSE,
  p.adjust.method = "bonferroni"
)

# non-parametric (Dunn test)
pairwise_comparisons(
  data = mtcars,
  x = cyl,
  y = wt,
  type = "nonparametric",
  paired = FALSE,
  p.adjust.method = "none"
)

# robust (Yuen's trimmed means *t*-test)
pairwise_comparisons(
  data = mtcars,
  x = cyl,
  y = wt,
  type = "robust",
  paired = FALSE,
  p.adjust.method = "fdr"
)

# Bayes Factor (Student's *t*-test)
pairwise_comparisons(
  data = mtcars,
  x = cyl,
  y = wt,
  type = "bayes",
```

paired = FALSE

#------------------- within-subjects design ----------------------------

# parametric (Student's *t*-test)
pairwise_comparisons(
  data = bugs_long,
  x = condition,
  y = desire,
  subject.id = subject,
  type = "parametric",
  paired = TRUE,
  p.adjust.method = "BH"
)

# non-parametric (Durbin-Conover test)
pairwise_comparisons(
  data = bugs_long,
  x = condition,
  y = desire,
  subject.id = subject,
  type = "nonparametric",
  paired = TRUE,
  p.adjust.method = "BY"
)

# robust (Yuen's trimmed means t-test)
pairwise_comparisons(
  data = bugs_long,
  x = condition,
  y = desire,
  subject.id = subject,
  type = "robust",
  paired = TRUE,
  p.adjust.method = "hommel"
)

# Bayes Factor (Student's *t*-test)
pairwise_comparisons(
  data = bugs_long,
  x = condition,
  y = desire,
  subject.id = subject,
  type = "bayes",
  paired = TRUE
)
Description

Preparing text to describe which $p$-value adjustment method was used

Usage

```r
p_adjust_text(p.adjust.method)
```

Arguments

- `p.adjust.method`
  Adjustment method for $p$-values for multiple comparisons. Possible methods are: "holm" (default), "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".

Value

Standardized text description for what method was used.

Examples

```r
p_adjust_text("none")
p_adjust_text("BY")
```

---

### stats_type_switch

*Switch the type of statistics.*

Description

Relevant mostly for `{ggstatsplot}` and `{statsExpressions}` packages, where different statistical approaches are supported via this argument: parametric, non-parametric, robust, and Bayesian. This switch function converts strings entered by users to a common pattern for convenience.

Usage

```r
stats_type_switch(type)
```

Arguments

- `type`  
  A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
  - "bayes"

You can specify just the initial letter.
tidy_model_expressions

Expressions with statistics for tidy regression data frames

Description

Expressions with statistics for tidy regression data frames

Usage

tidy_model_expressions(
  data,
  statistic = NULL,
  k = 2L,
  effsize.type = "omega",
  ...
)

Arguments

data A tidy data frame from regression model object (see statsExpressions::tidy_model_parameters()).
statistic Which statistic is to be displayed (either "t" or "f" or "z" or "chi") in the expression.
k Number of digits after decimal point (should be an integer) (Default: k = 2L).
effsize.type Type of effect size needed for parametric tests. The argument can be "eta" (partial eta-squared) or "omega" (partial omega-squared).
... Currently ignored.

Details

When any of the necessary numeric column values (estimate, statistic, p.value) are missing, for these rows, a NULL is returned instead of an expression with empty strings.

Note

This is an experimental function and may change in the future. Please do not use it yet in your workflow.

Examples

    stats_type_switch("p")
    stats_type_switch("bf")
Examples

```r
set.seed(123)

# tidy dataframe
df <- tidy_model_parameters(lm(wt ~ am * cyl, mtcars))

# create a column containing expressions
tidy_model_expressions(df, statistic = "t")
```

tidy_model_parameters  Convert parameters package output to tidyverse conventions

Description

Convert parameters package output to tidyverse conventions

Usage

```r
tidy_model_parameters(model, ...)
```

Arguments

- `model` Statistical Model.
- `...` Arguments passed to or from other methods. Non-documented arguments are digits, p_digits, ci_digits and footer_digits to set the number of digits for the output. group can also be passed to the print() method. See details in `print.parameters_model()` and 'Examples' in `model_parameters.default()`.

Examples

```r
model <- lm(mpg ~ wt + cyl, data = mtcars)
tidy_model_parameters(model)
```

two_sample_test  Two-sample tests

Description

The table below provides summary about:

- statistical test carried out for inferential statistics
- type of effect size estimate and a measure of uncertainty for this estimate
- functions used internally to compute these details

**between-subjects:**

Hypothesis testing
### two_sample_test

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>2</td>
<td>Student’s or Welch’s $t$-test</td>
<td>stats::t.test()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>2</td>
<td>Mann-Whitney $U$ test</td>
<td>stats::wilcox.test()</td>
</tr>
<tr>
<td>Robust</td>
<td>2</td>
<td>Yuen’s test for trimmed means</td>
<td>WRS2::yuen()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>2</td>
<td>Student’s $t$-test</td>
<td>BayesFactor::ttestBF()</td>
</tr>
</tbody>
</table>

#### Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>2</td>
<td>Cohen’s $d$, Hedge’s $g$</td>
<td>Yes</td>
<td>effectsize::cohens_d(), effectsize::hedges_g()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>2</td>
<td>$r$ (rank-biserial correlation)</td>
<td>Yes</td>
<td>effectsize::rank_biserial()</td>
</tr>
<tr>
<td>Robust</td>
<td>2</td>
<td>Algina-Keselman-Penfield robust standardized difference</td>
<td>Yes</td>
<td>WRS2::akp.e</td>
</tr>
<tr>
<td>Bayesian</td>
<td>2</td>
<td>difference</td>
<td>Yes</td>
<td>bayestestR::</td>
</tr>
</tbody>
</table>

#### within-subjects:

#### Hypothesis testing

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Test</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>2</td>
<td>Student’s $t$-test</td>
<td>stats::t.test()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>2</td>
<td>Wilcoxon signed-rank test</td>
<td>stats::wilcox.test()</td>
</tr>
<tr>
<td>Robust</td>
<td>2</td>
<td>Yuen’s test on trimmed means for dependent samples</td>
<td>WRS2::yuend()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>2</td>
<td>Student’s $t$-test</td>
<td>BayesFactor::ttestBF()</td>
</tr>
</tbody>
</table>

#### Effect size estimation

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of groups</th>
<th>Effect size</th>
<th>CI available?</th>
<th>Function used</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parametric</td>
<td>2</td>
<td>Cohen’s $d$, Hedge’s $g$</td>
<td>Yes</td>
<td>effectsize::cohens_d(), effectsize::hedges_g()</td>
</tr>
<tr>
<td>Non-parametric</td>
<td>2</td>
<td>$r$ (rank-biserial correlation)</td>
<td>Yes</td>
<td>effectsize::rank_biserial()</td>
</tr>
<tr>
<td>Robust</td>
<td>2</td>
<td>Algina-Keselman-Penfield robust standardized difference</td>
<td>Yes</td>
<td>WRS2::wmcpAKP()</td>
</tr>
<tr>
<td>Bayesian</td>
<td>2</td>
<td>difference</td>
<td>Yes</td>
<td>bayestestR::</td>
</tr>
</tbody>
</table>

#### Usage

```r
two_sample_test(
data, x, y, subject.id = NULL, type = "parametric", paired = FALSE, alternative = "two.sided", k = 2L, conf.level = 0.95, effsize.type = "g",
```
two_sample_test

```r
var.equal = FALSE,
bf.prior = 0.707,
tr = 0.2,
nboot = 100L,
...
```

**Arguments**

- **data** A data frame (or a tibble) from which variables specified are to be taken. Other data types (e.g., matrix, table, array, etc.) will **not** be accepted. Additionally, grouped data frames from `{dplyr}` should be ungrouped before they are entered as data.

- **x** The grouping (or independent) variable from `data`. In case of a repeated measures or within-subjects design, if `subject.id` argument is not available or not explicitly specified, the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is **not** sorted, the results *can* be inaccurate when there are more than two levels in `x` and there are NAs present. The data is expected to be sorted by user in subject-1, subject-2, ..., pattern.

- **y** The response (or outcome or dependent) variable from `data`.

- **subject.id** Relevant in case of a repeated measures or within-subjects design (paired = TRUE, i.e.), it specifies the subject or repeated measures identifier. **Important:** Note that if this argument is NULL (which is the default), the function assumes that the data has already been sorted by such an id by the user and creates an internal identifier. So if your data is **not** sorted and you leave this argument unspecified, the results *can* be inaccurate when there are more than two levels in `x` and there are NAs present.

- **type** A character specifying the type of statistical approach:
  - "parametric"
  - "nonparametric"
  - "robust"
  - "bayes"

  You can specify just the initial letter.

- **paired** Logical that decides whether the experimental design is repeated measures/within-subjects or between-subjects. The default is FALSE.

- **alternative** a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter.

- **k** Number of digits after decimal point (should be an integer) (Default: k = 2L).

- **conf.level** Scalar between 0 and 1. If unspecified, the defaults return 95% confidence/credible intervals (0.95).

- **effsize.type** Type of effect size needed for parametric tests. The argument can be "d" (for Cohen’s d) or "g" (for Hedge’s g).

- **var.equal** a logical variable indicating whether to treat the two variances as being equal. If TRUE then the pooled variance is used to estimate the variance otherwise the Welch (or Satterthwaite) approximation to the degrees of freedom is used.
bf.prior A number between 0.5 and 2 (default 0.707), the prior width to use in calculating Bayes factors and posterior estimates. In addition to numeric arguments, several named values are also recognized: "medium", "wide", and "ultrawide", corresponding to $r$ scale values of 1/2, $\sqrt{2}$/2, and 1, respectively. In case of an ANOVA, this value corresponds to scale for fixed effects.

tr Trim level for the mean when carrying out robust tests. In case of an error, try reducing the value of tr, which is by default set to 0.2. Lowering the value might help.

nboot Number of bootstrap samples for computing confidence interval for the effect size (Default: 100L).

... Currently ignored.

Value

The returned tibble data frame can contain some or all of the following columns (the exact columns will depend on the statistical test):

- statistic: the numeric value of a statistic
- df: the numeric value of a parameter being modeled (often degrees of freedom for the test)
- df.error and df: relevant only if the statistic in question has two degrees of freedom (e.g. anova)
- p.value: the two-sided $p$-value associated with the observed statistic
- method: the name of the inferential statistical test
- estimate: estimated value of the effect size
- conf.low: lower bound for the effect size estimate
- conf.high: upper bound for the effect size estimate
- conf.level: width of the confidence interval
- conf.method: method used to compute confidence interval
- conf.distribution: statistical distribution for the effect
- effectsize: the name of the effect size
- n.obs: number of observations
- expression: pre-formatted expression containing statistical details

For examples of dataframe outputs, see examples and this vignette.

Note that all examples are preceded by set.seed() calls for reproducibility.

Examples

```r
# for reproducibility
set.seed(123)
library(statsExpressions)
options(tibble.width = Inf, pillar.bold = TRUE, pillar.neg = TRUE)

# parametric -------------------------------------
# between-subjects design
two_sample_test(
    data = sleep,
    x = group,
    y = extra,
    type = "p"
)

# within-subjects design
two_sample_test(
    data = dplyr::filter(bugs_long, condition %in% c("HDHF", "HDLF")),
    x = condition,
    y = desire,
    paired = TRUE,
    subject.id = subject,
    type = "p"
)

# non-parametric ----------------------------------
# between-subjects design
two_sample_test(
    data = sleep,
    x = group,
    y = extra,
    type = "np"
)

# within-subjects design
two_sample_test(
    data = dplyr::filter(bugs_long, condition %in% c("HDHF", "HDLF")),
    x = condition,
    y = desire,
    paired = TRUE,
    subject.id = subject,
    type = "np"
)

# robust ----------------------------------
# between-subjects design
two_sample_test(
    data = sleep,
    x = group,
    y = extra,
    type = "r"
)

# within-subjects design
two_sample_test(
    data = dplyr::filter(bugs_long, condition %in% c("HDHF", "HDLF")),
    x = condition,
```r
y = desire,
paired = TRUE,
subject.id = subject,
type = "r"
)

#' # Bayesian ------------------------------

# between-subjects design
two_sample_test(
  data = sleep,
  x = group,
  y = extra,
  type = "bayes"
)

# within-subjects design
two_sample_test(
  data = dplyr::filter(bugs_long, condition %in% c("HDHF", "HDLF")),
  x = condition,
  y = desire,
  paired = TRUE,
  subject.id = subject,
  type = "bayes"
)
```
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