Package ‘rempsyc’

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Title Convenience Functions for Psychology

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Description Make your workflow faster and easier. Easily customizable plots (via ‘ggplot2’), nice APA tables (following the style of the *American Psychological Association*) exportable to Word (via ‘flextable’), easily run statistical tests or check assumptions, and automatize various other tasks.

License GPL (>= 3)

URL https://rempsyc.remi-theriault.com

BugReports https://github.com/rempsyc/rempsyc/issues

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best_duplicate \hspace{1cm}  Choose the best duplicate

\textbf{Description}

Chooses the best duplicate, based on the duplicate with the smallest number of missing values. In case of ties, it picks the first duplicate, as it is the one most likely to be valid and authentic, given practice effects.

\textbf{Usage}

\begin{verbatim}
best_duplicate(data, id, keep.rows = FALSE)
\end{verbatim}
**Arguments**

- `data`: The data frame.
- `id`: The ID variable for which to check for duplicates.
- `keep.rows`: Logical, whether to add a column at the beginning of the data frame with the original row indices.

**Details**

For the `easystats` equivalent, see: `datawizard::data_duplicated()`.

**Value**

A dataframe, containing only the "best" duplicates.

**Examples**

```r
df1 <- data.frame(
id = c(1, 2, 3, 1, 3),
item1 = c(NA, 1, 1, 2, 3),
item2 = c(NA, 1, 1, 2, 3),
item3 = c(NA, 1, 1, 2, 3)
)
best_duplicate(df1, id = "id", keep.rows = TRUE)
```

---

**cormatrix_excel**

**Easy export of correlation matrix to Excel**

**Description**

Easily output a correlation matrix and export it to Microsoft Excel, with the first row and column frozen, and correlation coefficients colour-coded based on effect size (0.0-0.2: small (no colour); 0.2-0.4: medium (pink/light blue); 0.4-1.0: large (red/dark blue)), following Cohen’s suggestions for small (.10), medium (.30), and large (.50) correlation sizes.

Based on the `correlation` and `openxlsx2` packages.

**Usage**

```r
cormatrix_excel(
data,
filename,
overwrite = TRUE,
p_adjust = "none",
print.mat = TRUE,
...)
```

**extract_duplicates**

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>The data frame</td>
</tr>
<tr>
<td>filename</td>
<td>Desired filename (path can be added beforehand but no need to specify extension).</td>
</tr>
<tr>
<td>overwrite</td>
<td>Whether to allow overwriting previous file.</td>
</tr>
<tr>
<td>p_adjust</td>
<td>Default p-value adjustment method (default is &quot;none&quot;, although <code>correlation::correlation()</code>'s default is &quot;holm&quot;)</td>
</tr>
<tr>
<td>print.mat</td>
<td>Logical, whether to also print the correlation matrix to console.</td>
</tr>
<tr>
<td>...</td>
<td>Parameters to be passed to the <code>correlation</code> package (see <code>correlation::correlation()</code>).</td>
</tr>
</tbody>
</table>

**Value**

A Microsoft Excel document, containing the colour-coded correlation matrix with significance stars, on the first sheet, and the colour-coded p-values on the second sheet.

**Author(s)**

Adapted from @JanMarvin (JanMarvin/openxlsx2#286) and the original `rempsys::cormatrix_excel`.

**Examples**

```
# Basic example

cormatrix_excel(mtcars, select = c("mpg", "cyl", "disp", "hp", "carb"), filename = "cormatrix1")
cormatrix_excel(iris, p_adjust = "none", filename = "cormatrix2")
cormatrix_excel(airquality, method = "spearman", filename = "cormatrix3")
```

---

**extract_duplicates**  Extract all duplicates

**Description**

Extract all duplicates, for visual inspection. Note that it also contains the first occurrence of future duplicates, unlike `duplicated()` or `dplyr::distinct()`. Also contains an additional column reporting the number of missing values for that row, to help in the decision-making when selecting which duplicates to keep.

**Usage**

```
extract_duplicates(data, id)
```
find_mad

Arguments

- **data**: The data frame.
- **id**: The ID variable for which to check for duplicates.

Details

For the easystats equivalent, see: `datawizard::data_unique()`.

Value

A dataframe, containing all duplicates.

Examples

```r
df1 <- data.frame(
  id = c(1, 2, 3, 1, 3),
  item1 = c(NA, 1, 1, 2, 3),
  item2 = c(NA, 1, 1, 2, 3),
  item3 = c(NA, 1, 1, 2, 3)
)

extract_duplicates(df1, id = "id")
# Filter to exclude duplicates
df2 <- df1[-c(1, 5), ]
df2
```

---

**find_mad**

*Identify outliers based on 3 MAD*

Description

Identify outliers based on 3 median absolute deviations (MAD) from the median.

Usage

```r
find_mad(data, col.list, ID = NULL, criteria = 3, mad.scores = TRUE)
```

Arguments

- **data**: The data frame.
- **col.list**: List of variables to check for outliers.
- **ID**: ID variable if you would like the outliers to be identified as such.
- **criteria**: How many MAD to use as threshold (similar to standard deviations)
- **mad.scores**: Logical, whether to output robust z (MAD) scores (default) or raw scores. Defaults to TRUE.
Details

The function internally use `scale_mad()` to "standardize" the data based on the MAD and median, and then check for any observation greater than the specified criteria (e.g., +/-3).

For the easystats equivalent, use: `performance::check_outliers(x, method = "zscore_robust, threshold = 3`).

Value

A list of dataframes of outliers per variable, with row numbers, based on the MAD. When printed, provides the number of outliers, selected variables, and any outlier flagged for more than one variable. More information can be obtained by using the `attributes()` function around the generated object.

References


Examples

```r
find_mad(
  data = mtcars,
  col.list = names(mtcars),
  criteria = 3
)

mtcars2 <- mtcars
mtcars2$car <- row.names(mtcars)
find_mad(
  data = mtcars2,
  col.list = names(mtcars),
  ID = "car",
  criteria = 3
)
```

format_value

<table>
<thead>
<tr>
<th>format_value</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Easily format p or r values</em></td>
</tr>
</tbody>
</table>

Description

Easily format p or r values. Note: converts to character class for use in figures or manuscripts to accommodate e.g., "< .001".
**format_value**

**Usage**

```r
format_value(value, type = "d", ...)

format_p(
  p,
  precision = 0.001,
  prefix = NULL,
  suffix = NULL,
  sign = FALSE,
  stars = FALSE
)

format_r(r, precision = 0.01)

format_d(d, precision = 0.01)
```

**Arguments**

- `value`: Value to be formatted, when using the generic `format_value()`.
- `type`: Specify r or p value.
- `...`: To specify precision level, if necessary, when using the generic `format_value()`.
  Simply add the precision argument.
- `p`: p value to format.
- `precision`: Level of precision desired, if necessary.
- `prefix`: To add a prefix before the value.
- `suffix`: To add a suffix after the value.
- `sign`: Logical. Whether to add an equal sign for p values higher or equal to .001.
- `stars`: Logical. Whether to add asterisks for significant p values.
- `r`: r value to format.
- `d`: d value to format.

**Details**

For the `easystats` equivalent, see: `insight::format_value()`.

**Value**

A formatted p, r, or d value.

**Examples**

```r
format_value(0.00041231, "p")
format_value(0.00041231, "r")
format_value(1.341231, "d")
format_p(0.0041231)
format_p(0.00041231)
```
install_if_not_installed

Description

Install package if not already installed

Usage

install_if_not_installed(pkgs)

Arguments

pkgs Packages to install if not already installed

get_dep_version

Get required version of specified package dependency

Description

Get required version of specified package dependency

Usage

get_dep_version(dep, pkg = utils::packageName())

Arguments

dep Dependency of the specified package to check
pkg Package to check the dependency from
Description

Test linear regression assumptions easily with a nice summary table.

Usage

nice_assumptions(model)

Arguments

model

The `lm()` object to be passed to the function.

Details

Interpretation: (p) values < .05 imply assumptions are not respected. Diagnostic is how many assumptions are not respected for a given model or variable.

Value

A dataframe, with p-value results for the Shapiro-Wilk, Breusch-Pagan, and Durbin-Watson tests, as well as a diagnostic column reporting how many assumptions are not respected for a given model. Shapiro-Wilk is set to NA if n < 3 or n > 5000.

See Also

Other functions useful in assumption testing: `nice_density`, `nice_normality`, `nice_qq`, `nice_varplot`, `nice_var`. Tutorial: [https://rempsyc.remi-theriault.com/articles/assumptions](https://rempsyc.remi-theriault.com/articles/assumptions)

Examples

```r
# Create a regression model (using data available in R by default)
model <- lm(mpg ~ wt * cyl + gear, data = mtcars)
nice_assumptions(model)

# Multiple dependent variables at once
model2 <- lm(qsec ~ disp + drat * carb, mtcars)
my.models <- list(model, model2)
nice_assumptions(my.models)
```
nice_contrasts

*Easy planned contrasts*

**Description**

Easily compute planned contrast analyses (pairwise comparisons similar to t-tests but more powerful when more than 2 groups), and format in publication-ready format. In this particular case, the confidence intervals are bootstrapped on chosen effect size (default to Cohen’s d).

**Usage**

```r
nice_contrasts(
  response,  
  group,     
  covariates = NULL, 
  data,      
  effect.type = "cohens.d", 
  bootstraps = 2000, 
  ...       
)
```

**Arguments**

- `response` The dependent variable.
- `group` The group for the comparison.
- `covariates` The desired covariates in the model.
- `data` The data frame.
- `effect.type` What effect size type to use. One of "cohens.d" (default), "akp.robust.d", "unstandardized", "hedges.g", "cohens.d.sigma", or "r".
- `bootstraps` The number of bootstraps to use for the confidence interval
- `...` Arguments passed to bootES::bootES.

**Details**

Statistical power is lower with the standard \( t \) test compared than it is with the planned contrast version for two reasons: a) the sample size is smaller with the \( t \) test, because only the cases in the two groups are selected; and b) in the planned contrast the error term is smaller than it is with the standard \( t \) test because it is based on all the cases (source).

The effect size and confidence interval are calculated via bootES::bootES, and correct for contrasts but not for covariates and other predictors. Because this method uses bootstrapping, it is recommended to set a seed before using for reproducibility reasons (e.g., sed.seet(100)).

Does not for the moment support nested comparisons for marginal means, only a comparison of all groups. For nested comparisons, please use emmeans::contrast() directly, or for the easystats equivalent, modelbased::estimate_contrasts().

When using `nice_lmer_contrasts()`, please use `as.factor()` outside the `lm()` formula, or it will lead to an error.
nice_contrasts

Value
A dataframe, with the selected dependent variable(s), comparisons of interest, degrees of freedom, t-values, p-values, Cohen’s d, and the lower and upper 95% confidence intervals of the effect size (i.e., dR).

See Also
nice_lm_contrasts, Tutorial: https://rempyc.remi-theriault.com/articles/contrasts

Examples

# Basic example
set.seed(100)
nice_contrasts(
  data = mtcars,
  response = "mpg",
  group = "cyl",
  bootstraps = 200
)

set.seed(100)
nice_contrasts(
  data = mtcars,
  response = "disp",
  group = "gear"
)

# Multiple dependent variables
set.seed(100)
nice_contrasts(
  data = mtcars,
  response = c("mpg", "disp", "hp"),
  group = "cyl"
)

# Adding covariates
set.seed(100)
nice_contrasts(
  data = mtcars,
  response = "mpg",
  group = "cyl",
  covariates = c("disp", "hp")
)

# Now supports more than 3 levels
mtcars2 <- mtcars
mtcars2$carb <- as.factor(mtcars2$carb)
set.seed(100)
nice_contrasts(
  data = mtcars,
  response = "mpg",
  group = "carb",
  bootstraps = 200
)
nice_density

Easy density plots

Description

Make nice density plots easily. Internally, uses na.rm = TRUE.

Usage

nice_density(
  data,
  variable,
  group = NULL,
  colours,
  ytitle = "Density",
  xtitle = variable,
  groups.labels = NULL,
  grid = TRUE,
  shapiro = FALSE,
  title = variable,
  histogram = FALSE,
  breaks.auto = FALSE,
  bins = 30
)

Arguments

data The data frame
variable The dependent variable to be plotted.
group The group by which to plot the variable.
colours Desired colours for the plot, if desired.
ytitle An optional y-axis label, if desired.
xtitle An optional x-axis label, if desired.
groups.labels The groups.labels (might rename to xlabels for consistency with other functions)
grid Logical, whether to keep the default background grid or not. APA style suggests not using a grid in the background, though in this case some may find it useful to more easily estimate the slopes of the different groups.
shapiro Logical, whether to include the p-value from the Shapiro-Wilk test on the plot.
nice_density

title The desired title of the plot. Can be put to NULL to remove.
histogram Logical, whether to add an histogram
breaks.auto If histogram = TRUE, then option to set bins/breaks automatically, mimicking the default behaviour of base R hist() (the Sturges method). Defaults to FALSE.
bins If histogram = TRUE, then option to change the default bin (30).

Value

A density plot of class ggplot, by group (if provided), along a reference line representing a matched normal distribution.

See Also

Other functions useful in assumption testing: nice_assumptions, nice_normality, nice_qq, nice_varplot, nice_var. Tutorial: https://rempsyc.remi-theriault.com/articles/assumptions

Examples

# Make the basic plot
nice_density(
  data = iris,
  variable = "Sepal.Length",
  group = "Species"
)

# Further customization
nice_density(
  data = iris,
  variable = "Sepal.Length",
  group = "Species",
  colours = c("#00BA38", "#619CFF", "#F8766D"),
  xtitle = "Sepal Length",
  ytitle = "Density (vs. Normal Distribution)",
  groups.labels = c(
    "(a) Setosa",
    "(b) Versicolor",
    "(c) Virginica"
  ),
  grid = FALSE,
  shapiro = TRUE,
  title = "Density (Sepal Length)",
  histogram = TRUE
)
nice_lm

Nice formatting of lm models

Description

Formats output of `lm()` model object for a publication-ready format.

Usage

```r
nice_lm(
  model,
  b.label = "b",
  standardize = FALSE,
  mod.id = TRUE,
  ci.alternative = "two.sided",
  ...
)
```

Arguments

- **model**: The model to be formatted.
- **b.label**: What to rename the default "b" column (e.g., to capital B if using standardized data for it to be converted to the Greek beta symbol in the `nice_table` function). Now attempts to automatically detect whether the variables were standardized, and if so, sets `b.label = "B"` automatically. Factor variables or dummy variables (only two numeric values) are ignored when checking for standardization. **This argument is now deprecated, please use argument `standardize` directly instead.**
- **standardize**: Logical, whether to standardize the data before refitting the model. If `TRUE`, automatically sets `b.label = "B"`. Defaults to `FALSE`. Note that if you have factor variables, these will be pseudo-betas, so these coefficients could be interpreted more like Cohen’s $d$.
- **mod.id**: Logical. Whether to display the model number, when there is more than one model.
- **ci.alternative**: Alternative for the confidence interval of the $sr^2$. It can be either "two.sided" (the default in this package), "greater", or "less".
- **...**: Further arguments to be passed to the `effectsize::r2_semipartial` function for the effect size.

Details

The effect size, $sr^2$ (semi-partial correlation squared, also known as delta R2), is computed through `effectsize::r2_semipartial`. Please read the documentation for that function, especially regarding the interpretation of the confidence interval. In `rempsyc`, instead of using the default one-sided alternative ("greater"), we use the two-sided alternative.
To interpret the sr2, use `effectsize::interpret_r2_semipartial()`. For the easystats equivalent, use `report::report()` on the `lm()` model object.

**Value**

A formatted dataframe of the specified lm model, with DV, IV, degrees of freedom, regression coefficient, t-value, p-value, and the effect size, the semi-partial correlation squared, and its confidence interval.

**See Also**


**Examples**

```r
# Make and format model
model <- lm(mpg ~ cyl + wt * hp, mtcars)
nice_lm(model)

# Make and format multiple models
model2 <- lm(qsec ~ disp + drat * carb, mtcars)
my.models <- list(model, model2)
x <- nice_lm(my.models)
x

# Get interpretations
cbind(x, Interpretation = effectsize::interpret_r2_semipartial(x$sr2))
```

---

**nice_lm_contrasts**  
*Easy planned contrasts using lm models*

**Description**

Easily compute planned contrast analyses (pairwise comparisons similar to t-tests but more powerful when more than 2 groups), and format in publication-ready format. In this particular case, the confidence intervals are bootstraped on chosen effect size (default to Cohen’s d).

**Usage**

```r
nice_lm_contrasts(
  model,  
  group,  
  data,  
  p_adjust = "none",  
)  
```
effect.type = "cohens.d",
bootstraps = 2000,
...
)

Arguments

model The model to be formatted.
group The group for the comparison.
data The data frame.
p_adjust Character: adjustment method (e.g., "bonferroni") – added to options
effect.type What effect size type to use. One of "cohens.d" (default), "akp.robust.d", "unstandardized", "hedges.g", "cohens.d.sigma", or "r".
bootstraps The number of bootstraps to use for the confidence interval
...
Arguments passed to bootES::bootES.

Details

Statistical power is lower with the standard t test compared than it is with the planned contrast version for two reasons: a) the sample size is smaller with the t test, because only the cases in the two groups are selected; and b) in the planned contrast the error term is smaller than it is with the standard t test because it is based on all the cases (source).

The effect size and confidence interval are calculated via bootES::bootES, and correct for contrasts but not for covariates and other predictors. Because this method uses bootstrapping, it is recommended to set a seed before using for reproducibility reasons (e.g., sed.set(100)).

Does not for the moment support nested comparisons for marginal means, only a comparison of all groups. For nested comparisons, please use emmeans::contrast() directly, or for the easystats equivalent, modelbased::estimate_contrasts().

When using nice_lm_contrasts(), please use as.factor() outside the lm() formula, or it will lead to an error.

Value

A dataframe, with the selected dependent variable(s), comparisons of interest, degrees of freedom, t-values, p-values, Cohen’s d, and the lower and upper 95% confidence intervals of the effect size (i.e., dR).

See Also

nice_contrasts, Tutorial: https://rempyc.remi-theriault.com/articles/contrasts

Examples

# Make and format model (group need to be a factor)
mtcars2 <- mtcars
mtcars2$cyl <- as.factor(mtcars2$cyl)
model <- lm(mpg ~ cyl + wt * hp, mtcars2)
set.seed(100)
nice_lm_contrasts(model, group = "cyl", data = mtcars, bootstraps = 500)

# Several models at once
mtcars2$gear <- as.factor(mtcars2$gear)
model2 <- lm(qsec ~ cyl, data = mtcars2)
my.models <- list(model, model2)
set.seed(100)
nice_lm_contrasts(my.models, group = "cyl", data = mtcars, bootstraps = 500)

# Now supports more than 3 levels
mtcars2$carb <- as.factor(mtcars2$carb)
model <- lm(mpg ~ carb + wt * hp, mtcars2)
set.seed(100)
nice_lm_contrasts(model, group = "carb", data = mtcars2, bootstraps = 500)

---

**nice_lm_slopes**

Nice formatting of simple slopes for lm models

**Description**

Extracts simple slopes from lm() model object and format for a publication-ready format.

**Usage**

```r
nice_lm_slopes(
  model,
  predictor,
  moderator,
  b.label = "b",
  standardize = FALSE,
  mod.id = TRUE,
  ci.alternative = "two.sided",
  ...
)
```

**Arguments**

- `model`: The model to be formatted.
- `predictor`: The independent variable.
- `moderator`: The moderating variable.
- `b.label`: What to rename the default "b" column (e.g., to capital B if using standardized data for it to be converted to the Greek beta symbol in the nice_table() function). Now attempts to automatically detect whether the variables were standardized, and if so, sets `b.label = "B"` automatically. Factor variables or dummy
variables (only two numeric values) are ignored when checking for standard-
ization. This argument is now deprecated, please use argument standardize
directly instead.

standardize Logical, whether to standardize the data before refitting the model. If TRUE, au-
tomatically sets b.label = "B". Defaults to FALSE. Note that if you have factor
variables, these will be pseudo-betas, so these coefficients could be interpreted
more like Cohen’s $d$.

mod.id Logical. Whether to display the model number, when there is more than one
model.

ci.alternative Alternative for the confidence interval of the sr2. It can be either "two.sided (the
default in this package), "greater", or "less".

... Further arguments to be passed to the lm() function for the models.

Details
The effect size, sr2 (semi-partial correlation squared, also known as delta R2), is computed through
effects::r2_semiPartial. Please read the documentation for that function, especially regarding
the interpretation of the confidence interval. In rempsyc, instead of using the default one-sided
alternative ("greater"), we use the two-sided alternative.
To interpret the sr2, use effects::interpret_r2_semiPartial().
For the easystats equivalent, use report::report() on the lm() model object.

Value
A formatted dataframe of the simple slopes of the specified lm model, with DV, levels of IV, degrees
of freedom, regression coefficient, t-value, p-value, and the effect size, the semi-partial correlation
squared, and its confidence interval.

See Also
Checking for moderation before checking simple slopes: nice_lm, nice_mod, nice_slopes. Tu-
torial: https://rempsyc.remi-theriault.com/articles/moderation

Examples

# Make and format model
model <- lm(mpg ~ gear * wt, mtcars)
nice_lm_slopes(model, predictor = "gear", moderator = "wt")

# Make and format multiple models
model2 <- lm(qsec ~ gear * wt, mtcars)
my.models <- list(model, model2)
x <- nice_lm_slopes(my.models, predictor = "gear", moderator = "wt")
x

# Get interpretations
cbind(x, Interpretation = effects::interpret_r2_semiPartial(x$sr2))
### nice_mod

**Easy moderations**

**Description**

Easily compute moderation analyses, with effect sizes, and format in publication-ready format.

**Usage**

```r
nice_mod(
  data,
  response,
  predictor,
  moderator,
  moderator2 = NULL,
  covariates = NULL,
  b.label = "b",
  standardize = TRUE,
  mod.id = TRUE,
  ci.alternative = "two.sided",
  ...
)
```

**Arguments**

- **data** The data frame
- **response** The dependent variable.
- **predictor** The independent variable.
- **moderator** The moderating variable.
- **moderator2** The second moderating variable, if applicable.
- **covariates** The desired covariates in the model.
- **b.label** What to rename the default "b" column (e.g., to capital B if using standardized data for it to be converted to the Greek beta symbol in the `nice_table()` function). Now attempts to automatically detect whether the variables were standardized, and if so, sets `b.label = "B"` automatically. Factor variables or dummy variables (only two numeric values) are ignored when checking for standardization. *This argument is now deprecated, please use argument `standardize` directly instead.*
- **standardize** Logical, whether to standardize the data before fitting the model. If `TRUE`, automatically sets `b.label = "B"`. Defaults to `TRUE`.
- **mod.id** Logical. Whether to display the model number, when there is more than one model.
- **ci.alternative** Alternative for the confidence interval of the sr2. It can be either "two.sided" (the default in this package), "greater", or "less".
- **...** Further arguments to be passed to the `lm()` function for the models.
Details

The effect size, sr2 (semi-partial correlation squared, also known as delta R2), is computed through `effectsize::r2_semipartial`. Please read the documentation for that function, especially regarding the interpretation of the confidence interval. In rempsyrc, instead of using the default one-sided alternative ("greater"), we use the two-sided alternative.

To interpret the sr2, use `effectsize::interpret_r2_semipartial()`.

For the easystats equivalent, use `report::report()` on the `lm()` model object.

Value

A formatted dataframe of the specified lm model, with DV, IV, degrees of freedom, regression coefficient, t-value, p-value, and the effect size, the semi-partial correlation squared, and its confidence interval.

See Also

Checking simple slopes after testing for moderation: `nice_slopes, nice_lm, nice_lm_slopes`.

Tutorial: https://rempsyc.remi-theriault.com/articles/moderation

Examples

```r
# Make the basic table
nice_mod(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt"
)

# Multiple dependent variables at once
nice_mod(
  data = mtcars,
  response = c("mpg", "disp", "hp"),
  predictor = "gear",
  moderator = "wt"
)

# Add covariates
nice_mod(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt",
  covariates = c("am", "vs")
)

# Three-way interaction
x <- nice_mod(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt",
  covariates = c("am", "vs")
)
```
```r
response = "mpg",
predictor = "gear",
moderator = "wt",
moderator2 = "am"
)
x

# Get interpretations
cbind(x, Interpretation = effectsize::interpret_r2_semipartial(x$sr2))
```

---

**nice_na**  
*Report missing values according to guidelines*

**Description**

Nicely reports NA values according to existing guidelines. This function reports both absolute and percentage values of specified column lists. Some authors recommend reporting item-level missing item per scale, as well as participant’s maximum number of missing items by scale. For example, Parent (2013) writes:

*I recommend that authors (a) state their tolerance level for missing data by scale or subscale (e.g., “We calculated means for all subscales on which participants gave at least 75% complete data”) and then (b) report the individual missingness rates by scale per data point (i.e., the number of missing values out of all data points on that scale for all participants) and the maximum by participant (e.g., “For Attachment Anxiety, a total of 4 missing data points out of 100 were observed, with no participant missing more than a single data point”).*

**Usage**

```r
nice_na(data, vars = NULL, scales = NULL)
```

**Arguments**

- **data**  
  The data frame.
- **vars**  
  Variable (or lists of variables) to check for NAs.
- **scales**  
  The scale names to check for NAs (single character string).

**Value**

A dataframe, with:

- **var**: variables selected
- **items**: number of items for selected variables
- **na**: number of missing cell values for those variables (e.g., 2 missing values for first participant + 2 missing values for second participant = total of 4 missing values)
• cells: total number of cells (i.e., number of participants multiplied by number of variables, items)

• na_percent: the percentage of missing values (number of missing cells, na, divided by total number of cells, cells)

• na_max: The amount of missing values for the participant with the most missing values for the selected variables

• na_max_percent: The amount of missing values for the participant with the most missing values for the selected variables, in percentage (i.e., na_max divided by the number of selected variables, items)

• all_na: the number of participants missing 100% of items for that scale (the selected variables)

References


Examples

# Use whole data frame
nice_na(airquality)

# Use selected columns explicitly
nice_na(airquality, vars = list(c("Ozone", "Solar.R", "Wind"), c("Temp", "Month", "Day")))

# If the questionnaire items start with the same name, e.g.,
set.seed(15)
fun <- function() {
c(sample(c(NA, 1:10), replace = TRUE), NA, NA, NA)
}
df <- data.frame(
  ID = c("idz", NA),
  open_1 = fun(), open_2 = fun(), open_3 = fun(),
  extrovert_1 = fun(), extrovert_2 = fun(), extrovert_3 = fun(),
  agreeable_1 = fun(), agreeable_2 = fun(), agreeable_3 = fun()
)

# One can list the scale names directly:
nice_na(df, scales = c("ID", "open", "extrovert", "agreeable"))
nice_normality

Description

Easily make nice per-group density and QQ plots through a wrapper around the ggplot2 and qqplotr packages.

Usage

nice_normality(
  data,
  variable,
  group = NULL,
  colours,
  groups.labels,
  grid = TRUE,
  shapiro = FALSE,
  title = NULL,
  histogram = FALSE,
  breaks.auto = FALSE,
  ...
)

Arguments

data The data frame.
variable The dependent variable to be plotted.
group The group by which to plot the variable.
colours Desired colours for the plot, if desired.
groups.labels How to label the groups.
grid Logical, whether to keep the default background grid or not. APA style suggests not using a grid in the background, though in this case some may find it useful to more easily estimate the slopes of the different groups.
shapiro Logical, whether to include the p-value from the Shapiro-Wilk test on the plot.
title An optional title, if desired.
histogram Logical, whether to add an histogram on top of the density plot.
breaks.auto If histogram = TRUE, then option to set bins/breaks automatically, mimicking the default behaviour of base R hist() (the Sturges method). Defaults to FALSE.
...

Further arguments from nice_qq() and nice_density() to be passed to nice_normality()

Value

A plot of classes patchwork and ggplot, containing two plots, resulting from nice_density and nice_qq.
See Also

Other functions useful in assumption testing: \texttt{nice_assumptions}, \texttt{nice_density}, \texttt{nice_qq}, \texttt{nice_var}, \texttt{nice_varplot}. Tutorial: \url{https://rempsyc.remi-theriault.com/articles/assumptions}

Examples

# Make the basic plot
nice_normality(
  data = iris,
  variable = "Sepal.Length",
  group = "Species"
)

# Further customization
nice_normality(
  data = iris,
  variable = "Sepal.Length",
  group = "Species",
  colours = c("#00BA38",
              "#619CFF",
              "#F8766D"),
  groups.labels = c("(a) Setosa",
                    "(b) Versicolor",
                    "(c) Virginica"),
  grid = FALSE,
  shapiro = TRUE
)

nice_qq \hspace{1cm} \textit{Easy QQ plots per group}

Description

Easily make nice per-group QQ plots through a wrapper around the \texttt{ggplot2} and \texttt{qqplotr} packages.

Usage

nice_qq(
  data,
  variable,
  group = NULL,
  colours,
  groups.labels = NULL,
)
Arguments

data The data frame.
variable The dependent variable to be plotted.
group The group by which to plot the variable.
colours Desired colours for the plot, if desired.
groups.labels How to label the groups.
grid Logical, whether to keep the default background grid or not. APA style suggests not using a grid in the background, though in this case some may find it useful to more easily estimate the slopes of the different groups.
shapiro Logical, whether to include the p-value from the Shapiro-Wilk test on the plot.
title An optional title, if desired.

Value

A qq plot of class ggplot, by group (if provided), along a reference interpretation helper, the 95% confidence band.

See Also

Other functions useful in assumption testing: nice_assumptions, nice_density, nice_normality, nice_var, nice_varplot. Tutorial: https://rempsyrc.remi-theriault.com/articles/assumptions

Examples

# Make the basic plot
nice_qq(
  data = iris,
  variable = "Sepal.Length",
  group = "Species"
)

# Further customization
nice_qq(
  data = iris,
  variable = "Sepal.Length",
  group = "Species",
  colours = c("#00BA38", "#619CFF", "#F8766D"),
  groups.labels = c("(a) Setosa", "(b) Versicolor", "(c) Virginica"),
  grid = FALSE,
  shapiro = TRUE,
  title = NULL
)
nice_randomize

Easily randomization

Description
Randomize easily with different designs.

Usage
nice_randomize(
  design = "between",
  Ncondition = 3,
  n = 9,
  condition.names = c("a", "b", "c"),
  col.names = c("id", "Condition")
)

Arguments
- **design**
The design: either between-subject (different groups) or within-subject (repeated-measures on same people).
- **Ncondition**
The number of conditions you want to randomize.
- **n**
The desired sample size. Note that it needs to be a multiple of your number of groups if you are using between.
- **condition.names**
The names of the randomized conditions.
- **col.names**
The desired additional column names for a runsheet.

Value
A dataframe, with participant ID and randomized condition, based on selected design.

See Also
- Tutorial: [https://rempsyc.remi-theriault.com/articles/randomize](https://rempsyc.remi-theriault.com/articles/randomize)

Examples
```r
# Specify design, number of conditions, number of participants, and names of conditions:
nice_randomize(
  design = "between", Ncondition = 4, n = 8,
  condition.names = c("BP", "CX", "PZ", "ZL")
)

# Within-Group Design
nice_randomize(
```
nice_reverse

```r
design = "within", Ncondition = 4, n = 6,
condition.names = c("SV", "AV", "ST", "AT")
)

# Make a quick runsheet
randomized <- nice_randomize(
  design = "within", Ncondition = 4, n = 128,
  condition.names = c("SV", "AV", "ST", "AT"),
col.names = c(
  "id", "Condition", "Date/Time",
  "SONA ID", "Age/Gd.", "Handedness",
  "Tester", "Notes"
)
)
head(randomized)
```

---

## nice_reverse

### Easily recode scores

**Description**

Easily recode scores (reverse-score), typically for questionnaire answers.

For the `easystats` equivalent, see: `datawizard::reverse()`.

**Usage**

```r
nice_reverse(x, max, min = 1)
```

**Arguments**

- `x` The score to reverse.
- `max` The maximum score on the scale.
- `min` The minimum score on the scale (optional unless it isn’t 1).

**Value**

A numeric vector, of reversed scores.

**Examples**

# Reverse score of 5 with a maximum score of 5
nice_reverse(5, 5)

# Reverse several scores at once
nice_reverse(1:5, 5)

# Reverse scores with maximum = 4 and minimum = 0
nice_reverse(1:4, 4, min = 0)
# Reverse scores with maximum = 3 and minimum = -3
nice_reverse(-3:3, 3, min = -3)

## Description

Make nice scatter plots easily.

## Usage

```r
nice_scatter(
  data,
  predictor,
  response,
  xtitle = predictor,
  ytitle = response,
  has.points = TRUE,
  has.jitter = FALSE,
  alpha = 0.7,
  has.line = TRUE,
  method = "lm",
  has.confband = FALSE,
  has.fullrange = FALSE,
  has.linetype = FALSE,
  has.shape = FALSE,
  xmin,
  xmax,
  xby = 1,
  ymin,
  ymax,
  yby = 1,
  has.legend = FALSE,
  legend.title = "",
  group = NULL,
  colours = "#619CFF",
  groups.order = "none",
  groups.labels = NULL,
  groups.alpha = NULL,
  has.r = FALSE,
  r.x = Inf,
  r.y = -Inf,
  has.p = FALSE,
  p.x = Inf,
)```
nice_scatter

p.y = -Inf
)
}

Arguments

data The data frame.
predictor The independent variable to be plotted.
response The dependent variable to be plotted.
xtitle An optional y-axis label, if desired.
ytitle An optional x-axis label, if desired.
has.points Whether to plot the individual observations or not.
has.jitter Alternative to has.points. "Jitters" the observations to avoid overlap (overplot-
ting). Use one or the other, not both.
alpha The desired level of transparency.
has.line Whether to plot the regression line(s).
method Which method to use for the regression line, either "lm" (default) or "loess".
has.confband Logical. Whether to display the confidence band around the slope.
has.fullrange Logical. Whether to extend the slope beyond the range of observations.
has.linetype Logical. Whether to change line types as a function of group.
has.shape Logical. Whether to change shape of observations as a function of group.
xmin The minimum score on the x-axis scale.
xmax The maximum score on the x-axis scale.
xby How much to increase on each "tick" on the x-axis scale.
ymin The minimum score on the y-axis scale.
ymax The maximum score on the y-axis scale.
yby How much to increase on each "tick" on the y-axis scale.
has.legend Logical. Whether to display the legend or not.
legend.title The desired legend title.
group The group by which to plot the variable
colours Desired colours for the plot, if desired.
groups.order Specifies the desired display order of the groups on the legend. Either provide
the levels directly, or a string: "increasing" or "decreasing", to order based on
the average value of the variable on the y axis, or "string.length", to order from
the shortest to the longest string (useful when working with long string names).
"Defaults to "none".

groups.labels Changes groups names (labels). Note: This applies after changing order of level.
groups.alpha The manually specified transparency desired for the groups slopes. Use only
when plotting groups separately.
has.r Whether to display the correlation coefficient, the r-value.
r.x The x-axis coordinates for the r-value.
nice_scatter

- `r.y`: The y-axis coordinates for the r-value.
- `has.p`: Whether to display the p-value.
- `p.x`: The x-axis coordinates for the p-value.
- `p.y`: The y-axis coordinates for the p-value.

**Value**

A scatter plot of class ggplot.

**See Also**


**Examples**

```r
# Make the basic plot
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg"
)

# Save a high-resolution image file to specified directory
ggplot2::ggsave("nicescatterplotthere.pdf", width = 7,
  height = 7, unit = "in", dpi = 300
) # change for your own desired path

# Change x- and y- axis labels
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  ytitle = "Miles/(US) gallon",
  xtitle = "Weight (1000 lbs)"
)

# Have points "jittered", loess method
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  has.jitter = TRUE,
  method = "loess"
)

# Change the transparency of the points
nice_scatter(
  data = mtcars,
  has.jitter = TRUE,
  alpha = 0.5
)```

```r```
nice_scatter

predictor = "wt",
response = "mpg",
alpha = 1
)

# Remove points
nice_scatter(
data = mtcars,
predictor = "wt",
response = "mpg",
has.points = FALSE,
has.jitter = FALSE
)

# Add confidence band
nice_scatter(
data = mtcars,
predictor = "wt",
response = "mpg",
has.confband = TRUE
)

# Set x- and y- scales manually
nice_scatter(
data = mtcars,
predictor = "wt",
response = "mpg",
xmin = 1,
xmax = 6,
xby = 1,
ymin = 10,
ymax = 35,
yby = 5
)

# Change plot colour
nice_scatter(
data = mtcars,
predictor = "wt",
response = "mpg",
colours = "blueviolet"
)

# Add correlation coefficient to plot and p-value
nice_scatter(
data = mtcars,
predictor = "wt",
response = "mpg",
has.r = TRUE,
has.p = TRUE
)

# Change location of correlation coefficient or p-value
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  has.r = TRUE,
  r.x = 4,
  r.y = 25,
  has.p = TRUE,
  p.x = 5,
  p.y = 20
)

# Plot by group
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl"
)

# Use full range on the slope/confidence band
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  has.fullrange = TRUE
)

# Remove lines
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  has.line = FALSE
)

# Change order of labels on the legend
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  groups.order = c(8, 4, 6)
)

# Change legend labels
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  }

groups.labels = c("Weak", "Average", "Powerful")
)
# Warning: This applies after changing order of level

# Add a title to legend
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  legend.title = "cylinders"
)

# Plot by group + manually specify colours
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  colours = c("burlywood", "darkgoldenrod", "chocolate")
)

# Plot by group + use different line types for each group
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  has.linetype = TRUE
)

# Plot by group + use different point shapes for each group
nice_scatter(
  data = mtcars,
  predictor = "wt",
  response = "mpg",
  group = "cyl",
  has.shape = TRUE
)

nice_slopes: Easy simple slopes

Description

Easily compute simple slopes in moderation analysis, with effect sizes, and format in publication-ready format.
Usage
	nice_slopes(
	
data,  # The data frame
	response,  # The dependent variable.
	predictor,  # The independent variable
	moderator,  # The moderating variable.
	moderator2 = NULL,  # The second moderating variable, if applicable. At this time, the second moder-

covariates = NULL,  # The desired covariates in the model.

b.label = "b",  # What to rename the default "b" column (e.g., to capital B if using standardized
standardize = TRUE,  # Logical, whether to standardize the data before fitting the model. If TRUE,
mod.id = TRUE,  # Logical. Whether to display the model number, when there is more than one

ci.alternative = "two.sided",  # Alternative for the confidence interval of the sr2. It can be either "two.sided (the
...
)

Arguments

data The data frame
response The dependent variable.
predictor The independent variable
moderator The moderating variable.
moderator2 The second moderating variable, if applicable. At this time, the second moder-
covariates The desired covariates in the model.
b.label What to rename the default "b" column (e.g., to capital B if using standardized
data for it to be converted to the Greek beta symbol in the nice_table() func-
standardize Logical, whether to standardize the data before fitting the model. If TRUE,
mod.id Logical. Whether to display the model number, when there is more than one

ci.alternative Alternative for the confidence interval of the sr2. It can be either "two.sided (the
... Further arguments to be passed to the lm() function for the models.

Details

The effect size, sr2 (semi-partial correlation squared, also known as delta R2), is computed through
effectsize::r2_semipartial. Please read the documentation for that function, especially regarding
the interpretation of the confidence interval. In rempsyc, instead of using the default one-sided
alternative ("greater"), we use the two-sided alternative.
To interpret the \( r^2 \), use `effectsize::interpret_r2_semipartial()`.

For the `easystats` equivalent, use `report::report()` on the `lm()` model object.

**Value**

A formatted dataframe of the simple slopes of the specified `lm` model, with DV, levels of IV, degrees of freedom, regression coefficient, t-value, p-value, and the effect size, the semi-partial correlation squared, and its confidence interval.

**See Also**


**Examples**

```r
# Make the basic table
nice_slopes(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt"
)

# Multiple dependent variables at once
nice_slopes(
  data = mtcars,
  response = c("mpg", "disp", "hp"),
  predictor = "gear",
  moderator = "wt"
)

# Add covariates
nice_slopes(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt",
  covariates = c("am", "vs")
)

# Three-way interaction (continuous moderator and binary
# second moderator required)
x <- nice_slopes(
  data = mtcars,
  response = "mpg",
  predictor = "gear",
  moderator = "wt",
  moderator2 = "am"
)
x
```
# Get interpretations
cbind(x, Interpretation = effectsize::interpret_r2_semipartial(x$sr2))

## Description

Make nice APA tables easily through a wrapper around the `flextable` package with sensical defaults and automatic formatting features.

## Usage

```r
nice_table(
  data,
  highlight = FALSE,
  stars = TRUE,
  italics,
  col.format.p,
  col.format.r,
  col.format.ci,
  format.custom,
  col.format.custom,
  width = NULL,
  broom = NULL,
  report = NULL,
  short = FALSE,
  title,
  note,
  separate.header
)
```

## Arguments

- **data**: The data frame, to be converted to a `flextable`. The data frame cannot have duplicate column names.
- **highlight**: Highlight rows with statistically significant results? Requires a column named "p" containing p-values. Can either accept logical (TRUE/FALSE) OR a numeric value for a custom critical p-value threshold (e.g., 0.10 or 0.001).
- **stars**: Logical. Whether to add asterisks for significant p values.
- **italics**: Which columns headers should be italic? Useful for column names that should be italic but that are not picked up automatically by the function. Select with numerical range, e.g., 1:3.
nice_table

**col.format.p** Applies p-value formatting to columns that cannot be named "p" (for example for a data frame full of p-values, also because it is not possible to have more than one column named "p"). Select with numerical range, e.g., 1:3.

**col.format.r** Applies r-value formatting to columns that cannot be named "r" (for example for a data frame full of r-values, also because it is not possible to have more than one column named "r"). Select with numerical range, e.g., 1:3.

**col.format.ci** Applies 95% confidence interval formatting to selected columns (e.g., when reporting more than one interval).

**format.custom** Applies custom formatting to columns selected via the `col.format.custom` argument. This is useful if one wants custom formatting other than for p- or r-values. It can also be used to transform (e.g., multiply) certain values or print a specific symbol along the values for instance.

**col.format.custom** Which columns to apply the custom function to. Select with numerical range, e.g., 1:3.

**width** Width of the table, in percentage of the total width, when exported e.g., to Word. For full width, use `width = 1`.

**broom** If providing a tidy table produced with the broom package, which model type to use if one wants automatic formatting (options are "t.test", "lm", "cor.test", and "wilcox.test").

**report** If providing an object produced with the report package, which model type to use if one wants automatic formatting (options are "t.test", "lm", and "cor.test").

**short** Logical. Whether to return an abbreviated version of the tables made by the report package.

**title** Optional, to add a table header, if desired.

**note** Optional, to add one or more table footnote (APA note), if desired.

**separate.header** Logical, whether to separate headers based on name delimiters (i.e., periods ".").

**Details**

The resulting `flextable` objects can be opened in Word with `print(table, preview ="docx")`, or saved to Word with the `flextable::save_as_docx()` function.

**Value**

An APA-formatted table of class "flextable" (and "nice_table").

**See Also**

Tutorial: [https://rempsyc.remi-theriault.com/articles/table](https://rempsyc.remi-theriault.com/articles/table)
Examples

```r
# Make the basic table
my_table <- nice_table(
  mtcars[1:3, ],
  title = c("Table 1", "Motor Trend Car Road Tests"),
  note = c(
    "The data was extracted from the 1974 Motor Trend US magazine.",
    "* p < .05, ** p < .01, *** p < .001"
  )
)
my_table

# Save table to word
mypath <- tempfile(fileext = ".docx")
flextable::save_as_docx(my_table, path = mypath)

# Publication-ready tables
mtcars.std <- lapply(mtcars, scale)
model <- lm(mpg ~ cyl + wt * hp, mtcars.std)
stats.table <- as.data.frame(summary(model)$coefficients)
CI <- confint(model)
stats.table <- cbind(
  row.names(stats.table),
  stats.table, CI
)
names(stats.table) <- c(
  "Term", "B", "SE", "t", "p",
  "CI_lower", "CI_upper"
)
nice_table(stats.table, highlight = TRUE)

# Test different column names
test <- head(mtcars)
names(test) <- c(
  "dR", "N", "M", "SD", "b", "np2",
  "ges", "p", "r", "R2", "sr2"
)
nice_table(test)

# Custom cell formatting (such as p or r)
nice_table(test[8:11], col.format.p = 2:4, highlight = .001)

nice_table(test[8:11], col.format.r = 1:4)

# Apply custom functions to cells
fun <- function(x) {
  x + 11.1
}
```
nice_table(test[8:11], col.format.custom = 2:4, format.custom = "fun")

fun <- function(x) {
  paste("x", x)
}

nice_table(test[8:11], col.format.custom = 2:4, format.custom = "fun")

# Separate headers based on periods
header.data <- structure(
  list(
    Variable = c(  
      "Sepal.Length",  
      "Sepal.Width", "Petal.Length"
    ),  
    setosa.M = c(  
      5.01, 3.43,  
      1.46
    ),  
    setosa.SD = c(0.35, 0.38, 0.17),  
    versicolor.M =  
      c(5.94, 2.77, 4.26),  
    versicolor.SD = c(0.52, 0.31, 0.47)
  ),  
  row.names = c(NA, -3L), class = "data.frame"
)

nice_table(header.data,  
  separate.header = TRUE,  
  italics = 2:4
)

---

**nice_t_test**

_Easy t-tests_

**Description**

Easily compute t-test analyses, with effect sizes, and format in publication-ready format. The 95% confidence interval is for the effect size, Cohen’s d, both provided by the `effectsize` package.

**Usage**

```r
nice_t_test(
  data,  
  response,  
  group = NULL,  
  correction = "none",  
  verbose = TRUE,  
  ...  
)
```

**Arguments**

- `data` The data frame.
response The dependent variable.
group The group for the comparison.
correction What correction for multiple comparison to apply, if any. Default is "none" and the only other option (for now) is "bonferroni".
verbose Whether to display the Welch test warning or not.
Further arguments to be passed to the `t.test()` function (e.g., to use Student instead of Welch test, to change from two-tail to one-tail, or to do a paired-sample t-test instead of independent samples).

Details

This function relies on the base R `t.test()` function, which uses the Welch t-test per default (see why here: https://daniellakens.blogspot.com/2015/01/always-use-welchs-t-test-instead-of.html). To use the Student t-test, simply add the following argument: `var.equal = TRUE`.

Note that for paired t tests, you need to use `paired = TRUE`, and you also need data in "long" format rather than wide format (like for the ToothGrowth data set). In this case, the group argument refers to the participant ID for example, so the same group/participant is measured several times, and thus has several rows. Note also that R >= 4.4.0 has stopped supporting the `paired` argument for the formula method used internally here.

For the easystats equivalent, use: `report::report()` on the `t.test()` object.

Value

A formatted dataframe of the specified model, with DV, degrees of freedom, t-value, p-value, the effect size, Cohen’s d, and its 95% confidence interval lower and upper bounds.

See Also

Tutorial: https://rempsyc.remi-theriault.com/articles/t-test

Examples

```r
# Make the basic table
nice_t_test(
  data = mtcars,
  response = "mpg",
  group = "am"
)

# Multiple dependent variables at once
nice_t_test(
  data = mtcars,
  response = names(mtcars)[1:7],
  group = "am"
)

# Can be passed some of the regular arguments
# of base [t.test()]
```
# Student t-test (instead of Welch)
nice_t_test(
    data = mtcars,
    response = "mpg",
    group = "am",
    var.equal = TRUE
)

# One-sided instead of two-sided
nice_t_test(
    data = mtcars,
    response = "mpg",
    group = "am",
    alternative = "less"
)

# One-sample t-test
nice_t_test(
    data = mtcars,
    response = "mpg",
    mu = 10
)

# Make sure cases appear in the same order for
# both levels of the grouping factor

---

### nice_var

**Obtain variance per group**

**Description**

Obtain variance per group as well as check for the rule of thumb of one group having variance four times bigger than any of the other groups. Variance ratio is calculated as Max / Min.

**Usage**

```r
nice_var(data, variable, group, criteria = 4)
```

**Arguments**

- `data` The data frame
- `variable` The dependent variable to be plotted.
- `group` The group by which to plot the variable.
- `criteria` Desired threshold if one wants something different than four times the variance.
42 nice_varplot

Value

A dataframe, with the values of the selected variables for each group, their max variance ratio (maximum variance divided by the minimum variance), the selected decision criterion, and whether the data are considered heteroscedastic according to the decision criterion.

See Also

Other functions useful in assumption testing: nice_assumptions, nice_density, nice_normality, nice_qq, nice_varplot. Tutorial: https://rempsyc.remi-theriault.com/articles/assumptions

Examples

# Make the basic table
nice_var(
    data = iris,
    variable = "Sepal.Length",
    group = "Species"
)

# Try on multiple variables
nice_var(
    data = iris,
    variable = names(iris[1:4]),
    group = "Species"
)

nice_varplot 

Attempt to visualize variance per group

Description

Attempt to visualize variance per group.

Usage

nice_varplot(
    data, 
    variable, 
    group, 
    colours, 
    groups.labels, 
    grid = TRUE, 
    shapiro = FALSE, 
    ytitle = variable
)
nice_varplot

Arguments

- **data**: The data frame
- **variable**: The dependent variable to be plotted.
- **group**: The group by which to plot the variable.
- **colours**: Desired colours for the plot, if desired.
- **groups.labels**: How to label the groups.
- **grid**: Logical, whether to keep the default background grid or not. APA style suggests not using a grid in the background, though in this case some may find it useful to more easily estimate the slopes of the different groups.
- **shapiro**: Logical, whether to include the p-value from the Shapiro-Wilk test on the plot.
- **ytitle**: An optional y-axis label, if desired.

Value

A scatter plot of class ggplot attempting to display the group variances. Also includes the max variance ratio (maximum variance divided by the minimum variance).

See Also

Other functions useful in assumption testing: `nice_assumptions`, `nice_density`, `nice_normality`, `nice_qq`, `nice_var`. Tutorial: https://rempsyc.remi-theriault.com/articles/assumptions

Examples

```r
# Make the basic plot
nice_varplot(
  data = iris,
  variable = "Sepal.Length",
  group = "Species"
)

# Further customization
nice_varplot(
  data = iris,
  variable = "Sepal.Length",
  group = "Species",
  colours = c("#00BA38",
              "#619CFF",
              "#F8766D"),
  ytitle = "Sepal Length",
  groups.labels = c("(a) Setosa",
                    "(b) Versicolor",
                    "(c) Virginica")
)
```
nice_violin  

Description

Make nice violin plots easily with 95% (possibly bootstrapped) confidence intervals.

Usage

nice_violin(
  data,
  response,
  group = NULL,
  boot = FALSE,
  bootstraps = 2000,
  colours,
  xlabels = NULL,
  ytitle = response,
  xtitle = NULL,
  has.ylabels = TRUE,
  has.xlabels = TRUE,
  comp1 = 1,
  comp2 = 2,
  signif_annotation = NULL,
  signif_yposition = NULL,
  signif_xmin = NULL,
  signif_xmax = NULL,
  ymin,
  ymax,
  yby = 1,
  CIcap.width = 0.1,
  obs = FALSE,
  alpha = 1,
  border.colour = "black",
  border.size = 2,
  has.d = FALSE,
  d.x = mean(c(comp1, comp2)) * 1.1,
  d.y = mean(data[[response]]) * 1.3,
  groups.order = "none",
  xlabels.angle = 0
)

Arguments

data The data frame.
response  The dependent variable to be plotted.
group  The group by which to plot the variable.
boot  Logical, whether to use bootstrapping for the confidence interval or not.
bootstraps  How many bootstraps to use.
colours  Desired colours for the plot, if desired.
xlabels  The individual group labels on the x-axis.
ytitle  An optional y-axis label, if desired.
xtitle  An optional x-axis label, if desired.
has.ylabels  Logical, whether the x-axis should have labels or not.
has.xlabels  Logical, whether the y-axis should have labels or not.
comp1  The first unit of a pairwise comparison, if the goal is to compare two groups. Automatically displays *, **, or *** depending on significance of the difference. Can take either a numeric value (based on the group number) or the name of the group directly. Must be provided along with argument comp2.
comp2  The second unit of a pairwise comparison, if the goal is to compare two groups. Automatically displays "", "", or "" depending on significance of the difference. Can take either a numeric value (based on the group number) or the name of the group directly. Must be provided along with argument comp1.
signif.annotation  Manually provide the required annotations/numbers of stars (as character strings). Useful if the automatic pairwise comparison annotation does not work as expected, or yet if one wants more than one pairwise comparison. Must be provided along with arguments signif_yposition, signif_xmin, and signif xmax.
signif.yposition  Manually provide the vertical position of the annotations/stars, based on the y-scale.
signif.xmin  Manually provide the first part of the horizontal position of the annotations/stars (start of the left-sided bracket), based on the x-scale.
signif.xmax  Manually provide the second part of the horizontal position of the annotations/stars (end of the right-sided bracket), based on the x-scale.
ymin  The minimum score on the y-axis scale.
ymax  The maximum score on the y-axis scale.
yby  How much to increase on each "tick" on the y-axis scale.
CIcap.width  The width of the confidence interval cap.
obsv  Logical, whether to plot individual observations or not. The type of plotting can also be specified, either "dotplot" (same as obs = TRUE for backward compatibility) or "jitter", useful when there are a lot of observations.
alpha  The transparency of the plot.
border.colour  The colour of the violins border.
border.size  The size of the violins border.
has.d  Whether to display the d-value.
d.x  The x-axis coordinates for the d-value.
d.y  The y-axis coordinates for the d-value.
groups.order  How to order the group factor levels on the x-axis. Either "increasing" or "decreasing", to order based on the value of the variable on the y axis, or "string.length", to order from the shortest to the longest string (useful when working with long string names). "Defaults to "none".
xlabels.angle  How much to tilt the labels of the x-axis. Useful when working with long string names. "Defaults to 0."

Details

Using boot = TRUE uses bootstrapping (for the confidence intervals only) with the BCa method, using the rcompanion_groupwiseMean function.

For the easystats equivalent, see: see::geom_violindot().

Value

A violin plot of class ggplot, by group.

See Also


Examples

# Make the basic plot
nice_violin(
  data = ToothGrowth,
  response = "len"
)

# Save a high-resolution image file to specified directory
ggplot2::ggsave("niceviolinplotthere.pdf", width = 7,
  height = 7, unit = "in", dpi = 300
)  # change for your own desired path

# Change x- and y- axes labels
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  ytitle = "Length of Tooth",
  xtitle = "Vitamin C Dosage"
)

# See difference between two groups
nice_violin(
nice_violin

data = ToothGrowth,
group = "dose",
response = "len",
comp1 = "0.5",
comp2 = "2"
)
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  comp1 = 2,
  comp2 = 3
)

# Compare all three groups
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  signif_annotation = c("\text{*}", "\text{**}", "\text{***}"),
  # manually enter the number of stars
  signif_yposition = c(30, 35, 40),
  # What height (y) should the stars appear
  signif_xmin = c(1, 2, 1),
  # Where should the left-sided brackets start (x)
  signif_xmax = c(2, 3, 3)
)

# Where should the right-sided brackets end (x)

# Set the colours manually
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  colours = c("darkseagreen", "cadetblue", "darkslateblue")
)

# Changing the names of the x-axis labels
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  xlabels = c("Low", "Medium", "High")
)

# Removing the x-axis or y-axis titles
nice_violin(
  data = ToothGrowth,
  group = "dose",
  response = "len",
  ytitle = NULL,
  xtitle = NULL
# Removing the x-axis or y-axis labels (for whatever purpose)
nice_violin(
    data = ToothGrowth,
    group = "dose",
    response = "len",
    has.ylabels = FALSE,
    has.xlabels = FALSE
)

# Set y-scale manually
nice_violin(
    data = ToothGrowth,
    group = "dose",
    response = "len",
    ymin = 5,
    ymax = 35,
    yby = 5
)

# Plotting individual observations
nice_violin(
    data = ToothGrowth,
    group = "dose",
    response = "len",
    obs = TRUE
)

# Micro-customizations
nice_violin(
    data = ToothGrowth,
    group = "dose",
    response = "len",
    CIcap.width = 0,
    alpha = .70,
    border.size = 1,
    border.colour = "white",
    comp1 = 1,
    comp2 = 2,
    has.d = TRUE
)
**Description**

Interpolating the Inclusion of the Other in the Self Scale (IOS; self-other merging) easily. The user provides the IOS score, from 1 to 7, and the function will provide a percentage of actual area overlap between the two circles (i.e., not linear overlap), so it is possible to say, e.g., that experimental group 1 had an average overlap of X% with the other person, whereas experimental group 2 had an average overlap of X% with the other person.

**Usage**

```r
overlap_circle(response, categories = c("Self", "Other"), scoring = "IOS")
```

**Arguments**

- **response**: The variable to plot: requires IOS scores ranging from 1 to 7 (when scoring = "IOS").
- **categories**: The desired category names of the two overlapping circles for display on the plot.
- **scoring**: One of c("IOS", "percentage", "direct"). If scoring = "IOS", response needs to be a value between 1 to 7. If set to "percentage" or "direct", responses need to be between 0 and 100. If set to "direct", must provide exactly three values that represent the area from the first circle, the middle overlapping area, and area from the second circle.

**Details**

The circles are generated through the `VennDiagram::draw.pairwise.venn()` function and the desired percentage overlap is passed to its `cross.area` argument ("The size of the intersection between the sets"). The percentage overlap values are interpolated from this reference grid: Score of 1 = 0%, 2 = 10%, 3 = 20%, 4 = 30%, 5 = 55%, 6 = 65%, 7 = 85%.

**Value**

A plot of class gList, displaying overlapping circles relative to the selected score.

**See Also**

Tutorial: [https://rempsyc.remi-theriault.com/articles/circles](https://rempsyc.remi-theriault.com/articles/circles)

For a javascript web plugin of a continuous version of the Inclusion of Other in the Self (IOS) task (instead of the pen and paper version), for experiments during data collection, rather than data analysis, please see: [https://github.com/jspsych/jspsych-contrib/tree/main/packages/plugin-ios](https://github.com/jspsych/jspsych-contrib/tree/main/packages/plugin-ios)

**Examples**

```r
# Score of 1 (0% overlap)
overlap_circle(1)
```

```r
# Score of 3.5 (25% overlap)
overlap_circle(3.5)
```
```

overlap_circle(3.5)

# Score of 6.84 (81.8% overlap)
overlap_circle(6.84)

# Changing labels
overlap_circle(3.12, categories = c("Humans", "Animals"))

# Saving to file (PDF or PNG)
plot <- overlap_circle(3.5)
 ggplot2::ggsave(plot,
   file = tempfile(fileext = ".pdf"), width = 7,
   height = 7, unit = "in", dpi = 300
)
# Change for your own desired path
```

---

**plot_outliers**

*Visually check outliers (dot plot)*

**Description**

Easily and visually check outliers through a dot plot with accompanying reference lines at +/- 3 MAD or SD. When providing a group, data are group-mean centered and standardized (based on MAD or SD); if no group is provided, data are simply standardized.

**Usage**

```r
plot_outliers(
  data,
  group = NULL,
  response,
  method = "mad",
  criteria = 3,
  colours,
  xlabels = NULL,
  ytitle = NULL,
  xtitle = NULL,
  has.ylabels = TRUE,
  has.xlabels = TRUE,
  ymin,
  ymax,
  yby = 1,
  ...
)
```
plot_outliers

Arguments

- **data**: The data frame.
- **group**: The group by which to plot the variable.
- **response**: The dependent variable to be plotted.
- **method**: Method to identify outliers, either (e.g., 3) median absolute deviations ("mad", default) or standard deviations ("sd").
- **criteria**: How many MADs (or standard deviations) to use as threshold (default is 3).
- **colours**: Desired colours for the plot, if desired.
- **xlabels**: The individual group labels on the x-axis.
- **ytitle**: An optional y-axis label, if desired.
- **xtitle**: An optional x-axis label, if desired.
- **has.ylabels**: Logical, whether the x-axis should have labels or not.
- **has.xlabels**: Logical, whether the y-axis should have labels or not.
- **ymin**: The minimum score on the y-axis scale.
- **ymax**: The maximum score on the y-axis scale.
- **yby**: How much to increase on each "tick" on the y-axis scale.
- **...**: Other arguments passed to ggplot2::geom_dotplot.

Value

A dot plot of class ggplot, by group.

See Also

Other functions useful in assumption testing: Tutorial: [https://rempyc.remi-theriault.com/articles/assumptions](https://rempyc.remi-theriault.com/articles/assumptions)

Examples

```r
# Make the basic plot
plot_outliers(
  airquality,
  group = "Month",
  response = "Ozone"
)

plot_outliers(
  airquality,
  response = "Ozone",
  method = "sd"
)
```
scale_mad

Description

Scale and center ("standardize") data based on the median absolute deviation (MAD).

Usage

scale_mad(x)

Arguments

x The vector to be scaled.

Details

The function subtracts the median to each observation, and then divides the outcome by the MAD. This is analogous to regular standardization which subtracts the mean to each observation, and then divides the outcome by the standard deviation.

For the easystats equivalent, use: datawizard::standardize(x, robust = TRUE).

Value

A numeric vector of standardized data.

References


Examples

scale_mad(mtcars$mpg)
Description

Winsorize (bring extreme observations to usually +/- 3 standard deviations) data based on median absolute deviations instead of standard deviations.

Usage

winsorize_mad(x, criteria = 3)

Arguments

- x: The vector to be winsorized based on the MAD.
- criteria: How many MAD to use as threshold (similar to standard deviations)

Details

For the easystats equivalent, use: datawizard::winsorize(x, method = "zscore", threshold = 3, robust = TRUE).

Value

A numeric vector of winsorized data.

References


Examples

winsorize_mad(mtcars$qsec, criteria = 2)
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