

Package ‘guideR’

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Type Package

Title Miscellaneous Statistical Functions Used in 'guide-R'

Version 0.5.1

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'guide-R : Guide pour l'analyse de données d'enquêtes avec R' available at
<<https://larmarange.github.io/guide-R/>>. 'guideR' implements miscellaneous
functions introduced in 'guide-R' to facilitate statistical analysis and
manipulation of survey data.

License GPL (>= 3)

URL <https://larmarange.github.io/guideR/>,
<https://github.com/larmarange/guideR>

BugReports <https://github.com/larmarange/guideR/issues>

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combine_answers	<i>Combine answers of a multiple answers question</i>
-----------------	---

Description

Considering a multiple answers question coded as several binary variables (one per item), create a new variable (list column or character) combining all positive answers. If defined, use variable labels (see examples).

Usage

```
combine_answers(data, answers, into, value = NULL, sep = NULL)
```

Arguments

data	A data frame, data frame extension (e.g. a tibble), or a survey design object.
answers	<tidy-select> List of variables identifying the different answers of the question.
into	Names of new variables to create as character vector.
value	Value indicating a positive answer. By default, will use the maximum observed value and will display a message.
sep	An optional character string to separate the results and return a character. If NULL, return a list column (see examples).

Note

If NA is observed for at least one item, return NA.

Examples

```
d <-
  dplyr::tibble(
    q1a = sample(c("y", "n"), size = 200, replace = TRUE),
    q1b = sample(c("y", "n", "n", NA), size = 200, replace = TRUE),
    q1c = sample(c("y", "y", "n"), size = 200, replace = TRUE),
    q1d = sample("n", size = 200, replace = TRUE)
  )

d |> combine_answers(q1a:q1d, into = "combined")
d |> combine_answers(q1a:q1d, into = "combined", sep = ", ", value = "y")
d |> combine_answers(q1a:q1d, into = "combined", sep = " | ", value = "n")
```

cut_quartiles

*Cut a continuous variable in quartiles***Description**

Convenient function to quickly cut a numeric vector into quartiles, i.e. by applying `cut(x, breaks = fivenum(x))`. Variable label is preserved by `cut_quartiles()`.

Usage

```
cut_quartiles(x, include.lowest = TRUE, ...)
```

Arguments

x a numeric vector which is to be converted to a factor by cutting.

include.lowest logical, indicating if an 'x[i]' equal to the lowest (or highest, for `right = FALSE`) 'breaks' value should be included.

... further arguments passed to `base::cut()`.

Examples

```
mtcars$mpg |> cut_quartiles() |> summary()
```

grouped_tbl_pivot_wider

Helpers for grouped tables generated with gtsummary

Description

A series of helpers for grouped tables generated by `tbl_regression()` in case of multinomial models, multi-components models or other grouped results. `grouped_tbl_pivot_wider()` allows to display results in a wide format, with one set of columns per group. `multinom_add_global_p_pivot_wider()` is a specific case for multinomial models, when displaying global p-values in a wide format: it calls `gtsummary::add_global_p()`, followed by `grouped_tbl_pivot_wider()`, and then keep only the last column with p-values (see examples). Finally, as grouped regression tables doesn't have exactly the same structure as ungrouped tables, functions as `gtsummary::bold_labels()` do not always work properly. If the grouped table is kept in a long format, `style_grouped_tbl()` could be use to improve the output by styling variable labels, levels and/or group names. **TO BE NOTED:** to style group names, `style_grouped_tbl()` convert the table into a gt object with `gtsummary::as_gt()`. This function should therefore be used last. If the table is intended to be exported to another format, do not use `style_grouped_tbl()`.

Usage

```
grouped_tbl_pivot_wider(x)

multinom_add_global_p_pivot_wider(
  x,
  ...,
  p_value_header = "**Likelihood-ratio test**"
)

style_grouped_tbl(
  x,
  bold_groups = TRUE,
  uppercase_groups = TRUE,
  bold_labels = FALSE,
  italicize_labels = TRUE,
  indent_labels = 4L,
  bold_levels = FALSE,
  italicize_levels = FALSE,
  indent_levels = 8L
)
```

Arguments

<code>x</code>	A grouped regression table generated with <code>gtsummary::tbl_regression()</code> .
<code>...</code>	Additional arguments passed to <code>gtsummary::add_global_p()</code> .
<code>p_value_header</code>	Header for the p-value column.

```

bold_groups      Bold group group names?
uppercase_groups  Convert group names to upper case?

bold_labels      Bold variable labels?
italicize_labels  Italicize variable labels?

indent_labels    Number of spaces to indent variable labels.

bold_levels      Bold levels?
italicize_levels  Italicize levels?

indent_levels    Number of spaces to indent levels.

```

Value

A gtsummary or a gt table.

Examples

```

mod <- nnet::multinom(
  grade ~ stage + marker + age,
  data = gtsummary::trial,
  trace = FALSE
)
tbl <- mod |> gtsummary::tbl_regression(exponentiate = TRUE)
tbl
tbl |> grouped_tbl_pivot_wider()

tbl |> multinom_add_global_p_pivot_wider() |> gtsummary::bold_labels()
tbl |> style_grouped_tbl()

```

install_dependencies *Install / Update project dependencies*

Description

This function uses `renv::dependencies()` to identify R package dependencies in a project and then calls `pak::pkg_install()` to install / update these packages. If some packages are not found, the function will install those available and returns a message indicated packages not installed/updated.

Usage

```
install_dependencies(dependencies = NULL, ask = TRUE)
```

Arguments

dependencies	An optional list of dependencies. If NULL, will be determined with <code>renv::dependencies()</code> . If equal to "old", will use the list returned by <code>utils::old.packages()</code> .
ask	Whether to ask for confirmation when installing a different version of a package that is already installed. Installations that only add new packages never require confirmation.

Value

(Invisibly) A data frame with information about the installed package(s).

Examples

```
## Not run:
install_dependencies()

## End(Not run)
```

is_different	<i>Comparison tests considering NA as values to be compared</i>
--------------	---

Description

`is_different()` and `is_equal()` performs comparison tests, considering NA values as legitimate values (see examples).

Usage

```
is_different(x, y)

is_equal(x, y)

cumdifferent(x)

num_cycle(x)
```

Arguments

x, y	Vectors to be compared.
------	-------------------------

Details

`cum_different()` allows to identify groups of continuous rows that have the same value. `num_cycle()` could be used to identify sub-groups that respect a certain condition (see examples).

`is_equal(x, y)` is equivalent to `(x == y & !is.na(x) & !is.na(y)) | (is.na(x) & is.na(y))`, and `is_different(x, y)` is equivalent to `(x != y & !is.na(x) & !is.na(y)) | xor(is.na(x), is.na(y))`.

Value

A vector of the same length as x.

Examples

```
v <- c("a", "b", NA)
is_different(v, "a")
is_different(v, NA)
is_equal(v, "a")
is_equal(v, NA)
d <- dplyr::tibble(group = c("a", "a", "b", "b", "a", "b", "c", "a"))
d |>
  dplyr::mutate(
    subgroup = cumdifferent(group),
    sub_a = num_cycle(group == "a")
  )
```

leading_zeros

Add leading zeros

Description

Add leading zeros

Usage

```
leading_zeros(x, left_digits = NULL, digits = 0, prefix = "", suffix = "", ...)
```

Arguments

x	a numeric vector
left_digits	number of digits before decimal point, automatically computed if not provided
digits	number of digits after decimal point
prefix, suffix	Symbols to display before and after value
...	additional parameters passed to base::formatC() , as <code>big.mark</code> or <code>decimal.mark</code>

Value

A character vector of the same length as x.

See Also

[base::formatC\(\)](#), [base::sprintf\(\)](#)

Examples

```
v <- c(2, 103.24, 1042.147, 12.4566, NA)
leading_zeros(v)
leading_zeros(v, digits = 1)
leading_zeros(v, left_digits = 6, big.mark = " ")
leading_zeros(c(0, 6, 12, 18), prefix = "M")
```

long_to_periods	<i>Transform a data frame from long format to period format</i>
-----------------	---

Description

Transform a data frame from long format to period format

Usage

```
long_to_periods(data, id, start, stop = NULL, by = NULL)
```

Arguments

data	A data frame, or a data frame extension (e.g. a tibble).
id	<tidy-select> Column containing individual ids
start	<tidy-select> Time variable indicating the beginning of each row
stop	<tidy-select> Optional time variable indicating the end of each row. If not provided, it will be derived from the dataset, considering that each row ends at the beginning of the next one.
by	<tidy-select> Co-variables to consider (optional)

Value

A tibble.

See Also

```
periods\_to\_long\(\)
```

Examples

```
d <- dplyr::tibble(
  patient = c(1, 2, 3, 3, 4, 4, 4),
  begin = c(0, 0, 0, 1, 0, 36, 39),
  end = c(50, 6, 1, 16, 36, 39, 45),
  covar = c("no", "no", "no", "yes", "no", "yes", "yes")
)
d

d |> long_to_periods(id = patient, start = begin, stop = end)
```

```
d |> long_to_periods(id = patient, start = begin, stop = end, by = covar)

# If stop not provided, it is deduced.
# However, it considers that observation ends at the last start time.
d |> long_to_periods(id = patient, start = begin)
```

`observed_vs_theoretical`*Plot observed vs predicted distribution of a fitted model*

Description

Plot observed vs predicted distribution of a fitted model

Usage

```
observed_vs_theoretical(model)
```

Arguments

<code>model</code>	A statistical model.
--------------------	----------------------

Details

Has been tested with `stats::lm()` and `stats::glm()` models. It may work with other types of models, but without any warranty.

Value

A ggplot2 plot.

Examples

```
# a linear model
mod <- lm(Sepal.Length ~ Sepal.Width + Species, data = iris)
mod |> observed_vs_theoretical()

# a logistic regression
mod <- glm(
  as.factor(Survived) ~ Class + Sex,
  data = titanic,
  family = binomial()
)
mod |> observed_vs_theoretical()
```

periods_to_long	<i>Transform a data frame from period format to long format</i>
-----------------	---

Description

Transform a data frame from period format to long format

Usage

```
periods_to_long(
  data,
  start,
  stop,
  time_step = 1,
  time_name = "time",
  keep = FALSE
)
```

Arguments

data	A data frame, or a data frame extension (e.g. a tibble).
start	<tidy-select> Time variable indicating the beginning of each row
stop	<tidy-select> Optional time variable indicating the end of each row. If not provided, it will be derived from the dataset, considering that each row ends at the beginning of the next one.
time_step	(numeric) Desired value for the time variable.
time_name	(character) Name of the time variable.
keep	(logical) Should start and stop variable be kept in the results?

Value

A tibble.

See Also

[long_to_periods\(\)](#)

Examples

```
d <- dplyr::tibble(
  patient = c(1, 2, 3, 3),
  begin = c(0, 2, 0, 3),
  end = c(6, 4, 2, 8),
  covar = c("no", "yes", "no", "yes")
)
d
```

```
d |> periods_to_long(start = begin, stop = end)
d |> periods_to_long(start = begin, stop = end, time_step = 5)
```

plot_inertia_from_tree

Plot inertia, absolute loss and relative loss from a classification tree

Description

Plot inertia, absolute loss and relative loss from a classification tree

Usage

```
plot_inertia_from_tree(tree, k_max = 15)

get_inertia_from_tree(tree, k_max = 15)
```

Arguments

tree	A dendrogram, i.e. an stats::hclust object, an FactoMineR::HCPC object or an object that can be converted to an stats::hclust object with stats::as.hclust() .
k_max	Maximum number of clusters to return / plot.

Value

A ggplot2 plot or a tibble.

Examples

```
hc <- hclust(dist(USArrests))
get_inertia_from_tree(hc)
plot_inertia_from_tree(hc)
```

plot_multiple_answers *Plot a multiple answers question*

Description

Considering a multiple answers question coded as several binary variables (one per answer), plot the proportion of positive answers. If `combine_answers = FALSE`, plot the proportion of positive answers of each item, separately. If `combine_answers = TRUE`, combine the different answers (see [combine_answers\(\)](#)) and plot the proportion of each combination ([ggupset](#) package required when `flip = FALSE`). See [proportion\(\)](#) for more details on the way proportions and confidence intervals are computed. By default, return a bar plot, but other geometries could be used (see examples). If defined, use variable labels (see examples).

Usage

```

plot_multiple_answers(
  data,
  answers = dplyr::everything(),
  value = NULL,
  by = NULL,
  combine_answers = FALSE,
  combine_sep = " | ",
  missing_label = " missing",
  none_label = "none",
  drop_na = FALSE,
  sort = c("none", "ascending", "descending", "degrees"),
  geom = "bar",
  ...,
  show_ci = TRUE,
  conf_level = 0.95,
  ci_color = "black",
  show_labels = TRUE,
  labels_labeller = scales::label_percent(1),
  labels_size = 3.5,
  labels_color = "black",
  flip = FALSE,
  return_data = FALSE
)

plot_multiple_answers_dodge(
  data,
  answers = dplyr::everything(),
  value = NULL,
  by,
  combine_answers = FALSE,
  combine_sep = " | ",
  missing_label = " missing",
  none_label = "none",
  drop_na = FALSE,
  sort = c("none", "ascending", "descending", "degrees"),
  geom = c("bar", "point"),
  width = 0.75,
  ...,
  show_ci = TRUE,
  conf_level = 0.95,
  ci_color = "black",
  show_labels = TRUE,
  labels_labeller = scales::label_percent(1),
  labels_size = 3.5,
  labels_color = "black",
  flip = FALSE
)

```

Arguments

data	A data frame, data frame extension (e.g. a tibble), or a survey design object.
answers	<code><tidy-select></code> List of variables identifying the different answers of the question.
value	Value indicating a positive answer. By default, will use the maximum observed value and will display a message.
by	<code><tidy-select></code> Optional list of variables to compare (using facets).
combine_answers	Should answers be combined? (see examples)
combine_sep	Character string to separate combined answers.
missing_label	When combining answers and <code>drop_na = FALSE</code> , label for missing values.
none_label	When combining answers and <code>flip = TRUE</code> , label when no item is selected.
drop_na	Should any observation with a least one NA value be dropped?
sort	Should answers be sorted according to their proportion? They could also be sorted by degrees (number of elements) when combining answers.
geom	Geometry to use for plotting proportions ("bar" by default).
...	Additional arguments passed to the geom defined by geom.
show_ci	Display confidence intervals?
conf_level	Confidence level for the confidence intervals.
ci_color	Color of the error bars representing confidence intervals.
show_labels	Display proportion labels?
labels_labeller	Labeller function for proportion labels.
labels_size	Size of proportion labels.
labels_color	Color of proportion labels.
flip	Flip x and y axis?
return_data	Return computed data instead of the plot?
width	Dodging width.

Note

If `drop_na = TRUE`, any observation with at least one NA value for one item will be dropped. If `drop_na = FALSE` and `combine_answers = FALSE`, NA values for a specific answer are excluded the denominator when computing proportions. Therefore, all proportions may be computed on different population sizes. If `drop_na = FALSE` and `combine_answers = TRUE`, any observation with at least one NA value will be labeled with `missing_label`.

Examples

```

d <-
  dplyr::tibble(
    q1a = sample(c("y", "n"), size = 200, replace = TRUE),
    q1b = sample(c("y", "n", "n", NA), size = 200, replace = TRUE),
    q1c = sample(c("y", "y", "n"), size = 200, replace = TRUE),
    q1d = sample("n", size = 200, replace = TRUE)
  )

d |> plot_multiple_answers(q1a:q1c)

d |>
  labelled::set_variable_labels(
    q1a = "apple",
    q1b = "banana",
    q1c = "chocolate",
    q1d = "Dijon mustard"
  ) |>
  plot_multiple_answers(
    value = "y",
    drop_na = TRUE,
    sort = "desc",
    fill = "lightblue",
    flip = TRUE
  )

d |>
  plot_multiple_answers(
    combine_answers = TRUE,
    value = "y",
    fill = "#DDCC77",
    drop_na = TRUE
  )

d |>
  plot_multiple_answers(
    combine_answers = TRUE,
    value = "y",
    flip = TRUE,
    mapping = ggplot2::aes(fill = prop),
    show.legend = FALSE
  ) +
  ggplot2::scale_fill_distiller(palette = "Spectral")

d$group <- sample(c("group A", "group B"), size = 200, replace = TRUE)
d |>
  plot_multiple_answers(
    answers = q1a:q1d,
    by = group,
    combine_answers = TRUE,
    sort = "degrees",
    value = "y",

```

```

    fill = "grey80"
  )

d |>
  plot_multiple_answers_dodge(q1a:q1d, by = group)
d |>
  plot_multiple_answers_dodge(q1a:q1d, by = group, flip = TRUE)
d |>
  plot_multiple_answers_dodge(q1a:q1d, by = group, combine_answers = TRUE)

```

plot_proportions	<i>Plot proportions by sub-groups</i>
------------------	---------------------------------------

Description

Plot one or several proportions (defined by logical conditions) by sub-groups. See [proportion\(\)](#) for more details on the way proportions and confidence intervals are computed. By default, return a bar plot, but other geometries could be used (see examples). `stratified_by()` is an helper function facilitating a stratified analyses (i.e. proportions by groups stratified according to a third variable, see examples). `dummy_proportions()` is an helper to easily convert a categorical variable into dummy variables and therefore showing the proportion of each level of the original variable (see examples).

Usage

```

plot_proportions(
  data,
  condition,
  by = NULL,
  drop_na_by = FALSE,
  convert_continuous = TRUE,
  geom = "bar",
  ...,
  show_overall = TRUE,
  overall_label = "Overall",
  show_ci = TRUE,
  conf_level = 0.95,
  ci_color = "black",
  show_pvalues = TRUE,
  pvalues_test = c("fisher", "chisq"),
  pvalues_labeller = scales::label_pvalue(add_p = TRUE),
  pvalues_size = 3.5,
  show_labels = TRUE,
  labels_labeller = scales::label_percent(1),
  labels_size = 3.5,
  labels_color = "black",

```

```

    show_overall_line = FALSE,
    overall_line_type = "dashed",
    overall_line_color = "black",
    overall_line_width = 0.5,
    facet_labeller = ggplot2::label_wrap_gen(width = 50, multi_line = TRUE),
    flip = FALSE,
    free_scale = FALSE,
    return_data = FALSE
)

stratified_by(condition, strata)

dummy_proportions(variable)

```

Arguments

<code>data</code>	A data frame, data frame extension (e.g. a tibble), or a survey design object.
<code>condition</code>	<code><data-masking></code> A condition defining a proportion, or a <code>dplyr::tibble()</code> defining several proportions (see examples).
<code>by</code>	<code><tidy-select></code> List of variables to group by (comparison is done separately for each variable).
<code>drop_na_by</code>	Remove NA values in by variables?
<code>convert_continuous</code>	Should continuous variables (with 5 unique values or more) be converted to quartiles (using <code>cut_quartiles()</code>)?
<code>geom</code>	Geometry to use for plotting proportions ("bar" by default).
<code>...</code>	Additional arguments passed to the geom defined by <code>geom</code> .
<code>show_overall</code>	Display "Overall" column?
<code>overall_label</code>	Label for the overall column.
<code>show_ci</code>	Display confidence intervals?
<code>conf_level</code>	Confidence level for the confidence intervals.
<code>ci_color</code>	Color of the error bars representing confidence intervals.
<code>show_pvalues</code>	Display p-values in the top-left corner?
<code>pvalues_test</code>	Test to compute p-values for data frames: "fisher" for <code>stats::fisher.test()</code> (with <code>simulate.p.value = TRUE</code>) or "chisq" for <code>stats::chisq.test()</code> . Has no effect on survey objects for those <code>survey::svychisq()</code> is used.
<code>pvalues_labeller</code>	Labeller function for p-values.
<code>pvalues_size</code>	Text size for p-values.
<code>show_labels</code>	Display proportion labels?
<code>labels_labeller</code>	Labeller function for proportion labels.
<code>labels_size</code>	Size of proportion labels.

labels_color	Color of proportion labels.
show_overall_line	Add an overall line?
overall_line_type	Line type of the overall line.
overall_line_color	Color of the overall line.
overall_line_width	Line width of the overall line.
facet_labeller	Labeller function for strip labels.
flip	Flip x and y axis?
free_scale	Allow y axis to vary between conditions?
return_data	Return computed data instead of the plot?
strata	Stratification variable
variable	Variable to be converted into dummy variables.

Examples

```
titanic |>
  plot_proportions(
    Survived == "Yes",
    overall_label = "All",
    labels_color = "white"
  )
```

```
titanic |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    fill = "lightblue"
  )
```

```
titanic |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    fill = "lightblue",
    flip = TRUE
  )
```

```
titanic |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    geom = "point",
    color = "red",
    size = 3,
  )
```

```

    show_labels = FALSE
  )

titanic |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    geom = "area",
    fill = "lightgreen",
    show_overall = FALSE
  )

titanic |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    geom = "line",
    color = "purple",
    ci_color = "darkblue",
    show_overall = FALSE
  )

titanic |>
  plot_proportions(
    Survived == "Yes",
    by = -Survived,
    mapping = ggplot2::aes(fill = variable),
    color = "black",
    show.legend = FALSE,
    show_overall_line = TRUE,
    show_pvalues = FALSE
  )

# defining several proportions

titanic |>
  plot_proportions(
    dplyr::tibble(
      Survived = Survived == "Yes",
      Male = Sex == "Male"
    ),
    by = c(Class),
    mapping = ggplot2::aes(fill = condition)
  )

titanic |>
  plot_proportions(
    dplyr::tibble(
      Survived = Survived == "Yes",
      Male = Sex == "Male"
    ),
    by = c(Class),
    mapping = ggplot2::aes(fill = condition),

```

```

    free_scale = TRUE
  )

iris |>
  plot_proportions(
    dplyr::tibble(
      "Long sepal" = Sepal.Length > 6,
      "Short petal" = Petal.Width < 1
    ),
    by = Species,
    fill = "palegreen"
  )

iris |>
  plot_proportions(
    dplyr::tibble(
      "Long sepal" = Sepal.Length > 6,
      "Short petal" = Petal.Width < 1
    ),
    by = Species,
    fill = "palegreen",
    flip = TRUE
  )

# works with continuous by variables
iris |>
  labelled::set_variable_labels(
    Sepal.Length = "Length of the sepal"
  ) |>
  plot_proportions(
    Species == "versicolor",
    by = dplyr::contains("leng"),
    fill = "plum",
    colour = "plum4"
  )

# works with survey object
titanic |>
  srvyr::as_survey() |>
  plot_proportions(
    Survived == "Yes",
    by = c(Class, Sex),
    fill = "darksalmon",
    color = "black",
    show_overall_line = TRUE,
    labels_labeller = scales::label_percent(.1)
  )

# stratified analysis
titanic |>
  plot_proportions(
    (Survived == "Yes") |> stratified_by(Sex),

```

```

    by = Class,
    mapping = ggplot2::aes(fill = condition)
  ) +
  ggplot2::theme(legend.position = "bottom") +
  ggplot2::labs(fill = NULL)

# Convert Class into dummy variables
titanic |>
  plot_proportions(
    dummy_proportions(Class),
    by = Sex,
    mapping = ggplot2::aes(fill = level)
  )

```

proportion

Compute proportions

Description

`proportion()` lets you quickly count observations (like `dplyr::count()`) and compute relative proportions. Proportions are computed separately by group (see examples).

Usage

```

proportion(data, ...)

## S3 method for class 'data.frame'
proportion(
  data,
  ...,
  .by = NULL,
  .na.rm = FALSE,
  .weight = NULL,
  .scale = 100,
  .sort = FALSE,
  .drop = FALSE,
  .drop_na_by = FALSE,
  .conf.int = FALSE,
  .conf.level = 0.95,
  .options = list(correct = TRUE)
)

## S3 method for class 'survey.design'
proportion(
  data,
  ...,
  .by = NULL,
  .na.rm = FALSE,

```

```

    .scale = 100,
    .sort = FALSE,
    .drop_na_by = FALSE,
    .conf.int = FALSE,
    .conf.level = 0.95,
    .options = NULL
  )

## Default S3 method:
proportion(
  data,
  ...,
  .na.rm = FALSE,
  .scale = 100,
  .sort = FALSE,
  .drop = FALSE,
  .conf.int = FALSE,
  .conf.level = 0.95,
  .options = list(correct = TRUE)
)
```

Arguments

<code>data</code>	A vector, a data frame, data frame extension (e.g. a tibble), or a survey design object.
<code>...</code>	<data-masking> Variable(s) for those computing proportions.
<code>.by</code>	<tidy-select> Optional additional variables to group by (in addition to those eventually previously declared using dplyr::group_by()).
<code>.na.rm</code>	Should NA values be removed (from variables declared in <code>...</code>)?
<code>.weight</code>	<data-masking> Frequency weights. Can be NULL or a variable.
<code>.scale</code>	A scaling factor applied to proportion. Use 1 for keeping proportions unchanged.
<code>.sort</code>	If TRUE, will show the highest proportions at the top.
<code>.drop</code>	If TRUE, will remove empty groups from the output.
<code>.drop_na_by</code>	If TRUE, will remove any NA values observed in the <code>.by</code> variables (or variables defined with dplyr::group_by()).
<code>.conf.int</code>	If TRUE, will estimate confidence intervals.
<code>.conf.level</code>	Confidence level for the returned confidence intervals.
<code>.options</code>	Additional arguments passed to stats::prop.test() or srvyr::survey_prop() .

Value

A tibble.

A tibble with one row per group.

Examples

```

# using a vector
titanic$Class |> proportion()

# univariable table
titanic |> proportion(Class)
titanic |> proportion(Class, .sort = TRUE)
titanic |> proportion(Class, .conf.int = TRUE)
titanic |> proportion(Class, .conf.int = TRUE, .scale = 1)

# bivariable table
titanic |> proportion(Class, Survived) # proportions of the total
titanic |> proportion(Survived, .by = Class) # row proportions
titanic |> # equivalent syntax
  dplyr::group_by(Class) |>
  proportion(Survived)

# combining 3 variables or more
titanic |> proportion(Class, Sex, Survived)
titanic |> proportion(Sex, Survived, .by = Class)
titanic |> proportion(Survived, .by = c(Class, Sex))

# missing values
dna <- titanic
dna$Survived[c(1:20, 500:530)] <- NA
dna |> proportion(Survived)
dna |> proportion(Survived, .na.rm = TRUE)

## SURVEY DATA -----

ds <- srvyr::as_survey(titanic)

# univariable table
ds |> proportion(Class)
ds |> proportion(Class, .sort = TRUE)
ds |> proportion(Class, .conf.int = TRUE)
ds |> proportion(Class, .conf.int = TRUE, .scale = 1)

# bivariable table
ds |> proportion(Class, Survived) # proportions of the total
ds |> proportion(Survived, .by = Class) # row proportions
ds |> dplyr::group_by(Class) |> proportion(Survived)

# combining 3 variables or more
ds |> proportion(Class, Sex, Survived)
ds |> proportion(Sex, Survived, .by = Class)
ds |> proportion(Survived, .by = c(Class, Sex))

# missing values
dsna <- srvyr::as_survey(dna)
dsna |> proportion(Survived)

```

```
dsna |> proportion(Survived, .na.rm = TRUE)
```

round_preserve_sum	<i>Round values while preserve their rounded sum in R</i>
--------------------	---

Description

Sometimes, the sum of rounded numbers (e.g., using `base::round()`) is not the same as their rounded sum.

Usage

```
round_preserve_sum(x, digits = 0)
```

Arguments

x	Numerical vector to sum.
digits	Number of decimals for rounding.

Details

This solution applies the following algorithm

- Round down to the specified number of decimal places
- Order numbers by their remainder values
- Increment the specified decimal place of values with k largest remainders, where k is the number of values that must be incremented to preserve their rounded sum

Value

A numerical vector of same length as x.

Source

<https://biostatmatt.com/archives/2902>

Examples

```
sum(c(0.333, 0.333, 0.334))
round(c(0.333, 0.333, 0.334), 2)
sum(round(c(0.333, 0.333, 0.334), 2))
round_preserve_sum(c(0.333, 0.333, 0.334), 2)
sum(round_preserve_sum(c(0.333, 0.333, 0.334), 2))
```

step_with_na

Apply step(), taking into account missing values

Description

When your data contains missing values, concerned observations are removed from a model. However, then at a later stage, you try to apply a descending stepwise approach to reduce your model by minimization of AIC, you may encounter an error because the number of rows has changed.

Usage

```
step_with_na(model, ...)

## Default S3 method:
step_with_na(model, ..., full_data = eval(model$call$data))

## S3 method for class 'svyglm'
step_with_na(model, ..., design)
```

Arguments

model	A model object.
...	Additional parameters passed to <code>stats::step()</code> .
full_data	Full data frame used for the model, including missing data.
design	Survey design previously passed to <code>survey::svyglm()</code> .

Details

`step_with_na()` applies the following strategy:

- recomputes the models using only complete cases;
- applies `stats::step()`;
- recomputes the reduced model using the full original dataset.

`step_with_na()` has been tested with `stats::lm()`, `stats::glm()`, `nnet::multinom()`, `survey::svyglm()` and `survival::coxph()`. It may be working with other types of models, but with no warranty.

In some cases, it may be necessary to provide the full dataset initially used to estimate the model.

`step_with_na()` may not work inside other functions. In that case, you may try to pass `full_data` to the function.

Value

The stepwise-selected model.

Examples

```

set.seed(42)
d <- titanic |>
  dplyr::mutate(
    Group = sample(
      c("a", "b", NA),
      dplyr::n(),
      replace = TRUE
    )
  )
mod <- glm(as.factor(Survived) ~ ., data = d, family = binomial())
# step(mod) should produce an error
mod2 <- step_with_na(mod, full_data = d)
mod2

## WITH SURVEY -----

library(survey)
ds <- d |>
  dplyr::mutate(Survived = as.factor(Survived)) |>
  srvyr::as_survey()
mods <- survey::svyglm(
  Survived ~ Class + Group + Sex,
  design = ds,
  family = quasibinomial()
)
mod2s <- step_with_na(mods, design = ds)
mod2s

```

titanic

Titanic data set in long format

Description

This titanic dataset is equivalent to `datasets::Titanic |> dplyr::as_tibble() |> tidyr::uncount(n)`.

Usage

```
titanic
```

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 2201 rows and 4 columns.

See Also

[datasets::Titanic](#)

unrowwise	<i>Remove row-wise grouping</i>
-----------	---------------------------------

Description

Remove row-wise grouping created with `dplyr::rowwise()` while preserving any other grouping declared with `dplyr::group_by()`.

Usage

```
unrowwise(data)
```

Arguments

data	A tibble.
------	-----------

Value

A tibble.

Examples

```
titanic |> dplyr::rowwise()
titanic |> dplyr::rowwise() |> unrowwise()

titanic |> dplyr::group_by(Sex, Class) |> dplyr::rowwise()
titanic |> dplyr::group_by(Sex, Class) |> dplyr::rowwise() |> unrowwise()
```

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