Package ‘gtsummary’

July 15, 2023

Title  Presentation-Ready Data Summary and Analytic Result Tables

Version  1.7.2

Description  Creates presentation-ready tables summarizing data sets, regression models, and more. The code to create the tables is concise and highly customizable. Data frames can be summarized with any function, e.g. mean(), median(), even user-written functions. Regression models are summarized and include the reference rows for categorical variables. Common regression models, such as logistic regression and Cox proportional hazards regression, are automatically identified and the tables are pre-filled with appropriate column headers.

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      https://www.danieldsjoberg.com/gtsummary/

BugReports  https://github.com/ddsjoberg/gtsummary/issues

Depends  R (>= 3.4)

Imports  broom (>= 1.0.1),
         broom.helpers (>= 1.13.0),
         cli (>= 3.1.1),
         dplyr (>= 1.1.1),
        forcats (>= 1.0.0),
         glue (>= 1.6.2),
        gt (>= 0.9.0),
        knitr (>= 1.37),
lifecycle (>= 1.0.1),
purrr (>= 1.0.1),
rlang (>= 1.0.3),
stringr (>= 1.4.0),
tibble (>= 3.2.1),
tidy (>= 1.1.4),
vctrs (>= 0.5.2)

Suggests  aod (>= 1.3.1),
broom.mixed (>= 0.2.9),
car (>= 3.0-11),
emprsk,
covr,
R topics documented:

- effectsize (>= 0.6.0),
- emmeans (>= 1.7.3),
- flextable (>= 0.8.1),
- geepack,
- ggstats (>= 0.2.1),
- Hmisc,
- huxtable (>= 5.4.0),
- insight (>= 0.15.0),
- kableExtra (>= 1.3.4),
- lme4,
- mgcv,
- mice (>= 3.10.0),
- nnet,
- officer,
- openxlsx,
- parameters (>= 0.20.2),
- parsnip (>= 0.1.7),
- rmarkdown,
- sandwich (>= 3.0.1),
- scales,
- smd (>= 0.6.6),
- spelling (>= 2.2),
- survey,
- survival (>= 3.2-11),
- testthat (>= 3.0.4),
- tidycmprsk (>= 0.1.2),
- workflows (>= 0.2.4)

VignetteBuilder knitr
RdMacros lifecycle
Encoding UTF-8
Language en-US
LazyData true
Roxygen list(markdown = TRUE)
RoxygenNote 7.2.3
Config/testthat/edition 3
Config/testthat/parallel true
Config/Needs/website lubridate

R topics documented:

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add_ci

Description
Add a new column with the confidence intervals for proportions, means, etc.

Usage

```r
add_ci(x, ...)
```

### S3 method for class 'tbl_summary'

```r
add_ci(
  x,
  method = NULL,
  include = everything(),
  statistic = NULL,
  conf.level = 0.95,
  style_fun = NULL,
  pattern = NULL,
  df = NULL,
  ...
)
```

### S3 method for class 'tbl_svysummary'

```r
add_ci(
  x,
  method = NULL,
  include = everything(),
  statistic = NULL,
  conf.level = 0.95,
  style_fun = NULL,
  pattern = NULL,
  df = NULL,
  ...
)
```
Arguments

- **x**: A `tbl_summary` or a `tbl_svysummary` object
- **method**: Confidence interval method. Default is `list(all_categorical() ~ "wilson", all_continuous() ~ "t.test")` for `tbl_summary` objects and `list(all_categorical() ~ "svyprop", all_continuous() ~ "svymean")` for `tbl_svysummary` objects. See details below.
- **include**: variables to include in the summary table. Default is `everything()`.
- **statistic**: Formula indicating how the confidence interval will be displayed. Default is `list(all_categorical() ~ "{conf.low}%, {conf.high}%", all_continuous() ~ "{conf.low}, {conf.high}"))`.
- **conf.level**: Confidence level. Default is 0.95.
- **style_fun**: Function to style upper and lower bound of confidence interval. Default is `list(all_categorical() ~ purrr::partial(style_sigfig, scale = 100), all_continuous() ~ style_sigfig)`.
- **pattern**: string indicating the pattern to use to merge the CI with the statistics cell. The default is `NULL`, where no columns are merged. The two columns that will be merged are the statistics column, represented by "{stat}" and the CI column represented by "{ci}", e.g. pattern = "{stat} ({ci})" will merge the two columns with the CI in parentheses.
- **df**: For `tbl_svysummary()`, the number of degrees of freedom used to estimate confidence intervals. By default, will use `survey::degf()`.

Value

gtsummary table

**method argument**

- **for tbl_summary tables**: Must be one of `c("wilson", "wilson.no.correct", "exact", "asymptotic")` for categorical variables, and `c("t.test", "wilcox.test")` for continuous variables.

- **for tbl_svysummary tables**: Must be one of `c("svyprop", "svyprop.logit", "svyprop.likelihood", "svyprop.asin", "svyprop.beta", "svyprop.mean", "svyprop.xlogit")` for categorical variables, and `c("svymean", "svymedian", "svymedian.mean", "svymedian.beta", "svymedian.xlogit", "svymedian.asin", "svymedian.score")` for continuous variables.

Confidence intervals for proportions are computed with `survey::svyciprop()`. See the help file of this function for details on the different methods available to compute CIs. The default method "svyprop" is equivalent to "svyprop.logit", corresponding to a call to `survey::svyciprop()` with method = "logit".

Confidence intervals for means (method "svymean") are computed using `confint(svymean())`. 

```r
add_ci
```
Confidence intervals for medians are computed with `survey::svyquantile()`.
See the help file of this function for details on the different methods available to compute CIs. The default method "svymedian" is equivalent to "svymedian.mean", corresponding to a call to `survey::svyquantile()` with method = "mean".

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

```r
# Example 1 ----------------------------------
add_ci_ex1 <-
  trial \%\%%
  select(marker, response, trt) \%\%
  tbl_summary(
    statistic = all_continuous() ~ "{mean} \{{sd}\}",
    missing = "no",
    statistic = all_continuous() ~ "{mean} \{{sd}\}
  ) \%\%
  add_ci()

# Example 2 ----------------------------------
add_ci_ex2 <-
  trial \%\%
  select(response, grade) \%\%
  tbl_summary(
    statistic = all_categorical() ~ "{p}\%",
    missing = "no"
  ) \%\%
  add_ci(pattern = "{stat} \{{ci}\}") \%\%
  modify_footnote(everything() ~ NA)

# Example 3 ----------------------------------
data(api, package = "survey")
add_ci_ex3 <-
  survey::svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc) \%\%
  tbl_svysummary(
    include = c(api00, hsg, stype),
    statistic = hsg ~ "{mean} \{{sd}\}",
  ) \%\%
  add_ci(
    method = api00 ~ "svymedian"
  )
```

---

`add_difference`  
Add difference between groups
add_difference

Description
Add the difference between two groups (typically mean difference), along with the difference confidence interval and p-value.

Usage
add_difference(
  x, test = NULL, group = NULL, adj.vars = NULL, test.args = NULL, conf.level = 0.95, include = everything(), pvalue_fun = NULL, estimate_fun = NULL)

Arguments
x "tbl_summary" or "tbl_svysummary" object
test List of formulas specifying statistical tests to perform for each variable, e.g. list(all_continuous() ~ "t.test"). Common tests include "t.test" or "ancova" for continuous data, and "prop.test" for dichotomous variables. See tests for details and more tests.
group Column name (unquoted or quoted) of an ID or grouping variable. The column can be used to calculate p-values with correlated data. Default is NULL. See tests for methods that utilize the group= argument.
adj.vars Variables to include in mean difference adjustment (e.g. in ANCOVA models)
test.args List of formulas containing additional arguments to pass to tests that accept arguments. For example, add an argument for all t-tests, use test.args = all_tests("t.test") ~ list(var.equal = TRUE)
conf.level Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval.
include Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is everything().
pvalue_fun Function to round and format p-values. Default is style_pvalue. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. pvalue_fun = function(x) style_pvalue(x, digits = 2) or equivalently, purrr::partial(style_pvalue, digits = 2)).
estimate_fun List of formulas specifying the formatting functions to round and format differences. Default is list(all_continuous() ~ style_sigfig, all_categorical() ~ function(x) paste0(style_sigfig(x * 100), "%")) Function to round and format difference. Default is style_sigfig()

Example Output
See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

# Example 1 ----------------------------------
add_difference_ex1 <-
trial %>%
  select(trt, age, marker, response, death) %>%
tbl_summary(
  by = trt,
  statistic =
    list(
      all_continuous() ~ "\{mean\} ({sd})",
      all_dichotomous() ~ "(p)\%"
    ),
  missing = "no"
) %>%
add_n() %>%
add_difference()

# Example 2 ----------------------------------
# ANCOVA adjusted for grade and stage
add_difference_ex2 <-
trial %>%
  select(trt, age, marker, grade, stage) %>%
tbl_summary(
  by = trt,
  statistic = list(all_continuous() ~ "\{mean\} ({sd})"),
  missing = "no",
  include = c(age, marker, trt)
) %>%
add_n() %>%
add_difference(adj.vars = c(grade, stage))

add_glance

Add Model Statistics

Description

Add model statistics returned from broom::glance(). Statistics can either be appended to the table (add_glance_table()), or added as a table source note (add_glance_source_note()).

Usage

add_glance_table(
x,
  include = everything(),
  label = NULL,
  fmt_fun = NULL,
  glance_fun = NULL
)
```r
add_glance_source_note(x, include = everything(), label = NULL, fmt_fun = NULL, glance_fun = NULL, text_interpret = c("md", "html"), sep1 = " = ", sep2 = "; ")
```

### Arguments

**x**

'tbl_regression' object

**include**

list of statistics to include in output. Must be column names of the tibble returned by `broom::glance()`. The include argument can also be used to specify the order the statistics appear in the table.

**label**

List of formulas specifying statistic labels, e.g. `list(r.squared ~ "R²", p.value ~ "p")`

**fmt_fun**

List of formulas where the LHS is a statistic and the RHS is a function to format/round the statistics. The default is to round the number of observations and degrees of freedom to the nearest integer, p-values are styled with `style_pvalue()` and the remaining statistics are styled with `style_sigfig(x, digits = 3)`

**glance_fun**

function that returns model statistics. Default is `broom::glance()` for most model objects, and `broom::glance(mice::pool())` for MICE 'mira' models. Custom functions must return a single row tibble.

**text_interpret**

String indicates whether source note text will be interpreted with `gt::md()` or `gt::html()`. Must be "md" (default) or "html".

**sep1**

Separator between statistic name and statistic. Default is " = ", e.g. "R² = 0.456"

**sep2**

Separator between statistics. Default is "; ".

### Value

gtsummary table

### Default Labels

The following statistics have set default labels when printed. When there is no default, the column name from `broom::glance()` is printed.

<table>
<thead>
<tr>
<th>Statistic Name</th>
<th>Default Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>r.squared</td>
<td>R²</td>
</tr>
<tr>
<td>adj.r.squared</td>
<td>Adjusted R²</td>
</tr>
<tr>
<td>p.value</td>
<td>p-value</td>
</tr>
<tr>
<td>logLik</td>
<td>Log-likelihood</td>
</tr>
<tr>
<td>statistic</td>
<td>Statistic</td>
</tr>
<tr>
<td>df.residual</td>
<td>Residual df</td>
</tr>
<tr>
<td>null.deviance</td>
<td>Null deviance</td>
</tr>
</tbody>
</table>
Tips

When combining `add_glance_table()` with `tbl_merge()`, the ordering of the model terms and the glance statistics may become jumbled. To re-order the rows with glance statistics on bottom, use the script below:

```r
tbl_merge(list(tbl1, tbl2)) %>%
  modify_table_body(~.x %>% arrange(row_type == "glance_statistic"))
```

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

```r
mod <- lm(age ~ marker + grade, trial) %>% tbl_regression()

# Example 1 ----------------------------------
add_glance_ex1 <-
  mod %>%
  add_glance_table(
    label = list(sigma ~ "\U03C3"),
    include = c(r.squared, AIC, sigma)
  )

# Example 2 ----------------------------------
add_glance_ex2 <-
  mod %>%
  add_glance_source_note(
    label = list(sigma ~ "\U03C3"),
    include = c(r.squared, AIC, sigma)
  )
```
add_global_p

Add the global p-values

Description

This function uses car::Anova() to calculate global p-values for model covariates. Output from tbl_regression and tbl_uvregression objects supported.

Usage

add_global_p(x, ...)

## S3 method for class 'tbl_regression'
add_global_p(
x, 
include = everything(),
type = NULL,
keep = FALSE,
anova_fun = NULL,
quiet = NULL,
...
)

## S3 method for class 'tbl_uvregression'
add_global_p(
x,
type = NULL,
include = everything(),
keep = FALSE,
anova_fun = NULL,
quiet = NULL,
...
)

Arguments

x Object with class 'tbl_regression' or 'tbl_uvregression'

... Additional arguments to be passed to car::Anova, aod::wald.test() or anova_fun (if specified)

include Variables to calculate global p-value for. Input may be a vector of quoted or unquoted variable names. Default is everything()

type Type argument passed to car::Anova(type=). Default is "III"

keep Logical argument indicating whether to also retain the individual p-values in the table output for each level of the categorical variable. Default is FALSE.

anova_fun Function that will be used in place of car::Anova() when specified to calculate the global p-values.

  • function must return a tibble matching the output of car::Anova() %>% broom::tidy() including a columns called "term" and "p.values"
**add_n.tbl_summary**

- function must accept arguments `anova_fun(x, ...)`, where `x` is a model object
- arguments passed in `...` will be passed to `anova_fun(...)`
- the `add_global_p(type=)` argument is ignored in `anova_fun=`
- a common function used here is `tidy_wald_test()`, a wrapper for `aod::wald.test()`

**quiet**

Logical indicating whether to print messages in console. Default is `FALSE`

**Example Output**

**Author(s)**

Daniel D. Sjoberg

**See Also**

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_uvregression tools: `add_q()`, `bold_italicize_labels_levels`, `inline_text.tbl_uvregression()`, `modify.tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_uvregression()`

Other tbl_regression tools: `add_q()`, `bold_italicize_labels_levels`, `combine_terms()`, `inline_text.tbl_regression()`, `modify.tbl_merge()`, `tbl_regression()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`

**Examples**

```r
# Example 1 ----------------------------------
tbl_lm_global_ex1 <-
  lm(marker ~ age + grade, trial) %>%
  tbl_regression() %>%
  add_global_p()

# Example 2 ----------------------------------
tbl_uv_global_ex2 <-
  trial[c("response", "trt", "age", "grade") ] %>%
  tbl_uvregression(
    method = glm,
    y = response,
    method.args = list(family = binomial),
    exponentiate = TRUE
  ) %>%
  add_global_p()
```

---

**add_n.tbl_summary**

*Add column with N*

**Description**

For each variable in a `tbl_summary` table, the `add_n` function adds a column with the total number of non-missing (or missing) observations
add_n.tbl_summary

Usage

## S3 method for class 'tbl_summary'
add_n(
    x,
    statistic = '{n}',
    col_label = '**N**',
    footnote = FALSE,
    last = FALSE,
    ...
)

## S3 method for class 'tbl_svysummary'
add_n(
    x,
    statistic = '{n}',
    col_label = '**N**',
    footnote = FALSE,
    last = FALSE,
    ...
)

Arguments

x Object with class tbl_summary from the tbl_summary function or with class tbl_svysummary from the tbl_svysummary function

statistic String indicating the statistic to report. Default is the number of non-missing observation for each variable, statistic = '{n}'. Other statistics available to report include:

- '{N_obs}' total number of observations,
- '{N_nonmiss}' number of non-missing observations,
- '{N_miss}' number of missing observations,
- '{p_nonmiss}' percent non-missing data,
- '{p_miss}' percent missing data

survey summaries also have the following unweighted statistics available:

- 'N_obs_unweighted', 'N_miss_unweighted', 'N_nonmiss_unweighted',
- 'p_miss_unweighted', 'p_nonmiss_unweighted'

The argument uses glue::glue syntax and multiple statistics may be reported, e.g. statistic = '{N_nonmiss} / {N.obs} ({p_nonmiss}%')

col_label String indicating the column label. Default is '**N**'

footnote Logical argument indicating whether to print a footnote clarifying the statistics presented. Default is FALSE

last Logical indicator to include N column last in table. Default is FALSE, which will display N column first.

... Not used

Value

A tbl_summary or tbl_svysummary object
Example Output

Author(s)
Daniel D. Sjoberg

See Also
Review list, formula, and selector syntax used throughout gtsummary

Other tbl_summary tools: add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(),
bold_italicize_labels_levels, inline_text, tbl_summary(), inline_text.tbl_survfit(),
modify, separate_p_footnotes(), tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(),
tbl_strata(), tbl_summary()

Other tbl_svysummary tools: add_overall(), add_p.tbl_svysummary(), add_q(), add_stat_label(),
modify, separate_p_footnotes(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(),
tbl_svysummary()

Examples

```r
# Example 1 ----------------------------------
tbl_n_ex <-
  trial[c("trt", "age", "grade", "response")] %>%
tbl_summary(by = trt) %>%
  add_n()
```

Description

[Maturing] For each survfit() object summarized with tbl_survfit() this function will add
the total number of observations in a new column.

Usage

```r
## S3 method for class 'tbl_survfit'
add_n(x, ...)
```

Arguments

- `x` object of class "tbl_survfit"
- `...` Not used

Example Output
add_nevent.tbl_survfit

See Also

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_survfit tools: add_nevent.tbl_survfit(), add_p.tbl_survfit(), modify.tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_survfit()

Examples

library(survival)
fit1 <- survfit(Surv(ttdeath, death) ~ 1, trial)
fit2 <- survfit(Surv(ttdeath, death) ~ trt, trial)

# Example 1 ----------------------------------
add_n.tbl_survfit_ex1 <-
  list(fit1, fit2) %>%
  tbl_survfit(times = c(12, 24)) %>%
  add_n()

Description

[Maturing] For each survfit() object summarized with tbl_survfit() this function will add the total number of events observed in a new column.

Usage

```r
## S3 method for class 'tbl_survfit'
add_nevent(x, ...)
```

Arguments

- `x`: object of class 'tbl_survfit'
- `...`: Not used

Example Output

See Also

Other tbl_survfit tools: add_n.tbl_survfit(), add_p.tbl_survfit(), modify, tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_survfit()
Examples

library(survival)
fit1 <- survfit(Surv(ttdeath, death) ~ 1, trial)
fit2 <- survfit(Surv(ttdeath, death) ~ trt, trial)

# Example 1 ----------------------------------
add_nevent.tbl_survfit_ex1 <-
  list(fit1, fit2) %>%
  tbl_survfit(times = c(12, 24)) %>%
  add_n() %>%
  add_nevent()

---

**add_nevent_regression**  
*Add event N to regression table*

Description

Add event N to regression table

Usage

```r
## S3 method for class 'tbl_regression'
add_nevent(x, location = NULL, ...)

## S3 method for class 'tbl_uvregression'
add_nevent(x, location = NULL, ...)
```

Arguments

- `x`  
a tbl_regression or tbl_uvregression table
- `location`  
location to place Ns. When "label" total Ns are placed on each variable’s label row. When "level" level counts are placed on the variable level for categorical variables, and total N on the variable’s label row for continuous.
- `...`  
Not used

Example Output

Examples

```r
# Example 1 ----------------------------------
add_nevent.tbl_regression_ex1 <-
  trial %>%
  select(response, trt, grade) %>%
  tbl_uvregression(
    y = response,
    method = glm,
    method.args = list(family = binomial),
```
# Example 2 ----------------------------------
add_nevent.tbl_regression_ex2 <- 
  glm(response ~ age + grade, trial, family = binomial) %>%
  tbl_regression(exponentiate = TRUE) %>%
  add_nevent(location = "level")

### Description
Add N to regression table

### Usage
```r
## S3 method for class 'tbl_regression'
add_n(x, location = NULL, ...)
```

```r
## S3 method for class 'tbl_uvregression'
add_n(x, location = NULL, ...)
```

### Arguments
- `x`: a tbl_regression or tbl_uvregression table
- `location`: location to place Ns. When "label" total Ns are placed on each variable’s label row. When "level" level counts are placed on the variable level for categorical variables, and total N on the variable’s label row for continuous.
- `...`: Not used

### Example Output

### Examples
```r
# Example 1 ----------------------------------
add_n.tbl_regression_ex1 <- 
  trial %>%
  select(response, age, grade) %>%
  tbl_uvregression(
    y = response,
    method = glm,
    method.args = list(family = binomial),
    hide_n = TRUE
  ) %>%
  add_n(location = "label")

# Example 2 ----------------------------------
add_n.tbl_regression_ex2 <- 
  glm(response ~ age + grade, trial, family = binomial) %>%
```
add_overall

```r
tbl_regression(exponentiate = TRUE) %>%
add_n(location = "level")
```

---

**add_overall**

**Add column with overall summary statistics**

**Description**

Adds a column with overall summary statistics to tables created by `tbl_summary`, `tbl_svysummary`, `tbl_continuous` or `tbl_custom_summary`.

**Usage**

```r
add_overall(x, ...)

## S3 method for class 'tbl_summary'
add_overall(
  x,
  last = FALSE,
  col_label = NULL,
  statistic = NULL,
  digits = NULL,
  ...
)

## S3 method for class 'tbl_svysummary'
add_overall(
  x,
  last = FALSE,
  col_label = NULL,
  statistic = NULL,
  digits = NULL,
  ...
)

## S3 method for class 'tbl_continuous'
add_overall(
  x,
  last = FALSE,
  col_label = NULL,
  statistic = NULL,
  digits = NULL,
  ...
)

## S3 method for class 'tbl_custom_summary'
add_overall(
  x,
  last = FALSE,
  col_label = NULL,
```
add_overall

```r
statistic = NULL,
digits = NULL,
...
)
```

**Arguments**

- `x` Object with class `tbl_summary` from the `tbl_summary` function, object with class `tbl_svysummary` from the `tbl_svysummary` function, object with class `tbl_continuous` from the `tbl_continuous` function or object with class `tbl_custom_summary` from the `tbl_custom_summary` function.
- `...` Not used
- `last` Logical indicator to display overall column last in table. Default is `FALSE`, which will display overall column first.
- `col_label` String indicating the column label. Default is "**Overall**, N = {N}"
- `statistic` Override the statistic argument in initial `tbl_*` function call. Default is `NULL`.
- `digits` Override the digits argument in initial `tbl_*` function call. Default is `NULL`.

**Value**

A `tbl_*` of same class as `x`

**Example Output**

**Author(s)**

Daniel D. Sjoberg

**See Also**

Other `tbl_summary` tools: `add_n.tbl_summary()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold.Italicize_labels_levels()`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Other `tbl_svysummary` tools: `add_n.tbl_summary()`, `add_p.tbl_svysummary()`, `add_q()`, `add_stat_label()`, `modify`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_stack()`, `tbl_strata()`, `tbl_svysummary()`

Other `tbl_continuous` tools: `add_p.tbl_continuous()`, `tbl_continuous()`

Other `tbl_custom_summary` tools: `continuous_summary()`, `proportion_summary()`, `ratio_summary()`, `tbl_custom_summary()`

**Examples**

```r
# Example 1 ----------------------------------
tbl_overall_ex1 <-
trial %>%
tbl_summary(include = c(age, grade), by = trt) %>%
add_overall()

# Example 2 ----------------------------------
```
tbl_overall_ex2 <-
  trial %>%
  tbl_summary(
    include = grade,
    by = trt,
    percent = "row",
    statistic = ""(p)"
  ) %>%
  add_overall(
    last = TRUE,
    statistic = "{p}% (n={n})",
    digits = c(1, 0)
  )

# Example 3 ----------------------------------

# Example 3 ----------------------------------

 tbl_overall_ex3 <-
  trial %>%
  tbl_continuous(
    variable = age,
    by = trt,
    include = grade
  ) %>%
  add_overall(last = TRUE)

===============================================

add_p.tbl_continuous P-values for tbl_continuous

Description

P-values for tbl_continuous

Usage

## S3 method for class 'tbl_continuous'
add_p(
  x,
  test = NULL,
pvalue_fun = NULL,
  include = everything(),
  test.args = NULL,
group = NULL,
  ...
)

Arguments

  x Object with class tbl_summary from the tbl_summary function

  test List of formulas specifying statistical tests to perform for each variable. Default is two-way ANOVA when by= is not NULL, and has the same defaults as add_p.tbl_continuous() when by = NULL. See tests for details, more tests, and instruction for implementing a custom test.
Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. `pvalue_fun = function(x) style_pvalue(x, digits = 2)` or equivalently, `purrr::partial(style_pvalue, digits = 2)`).

Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is `everything()`.

List of formulas containing additional arguments to pass to tests that accept arguments. For example, add an argument for all t-tests, use `test.args = all_tests("t.test") ~ list(var.equal = TRUE)`.

Column name (unquoted or quoted) of an ID or grouping variable. The column can be used to calculate p-values with correlated data. Default is `NULL`. See `tests` for methods that utilize the `group=` argument.

Not used

See Also

Other tbl_continuous tools: `add_overall()`, `tbl_continuous()`

Examples

```r
ddf_p_continuous_ex1 <-
tbl_continuous(
data = trial,
variable = age,
by = trt,
include = grade
) %>%
add_p()
```

Description

Calculate and add a p-value comparing the two variables in the cross table. Missing values are included in p-value calculations.

Usage

```r
## S3 method for class 'tbl_cross'
add_p(
  x,
  test = NULL,
pvalue_fun = NULL,
  source_note = NULL,
  test.args = NULL,
  ...)
```
Arguments

- **x**: Object with class `tbl_cross` from the `tbl_cross` function.
- **test**: A string specifying statistical test to perform. Default is "chisq.test" when expected cell counts >=5 and "fisher.test" when expected cell counts <5.
- **pvalue_fun**: Function to round and format p-value. Default is `style_pvalue`, except when `source_note = TRUE` when the default is `style_pvalue(x, prepend_p = TRUE)`.
- **source_note**: Logical value indicating whether to show p-value in the `{gt}` table source notes rather than a column.
- **test.args**: Named list containing additional arguments to pass to the test (if it accepts additional arguments). For example, add an argument for a chi-squared test with `test.args = list(correct = TRUE)`.

Examples

```r
# Example 1 ----------------------------------
add_p_cross_ex1 <-
  trial %>%
  tbl_cross(row = stage, col = trt) %>%
  add_p()

# Example 2 ----------------------------------
add_p_cross_ex2 <-
  trial %>%
  tbl_cross(row = stage, col = trt) %>%
  add_p(source_note = TRUE)
```

Author(s)

Karissa Whiting

See Also

Other `tbl_cross` tools: `inline_text.tbl_cross()`, `tbl_cross()`

Examples

```
# Example 1 ----------------------------------
add_p_cross_ex1 <-
  trial %>%
  tbl_cross(row = stage, col = trt) %>%
  add_p()

# Example 2 ----------------------------------
add_p_cross_ex2 <-
  trial %>%
  tbl_cross(row = stage, col = trt) %>%
  add_p(source_note = TRUE)
```

Description

Adds p-values to tables created by `tbl_summary` by comparing values across groups.
## Usage

```r
## S3 method for class 'tbl_summary'
add_p(
x,  
test = NULL,
pvalue_fun = NULL,
group = NULL,
include = everything(),
test.args = NULL,
...
)
```

### Arguments

- **x**: Object with class `tbl_summary` from the `tbl_summary` function.
- **test**: List of formulas specifying statistical tests to perform for each variable, e.g.,
  ```r
  list(all_continuous() ~ "t.test", all_categorical() ~ "fisher.test")
  ```
  Common tests include "t.test", "aov", "wilcox.test", "kruskal.test", "chisq.test", "fisher.test", and "lme4" (for clustered data). See `tests` for details, more tests, and instruction for implementing a custom test.
  Tests default to "kruskal.test" for continuous variables ("wilcox.test" when "by" variable has two levels), "chisq.test.no.correct" for categorical variables with all expected cell counts >=5, and "fisher.test" for categorical variables with any expected cell count <5.
- **pvalue_fun**: Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g.,
  ```r
  pvalue_fun = function(x) style_pvalue(x, digits = 2)
  ```
  or equivalently,
  ```r
  purrr::partial(style_pvalue, digits = 2)
  ```
- **group**: Column name (unquoted or quoted) of an ID or grouping variable. The column can be used to calculate p-values with correlated data. Default is `NULL`. See `tests` for methods that utilize the `group=` argument.
- **include**: Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is `everything()`.
- **test.args**: List of formulas containing additional arguments to pass to tests that accept arguments. For example, add an argument for all t-tests, use `test.args = all_tests("t.test") ~ list(var.equal = TRUE)`.
- **...**: Not used

### Value

A `tbl_summary` object

### Example Output

### Author(s)

Daniel D. Sjoberg, Emily C. Zabor
See Also

See tbl_summary vignette for detailed examples

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_q(), add_stat_label(), bold_italicize_labels_levels, inline_text.tbl_summary(), inline_text.tbl_survfit(), modify, separate_p_footnotes(), tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_summary()

Examples

# Example 1 ----------------------------------
add_p_ex1 <-
  trial[c("age", "grade", "trt")]
  tbl_summary(by = trt)
  add_p()

# Example 2 ----------------------------------
add_p_ex2 <-
  trial
  select(trt, age, marker)
  tbl_summary(by = trt, missing = "no")
  add_p(
    # perform t-test for all variables
    test = everything() ~ "t.test",
    # assume equal variance in the t-test
    test.args = all_tests("t.test") ~ list(var.equal = TRUE)
  )

add_p.tbl_survfit

Adds p-value to survfit table

Description

[Maturing] Calculate and add a p-value

Usage

## S3 method for class 'tbl_survfit'
add_p(
  x,
  test = "logrank",
  test.args = NULL,
  pvalue_fun = style_pvalue,
  include = everything(),
  quiet = NULL,
  ...
)
Arguments

- **x**: Object of class "tbl_survfit"
- **test**: String indicating test to use. Must be one of "logrank", "tarone", "survdiff", "petopeto_gehanwilcoxon", "coxph_lrt", "coxph_wald", "coxph_score". See details below
- **test.args**: List of formulas containing additional arguments to pass to tests that accept arguments. For example, add an argument for all t-tests, use `test.args = all_tests("t.test") ~ list(var.equal = TRUE)`
- **pvalue_fun**: Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. `pvalue_fun = function(x) style_pvalue(x, digits = 2)` or equivalently, `purrr::partial(style_pvalue, digits = 2)`).
- **include**: Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is `everything()`.
- **quiet**: Logical indicating whether to print messages in console. Default is `FALSE`
- **...**: Not used

**test argument**

The most common way to specify `test=` is by using a single string indicating the test name. However, if you need to specify different tests within the same table, the input in flexible using the list notation common throughout the gtsummary package. For example, the following code would call the log-rank test, and a second test of the G-rho family.

```r
... %>%
  add_p(test = list(trt ~ "logrank", grade ~ "survdiff"),
    test.args = grade ~ list(rho = 0.5))
```

**Example Output**

**See Also**

Other tbl_survfit tools: `add_n.tbl_survfit()`, `add_nevent.tbl_survfit()`, `modify.tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_survfit()`

**Examples**

```r
library(survival)

gts_survfit <-
  list(
    survfit(Surv(ttdeath, death) ~ grade, trial),
    survfit(Surv(ttdeath, death) ~ trt, trial)
  ) %>%
  tbl_survfit(times = c(12, 24))

# Example 1 ----------------------------------
add_p_tbl_survfit_ex1 <-
```
add_p.tbl_svysummary

# Example 2 ----------------------------------
# Pass "rho=" argument to "survdiff()"
gts_survfit %>%
  add_p()

add_p.tbl_survfit_ex2 <-
gts_survfit %>%
  add_p(test = "survdiff", test.args = list(rho = 0.5))

add_p.tbl_svysummary  Adds p-values to svysummary tables

Description

Adds p-values to tables created by tbl_svysummary by comparing values across groups.

Usage

## S3 method for class 'tbl_svysummary'
add_p(
  x,
  test = NULL,
  pvalue_fun = NULL,
  include = everything(),
  test.args = NULL,
  ...
)

Arguments

x  Object with class tbl_svysummary from the tbl_svysummary function
test List of formulas specifying statistical tests to perform, e.g. list(all_continuous() ~ "svy.t.test", all_categorical() ~ "svy.wald.test"). Options include
  • "svy.t.test" for a t-test adapted to complex survey samples (cf. survey::svytest),
  • "svy.wilcox.test" for a Wilcoxon rank-sum test for complex survey samples (cf. survey::svyranktest),
  • "svy.kruskal.test" for a Kruskal-Wallis rank-sum test for complex survey samples (cf. survey::svyranktest),
  • "svy.vanderwaerden.test" for a van der Waerden's normal-scores test for complex survey samples (cf. survey::svyranktest),
  • "svy.median.test" for a Mood's test for the median for complex survey samples (cf. survey::svyranktest),
  • "svy.chisq.test" for a Chi-squared test with Rao & Scott's second-order correction (cf. survey::svychisq),
  • "svy.adj.chisq.test" for a Chi-squared test adjusted by a design effect estimate (cf. survey::svychisq),
  • "svy.wald.test" for a Wald test of independence for complex survey samples (cf. survey::svychisq),
• "svy_adj.wald.test" for an adjusted Wald test of independence for complex survey samples (cf. survey::svychisq),
• "svy.lincom.test" for a test of independence using the exact asymptotic distribution for complex survey samples (cf. survey::svychisq),
• "svy.saddlepoint.test" for a test of independence using a saddlepoint approximation for complex survey samples (cf. survey::svychisq).
Tests default to "svy.wilcox.test" for continuous variables and "svy.chisq.test" for categorical variables.

pvalue_fun Function to round and format p-values. Default is style_pvalue. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. pvalue_fun = function(x) style_pvalue(x, digits = 2) or equivalently, purrr::partial(style_pvalue, digits = 2)).

include Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is everything().

test.args List of formulas containing additional arguments to pass to tests that accept arguments. For example, add an argument for all t-tests, use test.args = all_tests("t.test") ~ list(var.equal = TRUE)

... Not used

Value
A tbl_svysummary object

Example Output

Author(s)
Joseph Larmarange

See Also
Other tbl_svysummary tools: add_n.tbl_summary(), add_overall(), add_q(), add_stat_label(), modify, separate_p_footnotes(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_svysummary()

Examples

# Example 1 ----------------------------------
# A simple weighted dataset
add_p_svysummary_ex1 <-
survey::svydesign(~1, data = as.data.frame(Titanic), weights = ~Freq) %>%
tbl_svysummary(by = Survived, include = c(Sex, Age)) %>%
  add_p()

# A dataset with a complex design
data(api, package = "survey")
d_clust <- survey::svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc)

# Example 2 ----------------------------------
add_p_svysummary_ex2 <-
  tbl_svysummary(d_clust, by = both, include = c(api00, api99, both)) %>%
  add_p()

# Example 3 ----------------------------------
# change tests to svy t-test and Wald test
add_p_svysummary_ex3 <-
  tbl_svysummary(d_clust, by = both, include = c(cname, api00, api99, both)) %>%
  add_p(
    test = list(
      all_continuous() ~ "svy.t.test",
      all_categorical() ~ "svy.wald.test"
    )
  )

---------------------

add_q

Add a column of q-values to account for multiple comparisons

Description

Adjustments to p-values are performed with stats::p.adjust.

Usage

add_q(x, method = "fdr", pvalue_fun = NULL, quiet = NULL)

Arguments

x a gtsummary object

method String indicating method to be used for p-value adjustment. Methods from stats::p.adjust are accepted. Default is method = "fdr".

pvalue_fun Function to round and format p-values. Default is style_pvalue. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. pvalue_fun = function(x) style_pvalue(x, digits = 2) or equivalently, purrr::partial(style_pvalue, digits = 2)).

quiet Logical indicating whether to print messages in console. Default is FALSE

Example Output

Author(s)

Esther Drill, Daniel D. Sjoberg
**add_significance_stars**

**Description**

*Experimental* Add significance stars to estimates with small p-values

**Usage**

```
add_significance_stars(
  x,
  pattern = NULL,
)```

**See Also**

Other **tbl_summary** tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_stat_label()`, `bold_italicize_labels_levels()`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify()`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`.

Other **tbl_svysummary** tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_stat_label()`, `modify()`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_split()`, `tbl_strata()`, `tbl_svysummary()`.

Other **tbl_regression** tools: `add_global_p()`, `bold_italicize_labels_levels()`, `combine_terms()`, `inline_text.tbl_regression()`, `modify()`, `tbl_merge()`, `tbl_regression()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`.

Other **tbl_uvregression** tools: `add_global_p()`, `bold_italicize_labels_levels()`, `inline_text.tbl_uvregression()`, `modify()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_uvregression()`.

**Examples**

```
# Example 1 ----------------------------------
add_q_ex1 <-
  trial[c("trt", "age", "grade", "response")]
  %>%
  tbl_summary(by = trt) %>%
  add_p() %>%
  add_q()

# Example 2 ----------------------------------
add_q_ex2 <-
  trial[c("trt", "age", "grade", "response")]
  %>%
  tbl_uvregression(
    y = response,
    method = glm,
    method.args = list(family = binomial),
    exponentiate = TRUE
  ) %>%
  add_global_p() %>%
  add_q()
```
thresholds = c(0.001, 0.01, 0.05),
hide_ci = TRUE,
hide_p = inherits(x, c("tbl_regression", "tbl_uvregression")),
hide_se = FALSE
)

Arguments

x  a 'gtsummary' object with a 'p.value' column
pattern  glue-syntax string indicating what to display in formatted column. Default is "\{estimate\}\{stars\}" for regression summaries and "\{p.value\}\{stars\}" otherwise. A footnote is placed on the first column listed in the pattern. Other common patterns are "\{estimate\}\{stars\}\{\{conf.low\}, \{conf.high\}\}\{stars\}" and "\{estimate\} (\{conf.low\} to \{conf.high\})\{stars\}" thresholds  thresholds for significance stars. Default is c(0.001, 0.01, 0.05)
hide_ci  logical whether to hide confidence interval. Default is TRUE
hide_p  logical whether to hide p-value. Default is TRUE for regression summaries, and FALSE otherwise.
hide_se  logical whether to hide standard error. Default is FALSE

Future Updates

There are planned updates to the implementation of this function with respect to the pattern= argument. Currently, this function replaces the numeric estimate column, with a formatted character column following pattern=. Once gt::cols_merge() gains the rows= argument the implementation will be updated to use it, which will keep numeric columns numeric. For the vast majority of users, the planned change will be go unnoticed.

Example Output

Examples

```r
tbl <-
  lm(time ~ ph.ecog + sex, survival::lung) %>%
  tbl_regression(label = list(ph.ecog = "ECOG Score", sex = "Sex"))

# Example 1 ------------------------------
add_significance_stars_ex1 <-
  tbl %>%
  add_significance_stars(hide_ci = FALSE, hide_p = FALSE)

# Example 2 ------------------------------
add_significance_stars_ex2 <-
  tbl %>%
  add_significance_stars(
    pattern = "{estimate}\%(\{conf.low\}, \{conf.high\}){stars}"%,
    hide_ci = TRUE, hide_se = TRUE
  ) %>%
  modify_header(estimate ~ "**Beta (95\% CI)**") %>%
  modify_footnote(estimate ~ "CI = Confidence Interval", abbreviation = TRUE)
```
add_stat

# Example 3 ----------------------------------
# Use ' \n ' to put a line break between beta and SE
add_significance_stars_ex3 <-
tbl %>%
  add_significance_stars(
    hide_se = TRUE,
    pattern = "{estimate}{stars} \n({std.error})"
  ) %>%
  modify_header(estimate ~ "**Beta \n(SE)**") %>%
  modify_footnote(estimate ~ "SE = Standard Error", abbreviation = TRUE) %>%
  as_gt() %>%
  gt::fmt_markdown(columns = everything()) %>%
  gt::tab_style(
    style = "vertical-align:top",
    locations = gt::cells_body(columns = label)
  )

# Example 4 ----------------------------------
add_significance_stars_ex4 <-
lm(marker ~ stage + grade, data = trial) %>%
tbl_regression() %>%
  add_global_p() %>%
  add_significance_stars(
    hide_p = FALSE,
    pattern = "{p.value}{stars}"
  ) %>%
  as_gt() %>%
  gt::tab_style(
    style = "vertical-align:top",
    locations = gt::cells_body(columns = label)
  )

---

add_stat  
_Add a custom statistic column_

**Description**

*[Maturing]* The function allows a user to add a new column (or columns) of statistics to an existing `tbl_summary`, `tbl_svysummary`, or `tbl_continuous` object.

**Usage**

`add_stat(x, fns, location = NULL, ...)`

**Arguments**

- `x`  
  `tbl_summary`, `tbl_svysummary`, or `tbl_continuous` object

- `fns`  
  list of formulas indicating the functions that create the statistic. See details below.
location  list of formulas indicating the location the new statistics are placed. The RHS of the formula must be one of c("label", "level", "missing"). When "label", a single statistic is placed on the variable label row. When "level" the statistics are placed on the variable level rows. The length of the vector of statistics returned from the fns function must match the dimension of levels. Default is to place the new statistics on the label row.

DEPRECATED

Details

The returns from custom functions passed in fns= are required to follow a specified format. Each of these function will execute on a single variable.

1. Each function must return a tibble or a vector. If a vector is returned, it will be converted to a tibble with one column and number of rows equal to the length of the vector.
2. When location = "label", the returned statistic from the custom function must be a tibble with one row. When location = "level" the tibble must have the same number of rows as there are levels in the variable (excluding the row for unknown values).
3. Each function may take the following arguments: foo(data, variable, by, tbl, ...)  
   • data= is the input data frame passed to tbl_summary()  
   • variable= is a string indicating the variable to perform the calculation on. This is the variable in the label column of the table.  
   • by= is a string indicating the by variable from tbl_summary=, if present  
   • tbl= the original tbl_summary()/tbl_svysummary() object is also available to utilize

The user-defined does not need to utilize each of these inputs. It’s encouraged the user-defined function accept ... as each of the arguments will be passed to the function, even if not all inputs are utilized by the user's function, e.g. foo(data, variable, by, ...)  
   • Use modify_header() to update the column headers  
   • Use modify_fmt_fun() to update the functions that format the statistics  
   • Use modify_footnote() to add a explanatory footnote

If you return a tibble with column names p.value or q.value, default p-value formatting will be applied, and you may take advantage of subsequent p-value formatting functions, such as bold_p() or add_q().

To access the continuous variable in a tbl_continuous() table, use tbl$inputs$variable.

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

library(dplyr, warn.conflicts = FALSE)  
library(stringr)  
# Example 1 ----------------------------------  
# fn returns t-test pvalue  
my_ttest <- function(data, variable, by, ...) {
```r
add_stat

t.test(data[[variable]] ~ as.factor(data[[by]]))$p.value
}

add_stat_ex1 <-
trial %>%
select(trt, age, marker) %>%
tbl_summary(by = trt, missing = "no") %>%
add_stat(fns = everything() ~ my_ttest) %>%
modify_header(
  list(  
    add_stat_1 ~ "**p-value**",  
    all_stat_cols() ~ "**{level}**"
  )
)

# Example 2 ----------------------------------
# fn returns t-test test statistic and p-value
my_ttest2 <- function(data, variable, by, ...)
  t.test(data[[variable]] ~ as.factor(data[[by]])) %>%
  broom::tidy() %>%
  mutate(
    stat = str_glue("t={style_sigfig(statistic)}, {style_pvalue(p.value, prepend_p = TRUE)}")
  ) %>%
  pull(stat)

add_stat_ex2 <-
trial %>%
select(trt, age, marker) %>%
tbl_summary(by = trt, missing = "no") %>%
add_stat(fns = everything() ~ my_ttest2) %>%
modify_header(add_stat_1 ~ "**Treatment Comparison**")

# Example 3 ----------------------------------
# return test statistic and p-value is separate columns
my_ttest3 <- function(data, variable, by, ...)
  t.test(data[[variable]] ~ as.factor(data[[by]])) %>%
  broom::tidy() %>%
  select(statistic, p.value)

add_stat_ex3 <-
trial %>%
select(trt, age, marker) %>%
tbl_summary(by = trt, missing = "no") %>%
add_stat(fns = everything() ~ my_ttest3) %>%
modify_header(
  list(  
    statistic ~ "**t-statistic**",  
    p.value ~ "**p-value**"
  )
) %>%
modify_fmt_fun(
  list(  
    statistic ~ style_sigfig,  
    p.value ~ style_pvalue
  )
)
```

add_stat_label

Description
Adds labels describing the summary statistics presented for each variable in the tbl_summary / tbl_svysummary table.

Usage
add_stat_label(x, location = NULL, label = NULL)

Arguments
x Object with class tbl_summary from the tbl_summary function or with class tbl_svysummary from the tbl_svysummary function
location location where statistic label will be included. "row" (the default) to add the statistic label to the variable label row, and "column" adds a column with the statistic label.
label a list of formulas or a single formula updating the statistic label, e.g. label = all_categorical() ~ "No. (%)"

Value
A tbl_summary or tbl_svysummary object

Tips
When using add_stat_label(location='row') with subsequent tbl_merge(), it's important to have somewhat of an understanding of the underlying structure of the gtsummary table. add_stat_label(location='row') works by adding a new column called "stat_label" to x$table_body. The "label" and "stat_label" columns are merged when the gtsummary table is printed. The tbl_merge() function merges on the "label" column (among others), which is typically the first column you see in a gtsummary table. Therefore, when you want to merge a table that has run add_stat_label(location='row') you need to match the "label" column values before the "stat_column" is merged with it.

For example, the following two tables merge properly

```
tbl1 <- trial %>% select(age, grade) %>% tbl_summary() %>% add_stat_label()
tbl2 <- lm(marker ~ age + grade, trial) %>% tbl_regression()
tbl_merge(list(tbl1, tbl2))
```

Example Output

Author(s)
Daniel D. Sjoberg
add_vif

See Also

- Review list, formula, and selector syntax used throughout gtsummary
- Other tbl_summary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `bold_italicize_labels_levels()`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify()`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`
- Other tbl_svysummary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_q()`, `modify()`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()` `tbl_strata()`, `tbl_summary()`

Examples

```r
tbl <- trial %>%
    dplyr::select(trt, age, grade, response) %>%
    tbl_summary(by = trt)

# Example 1 ----------------------------------
# Add statistic presented to the variable label row
add_stat_label_ex1 <-
    tbl %>%
    add_stat_label(
        # update default statistic label for continuous variables
        label = all_continuous() ~ "med. (iqr)"
    )

# Example 2 ----------------------------------
add_stat_label_ex2 <-
    tbl %>%
    add_stat_label(
        # add a new column with statistic labels
        location = "column"
    )

# Example 3 ----------------------------------
add_stat_label_ex3 <-
    trial %>%
    select(age, grade, trt) %>%
    tbl_summary(
        by = trt,
        type = all_continuous() ~ "continuous2",
        statistic = all_continuous() ~ c("{mean} ({sd})", "{min} - {max}"),
    ) %>%
    add_stat_label(label = age ~ c("Mean (SD)", "Min - Max")
```

---

**Description**

[Maturing] Add the variance inflation factor (VIF) or generalized VIF (GVIF) to the regression table. Function uses `car::vif()` to calculate the VIF.
Usage

```
add_vif(x, statistic = NULL, estimate_fun = NULL)
```

Arguments

- `x`: `tbl_regression` object
- `statistic`: "VIF" (variance inflation factors, for models with no categorical terms) or one of combination of "GVIF" (generalized variance inflation factors), "aGVIF" adjusted GVIF, i.e. $GVIF^{1/(2*df)}$ and/or "df" (degrees of freedom). See `car::vif()` for details.
- `estimate_fun`: Default is `style_sigfig()`.

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

```
# Example 1 ----------------------------------
add_vif_ex1 <-
  lm(age ~ grade + marker, trial) %>%
  tbl_regression() %>%
  add_vif()

# Example 2 ----------------------------------
add_vif_ex2 <-
  lm(age ~ grade + marker, trial) %>%
  tbl_regression() %>%
  add_vif(c("aGVIF", "df"))
```

---

### as_flex_table

Convert `gtsummary` object to a `flextable` object

Description

Function converts a `gtsummary` object to a `flextable` object. A user can use this function if they wish to add customized formatting available via the `flextable` functions. The `flextable` output is particularly useful when combined with R markdown with Word output, since the `gt` package does not support Word.

Usage

```
as_flex_table(x, include = everything(), return_calls = FALSE, ...)
```
Arguments

- **x**: Object created by a function from the gtsummary package (e.g. `tbl_summary` or `tbl_regression`)
- **include**: Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is `everything()`.
- **return_calls**: Logical. Default is `FALSE`. If `TRUE`, the calls are returned as a list of expressions.
- **...**: Not used

Details

The `as_flex_table()` function supports bold and italic markdown syntax in column headers and spanning headers (`**` and `_` only). Text wrapped in double stars (`**bold**`) will be made bold, and text between single underscores (`_italic_`) will be made italic. No other markdown syntax is supported and the double-star and underscore cannot be combined. To further style your table, you may convert the table to flextable with `as_flex_table()`, then utilize any of the flextable functions.

Value

A flextable object

Example Output

Author(s)

Daniel D. Sjoberg

See Also

Other gtsummary output types: `as_gt()`, `as_hux_table()`, `as_kable_extra()`, `as_kable()`, `as_tibble.gtsummary()`

Examples

```r
as_flex_table_ex1 <-
trial %>%
  select(trt, age, grade) %>%
  tbl_summary(by = trt) %>%
  add_p() %>%
  as_flex_table()
```
Description

Function converts a gtsummary object to a "gt_tbl" object, that is, a table created with gt::gt(). Function is used in the background when the results are printed or knit. A user can use this function if they wish to add customized formatting available via the gt package.

Review the tbl_summary vignette or tbl_regression vignette for detailed examples in the ‘Advanced Customization’ section.

Usage

as_gt(x, include = everything(), return_calls = FALSE, ...)

Arguments

x Object created by a function from the gtsummary package (e.g. tbl_summary or tbl_regression)
include Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is everything().
return_calls Logical. Default is FALSE. If TRUE, the calls are returned as a list of expressions.
... Arguments passed on to gt::gt

Value

A gt_tbl object

Example Output

Author(s)

Daniel D. Sjoberg

See Also

Other gtsummary output types: as_flex_table(), as_hux_table(), as_kable_extra(), as_kable(), as_tibble.gtsummary()

Examples

# Example 1 ----------------------------------
as_gt_ex1 <-
trial[c("trt", "age", "response", "grade")]
%>%
tbl_summary(by = trt)
%>%
as_gt()
as_hux_table

Convert gtsummary object to a huxtable object

Description

Function converts a gtsummary object to a huxtable object. A user can use this function if they wish to add customized formatting available via the huxtable functions. The huxtable package supports output to PDF via LaTeX, as well as HTML and Word.

Usage

as_hux_table(
  x,
  include = everything(),
  return_calls = FALSE,
  strip_md_bold = FALSE
)

as_hux_xlsx(x, file, include = everything(), bold_header_rows = TRUE)

Arguments

x  
Object created by a function from the gtsummary package (e.g. tbl_summary or tbl_regression)

include  
Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is everything().

return_calls  
Logical. Default is FALSE. If TRUE, the calls are returned as a list of expressions.

strip_md_bold  
DEPRECATED

file  
File path for the output.

bold_header_rows  
logical indicating whether to bold header rows. Default is TRUE

Value

A huxtable object

Excel Output

Use the as_hux_xlsx() function to save a copy of the table in an excel file. The file is saved using huxtable::quick_xlsx().

Author(s)

David Hugh-Jones, Daniel D. Sjoberg

See Also

Other gtsummary output types: as_flex_table(), as_gt(), as_kable_extra(), as_kable(), as_tibble.gtsummary()
Examples

```r
trial %>%
  dplyr::select(trt, age, grade) %>%
  tbl_summary(by = trt) %>%
  add_p() %>%
  as_hux_table()
```

Description

Output from `knitr::kable` is less full featured compared to summary tables produced with `gt`. For example, kable summary tables do not include indentation, footnotes, or spanning header rows.

Line breaks (`\n`) are removed from column headers and table cells.

Usage

```r
as_kable(x, ..., include = everything(), return_calls = FALSE)
```

Arguments

- `x` Object created by a function from the gtsummary package (e.g. `tbl_summary` or `tbl_regression`)
- `...` Additional arguments passed to `knitr::kable`
- `include` Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is `everything()`.
- `return_calls` Logical. Default is `FALSE`. If `TRUE`, the calls are returned as a list of expressions.

Details

Tip: To better distinguish variable labels and level labels when indenting is not supported, try `bold_labels()` or `italicize_levels()`.

Value

A `knitr_kable` object

Author(s)

Daniel D. Sjoberg

See Also

Other gtsummary output types: `as_flex_table()`, `as_gt()`, `as_hux_table()`, `as_kable_extra()`, `as_tibble.gtsummary()`
Examples

```r
trial %>%
  tbl_summary(by = trt) %>%
  bold_labels() %>%
  as_kable()
```

**as_kable_extra**  
Convert gtsummary object to a kableExtra object

**Description**

Function converts a gtsummary object to a knitr_kable + kableExtra object. This allows the customized formatting available via knitr::kable() and kableExtra; as_kable_extra() supports arguments in knitr::kable(). as_kable_extra() output via gtsummary supports bold and italic cells for table bodies. Users are encouraged to leverage as_kable_extra() for enhanced pdf printing; for html output options there is better support via as_gt().

**Usage**

```r
as_kable_extra(
  x,
  escape = FALSE,
  format = NULL,
  ...,
  include = everything(),
  addtl_fmt = TRUE,
  return_calls = FALSE
)
```

**Arguments**

- `x`  
  Object created by a function from the gtsummary package (e.g. `tbl_summary` or `tbl_regression`)
- `format`, `escape`, `...`  
  Arguments passed to knitr::kable(). Default is `escape = FALSE`, and the format is auto-detected.
- `include`  
  Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is `everything()`.
- `addtl_fmt`  
  Logical indicating whether to include additional formatting. Default is `TRUE`. This is primarily used to escape special characters, convert markdown to LaTeX, and remove line breaks from the footnote.
- `return_calls`  
  Logical. Default is `FALSE`. If `TRUE`, the calls are returned as a list of expressions.

**Value**

A kableExtra table
This section shows options intended for use with `output: pdf_document` in yaml of `.Rmd`. When the default values of `as_kable_extra(escape = FALSE, addtl_fmt = TRUE)` are utilized, the following formatting occurs.

- Markdown bold, italic, and underline syntax in the headers, spanning headers, caption, and footnote will be converted to escaped LaTeX code
- Special characters in the table body, headers, spanning headers, caption, and footnote will be escaped with `.escape_latex()` or `.escape_latex2()`
- The "\n" symbol will be recognized as a line break in the table headers, spanning headers, caption, and the table body
- The "\n" symbol is removed from the footnotes

To suppress these additional formats, set `as_kable_extra(addtl_fmt = FALSE)` Additional styling is available with `kableExtra::kable_styling()` as shown in Example 2, which implements row striping and repeated column headers in the presence of page breaks.

HTML

This section discusses options intended for use with `output: html_document` in yaml of `.Rmd`. When the default values of `as_kable_extra(escape = FALSE, addtl_fmt = TRUE)` are utilized, the following formatting occurs.

- The default markdown syntax in the headers and spanning headers is removed
- Special characters in the table body, headers, spanning headers, caption, and footnote will be escaped with `.escape_html()`
- The "\n" symbol is removed from the footnotes

To suppress the additional formatting, set `as_kable_extra(addtl_fmt = FALSE)`

Example Output

Author(s)

Daniel D. Sjoberg

See Also

Other gtsummary output types: `as_flex_table()`, `as_gt()`, `as_hux_table()`, `as_kable()`, `as_tibble.gtsummary()`

Examples

```r
# basic gtsummary tbl to build upon
as_kable_extra_base <-
  trial %>%
  select(trt, age, stage) %>%
  tbl_summary(by = trt) %>%
  bold_labels()

# Example 1 (PDF via LaTeX) ---------------------
```
# add linebreak in table header with '
'

```r
as_kable_extra_ex1_pdf <-
  as_kable_extra_base %>%
  modify_header(all_stat_cols() ~ "**{level}**
  *N = {n}*") %>%
  as_kable_extra()
```

# Example 2 (PDF via LaTeX) ---------------------
# additional styling in `knitr::kable()` and with
# call to `kableExtra::kable_styling()`

```r
as_kable_extra_ex2_pdf <-
  as_kable_extra_base %>%
  as_kable_extra(  
    booktabs = TRUE,  
    longtable = TRUE,  
    linesep = ""  
  ) %>%
  kableExtra::kable_styling(  
    position = "left",  
    latex_options = c("striped", "repeat_header"),  
    stripe_color = "gray!15"  
  )
```

---

**as_tibble.gtsummary**  
*Convert gtsummary object to a tibble*

**Description**

Function converts a gtsummary object to a tibble.

**Usage**

```r
## S3 method for class 'gtsummary'
as_tibble(
  x,  
  include = everything(),  
  col_labels = TRUE,  
  return_calls = FALSE,  
  fmt_missing = FALSE,  
  ...  
)
```

```r
## S3 method for class 'gtsummary'
as.data.frame(...)
```

**Arguments**

- **x**  
  Object created by a function from the gtsummary package (e.g. tbl_summary or tbl_regression)

- **include**  
  Commands to include in output. Input may be a vector of quoted or unquoted names. tidyselect and gtsummary select helper functions are also accepted. Default is everything().
bold_italicize_labels_levels

boldLabels(x) italicizeLabels(x) boldLevels(x) italicizeLevels(x)

## S3 method for class 'gtsummary'
bold_labels(x)

Description

Bold or Italicize labels or levels in gtsummary tables

Usage

bold_labels(x)

italicize_labels(x)

bold_levels(x)

italicize_levels(x)

# # S3 method for class 'gtsummary'
bold_labels(x)
## S3 method for class 'gtsummary'
bold_levels(x)

## S3 method for class 'gtsummary'
italicize_labels(x)

## S3 method for class 'gtsummary'
italicize_levels(x)

## S3 method for class 'tbl_cross'
bold_labels(x)

## S3 method for class 'tbl_cross'
bold_levels(x)

## S3 method for class 'tbl_cross'
italicize_labels(x)

## S3 method for class 'tbl_cross'
italicize_levels(x)

Arguments

x Object created using gtsummary functions

Value

Functions return the same class of gtsummary object supplied

Example Output

Author(s)

Daniel D. Sjoberg

See Also

Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(), inline_text.tbl_summary(), inline_text.tbl_survfit(), modify, separate_p_footnotes, tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_summary()

Other tbl_regression tools: add_global_p(), add_q(), combine_terms(), inline_text.tbl_regression(), modify, tbl_merge(), tbl_regression(), tbl_split(), tbl_stack(), tbl_strata()

Other tbl_uvregression tools: add_global_p(), add_q(), inline_text.tbl_uvregression(), modify, tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_uvregression()

Examples

# Example 1 ----------------------------------
tbl_bold_ital_ex1 <-
  trial[c("trt", "age", "grade")]
  tbl_summary() %>%
  bold_labels() %>%
bold_p

**Bold significant p-values or q-values**

**Description**

Bold values below a chosen threshold (e.g. <0.05) in a gtsummary tables.

**Usage**

```r
bold_p(x, t = 0.05, q = FALSE)
```

**Arguments**

- `x`: Object created using gtsummary functions
- `t`: Threshold below which values will be bold. Default is 0.05.
- `q`: Logical argument. When TRUE will bold the q-value column rather than the p-values. Default is FALSE.

**Example Output**

**Author(s)**

Daniel D. Sjoberg, Esther Drill

**Examples**

```r
# Example 1 ----------------------------------
bold_p_ex1 <-
  trial[c("age", "grade", "response", "trt")]
  %>%
  tbl_summary(by = trt)
  %>%
  add_p()
  %>%
  bold_p(t = 0.65)

# Example 2 ----------------------------------
bold_p_ex2 <-
  glm(response ~ trt + grade, trial, family = binomial(link = "logit"))
  %>%
  tbl_regression(exponentiate = TRUE)
  %>%
  bold_p(t = 0.65)
```
**combine_terms**

**Combine terms in a regression model**

**Description**

The function combines terms from a regression model, and replaces the terms with a single row in the output table. The p-value is calculated using `stats::anova()`.

**Usage**

```r
combine_terms(x, formula_update, label = NULL, quiet = NULL, ...)
```

**Arguments**

- `x` a `tbl_regression` object
- `formula_update` formula update passed to the `stats::update`. This updated formula is used to construct a reduced model, and is subsequently passed to `stats::anova()` to calculate the p-value for the group of removed terms. See the `stats::update` help file for proper syntax. function's `formula =` argument
- `label` Option string argument labeling the combined rows
- `quiet` Logical indicating whether to print messages in console. Default is `FALSE`
- `...` Additional arguments passed to `stats::anova`

**Value**

`tbl_regression` object

**Example Output**

**Author(s)**

Daniel D. Sjoberg

**See Also**

Review list, formula, and selector syntax used throughout gtsummary

Other `tbl_regression` tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels`, `inline_text.tbl_regression()`, `modify`, `tbl_merge()`, `tbl_regression()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`

**Examples**

```r
# Example 1 ----------------------------------
# Logistic Regression Example, LRT p-value
combine_terms_ex1 <-
  glm(
    response ~ marker + I(marker^2) + grade,
    trial[c("response", "marker", "grade")] %>% na.omit(), # keep complete cases only!
    family = binomial
```
```r
) %>%
tbl_regression(label = grade ~ "Grade", exponentiate = TRUE) %>%
# collapse non-linear terms to a single row in output using anova
combine_terms(
  formula_update = . ~ . - marker - I(marker^2),
  label = "Marker (non-linear terms),
  test = "LRT"
)
```

### continuous_summary

**Summarize a continuous variable**

#### Description

[Experimental] This helper, to be used with `tbl_custom_summary()`, creates a function summarizing a continuous variable.

#### Usage

```r
continuous_summary(variable)
```

#### Arguments

- `variable`  
  String indicating the name of the variable to be summarized. This variable should be continuous.

#### Details

When using `continuous_summary`, you can specify in the `statistic=` argument of `tbl_custom_summary()` the same continuous statistics than in `tbl_summary()`. See the `statistic argument` section of the help file of `tbl_summary()`.

#### Example Output

#### Author(s)

Joseph Larmarange

#### See Also

Other `tbl_custom_summary` tools: `add_overall()`, `proportion_summary()`, `ratio_summary()`, `tbl_custom_summary()`
Examples

# Example 1 ----------------------------------
continuous_summary_ex1 <-
  trial %>%
  tbl_custom_summary(
    include = c("stage", "grade"),
    by = "trt",
    stat_fns = ~ continuous_summary("age"),
    statistic = ~(\(median\) \{p25\}-{p75\}),
    overall_row = TRUE,
    overall_row_label = "All stages & grades"
  ) %>%
  modify_footnote(
    update = all_stat_cols() ~ "Median age (IQR)"
  )


custom_tidders  Collection of custom tidiers

Description

[Maturing] Collection of tidiers that can be utilized in gtsummary. See details below.

Usage

tidy_standardize(
  x,
  exponentiate = FALSE,
  conf.level = 0.95,
  conf.int = TRUE,
  ..., 
  quiet = FALSE
)

tidy_bootstrap(
  x,
  exponentiate = FALSE,
  conf.level = 0.95,
  conf.int = TRUE,
  ..., 
  quiet = FALSE
)

tidy_robust(
  x,
  exponentiate = FALSE,
  conf.level = 0.95,
  conf.int = TRUE,
  vcov = NULL,
  vcov_args = NULL,
  ...,
quiet = FALSE
)
pool_and_tidy_mice(x, pool.args = NULL, ..., quiet = FALSE)
tidy_gam(x, conf.int = FALSE, exponentiate = FALSE, conf.level = 0.95, ...)
tidy_wald_test(x, tidy_fun = NULL, ...)

Arguments

- **x**: a regression model object
- **exponentiate**: Logical indicating whether or not to exponentiate the coefficient estimates. This is typical for logistic and multinomial regressions, but a bad idea if there is no log or logit link. Defaults to FALSE.
- **conf.level**: The confidence level to use for the confidence interval if conf.int = TRUE. Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval.
- **conf.int**: Logical indicating whether or not to include a confidence interval in the tidied output. Defaults to FALSE.
- **...**: arguments passed to method:
  - pool_and_tidy_mice(): mice::tidy(x, ...)
  - tidy_standardize(): parameters::standardize_parameters(x, ...)
  - tidy_bootstrap(): parameters::bootstrap_parameters(x, ...)
  - tidy_robust(): parameters::model_parameters(x, ...)
- **quiet**: Logical indicating whether to print messages in console. Default is FALSE
- **vcov, vcov_args**: arguments passed to parameters::model_parameters(). At least one of these arguments must be specified.
- **pool.args**: named list of arguments passed to mice::pool() in pool_and_tidy_mice(). Default is NULL
- **tidy_fun**: Option to specify a particular tidier function for the model. Default is to use broom::tidy(), but if an error occurs then tidying of the model is attempted with parameters::model_parameters(), if installed.

Regression Model Tidiers

These tidiers are passed to tbl_regression() and tbl_uvregression() to obtain modified results.

- **tidy_standardize()** tidier to report standardized coefficients. The parameters package includes a wonderful function to estimate standardized coefficients. The tidier uses the output from parameters::standardize_parameters(), and merely takes the result and puts it in broom::tidy() format.
- **tidy_bootstrap()** tidier to report bootstrapped coefficients. The parameters package includes a wonderful function to estimate bootstrapped coefficients. The tidier uses the output from parameters::bootstrap_parameters(test = "p"), and merely takes the result and puts it in broom::tidy() format.
custom_tidiers

• tidy_robust() tidier to report robust standard errors, confidence intervals, and p-values. The `parameters` package includes a wonderful function to calculate robust standard errors, confidence intervals, and p-values. The tidier uses the output from `parameters::model_parameters()`, and merely takes the result and puts it in `broom::tidy()` format. To use this function with `tbl_regression()`, pass a function with the arguments for `tidy_robust()` populated. This is easily done using `purrr::partial()` e.g. `tbl_regression(tidy_fun = partial(tidy_robust, vcov = "CL"))

• pool_and_tidy_mice() tidier to report models resulting from multiply imputed data using the mice package. Pass the mice model object before the model results have been pooled. See example.

Other Tidiers

• tidy_wald_test() tidier to report Wald p-values, wrapping the `aod::wald.test()` function. Use this tidier with `add_global_p(anova_fun = tidy_wald_test)`

Example Output

Examples

```r
# Example 1 ----------------------------------
mod <- lm(age ~ marker + grade, trial)

tbl_stnd <- tbl_regression(mod, tidy_fun = tidy_standardize)
tbl <- tbl_regression(mod)

tidy_standardize_ex1 <-
tbl_merge(
  list(tbl_stnd, tbl),
  tab_spanner = c("**Standardized Model**", "**Original Model**")
)

# Example 2 ----------------------------------
# use "posthoc" method for coef calculation

# Example 3 ----------------------------------
# Multiple Imputation using the mice package

set.seed(1123)

pool_and_tidy_mice_ex3 <-
suppressWarnings(mice::mice(trial, m = 2)) %>%
  with(lm(age ~ marker + grade)) %>%
tbl_regression()
```
inline_text.tbl_cross  Report statistics from cross table inline

Description

[Maturing] Extracts and returns statistics from a tbl_cross object for inline reporting in an R markdown document. Detailed examples in the inline_text vignette

Usage

## S3 method for class 'tbl_cross'
inline_text(x, col_level = NULL, row_level = NULL, pvalue_fun = NULL, ...)

Arguments

- x: gtsummary object
- variable: Variable name of statistic to present
- level: Level of the variable to display for categorical variables. Default is NULL
- column: Column name to return from x$table_body.
- pattern: String indicating the statistics to return. Uses glue::glue formatting. Default is NULL
- ...: Not used

column + pattern

Some gtsummary tables report multiple statistics in a single cell, e.g. "(mean) ((sd))" in tbl_summary() or tbl_svysummary(). We often need to report just the mean or the SD, and that can be accomplished by using both the column= and pattern= arguments. When both of these arguments are specified, the column argument selects the column to report statistics from, and the pattern argument specifies which statistics to report, e.g. inline_text(x, column = "stat_1", pattern = "{mean}") reports just the mean from a tbl_summary().
Arguments

- `x`: a `tbl_cross` object
- `col_level`: Level of the column variable to display. Default is NULL. Can also specify "p.value" for the p-value and "stat_0" for Total column.
- `row_level`: Level of the row variable to display. Can also specify the 'Unknown' row. Default is NULL.
- `pvalue_fun`: Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. `pvalue_fun = function(x) style_pvalue(x, digits = 2)` or equivalently, `purrr::partial(style_pvalue, digits = 2)`).
- `...`: Not used.

Value

A string reporting results from a `gtsummary` table.

See Also

Other `tbl_cross` tools: `add_p.tbl_cross()`, `tbl_cross()`

Examples

```r
tbl_cross <-
  tbl_cross(trial, row = trt, col = response) %>%
  add_p()

inline_text(tbl_cross, row_level = "Drug A", col_level = "1")
inline_text(tbl_cross, row_level = "Total", col_level = "1")
inline_text(tbl_cross, col_level = "p.value")
```
estimate_fun = NULL,
pvalue_fun = NULL,
...
)

Arguments

x Object created from tbl_regression

variable Variable name of statistics to present

level Level of the variable to display for categorical variables. Default is NULL, returning the top row in the table for the variable.

pattern String indicating the statistics to return. Uses glue::glue formatting. Default is 

"\{estimate\} (\{conf.level\}% CI \{conf.low\}, \{conf.high\}; \{p.value\})\". All columns from x$table_body are available to print as well as the confidence level (conf.level). See below for details.

estimate_fun function to style model coefficient estimates. Columns 'estimate', 'conf.low', and 'conf.high' are formatted. Default is x$inputs$estimate_fun

pvalue_fun function to style p-values and/or q-values. Default is function(x) style_pvalue(x, prepend_p = TRUE)

... Not used

Value

A string reporting results from a gtsummary table

pattern argument

The following items (and more) are available to print. Use print(x$table_body) to print the table the estimates are extracted from.

• \{estimate\} coefficient estimate formatted with 'estimate_fun'
• \{conf.low\} lower limit of confidence interval formatted with 'estimate_fun'
• \{conf.high\} upper limit of confidence interval formatted with 'estimate_fun'
• \{p.value\} p-value formatted with 'pvalue_fun'
• \{N\} number of observations in model
• \{label\} variable/variable level label

Author(s)

Daniel D. Sjoberg

See Also

Other tbl_regression tools: add_global_p(), add_q(), bold_italicize_labels_levels, combine_terms(), modify, tbl_merge(), tbl_regression(), tbl_split(), tbl_stack(), tbl_strata()
Examples

```r
inline_text_ex1 <-
  glm(response ~ age + grade, trial, family = binomial(link = "logit")) %>%
  tbl_regression(exponentiate = TRUE)
inline_text(inline_text_ex1, variable = age)
inline_text(inline_text_ex1, variable = grade, level = "III")
```

inline_text.tbl_summary

Report statistics from summary tables inline

Description

Extracts and returns statistics from a `tbl_summary` object for inline reporting in an R markdown document. Detailed examples in the `inline_text` vignette

Usage

```r
## S3 method for class 'tbl_summary'
inline_text(
  x,
  variable,
  column = NULL,
  level = NULL,
  pattern = NULL,
  pvalue_fun = NULL,
  ...
)

## S3 method for class 'tbl_svysummary'
inline_text(
  x,
  variable,
  column = NULL,
  level = NULL,
  pattern = NULL,
  pvalue_fun = NULL,
  ...
)
```

Arguments

- `x` Object created from `tbl_summary`
- `variable` Variable name of statistic to present
- `column` Column name to return from `x$table_body`. Can also pass the level of a by variable.
- `level` Level of the variable to display for categorical variables. Can also specify the 'Unknown' row. Default is `NULL`
pattern: String indicating the statistics to return. Uses glue::glue formatting. Default is pattern shown in tbl_summary() output.

pvalue_fun: Function to round and format p-values. Default is style_pvalue. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. pvalue_fun = function(x) style_pvalue(x, digits = 2) or equivalently, purrr::partial(style_pvalue, digits = 2)).

Value
A string reporting results from a gtsummary table.

Author(s)
Daniel D. Sjoberg

See Also
Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(), bold_italicize_labels_levels, inline_text.tbl_survfit(), modify, separate_p_footnotes(), tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata(), tbl_summary()

Examples
```r
t1 <- trial[, c("trt", "grade")]
  tbl_summary(by = trt)
  add_p()
  inline_text(t1, variable = grade, level = "I", column = "Drug A", pattern = "(n)/(N) ({p})%")
  inline_text(t1, variable = grade, column = "p.value")
```

Description


Usage
```r
## S3 method for class 'tbl_survfit'
inline_text(  
  x,  
  variable = NULL,  
  level = NULL,  
  pattern = NULL,  
  time = NULL,  
  prob = NULL,
)```
Arguments

- **x**: Object created from `tbl_survfit`
- **variable**: Variable name of statistic to present.
- **level**: Level of the variable to display for categorical variables. Can also specify the 'Unknown' row. Default is NULL
- **pattern**: String indicating the statistics to return.
- **time**: time for which to return survival probabilities.
- **prob**: probability with values in (0,1)
- **column**: column to print from `x$table_body`. Columns may be selected with `time=` or `prob=` as well.
- **estimate_fun**: Function to round and format estimate and confidence limits. Default is the same function used in `tbl_survfit()`
- **pvalue_fun**: Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. `pvalue_fun = function(x) style_pvalue(x, digits = 2)` or equivalently, `purrr::partial(style_pvalue, digits = 2)`).

Value

A string reporting results from a gtssummary table

Author(s)

Daniel D. Sjoberg

See Also

Other `tbl_summary` tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold_italicize_labels_levels`, `inline_text.tbl_summary()`, `modify`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Examples

```r
library(survival)
# fit survfit
fit1 <- survfit(Surv(ttdeath, death) ~ trt, trial)
fit2 <- survfit(Surv(ttdeath, death) ~ 1, trial)

# summarize survfit objects
tbl1 <-
  tbl_survfit(
    fit1,
```
\begin{verbatim}
times = c(12, 24),
label = "Treatment",
label_header = "**{time} Month**"
) %>%
add_p()

tbl2 <-
tbl_survfit(
  fit2,
  probs = 0.5,
  label_header = "**Median Survival**"
)

# report results inline
inline_text(tbl1, time = 24, level = "Drug B")
inline_text(tbl1, column = p.value)
inline_text(tbl2, prob = 0.5)
\end{verbatim}

\section*{inline_text.tbl_uvregression}

\textit{Report statistics from regression summary tables inline}

\subsection*{Description}
Extracts and returns statistics from a table created by the \texttt{tbl_uvregression} function for inline reporting in an R markdown document. Detailed examples in the \texttt{inline_text} vignette.

\subsection*{Usage}

\begin{verbatim}
## S3 method for class 'tbl_uvregression'
inline_text(
  x,
  variable,
  level = NULL,
  pattern = "{estimate} ({conf.level*100}% CI {conf.low}, {conf.high}; {p.value})",
  estimate_fun = NULL,
  pvalue_fun = NULL,
  ...
)
\end{verbatim}

\subsection*{Arguments}

\begin{description}
\item \textbf{x} Object created from \texttt{tbl_uvregression}
\item \textbf{variable} Variable name of statistics to present
\item \textbf{level} Level of the variable to display for categorical variables. Default is \texttt{NULL}, returning the top row in the table for the variable.
\item \textbf{pattern} String indicating the statistics to return. Uses \texttt{glue::glue} formatting. Default is "\{estimate\} \{(conf.level*100)% CI \{conf.low\}, \{conf.high\}; \{p.value\}". All columns from \texttt{x$table_body} are available to print as well as the confidence level (\texttt{conf.level}). See below for details.
\end{description}
Value

A string reporting results from a gtsummary table

pattern argument

The following items (and more) are available to print. Use `print(x$table_body)` to print the table the estimates are extracted from.

- `{estimate}` coefficient estimate formatted with `estimate_fun`
- `{conf.low}` lower limit of confidence interval formatted with `estimate_fun`
- `{conf.high}` upper limit of confidence interval formatted with `estimate_fun`
- `{p.value}` p-value formatted with `pvalue_fun`
- `{N}` number of observations in model
- `{label}` variable/variable level label

Author(s)

Daniel D. Sjoberg

See Also

Other tbl_uvregression tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels`, `modify()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_uvregression()

Examples

```r
inline_text_ex1 <-
  trial[c("response", "age", "grade")]%>%
tbl_uvregression(
    method = glm,
    method.args = list(family = binomial),
    y = response,
    exponentiate = TRUE
  )

inline_text(inline_text_ex1, variable = age)
inline_text(inline_text_ex1, variable = grade, level = "III")
```
### Description

These functions assist with updating or adding column headers (`modify_header()`), footnotes (`modify_footnote()`), spanning headers (`modify_spanning_header()`), and table captions (`modify_caption()`). Use `show_header_names()` to learn the column names.

### Usage

```r
modify_header(
  x,
  update = NULL,
  ...,                
  text_interpret = c("md", "html"),
  quiet = NULL,
  stat_by = NULL
)
```

```r
modify_footnote(
  x,
  update = NULL,
  ...,                
  abbreviation = FALSE,
  text_interpret = c("md", "html"),
  quiet = NULL
)
```

```r
modify_spanning_header(
  x,
  update = NULL,
  ...,                
  text_interpret = c("md", "html"),
  quiet = NULL
)
```

```r
modify_caption(x, caption, text_interpret = c("md", "html"))
```

```r
show_header_names(x = NULL, include_example = TRUE, quiet = NULL)
```

### Arguments

- `x`: a `gtsummary` object
- `update, ...`: use these arguments to assign updates to headers, spanning headers, and footnotes. See examples below.
  - `update` expects a list of assignments, with the variable name or selector on the LHS of the formula, and the updated string on the RHS. Also accepts a named list.
• ... pass individual updates outside of a list, e.g., 

\[
\text{modify\_header}(p.\text{value} = "**P**", \text{all\_stat\_cols()} \sim "**\{level\}**")
\]

Use the \texttt{show\_header\_names()} to see the column names that can be modified.

\begin{itemize}
\item \texttt{text\_interpret}: String indicates whether text will be interpreted with \texttt{gt::md()} or \texttt{gt::html()}. Must be "md" (default) or "html".
\item \texttt{quiet}: Logical indicating whether to print messages in console. Default is \texttt{FALSE}.
\item \texttt{stat\_by}: DEPRECATED, use update = \texttt{all\_stat\_cols() \sim "<label>"} instead.
\item \texttt{abbreviation}: Logical indicating if an abbreviation is being updated.
\item \texttt{caption}: a string of the table caption/title
\item \texttt{include\_example}: logical whether to include print of \texttt{modify\_header()} example
\end{itemize}

**Value**

Updated \texttt{gtsummary} object

**\texttt{tbl\_summary()}, \texttt{tbl\_svysummary()}, and \texttt{tbl\_cross()}**

When assigning column headers, footnotes, spanning headers, and captions for these \texttt{gtsummary} tables, you may use \{N\} to insert the number of observations. \texttt{tbl\_svysummary} objects additionally have \{N\_unweighted\} available.

When there is a stratifying \texttt{by=} argument present, the following fields are additionally available to stratifying columns: \{level\}, \{n\}, and \{p\} (\{n\_unweighted\} and \{p\_unweighted\} for \texttt{tbl\_svysummary} objects).

Syntax follows \texttt{glue::glue()}, e.g., \texttt{all\_stat\_cols() \sim "**\{level\}**\ N = \{n\}"}.

**\texttt{tbl\_regression()}**

When assigning column headers for \texttt{tbl\_regression} tables, you may use \{N\} to insert the number of observations, and \{N\_event\} for the number of events (when applicable).

**Captions**

Captions are assigned based on output type.

\begin{itemize}
\item \texttt{gt::gt(caption=)}
\item \texttt{flextable::set\_caption(caption=)}
\item \texttt{huxtable::set\_caption(value=)}
\item \texttt{knitr::kable(caption=)}
\end{itemize}

**Example Output**

**Author(s)**

Daniel D. Sjoberg
modify

See Also

Other tbl_summary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold_italicize_labels_levels()`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Other tbl_svysummary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_q()`, `add_stat_label()`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_svysummary()`

Other tbl_regression tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels()`, `combine_terms()`, `inline_text.tbl_regression()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_regression()`

Other tbl_uvregression tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels()`, `inline_text.tbl_uvregression()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_uvregression()`

Other tbl_survfit tools: `add_n.tbl_survfit()`, `add_nevent.tbl_survfit()`, `add_p.tbl_survfit()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_survfit()`

Examples

# create summary table
tbl <- trial[c("age", "grade", "trt")]
   %>%
   tbl_summary(by = trt, missing = "no")
   %>%
   add_p()

# print the column names that can be modified
show_header_names(tbl)

# Example 1 -------------------------------
# updating column headers, footnote, and table caption
modify_ex1 <- tbl %>%
   modify_header(label = "**Variable**", p.value = "**P**") %>%
   modify_footnote(all_stat_cols() ~ "median (IQR) for Age; n (%) for Grade") %>%
   modify_caption("**Patient Characteristics** (N = {N})")

# Example 2 -------------------------------
# updating headers, remove all footnotes, add spanning header
modify_ex2 <- tbl %>%
   modify_header(all_stat_cols() ~ "**(level)**, N = {n} ((style_percent(p))\%") %>%
   modify_footnote(everything() ~ NA, abbreviation = TRUE) %>%
   modify_caption("**Treatment Received**")

# Example 3 -------------------------------
# updating an abbreviation in table footnote
modify_ex3 <-
   glm(response ~ age + grade, trial, family = binomial)
   %>%
   tbl_regression(exponentiate = TRUE)
   %>%
   modify_footnote(ci = "CI = Credible Interval", abbreviation = TRUE)
**modify_column_alignment**

*Modify Column Alignment*

**Description**

*[Maturing]* Update column alignment/justification in a gtsummary table.

**Usage**

```r
modify_column_alignment(x, columns, align = c("left", "right", "center"))
```

**Arguments**

- `x` gtsummary object
- `columns` vector or selector of columns in `x$table_body`
- `align` string indicating alignment of column, must be one of c("left", "right", "center")

**See Also**

Review list, formula, and selector syntax used throughout gtsummary

Other Advanced modifiers: `modify_column_hide()`, `modify_column_indent()`, `modify_column_merge()`, `modify_fmt_fun()`, `modify_table_body()`, `modify_table_styling()`

**Examples**

```r
# Example 1 ----------------------------------
tbl <-
  lm(age ~ marker + grade, trial) %>%
  tbl_regression() %>%
  modify_column_alignment(columns = everything(), align = "left")
```

---

**modify_column_hide**

*Modify Hidden Columns*

**Description**

*[Maturing]* Use these functions to hide or unhide columns in a gtsummary table.

**Usage**

```r
modify_column_hide(x, columns)
modify_column_unhide(x, columns)
```
modify_column_indent

Arguments

x  gtsummary object
columns  vector or selector of columns in x$table_body

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary
Other Advanced modifiers: modify_column_alignment(), modify_column_indent(), modify_column_merge(), modify_fmt_fun(), modify_table_body(), modify_table_styling()

Examples

# Example 1 ----------------------------------
# hide 95% CI, and replace with standard error
modify_column_hide_ex1 <-
  lm(age ~ marker + grade, trial) %>%
  tbl_regression() %>%
  modify_column_hide(columns = ci) %>%
  modify_column_unhide(columns = std.error)

modify_column_indent  Add/Remove Indentation

Description

Add/Remove Indentation

Usage

modify_column_indent(
  x,
  columns,
  rows = NULL,
  double_indent = FALSE,
  undo = FALSE
)

Arguments

x  a gtsummary table
columns  columns to add indentation to
rows  predicate expression to select rows to indent. Default is NULL, indicating all rows
double_indent  logical indicating whether to double indent the cells. Default is FALSE for a single indentation
undo  logical indicating whether an indentation should be removed/undone. Default is FALSE
modify_column_merge

Value

a gtsummary table

See Also

Other Advanced modifiers: modify_column_alignment(), modify_column_hide(), modify_column_merge(), modify_fmt_fun(), modify_table_body(), modify_table_styling()

Examples

# remove indentation from `tbl_summary()`
modify_column_indent_ex1 <-
  trial %>%
  select(grade) %>%
  tbl_summary() %>%
  modify_column_indent(columns = label, undo = TRUE)

modify_column_merge  Modify Column Merging

Description

[Experimental] Merge two or more columns in a gtsummary table. Use show_header_names() to print underlying column names.

Usage

modify_column_merge(x, pattern, rows = NULL)

Arguments

x  gtsummary object
pattern  glue syntax string indicating how to merge columns in x$table_body. For example, to construct a confidence interval use “{conf.low}, {conf.high}”.
rows  predicate expression to select rows in x$table_body. Can be used to style footnote, formatting functions, missing symbols, and text formatting. Default is NULL. See details below.

Value

gtsummary table

Details

1. Calling this function merely records the instructions to merge columns. The actual merging occurs when the gtsummary table is printed or converted with a function like as_gt().
2. Because the column merging is delayed, it is recommended to perform major modifications to the table, such as those with tbl_merge() and tbl_stack(), before assigning merging instructions. Otherwise, unexpected formatting may occur in the final table.
modify_column_merge

3. If this functionality is used in conjunction with tbl_stack() (which includes tbl_uvregression()), there is potential issue with printing. When columns are stack AND when the column-merging is defined with a quosure, you may run into issues due to the loss of the environment when 2 or more quosures are combined. If the expression version of the quosure is the same as the quosure (i.e. no evaluated objects), there should be no issues. Regardless, this argument is used internally with care, and it is not recommended for users.

Future Updates

There are planned updates to the implementation of this function with respect to the pattern= argument. Currently, this function replaces a numeric column with a formatted character column following pattern=. Once gt::cols_merge() gains the rows= argument the implementation will be updated to use it, which will keep numeric columns numeric. For the vast majority of users, the planned change will be go unnoticed.

If this functionality is used in conjunction with tbl_stack() (which includes tbl_uvregression()), there is potential issue with printing. When columns are stack AND when the column-merging is defined with a quosure, you may run into issues due to the loss of the environment when 2 or more quosures are combined. If the expression version of the quosure is the same as the quosure (i.e. no evaluated objects), there should be no issues. Regardless, this argument is used internally with care, and it is not recommended for users.

Example Output

See Also

Other Advanced modifiers: modify_column_alignment(), modify_column_hide(), modify_column_indent(), modify_fmt_fun(), modify_table_body(), modify_table_styling()

Examples

# Example 1 ----------------------------------
modify_column_merge_ex1 <-
trial %>%
  select(age, marker, trt) %>%
  tbl_summary(by = trt, missing = "no") %>%
  add_p(all_continuous() ~ "t.test", pvalue_fun = ~ style_pvalue(., prepend_p = TRUE)) %>%
  modify_fmt_fun(statistic ~ style_sigfig) %>%
  modify_column_merge(pattern = "t = {statistic}; {p.value}")) %>%
  modify_header(statistic ~ "**t-test**")

# Example 2 ----------------------------------
modify_column_merge_ex2 <-
lm(marker ~ age + grade, trial) %>%
tbl_regression() %>%
modify_column_merge(
  pattern = "{estimate} ({ci})",
  rows = !is.na(estimate)
)
modify_fmt_fun

Modify Formatting Functions

Description

[Maturing] Use this function to update the way numeric columns and rows of $.table_body are formatted

Usage

modify_fmt_fun(x, update, rows = NULL)

Arguments

x         gtsummary object
update    list of formulas or a single formula specifying the updated formatting function. The LHS specifies the column(s) to be updated, and the RHS is the updated formatting function.
rows      predicate expression to select rows in x$table_body. Default is NULL. See details below.

Example Output

rows argument

The rows argument accepts a predicate expression that is used to specify rows to apply formatting. The expression must evaluate to a logical when evaluated in x$table_body. For example, to apply formatting to the age rows pass rows = variable == "age". A vector of row numbers is NOT acceptable.

A couple of things to note when using the rows= argument.

1. You can use saved objects to create the predicate argument, e.g. rows = variable == letters[1].
2. The saved object cannot share a name with a column in x$table_body. The reason for this is that in tbl_merge() the columns are renamed, and the renaming process cannot disambiguate the variable column from an external object named variable in the following expression rows = .data$variable = .env$variable.

See Also

Review list, formula, and selector syntax used throughout gtsummary

Other Advanced modifiers: modify_column_alignment(), modify_column_hide(), modify_column_indent(), modify_column_merge(), modify_table_body(), modify_table_styling()
Examples

# Example 1 ----------------------------------
# show 'grade' p-values to 3 decimal places
modify_fmt_fun_ex1 <-
  lm(age ~ marker + grade, trial) %>%
  tbl_regression() %>%
  modify_fmt_fun(
    update = p.value ~ function(x) style_pvalue(x, digits = 3),
    rows = variable == "grade"
  )

Description

[Maturing] Function is for advanced manipulation of gtsummary tables. It allow users to modify the $.table_body data frame included in each gtsummary object.

If a new column is added to the table, default printing instructions will then be added to $.table_styling. By default, columns are hidden. To show a column, add a column header with modify_header().

Usage

modify_table_body(x, fun, ...)

Arguments

x  gtsummary object

fun  A function or formula. If a function, it is used as is. If a formula, e.g. fun = ~ .x %>% arrange(variable), it is converted to a function. The argument passed to fun= is x$table_body.

...  Additional arguments passed on to the mapped function

Example Output

See Also

modify_table_styling()

See gtsummary internals vignette

Other Advanced modifiers: modify_column_alignment(), modify_column_hide(), modify_column_indent(), modify_column_merge(), modify_fmt_fun(), modify_table_styling()
Examples

# Example 1 --------------------------------
# Add number of cases and controls to regression table
modify_table_body_ex1 <-
  trial %>%
  select(response, age, marker) %>%
  tbl_uvregression(
    y = response,
    method = glm,
    method.args = list(family = binomial),
    exponentiate = TRUE,
    hide_n = TRUE
  ) %>%
  # adding number of non-events to table
  modify_table_body(
    .x %>%
    dplyr::mutate(N_nonevent = N_obs - N_event) %>%
    dplyr::relocate(c(N_event, N_nonevent), .before = estimate)
  ) %>%
  # assigning header labels
  modify_header(N_nonevent = "**Control N**", N_event = "**Case N**") %>%
  modify_fmt_fun(c(N_event, N_nonevent) ~ style_number)

modify_table_styling

Modify Table Styling

Description

This is a function meant for advanced users to gain more control over the characteristics of the resulting gtsummary table by directly modifying .$table_styling

Usage

modify_table_styling(
  x,
  columns,
  rows = NULL,
  label = NULL,
  spanning_header = NULL,
  hide = NULL,
  footnote = NULL,
  footnote_abbrev = NULL,
  align = NULL,
  missing_symbol = NULL,
  fmt_fun = NULL,
  text_format = NULL,
  undo_text_format = FALSE,
  text_interpret = c("md", "html"),
  cols_merge_pattern = NULL
)
modify_table_styling

Arguments

- **x**: gtsummary object
- **columns**: vector or selector of columns in x$table_body
- **rows**: predicate expression to select rows in x$table_body. Can be used to style footnote, formatting functions, missing symbols, and text formatting. Default is NULL. See details below.
- **label**: string of column label(s)
- **spanning_header**: string with text for spanning header
- **hide**: logical indicating whether to hide column from output
- **footnote**: string with text for footnote
- **footnote_abbrev**: string with abbreviation definition, e.g. "CI = Confidence Interval"
- **align**: string indicating alignment of column, must be one of c("left", "right", "center")
- **missing_symbol**: string indicating how missing values are formatted.
- **fmt_fun**: function that formats the statistics in the columns/rows in columns= and rows=
- **text_format**: string indicated which type of text formatting to apply to the rows and columns. Must be one of c("bold", "italic", "indent", "indent2"). Do not assign both "indent" and "indent2" to the same cell.
- **undo_text_format**: rarely used. Logical that undoes the indent, bold, and italic styling when TRUE
- **text_interpret**: string, must be one of "md" or "html"
- **cols_merge_pattern**: [Experimental] glue-syntax string indicating how to merge columns in x$table_body. For example, to construct a confidence interval use "{conf.low}, {conf.high}". The first column listed in the pattern string must match the single column name passed in columns=.

Details

Review the gtsummary definition vignette for information on .$table_styling objects.

rows argument

The rows argument accepts a predicate expression that is used to specify rows to apply formatting. The expression must evaluate to a logical when evaluated in x$table_body. For example, to apply formatting to the age rows pass rows = variable == "age". A vector of row numbers is NOT acceptable.

A couple of things to note when using the rows= argument.

1. You can use saved objects to create the predicate argument, e.g. rows = variable == letters[1].
2. The saved object cannot share a name with a column in x$table_body. The reason for this is that in tbl_merge() the columns are renamed, and the renaming process cannot disambiguate the variable column from an external object named variable in the following expression rows = .data$variable = .env$variable.
There are planned updates to the implementation of column merging. Currently, this function replaces the numeric column with a formatted character column following `cols_merge_pattern=`. Once `gt::cols_merge()` gains the `rows=` argument the implementation will be updated to use it, which will keep numeric columns numeric. For the vast majority of users, the planned change will be go unnoticed.

If this functionality is used in conjunction with `tbl_stack()` (which includes `tbl_uvregression()`), there is potential issue with printing. When columns are stack AND when the column-merging is defined with a quosure, you may run into issues due to the loss of the environment when 2 or more quosures are combined. If the expression version of the quosure is the same as the quosure (i.e. no evaluated objects), there should be no issues. Regardless, this argument is used internally with care, and it is not recommended for users.

See Also

modify_table_body()

See `gtsummary internals vignette`

Review list, formula, and selector syntax used throughout gtsummary

Other Advanced modifiers: modify_column_alignment(), modify_column_hide(), modify_column_indent(), modify_column_merge(), modify_fmt_fun(), modify_table_body()

---

**plot**

*Plot Regression Coefficients*

**Description**

The `plot()` function extracts `x$table_body` and passes the it to `ggstats::ggcoef_plot()` along with a formatting options.

**Usage**

```r
## S3 method for class 'tbl_regression'
plot(x, remove_header_rows = TRUE, remove_reference_rows = FALSE, ...)

## S3 method for class 'tbl_uvregression'
plot(x, remove_header_rows = TRUE, remove_reference_rows = FALSE, ...)
```

**Arguments**

- `x` `tbl_regression` or `tbl_uvregression` object
- `remove_header_rows` logical indicating whether to remove header rows for categorical variables. Default is `TRUE`.
- `remove_reference_rows` logical indicating whether to remove reference rows for categorical variables. Default is `FALSE`.
- `...` arguments passed to `ggstats::ggcoef_plot(...)`
proportion_summary

Details

[Experimental]

Value

a ggplot

Examples

```r
glm(response ~ marker + grade, trial, family = binomial) %>%
  tbl_regression(
    add_estimate_to_reference_rows = TRUE,
    exponentiate = TRUE
  ) %>%
plot()
```

Description

[Experimental] This helper, to be used with `tbl_custom_summary()`, creates a function computing a proportion and its confidence interval.

Usage

```r
proportion_summary(
  variable,
  value,
  weights = NULL,
  na.rm = TRUE,
  conf.level = 0.95,
  method = c("wilson", "wilson.no.correct", "exact", "asymptotic")
)
```

Arguments

- **variable**: String indicating the name of the variable from which the proportion will be computed.
- **value**: Value (or list of values) of `variable` to be taken into account in the numerator.
- **weights**: Optional string indicating the name of a weighting variable. If `NULL`, all observations will be assumed to have a weight equal to 1.
- **na.rm**: Should missing values be removed before computing the proportion? (default is `TRUE`)?
- **conf.level**: Confidence level for the returned confidence interval. Must be strictly greater than 0 and less than 1. Default to 0.95, which corresponds to a 95 percent confidence interval.
- **method**: Confidence interval method. Must be one of c("wilson", "wilson.no.correct", "exact", "asymptotic"). See details below.
Details

Computed statistics:

- \( n \) numerator, (weighted) number of observations equal to values
- \( N \) denominator, (weighted) number of observations
- \( prop \) proportion, i.e. \( n/N \)
- \( \text{conf.low} \) lower confidence interval
- \( \text{conf.high} \) upper confidence interval

Methods \( \text{c(} \text{"wilson"}, \text{"wilson.no.correct"} \) \) are calculated with \( \text{stats::prop.test()} \) (with \( \text{correct} = \text{c(TRUE, FALSE)} \)). The default method, "wilson", includes the Yates continuity correction. Methods \( \text{c(} \text{"exact"}, \text{"asymptotic"} \) \) are calculated with \( \text{Hmisc::binconf()} \) and the corresponding method.

Example Output

Author(s)

Joseph Larmarange

See Also

Other tbl_custom_summary tools: \text{add_overall()}, \text{continuous_summary()}, \text{ratio_summary()}, \text{tbl_custom_summary()}

Examples

# Example 1 -------------------------------
proportion_summary_ex1 <-
Titanic %>%
  as.data.frame() %>%
tbl_custom_summary(
  include = c("Age", "Class"),
  by = "Sex",
  stat_fns = ~ proportion_summary("Survived", "Yes", weights = "Freq"),
  statistic = "\{prop\}% \{n\}/\{N\} \{conf.low}\,\{conf.high\}\",
  digits = ~ list(
    function(x) {
      style_percent(x, digits = 1)
    },
    0, 0, style_percent, style_percent
  ),
  overall_row = TRUE,
  overall_row_last = TRUE
) %>%
bold_labels() %>%
modify_footnote(
  update = all_stat_cols() ~ "Proportion (%) of survivors \{n\}/\{N\} [95% CI]"
)
ratio_summary  

Summarize the ratio of two variables

Description

[Experimental] This helper, to be used with tbl_custom_summary(), creates a function computing the ratio of two continuous variables and its confidence interval.

Usage

ratio_summary(numerator, denominator, na.rm = TRUE, conf.level = 0.95)

Arguments

- **numerator**: String indicating the name of the variable to be summed for computing the numerator.
- **denominator**: String indicating the name of the variable to be summed for computing the denominator.
- **na.rm**: Should missing values be removed before summing the numerator and the denominator? (default is TRUE)
- **conf.level**: Confidence level for the returned confidence interval. Must be strictly greater than 0 and less than 1. Default to 0.95, which corresponds to a 95 percent confidence interval.

Details

Computed statistics:

- `{num}` sum of the variable defined by numerator
- `{denom}` sum of the variable defined by denominator
- `{ratio}` ratio of num by denom
- `{conf.low}` lower confidence interval
- `{conf.high}` upper confidence interval

Confidence interval is computed with stats::poisson.test(), if and only if num is an integer.

Example Output

Author(s)

Joseph Larmarange

See Also

Other tbl_custom_summary tools: add_overall(), continuous_summary(), proportion_summary(), tbl_custom_summary()
Examples

# Example 1 ----------------------------------
ratio_summary_ex1 <-
trial %>%
tbl_custom_summary(
  include = c("stage", "grade"),
  by = "trt",
  stat_fns = ~ ratio_summary("response", "ttdeath"),
  statistic = "{ratio} [{conf.low}; {conf.high}] ({num}/{denom})",
  digits = ~ c(3, 2, 2, 0, 0),
  overall_row = TRUE,
  overall_row_label = "All stages & grades"
) %>%
bold_labels() %>%
modify_footnote(
  update = all_stat_cols() ~ "Ratio [95% CI] (n/N)"
)

remove_row_type  Remove rows by type

Description

Removes either the header, reference, or missing rows from a gtsummary table.

Usage

remove_row_type(
  x,
  variables = everything(),
  type = c("header", "reference", "missing", "level", "all"),
  level_value = NULL
)

Arguments

x  gtsummary object
variables  variables to to remove rows from. Default is everything()
type  type of row to remove. Must be one of
level_value  When type='level' you can specify the character value of the level to remove. When NULL all levels are removed. c("header", "reference", "missing")

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary
**Examples**

# Example 1 ----------------------------------
library(dplyr, warn.conflicts = FALSE, quietly = TRUE)
remove_row_type_ex1 <-
  trial %>%
  select(trt, age) %>%
  mutate(
    age60 = case_when(age < 60 ~ "<60", age >= 60 ~ "60+")
  ) %>%
  tbl_summary(by = trt, missing = "no") %>%
  remove_row_type(age60, type = "header")

**select_helpers**

Select helper functions

**Description**

Set of functions to supplement the tidyselect set of functions for selecting columns of data frames (and other items as well).

- all_continuous() selects continuous variables
- all_continuous2() selects only type "continuous2"
- all_categorical() selects categorical (including "dichotomous") variables
- all_dichotomous() selects only type "dichotomous"
- all_tests() selects variables by the name of the test performed
- all_stat_cols() selects columns from tbl_summary/tbl_svysummary object with summary statistics (i.e. "stat_0", "stat_1", "stat_2", etc.)
- all_interaction() selects interaction terms from a regression model
- all_intercepts() selects intercept terms from a regression model
- all_contrasts() selects variables in regression model based on their type of contrast

**Usage**

all_continuous(continuous2 = TRUE)

all_continuous2()

all_categorical(dichotomous = TRUE)

all_dichotomous()

all_tests(tests = NULL)

all_stat_cols(stat_0 = TRUE)

all_interaction()

all_intercepts()

all_contrasts(contrasts_type = NULL)
Arguments

- `continuous2`: Logical indicating whether to include continuous2 variables. Default is TRUE
- `dichotomous`: Logical indicating whether to include dichotomous variables. Default is TRUE
- `tests`: string indicating the test type of the variables to select, e.g. select all variables being compared with "t.test"
- `stat_0`: When FALSE, will not select the "stat_0" column. Default is TRUE
- `contrasts_type`: type of contrast to select. When NULL, all variables with a contrast will be selected. Default is NULL. Select among contrast types c("treatment", "sum", "poly", "helmert", "other")

Value

A character vector of column names selected

Example Output

See Also

Review list, formula, and selector syntax used throughout gtsummary

Examples

```r
select_ex1 <-
  trial %>%
  select(age, response, grade) %>%
  tbl_summary(
    statistic = all_continuous() ~ "{mean} ({sd})",
    type = all_dichotomous() ~ "categorical"
  )
```

Description

[Experimental] The usual presentation of footnotes for p-values on a gtsummary table is to have a single footnote that lists all statistical tests that were used to compute p-values on a given table. The `separate_p_footnotes()` function separates aggregated p-value footnotes to individual footnotes that denote the specific test used for each of the p-values.

Usage

```r
separate_p_footnotes(x)
```

Arguments

- `x`: object with class "tbl_summary" or "tbl_svysummary"
**set_gtsummary_theme**

### Example Output

**See Also**

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_summary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold_italicize_labels_levels`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify.tbl_custom_summary()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Other tbl_svysummary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_q()`, `add_stat_label()`, `modify.tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

### Examples

```r
# Example 1 ----------------------------------
separate_p_footnotes_ex1 <-
  trial %>%
  select(trt, age, grade) %>%
  tbl_summary(by = trt) %>%
  add_p() %>%
  separate_p_footnotes()
```

### Description

[Maturing] Functions to set, reset, get, and evaluate with gtsummary themes.

- `set_gtsummary_theme()` set a theme
- `reset_gtsummary_theme()` reset themes
- `get_gtsummary_theme()` get a named list with all active theme elements
- `with_gtsummary_theme()` evaluate an expression with a theme temporarily set
- `check_gtsummary_theme()` checks if passed theme is valid

### Usage

```r
set_gtsummary_theme(x, quiet = NULL)
reset_gtsummary_theme()
get_gtsummary_theme()

with_gtsummary_theme(
  x,
  expr,
  env = rlang::caller_env(),
  msg_ignored_elements = NULL
)

check_gtsummary_theme(x)
```
Arguments

- **x**: A named list defining a gtsummary theme.
- **quiet**: Logical indicating whether to print messages in console. Default is `FALSE`.
- **expr**: Expression to be evaluated with the theme specified in `x`.
- **env**: The environment in which to evaluate `expr`.
- **msg_ignored_elements**: Default is NULL with no message printed. Pass a string that will be printed with `cli::cli_alert_info()`. The `{elements}` object contains vector of theme elements that will be overwritten and ignored.

Details

The default formatting and styling throughout the gtsummary package are taken from the published reporting guidelines of the top four urology journals: European Urology, The Journal of Urology, Urology and the British Journal of Urology International. Use this function to change the default reporting style to match another journal, or your own personal style.

Example Output

See Also

- Themes vignette
- Available gtsummary themes

Examples

```r
# Setting JAMA theme for gtsummary
set_gtsummary_theme(theme_gtsummary_journal("jama"))

# Themes can be combined by including more than one
set_gtsummary_theme(theme_gtsummary_compact())

set_gtsummary_theme_ex1 <-
  trial %>%
  dplyr::select(age, grade, trt) %>%
  tbl_summary(by = trt) %>%
  add_stat_label() %>%
  as_gt()

# reset gtsummary theme
reset_gtsummary_theme()
```
sort_filter_p  Sort and filter variables in table by p-values

Description
Sort and filter variables in table by p-values

Usage
sort_p(x, q = FALSE)
filter_p(x, q = FALSE, t = 0.05)

Arguments
x An object created using gtsummary functions
q Logical argument. When TRUE will the q-value column is used
t p-values/q-values less than or equal to this threshold will be retained. Default is 0.05

Example Output

Author(s)
Karissa Whiting, Daniel D. Sjoberg

Examples
# Example 1 ----------------------------------
sort_filter_p_ex1 <-
  trial %>%
  select(age, grade, response, trt) %>%
  tbl_summary(by = trt) %>%
  add_p() %>%
  filter_p(t = 0.8) %>%
  sort_p()

# Example 2 ----------------------------------
sort_p_ex2 <-
  glm(response ~ trt + grade, trial, family = binomial(link = "logit")) %>%
  tbl_regression(exponentiate = TRUE) %>%
  sort_p()
### Description

Style numbers

### Usage

```r
style_number(
  x,
  digits = 0,
  big.mark = NULL,
  decimal.mark = NULL,
  scale = 1,
  ...
)
```

### Arguments

- **x**: Numeric vector
- **digits**: Integer or vector of integers specifying the number of decimals to round x=`. When vector is passed, each integer is mapped 1:1 to the numeric values in x
- **big.mark**: Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is ",", except when decimal.mark = "", when the default is a space.
- **decimal.mark**: The character to be used to indicate the numeric decimal point. Default is "." or getOption("OutDec")
- **scale**: A scaling factor: x will be multiplied by scale before formatting.
- **...**: Other arguments passed on to base::format()

### Value

formatted character vector

### See Also

Other style tools: `style_percent()`, `style_pvalue()`, `style_ratio()`, `style_sigfig()`

### Examples

```r
c(0.111, 12.3) %>% style_number(digits = 1)
c(0.111, 12.3) %>% style_number(digits = c(1, 0))
```
style_percent

Style percentages

Description

Style percentages

Usage

style_percent(
  x, 
  symbol = FALSE, 
  digits = 0, 
  big.mark = NULL, 
  decimal.mark = NULL, 
  ... 
)

Arguments

x numeric vector of percentages
symbol Logical indicator to include percent symbol in output. Default is FALSE.
digits number of digits to round large percentages (i.e. greater than 10%). Smaller percentages are rounded to digits + 1 places. Default is 0
big.mark Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is "", except when decimal.mark = "," when the default is a space.
decimal.mark The character to be used to indicate the numeric decimal point. Default is "." or getOption("OutDec")
...

Value

A character vector of styled percentages

Author(s)

Daniel D. Sjoberg

See Also

See Table Gallery vignette for example

Other style tools: style_number(), style_pvalue(), style_ratio(), style_sigfig()

Examples

percent_vals <- c(-1, 0, 0.0001, 0.005, 0.01, 0.10, 0.45356, 0.99, 1.45)
style_percent(percent_vals)
style_percent(percent_vals, symbol = TRUE, digits = 1)
Style p-values

**Usage**

```r
style_pvalue(
  x,
  digits = 1,
  prepend_p = FALSE,
  big.mark = NULL,
  decimal.mark = NULL,
  ...  
)
```

**Arguments**

- `x` Numeric vector of p-values.
- `digits` Number of digits large p-values are rounded. Must be 1, 2, or 3. Default is 1.
- `prepend_p` Logical. Should 'p=' be prepended to formatted p-value. Default is FALSE
- `big.mark` Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is ",", except when `decimal.mark` = "", when the default is a space.
- `decimal.mark` The character to be used to indicate the numeric decimal point. Default is "." or `getOption("OutDec")`
- `...` Other arguments passed on to `base::format()`

**Value**

A character vector of styled p-values

**Author(s)**

Daniel D. Sjoberg

**See Also**

See tbl_summary vignette for examples

Other style tools: `style_number()`, `style_percent()`, `style_ratio()`, `style_sigfig()`

**Examples**

```r
pvals <- c(
  1.5, 1, 0.999, 0.5, 0.25, 0.2, 0.197, 0.12, 0.10, 0.0999, 0.06,
  0.03, 0.002, 0.001, 0.00099, 0.0002, 0.00002, -1
)
style_pvalue(pvals)
style_pvalue(pvals, digits = 2, prepend_p = TRUE)
```
**style_ratio**

*Style significant figure-like rounding for ratios*

**Description**

When reporting ratios, such as relative risk or an odds ratio, we’ll often want the rounding to be similar on each side of the number 1. For example, if we report an odds ratio of 0.95 with a confidence interval of 0.70 to 1.24, we would want to round to two decimal places for all values. In other words, 2 significant figures for numbers less than 1 and 3 significant figures 1 and larger. `style_ratio()` performs significant figure-like rounding in this manner.

**Usage**

```r
style_ratio(x, digits = 2, big.mark = NULL, decimal.mark = NULL, ...)
```

**Arguments**

- `x` Numeric vector
- `digits` Integer specifying the number of significant digits to display for numbers below 1. Numbers larger than 1 will be be digits + 1. Default is `digits = 2`.
- `big.mark` Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is `""`, except when `decimal.mark = ""` when the default is a space.
- `decimal.mark` The character to be used to indicate the numeric decimal point. Default is `"."` or `getOption("OutDec")`
- `...` Other arguments passed on to `base::format()`

**Value**

A character vector of styled ratios

**Author(s)**

Daniel D. Sjoberg

**See Also**

Other style tools: `style_number()`, `style_percent()`, `style_pvalue()`, `style_sigfig()`

**Examples**

```r
x <- c(0.123, 0.9, 1.1234, 12.345, 101.234, -0.123, -0.9, -1.1234, -12.345, -101.234)
> style_ratio(x)
```

**style_sigfig**

**Style significant figure-like rounding**

**Description**

Converts a numeric argument into a string that has been rounded to a significant figure-like number. Scientific notation output is avoided, however, and additional significant figures may be displayed for large numbers. For example, if the number of significant digits requested is 2, 123 will be displayed (rather than 120 or 1.2x10^2).

**Usage**

```r
style_sigfig(
  x,
  digits = 2,
  scale = 1,
  big.mark = NULL,
  decimal.mark = NULL,
  ...
)
```

**Arguments**

- **x** Numeric vector
- **digits** Integer specifying the minimum number of significant digits to display
- **scale** A scaling factor: x will be multiplied by scale before formatting.
- **big.mark** Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is ",", except when `decimal.mark = ""," when the default is a space.
- **decimal.mark** The character to be used to indicate the numeric decimal point. Default is "." or `getOption("OutDec")`
- **...** Other arguments passed on to `base::format()`

**Value**

A character vector of styled numbers

**Details**

- Scientific notation output is avoided.
- If 2 significant figures are requested, the number is rounded to no more than 2 decimal places. For example, a number will be rounded to 2 decimals places when `abs(x) < 1`, 1 decimal place when `abs(x) >= 1 & abs(x) < 10`, and to the nearest integer when `abs(x) >= 10`.
- Additional significant figures may be displayed for large numbers. For example, if the number of significant digits requested is 2, 123 will be displayed (rather than 120 or 1.2x10^2).

**Author(s)**

Daniel D. Sjoberg
See Also

Other style tools: `style_number()`, `style_percent()`, `style_pvalue()`, `style_ratio()

Examples

c(0.123, 0.9, 1.1234, 12.345, -0.123, -0.9, -1.1234, -132.345, NA, -0.001) %>%
  style_sigfig()

---

**tbl_butcher**

*Reduce size of gtsummary objects*

Description

Some gtsummary objects can become large and the size becomes cumbersome when working with the object. The function removes all elements from a gtsummary object, except those required to print the table. This may result in gtsummary functions that add information or modify the table, such as `add_global_p()`, will no longer execute after the excess elements have been removed (aka butchered). Of note, the majority of `inline_text()` calls will continue to execute properly.

Usage

tbl_butcher(x, include = NULL)

Arguments

- **x**
  - a gtsummary object
- **include**
  - names of additional elements to retain in the gtsummary object. `c("table_body", "table_styling")` will always be retained. Default is NULL

Value

a gtsummary object

Examples

tbl_large <-
  trial %>%
  tbl_uvregression(
    y = age, 
    method = lm
  )

tbl_butchered <-
  tbl_large %>%
  tbl_butcher()

  # size comparison
  object.size(tbl_large) %>% format(units = "Mb")
  object.size(tbl_butchered) %>% format(units = "Mb")
**tbl_continuous**

Summarize a continuous variable

---

**Description**

*Experimental* Summarize a continuous variable by one or more categorical variables

**Usage**

```r
tbl_continuous(
  data,
  variable,
  include = everything(),
  digits = NULL,
  by = NULL,
  statistic = NULL,
  label = NULL
)
```

**Arguments**

- `data` A data frame
- `variable` Variable name of the continuous column to be summarized
- `include` variables to include in the summary table. Default is `everything()`
- `digits` List of formulas specifying the number of decimal places to round continuous summary statistics. If not specified, an appropriate number of decimals to round statistics will be guessed based on the the variable’s distribution.
- `by` A column name (quoted or unquoted) in `data`. Summary statistics will be calculated separately for each level of the `by` variable (e.g. `by = trt`). If `NULL`, summary statistics are calculated using all observations. To stratify a table by two or more variables, use `tbl_strata()`
- `statistic` List of formulas specifying types of summary statistics to display for each variable. The default is `everything() ~ {median} ((p25), (p75))`
- `label` List of formulas specifying variables labels, e.g. `list(age ~ "Age", stage ~ "Path T Stage")`. If a variable’s label is not specified here, the label attribute (`attr(data$age, "label")`) is used. If attribute label is `NULL`, the variable name will be used.

**Value**

a `gtsummary` table

**Example Output**

---

**See Also**

Review list, formula, and selector syntax used throughout `gtsummary` Other `tbl_continuous` tools: `add_overall()`, `add_p.tbl_continuous()`
Examples

# Example 1 ----------------------------------
expr1 <-
tbl_continuous(
  data = trial,
  variable = age,
  by = trt,
  include = grade
)

# Example 2 ----------------------------------
expr2 <-
tbl_continuous(
  data = trial,
  variable = age,
  include = c(trt, grade)
)

tbl_cross  
Create a cross table of summary statistics

Description

The function creates a cross table of two categorical variables.

Usage

tbl_cross(
  data, 
  row = NULL, 
  col = NULL, 
  label = NULL, 
  statistic = NULL, 
  digits = NULL, 
  percent = c("none", "column", "row", "cell"), 
  margin = c("column", "row"), 
  missing = c("ifany", "always", "no"), 
  missing_text = "Unknown", 
  margin_text = "Total"
)

Arguments

data  
A data frame

row  
A column name in data= to be used for the rows of cross table.

col  
A column name in data= to be used for the columns of cross table.

label  
List of formulas specifying variables labels, e.g. list(age ~ "Age", stage ~ "Path T Stage"). If a variable’s label is not specified here, the label attribute (attr(data$age, "label")). If attribute label is NULL, the variable name will be used.
### statistic
A string with the statistic name in curly brackets to be replaced with the numeric statistic (see glue::glue). The default is \{n\}. If percent argument is "column", "row", or "cell", default is "{n} \{(p)\}%\)".

### digits
Specifies the number of decimal places to round the summary statistics. By default integers are shown to the zero decimal places, and percentages are formatted with style_percent(). If you would like to modify either of these, pass a vector of integers indicating the number of decimal places to round the statistics. For example, if the statistic being calculated is "\{(n) \{(p)\}\}%" and you want the percent rounded to 2 decimal places use digits = c(0, 2). User may also pass a styling function: digits = style_sigfig

### percent
Indicates the type of percentage to return. Must be one of "none", "column", "row", or "cell". Default is "cell" when \(N\) or \(p\) is used in statistic.

### margin
Indicates which margins to add to the table. Default is c("row", "column"). Use margin = NULL to suppress both row and column margins.

### missing
Indicates whether to include counts of NA values in the table. Allowed values are "no" (never display NA values), "ifany" (only display if any NA values), and "always" (includes NA count row for all variables). Default is "ifany".

### missing_text
String to display for count of missing observations. Default is "Unknown".

### margin_text
Text to display for margin totals. Default is "Total"

#### Value
A tbl_cross object

#### Example Output

#### Author(s)
Karissa Whiting, Daniel D. Sjoberg

#### See Also
Review list, formula, and selector syntax used throughout gtsummary

Other tbl_cross tools: add_p.tbl_cross(), inline_text.tbl_cross()

#### Examples

```r
# Example 1 ----------------------------------
tbl_cross_ex1 <-
trial %>%
tbl_cross(row = trt, col = response) %>%
bold_labels()

# Example 2 ----------------------------------
tbl_cross_ex2 <-
trial %>%
tbl_cross(row = stage, col = trt, percent = "cell") %>%
add_p() %>%
bold_labels()
```
Create a table of summary statistics using a custom summary function

Description

[Experimental] The tbl_custom_summary() function calculates descriptive statistics for continuous, categorical, and dichotomous variables. This function is similar to tbl_summary() but allows you to provide a custom function in charge of computing the statistics (see Details).

Usage

tbl_custom_summary(
  data,
  by = NULL,
  label = NULL,
  stat_fns,
  statistic,
  digits = NULL,
  type = NULL,
  value = NULL,
  missing = NULL,
  missing_text = NULL,
  include = everything(),
  overall_row = FALSE,
  overall_row_last = FALSE,
  overall_row_label = NULL
)

Arguments

data A data frame
by A column name (quoted or unquoted) in data. Summary statistics will be calculated separately for each level of the by variable (e.g. by = trt). If NULL, summary statistics are calculated using all observations. To stratify a table by two or more variables, use tbl_strata()
label List of formulas specifying variables labels, e.g. list(age ~ "Age", stage ~ "Path T Stage"). If a variable’s label is not specified here, the label attribute (attr(data$age, "label")) is used. If attribute label is NULL, the variable name will be used.
stat_fns Formula or list of formulas specifying the function to be used to compute the statistics (see below for details and examples). You can also use dedicated helpers such as continuous_summary(), ratio_summary() or proportion_summary().
statistic List of formulas specifying the glue::glue() pattern to display the statistics for each variable. The statistics should be returned by the functions specified in stat_fns (see below for details and examples).
digits List of formulas specifying the number of decimal places to round summary statistics. If not specified, tbl_summary guesses an appropriate number of decimals to round statistics. When multiple statistics are displayed for a single variable, supply a vector rather than an integer. For example, if the statistic being calculated is "{mean} \{sd\}" and you want the mean rounded to 1 decimal
place, and the SD to 2 use digits = list(age ~ c(1, 2)). User may also pass a styling function: digits = age ~ style_s

type
List of formulas specifying variable types. Accepted values are c("continuous", "continuous2", "categorical", "dichotomous"), e.g. type = list(age ~ "continuous", female ~ "dichotomous"). If type not specified for a variable, the function will default to an appropriate summary type. See below for details.

value
List of formulas specifying the value to display for dichotomous variables. gt-summary selectors, e.g. all_dichotomous(), cannot be used with this argument. See below for details.

missing
Indicates whether to include counts of NA values in the table. Allowed values are "no" (never display NA values), "ifany" (only display if any NA values), and "always" (includes NA count row for all variables). Default is "ifany".

missing_text
String to display for count of missing observations. Default is "Unknown".

include
variables to include in the summary table. Default is everything().

overall_row
Logical indicator to display an overall row. Default is FALSE. Use add_overall() to add an overall column.

overall_row_last
Logical indicator to display overall row last in table. Default is FALSE, which will display overall row first.

overall_row_label
String indicating the overall row label. Default is "Overall".

Value
A tbl_custom_summary and tbl_summary object

Similarities with tbl_summary()

Please refer to the help file of tbl_summary() regarding the use of select helpers, and arguments include, by, type, value, digits, missing and missing_text.

stat_fns argument

The stat_fns argument specify the custom function(s) to be used for computing the summary statistics. For example, stat_fns = everything() ~ foo.

Each function may take the following arguments: foo(data, full_data, variable, by, type, ...

• data= is the input data frame passed to tbl_custom_summary(), subset according to the level of by or variable if any, excluding NA values of the current variable
• full_data= is the full input data frame passed to tbl_custom_summary()
• variable= is a string indicating the variable to perform the calculation on
• by= is a string indicating the by variable from tbl_custom_summary=, if present
• type= is a string indicating the type of variable (continuous, categorical, ...)
• stat_display= a string indicating the statistic to display (for the statistic argument, for that variable)

The user-defined does not need to utilize each of these inputs. It’s encouraged the user-defined function accept ... as each of the arguments will be passed to the function, even if not all inputs are utilized by the user’s function, e.g. foo(data, ...) (see examples).

The user-defined function should return a one row dplyr::tibble() with one column per summary statistics (see examples).
statistic argument

The statistic argument specifies the statistics presented in the table. The input is a list of formulas that specify the statistics to report. For example, `statistic = list(age ~ "{mean} ({sd})")`. A statistic name that appears between curly brackets will be replaced with the numeric statistic (see `glue::glue()`). All the statistics indicated in the statistic argument should be returned by the functions defined in the `stat_fns` argument.

When the summary type is "continuous2", pass a vector of statistics. Each element of the vector will result in a separate row in the summary table.

For both categorical and continuous variables, statistics on the number of missing and non-missing observations and their proportions are also available to display.

- \{N\_obs\} total number of observations
- \{N\_miss\} number of missing observations
- \{N\_nonmiss\} number of non-missing observations
- \{p\_miss\} percentage of observations missing
- \{p\_nonmiss\} percentage of observations not missing

Note that for categorical variables, \{N\_obs\}, \{N\_miss\} and \{N\_nonmiss\} refer to the total number, number missing and number non missing observations in the denominator, not at each level of the categorical variable.

It is recommended to use `modify_footnote()` to properly describe the displayed statistics (see examples).

Caution

The returned table is compatible with all `gtsummary` features applicable to a `tbl_summary` object, like `add_overall()`, `modify_footnote()` or `bold_labels()`.

However, some of them could be inappropriate in such case. In particular, `add_p()` do not take into account the type of displayed statistics and always return the p-value of a comparison test of the current variable according to the by groups, which may be incorrect if the displayed statistics refer to a third variable.

Example Output

Author(s)

Joseph Larmarange

See Also

Review list, formula, and selector syntax used throughout `gtsummary`

Other `tbl_summary` tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold_italicize_labels_levels()`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify()`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Other `tbl_custom_summary` tools: `add_overall()`, `continuous_summary()`, `proportion_summary()`, `ratio_summary()`
Examples

# Example 1 ----------------------------------
my_stats <- function(data, ...) {
  marker_sum <- sum(data$marker, na.rm = TRUE)
  mean_age <- mean(data$age, na.rm = TRUE)
  dplyr::tibble(
    marker_sum = marker_sum,
    mean_age = mean_age
  )
}

my_stats(trial)

tbl_custom_summary_ex1 <-
  trial %>%
  tbl_custom_summary(
    include = c("stage", "grade"),
    by = "trt",
    stat_fns = everything() ~ my_stats,
    statistic = everything() ~ "A: {mean_age} - S: {marker_sum}"
  ) %>%
  add_overall(last = TRUE) %>%
  modify_footnote(
    update = all_stat_cols() ~ "A: mean age - S: sum of marker"
  ) %>%
  bold_labels()

# Example 2 ----------------------------------
# Use `data[[variable]]` to access the current variable
mean_ci <- function(data, variable, ...) {
  test <- t.test(data[[variable]])
  dplyr::tibble(
    mean = test$estimate,
    conf.low = test$conf.int[1],
    conf.high = test$conf.int[2]
  )
}

tbl_custom_summary_ex2 <-
  trial %>%
  tbl_custom_summary(
    include = c("marker", "ttdeath"),
    by = "trt",
    stat_fns = ~mean_ci,
    statistic = ~"(mean) [{conf.low}; {conf.high}]"
  ) %>%
  add_overall(last = TRUE) %>%
  modify_footnote(
    update = all_stat_cols() ~ "mean [95% CI]"
  )

# Example 3 ----------------------------------
# Use "full_data" to access the full datasets
# Returned statistic can also be a character

diff_to_great_mean <- function(data, full_data, ...) {
  mean <- mean(data$marker, na.rm = TRUE)
  great_mean <- mean(full_data$marker, na.rm = TRUE)
  diff <- mean - great_mean
  dplyr::tibble(
    mean = mean,
    great_mean = great_mean,
    diff = diff,
    level = ifelse(diff > 0, "high", "low")
  )
}

tbl_custom_summary_ex3 <-
  trial %>%
  tbl_custom_summary(
    include = c("grade", "stage"),
    by = "trt",
    stat_fns = ~diff_to_great_mean,
    statistic = ~"{mean} ({level}, diff: {diff})",
    overall_row = TRUE
  ) %>%
  bold_labels()

---

**tbl_merge**

_Merge two or more gtsummary objects_

**Description**

Merges two or more tbl_regression, tbl_uvregression, tbl_stack, tbl_summary, or tbl_svysummary objects and adds appropriate spanning headers.

**Usage**

```r
tbl_merge(tbls, tab_spanner = NULL)
```

**Arguments**

- **tbls** List of gtsummary objects to merge
- **tab_spanner** Character vector specifying the spanning headers. Must be the same length as tbls. The strings are interpreted with gt::md. Must be same length as tbls argument. Default is NULL, and places a default spanning header. If FALSE, no header will be placed.

**Value**

A tbl_merge object

**Example Output**
Author(s)

Daniel D. Sjoberg

See Also

tbl_stack

Other tbl_regression tools: add_global_p(), add_q(), bold_italicize_labels_levels, combine_terms(), inline_text.tbl_regression(), modify.tbl_regression(), tbl_split(), tbl_stack(), tbl_strata()

Other tbl_uvregression tools: add_global_p(), add_q(), bold_italicize_labels_levels, inline_text.tbl_uvregression(), modify.tbl_split(), tbl_stack(), tbl_strata(), tbl_uvregression()

Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(), bold_italicize_labels_levels, inline_text.tbl_summary(), inline_text.tbl_survfit(), modify, separate_p_footnotes(), tbl_custom_summary(), tbl_split(), tbl_stack(), tbl_strata(), tbl_summary()

Other tbl_survfit tools: add_n.tbl_survfit(), add_nevent.tbl_survfit(), add_p.tbl_survfit(), modify.tbl_split(), tbl_stack(), tbl_strata(), tbl_survfit()

Other tbl_svysummary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_svysummary(), add_q(), add_stat_label(), modify, separate_p_footnotes(), tbl_split(), tbl_stack(), tbl_strata(), tbl_svysummary()

Examples

# Example 1 ----------------------------------
# Side-by-side Regression Models
library(survival)
t1 <-
  glm(response ~ trt + grade + age, trial, family = binomial) %>%
  tbl_regression(exponentiate = TRUE)
t2 <-
  coxph(Surv(ttdeath, death) ~ trt + grade + age, trial) %>%
  tbl_regression(exponentiate = TRUE)
tbl_merge_ex1 <-
tbl_merge(
  tbls = list(t1, t2),
  tab_spanner = c("**Tumor Response**", "**Time to Death**")
)

# Example 2 ----------------------------------
# Descriptive statistics alongside univariate regression, with no spanning header
t3 <-
  trial[c("age", "grade", "response")]
  tbl_summary(missing = "no") %>%
  add_n() %>%
  modify_header(stat_0 ~ "**Summary Statistics**")
t4 <-
  tbl_uvregression(
    trial[c("ttdeath", "death", "age", "grade", "response")],
    method = coxph,
    y = Surv(ttdeath, death),
    exponentiate = TRUE,
    hide_n = TRUE
)
tbl_regression

Display regression model results in table

Description

This function takes a regression model object and returns a formatted table that is publication-ready. The function is highly customizable allowing the user to obtain a bespoke summary table of the regression model results. Review the tbl_regression vignette for detailed examples.

Usage

tbl_regression(x, ...)

## Default S3 method:
tbl_regression(
  x,
  label = NULL,
  exponentiate = FALSE,
  include = everything(),
  show_single_row = NULL,
  conf.level = NULL,
  intercept = FALSE,
  estimate_fun = NULL,
  pvalue_fun = NULL,
  tidy_fun = NULL,
  add_estimate_to_reference_rows = FALSE,
  conf.int = NULL,
  ...
)

Arguments

x Regression model object

... [Experimental] Additional arguments passed to broom.helpers::tidy_plus_plus(). See ?tidy_plus_plus_dots for details.

label List of formulas specifying variables labels, e.g. list(age ~ "Age", stage ~ "Path T Stage")

exponentiate Logical indicating whether to exponentiate the coefficient estimates. Default is FALSE.

include Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is everything().

show_single_row By default categorical variables are printed on multiple rows. If a variable is dichotomous (e.g. Yes/No) and you wish to print the regression coefficient on a single row, include the variable name(s) here--quoted and unquoted variable name accepted.
conf.level  Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval.

intercept  Logical argument indicating whether to include the intercept in the output. Default is FALSE.

estimate_fun  Function to round and format coefficient estimates. Default is style_sigfig when the coefficients are not transformed, and style_ratio when the coefficients have been exponentiated.

pvalue_fun  Function to round and format p-values. Default is style_pvalue. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. pvalue_fun = function(x) style_pvalue(x, digits = 2) or equivalently, purrr::partial(style_pvalue, digits = 2)).

tidy_fun  Option to specify a particular tidier function for the model. Default is to use broom::tidy(), but if an error occurs then tidying of the model is attempted with parameters::model_parameters(), if installed.

add_estimate_to_reference_rows  Add a reference value. Default is FALSE.

conf.int  Logical indicating whether or not to include a confidence interval in the output. Defaults to TRUE.

Value

A tbl_regression object

Methods

The default method for tbl_regression() model summary uses broom::tidy(x) to perform the initial tidying of the model object. There are, however, a few models that use modifications.

- "parsnip/workflows": If the model was prepared using parsnip/workflows, the original model fit is extracted and the original x= argument is replaced with the model fit. This will typically go unnoticed; however, if you’ve provided a custom tidier in tidy_fun= the tidier will be applied to the model fit object and not the parsnip/workflows object.

- "survreg": The scale parameter is removed, broom::tidy(x) %>% dplyr::filter(term != "Log(scale)"

- "multinom": This multinomial outcome is complex, with one line per covariate per outcome (less the reference group)

- "gam": Uses the internal tidier tidy_gam() to print both parametric and smooth terms.

- "tidyrr": Uses the tidier tidycmprsk::tidy() to print the model terms.

- "lmerMod", "glmerMod", "glmmTMB", "glmmadmb", "stanreg", "brmsfit": These mixed effects models use broom.mixed::tidy(x, effects = "fixed"). Specify tidy_fun = broom.mixed::tidy to print the random components.

Example Output

Author(s)

Daniel D. Sjoberg
See Also

See tbl_regression vignette for detailed examples

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_regression tools: add_global_p(), add_q(), bolditalicize_labels_levels, combine_terms(), inline_text.tbl_regression(), modify, tbl_merge(), tbl_split(), tbl_stack(), tbl_strata()

Examples

# Example 1 ----------------------------------
library(survival)
tbl_regression_ex1 <-
  coxph(Surv(ttdeath, death) ~ age + marker, trial) %>%
  tbl_regression(exponentiate = TRUE)

# Example 2 ----------------------------------
tbl_regression_ex2 <-
  glm(response ~ age + grade, trial, family = binomial(link = "logit")) %>%
  tbl_regression(exponentiate = TRUE)

# Example 3 ----------------------------------
# round all estimates to 3 decimal places
suppressMessages(library(lme4))
tbl_regression_ex3 <-
  lmer(hp ~ am + (1 | gear), data = mtcars) %>%
  tbl_regression(estimate_fun = function(x) style_number(x, digits = 3))

tbl_split

Split gtsummary table

Description

[Experimental] The tbl_split function splits a single gtsummary table into multiple tables. Updates to the print method are expected.

Usage

tbl_split(x, ...)

## S3 method for class 'gtsummary'
tbl_split(x, variables, ...)

## S3 method for class 'tbl_split'
print(x, ...)

Arguments

x  gtsummary table
...
not used
variables  variables at which to split the gtsummary table rows (tables will be separated after each of these variables)
Value
tbl_split object

See Also
Other tbl_regression tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels`, `combine_terms()`, `inline_text.tbl_regression()`, `modify.tbl_merge()`, `tbl_regression()`, `tbl_stack()`

Other tbl_uvregression tools: `add_global_p()`, `add_q()`, `bold_italicize_labels_levels`, `inline_text.tbl_uvregression()`, `modify.tbl_merge()`, `tbl_stack()`

Other tbl_summary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_summary()`, `add_q()`, `add_stat_label()`, `bold_italicize_labels_levels`, `inline_text.tbl_summary()`, `inline_text.tbl_survfit()`, `modify`, `separate_p_footnotes()`, `tbl_custom_summary()`, `tbl_merge()`, `tbl_stack()`, `tbl_strata()`, `tbl_summary()`

Other tbl_survfit tools: `add_n.tbl_survfit()`, `add_nevent.tbl_survfit()`, `add_p.tbl_survfit()`, `modify`, `tbl_merge()`, `tbl_stack()`, `tbl_strata()`, `tbl_survfit()`

Other tbl_svysummary tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_q()`, `add_stat_label()`, `modify`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_stack()`, `tbl_strata()`, `tbl_svysummary()`

Examples
tbl <-
tbl_summary(trial) %>%
tbl_split(variables = c(marker, grade))

---

**tbl_stack**

*Stacks two or more gtsummary objects*

**Description**

Assists in patching together more complex tables. `tbl_stack()` appends two or more `tbl_regression`, `tbl_summary`, `tbl_svysummary`, or `tbl_merge` objects. Column attributes, including number formatting and column footnotes, are retained from the first passed gtsummary object.

**Usage**

```
tbl_stack(tbls, group_header = NULL, quiet = NULL)
```

**Arguments**

- **tbls** List of gtsummary objects
- **group_header** Character vector with table headers where length matches the length of `tbls`
- **quiet** Logical indicating whether to print messages in console. Default is `FALSE`

**Value**

A `tbl_stack` object
Example Output

Author(s)
Daniel D. Sjoberg

See Also
- tbl_merge

Other tbl_summary tools:
  - add_n.tbl_summary()
  - add_overall()
  - add_p.tbl_summary()
  - add_q()
  - add_stat_label()
  - bold_italicize_labels_levels
  - inline_text.tbl_summary()
  - inline_text.tbl_survfit()
  - modify
  - separate_p_footnotes()
  - tbl_custom_summary()
  - tbl_merge()
  - tbl_split()
  - tbl_strata()

Other tbl_svysummary tools:
  - add_n.tbl_summary()
  - add_overall()
  - add_p.tbl_svysummary()
  - add_q()
  - add_stat_label()
  - modify
  - separate_p_footnotes()
  - tbl_merge()
  - tbl_split()
  - tbl_strata()

Other tbl_regression tools:
  - add_global_p()
  - add_q()
  - bold_italicize_labels_levels
  - combine_terms()
  - inline_text.tbl_regression()
  - modify
  - tbl_merge()
  - tbl_regression()
  - tbl_split()
  - tbl_strata()

Other tbl_uvregression tools:
  - add_global_p()
  - add_q()
  - bold_italicize_labels_levels
  - inline_text.tbl_uvregression()
  - modify
  - tbl_merge()
  - tbl_split()
  - tbl_strata()
  - tbl_uvregression()

Other tbl_survfit tools:
  - add_n.tbl_survfit()
  - add_nevent.tbl_survfit()
  - add_p.tbl_survfit()
  - modify
  - tbl_merge()
  - tbl_split()
  - tbl_strata()
  - tbl_survfit()

Examples

# Example 1 ----------------------------------
# stacking two tbl_regression objects

t1 <-
  glm(response ~ trt, trial, family = binomial) %>%
  tbl_regression(
    exponentiate = TRUE,
    label = list(trt ~ "Treatment (unadjusted)")
  )

t2 <-
  glm(response ~ trt + grade + stage + marker, trial, family = binomial) %>%
  tbl_regression(
    include = "trt",
    exponentiate = TRUE,
    label = list(trt ~ "Treatment (adjusted)")
  )

tbl_stack_ex1 <- tbl_stack(list(t1, t2))

# Example 2 ----------------------------------
# stacking two tbl_merge objects

t3 <-
  coxph(Surv(ttdeath, death) ~ trt, trial) %>%
  tbl_regression(
    exponentiate = TRUE,
    label = list(trt ~ "Treatment (unadjusted)")
  )
t4 <-
  coxph(Surv(ttdeath, death) ~ trt + grade + stage + marker, trial) %>%
  tbl_regression(
    include = "trt",
    exponentiate = TRUE,
    label = list(trt ~ "Treatment (adjusted)"
  )
)

# first merging, then stacking
row1 <- tbl_merge(list(t1, t3), tab_spanner = c("Tumor Response", "Death"))
row2 <- tbl_merge(list(t2, t4))
tbl_stack_ex2 <-
  tbl_stack(list(row1, row2), group_header = c("Unadjusted Analysis", "Adjusted Analysis"))

---

tbl_strata

Stratified gtsummary tables

Description

[Maturing] Build a stratified gtsummary table. Any gtsummary table that accepts a data frame as its first argument can be stratified.

- In tbl_strata(), the stratified or subset data frame is passed to the function in .tbl_fun=, e.g. purrr::map(data, .tbl_fun).
- In tbl_strata2(), both the stratified data frame and the strata level are passed to .tbl_fun=, e.g. purrr::map2(data, strata, .tbl_fun)

Usage

tbl_strata(
data,
strata,
.tbl_fun,
...
.sep = "", 
.combine_with = c("tbl_merge", "tbl_stack"),
.combine_args = NULL,
.header = ifelse(.combine_with == "tbl_merge", "**(strata)**", "{strata}"),
.stack_group_header = NULL,
.quiet = NULL
)

tbl_strata2(
data,
strata,
.tbl_fun,
...
.sep = "", 

Arguments

data a data frame or survey object
strata character vector or tidy-selector of columns in data to stratify results by
.tbl_fun A function or formula. If a function, it is used as is. If a formula, e.g. ~ .x %>%
tbl_summary() %>% add_p(), it is converted to a function. The stratified data
frame is passed to this function.
... Additional arguments passed on to the .tbl_fun function.
.sep when more than one stratifying variable is passed, this string is used to separate
the levels in the spanning header. Default is ", "
.combine_with One of c("tbl_merge", "tbl_stack"). Names the function used to combine
the stratified tables.
.combine_args named list of arguments that are passed to function specified in .combine_with=
.header String indicating the headers that will be placed. Default is "**(strata)**" when
.combine_with = "tbl_merge" and "{strata}" when .combine_with
= "tbl_stack". Items placed in curly brackets will be evaluated according to
\texttt{glue::glue()} syntax.
  • strata stratum levels
  • n N within stratum
  • N Overall N
The evaluated value of .header= is also available within tbl_strata2(.tbl_fun=)
.stack_group_header DEPRECATED.
.quiet Logical indicating whether to print messages in console. Default is FALSE

Tips

• 
  • \texttt{tbl_summary()} 
    -- The number of digits continuous variables are rounded to is determined separately within
each stratum of the data frame. Set the \texttt{digits=} argument to ensure continuous variables
are rounded to the same number of decimal places.
    -- If some levels of a categorical variable are unobserved within a stratum, convert the vari-
able to a factor to ensure all levels appear in each stratum’s summary table.

Example Output

Author(s)

Daniel D. Sjoberg
See Also

Other tbl_regression tools: add_global_p(), add_q(), bold_italicize_labels_levels, combine_terms(), inline_text.tbl_regression(), modify.tbl_merge(), tbl_regression(), tbl_split(), tbl_stack()

Other tbl_uvregression tools: add_global_p(), add_q(), bold_italicize_labels_levels, inline_text.tbl_uvregrgression(), modify.tbl_merge(), tbl_split(), tbl_stack(), tbl_uvregrgression()

Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(), bold_italicize_labels_levels, inline_text.tbl_summary(), inline_text.tbl_survfit(), modify.separate_p_footnotes(), tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(), tbl_summary()

Other tbl_survfit tools: add_n.tbl_survfit(), add_nevent.tbl_survfit(), add_p.tbl_survfit(), modify.tbl_merge(), tbl_split(), tbl_stack(), tbl_survfit()

Other tbl_svysummary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_svysummary(), add_q(), add_stat_label(), modify.separate_p_footnotes(), tbl_merge(), tbl_split(), tbl_stack(), tbl_svysummary()

Examples

# Example 1 ----------------------------------
# Example 2 ----------------------------------
Create a table of summary statistics

Description

The tbl_summary function calculates descriptive statistics for continuous, categorical, and dichotomous variables. Review the tbl_summary vignette for detailed examples.

Usage

```r
tbl_summary(
  data, 
  by = NULL, 
  label = NULL, 
  statistic = NULL, 
  digits = NULL, 
  type = NULL, 
  value = NULL, 
  missing = NULL, 
  missing_text = NULL, 
  sort = NULL, 
  percent = NULL, 
  include = everything()
)
```

Arguments

- **data**: A data frame
- **by**: A column name (quoted or unquoted) in `data`. Summary statistics will be calculated separately for each level of the `by` variable (e.g. `by = trt`). If `NULL`, summary statistics are calculated using all observations. To stratify a table by two or more variables, use `tbl_strata()`.
- **label**: List of formulas specifying variables labels, e.g. `list(age ~ "Age", stage ~ "Path T Stage")`. If a variable's label is not specified here, the label attribute (`attr(data$age, "label")`) is used. If attribute label is `NULL`, the variable name will be used.
- **statistic**: List of formulas specifying types of summary statistics to display for each variable. The default is `list(all_continuous() ~ "(median) {(p25}, (p75)})`, `all_categorical() ~ "(n) {{p}%}\)`. See below for details.
- **digits**: List of formulas specifying the number of decimal places to round summary statistics. If not specified, `tbl_summary` guesses an appropriate number of decimals to round statistics. When multiple statistics are displayed for a single variable, supply a vector rather than an integer. For example, if the statistic being calculated is "{mean} {{sd)}" and you want the mean rounded to 1 decimal place, and the SD to 2 use `digits = list(age ~ c(1, 2))`. User may also pass a styling function: `digits = age ~ style_sigfig`
- **type**: List of formulas specifying variable types. Accepted values are `c("continuous", "continuous2", "categorical", "dichotomous")`. e.g. `type = list(age ~ "continuous", female ~ "dichotomous")`. If type not specified for a variable, the function will default to an appropriate summary type. See below for details.
**Value**

A tbl_summary object

**select helpers**

Select helpers from the tidyselect\ package and \gtsummary\ package are available to modify default behavior for groups of variables. For example, by default continuous variables are reported with the median and IQR. To change all continuous variables to mean and standard deviation use `statistic = list(all_continuous() ~ "{mean} ({sd})")`. All columns with class logical are displayed as dichotomous variables showing the proportion of events that are TRUE on a single row. To show both rows (i.e. a row for TRUE and a row for FALSE) use `type = list(where(is.logical) ~ "categorical")`. The select helpers are available for use in any argument that accepts a list of formulas (e.g. statistic, type, digits, value, sort, etc.) Read more on the syntax used through the package.

**type argument**

The tbl_summary() function has four summary types:

- "continuous" summaries are shown on a single row. Most numeric variables default to summary type continuous.
- "continuous2" summaries are shown on 2 or more rows
- "categorical" multi-line summaries of nominal data. Character variables, factor variables, and numeric variables with fewer than 10 unique levels default to type categorical. To change a numeric variable to continuous that defaulted to categorical, use `type = list(varname ~ "continuous")`
- "dichotomous" categorical variables that are displayed on a single row, rather than one row per level of the variable. Variables coded as TRUE/FALSE, 0/1, or yes/no are assumed to be dichotomous, and the TRUE, 1, and yes rows are displayed. Otherwise, the value to display must be specified in the value argument, e.g. `value = list(varname ~ "level to show")`
**statistic argument**

The statistic argument specifies the statistics presented in the table. The input is a list of formulas that specify the statistics to report. For example, `statistic = list(age ~ "{mean} (sd)")` would report the mean and standard deviation for age; `statistic = list(all_continuous() ~ "{mean} (sd)")` would report the mean and standard deviation for all continuous variables. A statistic name that appears between curly brackets will be replaced with the numeric statistic (see `glue::glue`).

For categorical variables the following statistics are available to display.

- `{n}` frequency
- `{N}` denominator, or cohort size
- `{p}` formatted percentage

For continuous variables the following statistics are available to display.

- `{median}` median
- `{mean}` mean
- `{sd}` standard deviation
- `{var}` variance
- `{min}` minimum
- `{max}` maximum
- `{sum}` sum
- `{p##}` any integer percentile, where ## is an integer from 0 to 100
- `{foo}` any function of the form `foo(x)` is accepted where x is a numeric vector

When the summary type is "continuous2", pass a vector of statistics. Each element of the vector will result in a separate row in the summary table.

For both categorical and continuous variables, statistics on the number of missing and non-missing observations and their proportions are available to display.

- `{N_obs}` total number of observations
- `{N_miss}` number of missing observations
- `{N_nonmiss}` number of non-missing observations
- `{p_miss}` percentage of observations missing
- `{p_nonmiss}` percentage of observations not missing

Note that for categorical variables, `{N_obs}`, `{N_miss}` and `{N_nonmiss}` refer to the total number, number missing and number non missing observations in the denominator, not at each level of the categorical variable.

**Example Output**

**Author(s)**

Daniel D. Sjoberg
See Also
See tbl_summary vignette for detailed tutorial
See table gallery for additional examples
Review list, formula, and selector syntax used throughout gtsummary
Other tbl_summary tools: add_n.tbl_summary(), add_overall(), add_p.tbl_summary(), add_q(), add_stat_label(), bold_italicize_labels_levels, inline_text.tbl_summary(), inline_text.tbl_survfit(), modify, separate_p_footnotes(), tbl_custom_summary(), tbl_merge(), tbl_split(), tbl_stack(), tbl_strata()

Examples

```
# Example 1 ----------------------------------
tbl_summary_ex1 <-
  trial %>%
    select(age, grade, response) %>%
    tbl_summary()
# Example 2 ----------------------------------
tbl_summary_ex2 <-
  trial %>%
    select(age, grade, response, trt) %>%
    tbl_summary(
      by = trt,
      label = list(age ~ "Patient Age"),
      statistic = list(all_continuous() ~ "\{mean\} \{sd\}"),
      digits = list(age ~ c(0, 1))
    )
# Example 3 ----------------------------------
# for convenience, you can also pass named lists to any arguments
# that accept formulas (e.g label, digits, etc.)
tbl_summary_ex3 <-
  trial %>%
    select(age, trt) %>%
    tbl_summary(
      by = trt,
      label = list(age = "Patient Age")
    )
# Example 4 ----------------------------------
# multi-line summaries of continuous data with type 'continuous2'
tbl_summary_ex4 <-
  trial %>%
    select(age, marker) %>%
    tbl_summary(
      type = all_continuous() ~ "continuous2",
      statistic = all_continuous() ~ "\{median\} \{p25\}, \{p75\}" ~ "\{min\}, \{max\}"
    )
```
### tbl_survfit

**Description**

[Maturing] Function takes a `survfit` object as an argument, and provides a formatted summary table of the results

**Usage**

```r
tbl_survfit(x, ...)
```

#### S3 method for class 'list'

```r
tbl_survfit(
  x,
  times = NULL,
  probs = NULL,
  statistic = NULL,
  label = NULL,
  label_header = NULL,
  estimate_fun = NULL,
  missing = NULL,
  conf.level = 0.95,
  reverse = FALSE,
  quiet = NULL,
  ...
)
```

#### S3 method for class 'survfit'

```r
tbl_survfit(x, ...)
```

#### S3 method for class 'data.frame'

```r
tbl_survfit(x, y, include = everything(), ...)
```

**Arguments**

- **x**
  - a `survfit` object, list of `survfit` objects, or a data frame. If a data frame is passed, a list of `survfit` objects is constructed using each variable as a stratifying variable.
- **...**
  - For `tbl_survfit.data.frame()` and `tbl_survfit.survfit()` the arguments are passed to `tbl_survfit.list()`. They are not used when `tbl_survfit.list()` is called directly.
- **times**
  - numeric vector of times for which to return survival probabilities.
- **probs**
  - numeric vector of probabilities with values in (0,1) specifying the survival quantiles to return.
- **statistic**
  - string defining the statistics to present in the table. Default is “{estimate} ({conf.low}, {conf.high})”
- **label**
  - List of formulas specifying variables labels, e.g. `list(age ~ "Age, yrs", stage ~ "Path T Stage")`, or a string for a single variable table.
tbl_survfit

- **label_header**: string specifying column labels above statistics. Default is "\(\{\text{prob}\}\) Percentile" for survival percentiles, and "Time \(\{\text{time}\}\)" for n-year survival estimates.
- **estimate_fun**: function to format the Kaplan-Meier estimates. Default is `style_percent()` for survival probabilities and `style_sigfig` for survival times.
- **missing**: text to fill when estimate is not estimable. Default is "--".
- **conf.level**: confidence level for confidence intervals. Default is 0.95.
- **reverse**: flip the probability reported, i.e. \(1 - \text{estimate}\). Default is `FALSE`. Does not apply to survival quantile requests.
- **quiet**: logical indicating whether to print messages in console. Default is `FALSE`.
- **y**: outcome call, e.g. \(y = \text{Surv(ttdeath, death)}\).
- **include**: variable to include as stratifying variables.

**Example Output**

**Author(s)**

Daniel D. Sjoberg

**See Also**

Review list, formula, and selector syntax used throughout gtsummary.

Other tbl_survfit tools: `add_n.tbl_survfit()`, `add_nevent.tbl_survfit()`, `add_p.tbl_survfit()`, `modify.tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`

**Examples**

```r
library(survival)

# Example 1 ----------------------------------
# Pass single survfit() object
tbl_survfit_ex1 <- tbl_survfit(
  survfit(Surv(ttdeath, death) ~ trt, trial),
  times = c(12, 24),
  label_header = "**{time} Month**"
)

# Example 2 ----------------------------------
# Pass a data frame
tbl_survfit_ex2 <- tbl_survfit(
  trial,
  y = Surv(ttdeath, death),
  include = c(trt, grade),
  probs = 0.5,
  label_header = "**Median Survival**"
)

# Example 3 ----------------------------------
# Pass a list of survfit() objects
tbl_survfit_ex3 <-
  list(
    survfit(Surv(ttdeath, death) ~ 1, trial),
```


```r
# Example 4 Competing Events Example ---------
# adding a competing event for death (cancer vs other causes)
set.seed(1123)
library(dplyr, warn.conflicts = FALSE, quietly = TRUE)
trial2 <- trial %>%
  mutate(
    death_cr = case_when(
      death == 0 ~ "censor",
      runif(n()) < 0.5 ~ "death from cancer",
      TRUE ~ "death other causes"
    )
  )
survfit_cr_ex4 <-
  survfit(Surv(ttdeath, death_cr) ~ grade, data = trial2)
  tbl_survfit(times = c(12, 24), label = "Tumor Grade")
```

---

### tbl_svysummary

Create a table of summary statistics from a survey object

**Description**

The `tbl_svysummary` function calculates descriptive statistics for continuous, categorical, and dichotomous variables taking into account survey weights and design. It is similar to `tbl_summary()`.

**Usage**

```r
tbl_svysummary(
  data,
  by = NULL,
  label = NULL,
  statistic = NULL,
  digits = NULL,
  type = NULL,
  value = NULL,
  missing = NULL,
  missing_text = NULL,
  sort = NULL,
  percent = NULL,
  include = everything()
)
```

**Arguments**

- `data` A survey object created with `survey::svydesign()`
- `by` A column name (quoted or unquoted) in `data`. Summary statistics will be calculated separately for each level of the `by` variable (e.g. `by = trt`). If `NULL`, summary statistics are calculated using all observations. To stratify a table by two or more variables, use `tbl_strata()`
label
List of formulas specifying variables labels, e.g. `list(age ~ "Age", stage ~ "Path T Stage")`. If a variable’s label is not specified here, the label attribute (`attr(data$age, "label")`) is used. If attribute label is NULL, the variable name will be used.

statistic
List of formulas specifying types of summary statistics to display for each variable. The default is `list(all_continuous() ~ \(\{\text{median}\}\ (\{p25\}, \{p75\})\), all_categorical() ~ \(\{n\}\ (\{p\}\%))\)`. See below for details.

digits
List of formulas specifying the number of decimal places to round summary statistics. If not specified, `tbl_summary` guesses an appropriate number of decimals to round statistics. When multiple statistics are displayed for a single variable, supply a vector rather than an integer. For example, if the statistic being calculated is \(\{\text{mean}\}\ (\{sd\})\) and you want the mean rounded to 1 decimal place, and the SD to 2 use `digits = list(age ~ c(1, 2))`. User may also pass a styling function: `digits = age ~ style_sigfig`

type
List of formulas specifying variable types. Accepted values are `c("continuous", "continuous2", "categorical", "dichotomous")`, e.g. `type = list(age ~ "continuous", female ~ "dichotomous")`. If type not specified for a variable, the function will default to an appropriate summary type. See below for details.

value
List of formulas specifying the value to display for dichotomous variables. gt-summary selectors, e.g. `all_dichotomous()`, cannot be used with this argument. See below for details.

missing
Indicates whether to include counts of NA values in the table. Allowed values are "no" (never display NA values), "ifany" (only display if any NA values), and "always" (includes NA count row for all variables). Default is "ifany".

missing_text
String to display for count of missing observations. Default is "Unknown".

sort
List of formulas specifying the type of sorting to perform for categorical data. Options are frequency where results are sorted in descending order of frequency and alphanumeric, e.g. `sort = list(everything() ~ "frequency")`

percent
Indicates the type of percentage to return. Must be one of "column", "row", or "cell". Default is "column".

include
variables to include in the summary table. Default is `everything()`

Value
A `tbl_svysummary` object

statistic argument
The statistic argument specifies the statistics presented in the table. The input is a list of formulas that specify the statistics to report. For example, `statistic = list(age ~ \("\{\text{mean}\}\ (\{sd\})\")`) would report the mean and standard deviation for age; `statistic = list(all_continuous() ~ \("\{\text{mean}\}\ (\{sd\})\")`) would report the mean and standard deviation for all continuous variables. A statistic name that appears between curly brackets will be replaced with the numeric statistic (see `glue::glue`).

For categorical variables the following statistics are available to display.

- \(\{n\}\) frequency
- \(\{N\}\) denominator, or cohort size
- \(\{p\}\) percentage
• `{p.std.error}` standard error of the sample proportion computed with `survey::svymean()`

• `{deff}` design effect of the sample proportion computed with `survey::svymean()`

• `{n_unweighted}` unweighted frequency

• `{N_unweighted}` unweighted denominator

• `{p_unweighted}` unweighted formatted percentage

For continuous variables the following statistics are available to display.

• `{median}` median

• `{mean}` mean

• `{mean.std.error}` standard error of the sample mean computed with `survey::svymean()`

• `{deff}` design effect of the sample mean computed with `survey::svymean()`

• `{sd}` standard deviation

• `{var}` variance

• `{min}` minimum

• `{max}` maximum

• `{p##}` any integer percentile, where `##` is an integer from 0 to 100

• `{sum}` sum

Unlike `tbl_summary()`, it is not possible to pass a custom function.

For both categorical and continuous variables, statistics on the number of missing and non-missing observations and their proportions are available to display.

• `{N_obs}` total number of observations

• `{N_miss}` number of missing observations

• `{N_nonmiss}` number of non-missing observations

• `{p_miss}` percentage of observations missing

• `{p_nonmiss}` percentage of observations not missing

• `{N_obs_unweighted}` unweighted total number of observations

• `{N_miss_unweighted}` unweighted number of missing observations

• `{N_nonmiss_unweighted}` unweighted number of non-missing observations

• `{p_miss_unweighted}` unweighted percentage of observations missing

• `{p_nonmiss_unweighted}` unweighted percentage of observations not missing

Note that for categorical variables, `{N_obs}`, `{N_miss}` and `{N_nonmiss}` refer to the total number, number missing and number non missing observations in the denominator, not at each level of the categorical variable.

**Example Output**
**type argument**

The `tbl_summary()` function has four summary types:
- "continuous" summaries are shown on a single row. Most numeric variables default to summary type continuous.
- "continuous2" summaries are shown on 2 or more rows
- "categorical" multi-line summaries of nominal data. Character variables, factor variables, and numeric variables with fewer than 10 unique levels default to type categorical. To change a numeric variable to continuous that defaulted to categorical, use `type = list(varname ~ "continuous")`
- "dichotomous" categorical variables that are displayed on a single row, rather than one row per level of the variable. Variables coded as TRUE/FALSE, 0/1, or yes/no are assumed to be dichotomous, and the TRUE, 1, and yes rows are displayed. Otherwise, the value to display must be specified in the `value` argument, e.g. `value = list(varname ~ "level to show")`

**select helpers**

Select helpers from the `tidyselect` package and `gtsummary` package are available to modify default behavior for groups of variables. For example, by default continuous variables are reported with the median and IQR. To change all continuous variables to mean and standard deviation use `statistic = list(all_continuous() ~ "(mean) (sd)")`. All columns with class logical are displayed as dichotomous variables showing the proportion of events that are TRUE on a single row. To show both rows (i.e. a row for TRUE and a row for FALSE) use `type = list(where(is.logical) ~ "categorical")`. The select helpers are available for use in any argument that accepts a list of formulas (e.g. `statistic`, `type`, `digits`, `value`, `sort`, etc.) Read more on the syntax used through the package.

**Author(s)**

Joseph Larmarange

**See Also**

Review `list`, `formula`, and `selector syntax` used throughout gtsummary

Other `tbl_svysummary` tools: `add_n.tbl_summary()`, `add_overall()`, `add_p.tbl_svysummary()`, `add_q()`, `add_stat_label()`, `modify`, `separate_p_footnotes()`, `tbl_merge()`, `tbl_split()`, `tbl_stack()`, `tbl_strata()`

**Examples**

# A simple weighted dataset
tbl_svysummary_ex1 <-
survey::svydesign(~1, data = as.data.frame(Titanic), weights = ~Freq) %>%
tbl_svysummary(by = Survived, percent = "row", include = c(Class, Age))

# Example 2 -------------------------------
# A dataset with a complex design
data(api, package = "survey")
tbl_svysummary_ex2 <-
survey::svydesign(id = ~dnum, weights = ~pw, data = apiclus1, fpc = ~fpc) %>%
tbl_svysummary(by = "both", include = c(api00, stype))
This function estimates univariate regression models and returns them in a publication-ready table. It can create univariate regression models holding either a covariate or outcome constant.

For models holding outcome constant, the function takes as arguments a data frame, the type of regression model, and the outcome variable \( y \). Each column in the data frame is regressed on the specified outcome. The `tbl_uvregression` function arguments are similar to the `tbl_regression` arguments. Review the `tbl_uvregression` vignette for detailed examples.

You may alternatively hold a single covariate constant. For this, pass a data frame, the type of regression model, and a single covariate in the `x=` argument. Each column of the data frame will serve as the outcome in a univariate regression model. Take care using the `x` argument that each of the columns in the data frame are appropriate for the same type of model, e.g. they are all continuous variables appropriate for `lm`, or dichotomous variables appropriate for logistic regression with `glm`.

**Usage**

```r
tbl_uvregression(
  data,
  method,
  y = NULL,
  x = NULL,
  method.args = NULL,
  exponentiate = FALSE,
  label = NULL,
  include = everything(),
  tidy_fun = NULL,
  hide_n = FALSE,
  show_single_row = NULL,
  conf.level = NULL,
  estimate_fun = NULL,
  pvalue_fun = NULL,
  formula = "{y} ~ {x}",
  add_estimate_to_reference_rows = NULL,
  conf.int = NULL,
...
)
```

**Arguments**

- `data` Data frame to be used in univariate regression modeling. Data frame includes the outcome variable(s) and the independent variables. Survey design objects are also accepted.
- `method` Regression method (e.g. `lm`, `glm`, `survival::coxph`, `survey::svyglm`, and more).
- `y` Model outcome (e.g. \( y = \) recurrence or \( y = \) \texttt{Surv(time, recur)}). All other column in data will be regressed on \( y \). Specify one and only one of \( y \) or \( x \)
### tbl_uvregression

| x | Model covariate (e.g. \( x = \text{trt} \)). All other columns in data will serve as the outcome in a regression model with \( x \) as a covariate. Output table is best when \( x \) is a continuous or dichotomous variable displayed on a single row. Specify one and only one of \( y \) or \( x \) |
| method.args | List of additional arguments passed on to the regression function defined by method. |
| exponentiate | Logical indicating whether to exponentiate the coefficient estimates. Default is FALSE. |
| label | List of formulas specifying variables labels, e.g. `list(age ~ "Age", stage ~ "Path T Stage")` |
| include | Variables to include in output. Input may be a vector of quoted variable names, unquoted variable names, or tidyselect select helper functions. Default is `everything()` |
| tidy_fun | Option to specify a particular tidier function for the model. Default is to use `broom::tidy()`, but if an error occurs then tidying of the model is attempted with `parameters::model_parameters()`, if installed. |
| hide_n | Hide N column. Default is FALSE |
| show_single_row | By default categorical variables are printed on multiple rows. If a variable is dichotomous (e.g. Yes/No) and you wish to print the regression coefficient on a single row, include the variable name(s) here–quoted and unquoted variable name accepted. |
| conf.level | Must be strictly greater than 0 and less than 1. Defaults to 0.95, which corresponds to a 95 percent confidence interval. |
| estimate_fun | Function to round and format coefficient estimates. Default is `style_sigfig` when the coefficients are not transformed, and `style_ratio` when the coefficients have been exponentiated. |
| pvalue_fun | Function to round and format p-values. Default is `style_pvalue`. The function must have a numeric vector input (the numeric, exact p-value), and return a string that is the rounded/formatted p-value (e.g. `pvalue_fun = function(x) style_pvalue(x, digits = 2)` or equivalently, `purrr::partial(style_pvalue, digits = 2)`). |
| formula | String of the model formula. Uses `glue::glue` syntax. Default is "\((y) \sim (x)\)". where \((y)\) is the dependent variable, and \((x)\) represents a single covariate. For a random intercept model, the formula may be `formula = "(y) \sim (x) + (1 | gear)"`. |
| add_estimate_to_reference_rows | add a reference value. Default is FALSE |
| conf.int | Logical indicating whether or not to include a confidence interval in the output. Defaults to TRUE. |
| ... | [Experimental] Additional arguments passed to `broom.helpers::tidy_plus_plus()`. See `?tidy_plus_plus_dots` for details. |

### Value

A `tbl_uvregression` object

### Example Output
Methods

The default method for tbl_regression() model summary uses broom::tidy(x) to perform the initial tidying of the model object. There are, however, a few models that use modifications.

- "parsnip/workflows": If the model was prepared using parsnip/workflows, the original model fit is extracted and the original x= argument is replaced with the model fit. This will typically go unnoticed; however, if you’ve provided a custom tidier in tidy_fun= the tidier will be applied to the model fit object and not the parsnip/workflows object.
- "survreg": The scale parameter is removed, broom::tidy(x) %>% dplyr::filter(term != "Log(scale)"
- "multinom": This multinomial outcome is complex, with one line per covariate per outcome (less the reference group)
- "gam": Uses the internal tidier tidy_gam() to print both parametric and smooth terms.
- "tidycrr": Uses the tidier tidycmprsk::tidy() to print the model terms.
- "lmerMod", "glmerMod", "glmmTMB", "glmmadmb", "stanreg", "brmsfit": These mixed effects models use broom.mixed::tidy(x, effects = "fixed"). Specify tidy_fun = broom.mixed::tidy to print the random components.

Author(s)

Daniel D. Sjoberg

See Also

See tbl_regression vignette for detailed examples

Review list, formula, and selector syntax used throughout gtsummary

Other tbl_uvregression tools: add_global_p(), add_q(), bold_italicize_labels_levels, inline_text.tbl_uvregression(), modify, tbl_merge(), tbl_split(), tbl_stack(), tbl_strata()

Examples

```r
# Example 1 ----------------------------------
tbl_uv_ex1 <-
  tbl_uvregression(
    trial[, c("response", "age", "grade")],
    method = glm,
    y = response,
    method.args = list(family = binomial),
    exponentiate = TRUE
  )

# Example 2 ----------------------------------
# rounding pvalues to 2 decimal places
library(survival)
tbl_uv_ex2 <-
  tbl_uvregression(
    trial[, c("ttdeath", "death", "age", "grade", "response")],
    method = coxph,
    y = Surv(ttdeath, death),
    exponentiate = TRUE,
    pvalue_fun = function(x) style_pvalue(x, digits = 2)
  )
```
Available gtsummary themes

Description

[Maturing] The following themes are available to use within the gtsummary package. Print theme elements with `theme_gtsummary_journal(set_theme = FALSE) %>% print()`. Review the themes vignette for details.

Usage

```r
theme_gtsummary_journal(
  journal = c("jama", "lancet", "nejm", "qjecon"),
  set_theme = TRUE
)
```

```r
theme_gtsummary_compact(set_theme = TRUE, font_size = NULL)
```

```r
theme_gtsummary_printer(
  print_engine = c("gt", "kable", "kable_extra", "flextable", "huxtable", "tibble"),
  set_theme = TRUE
)
```

```r
theme_gtsummary_language(
  language = c("de", "en", "es", "fr", "gu", "hi", "is", "ja", "kr", "mr", "nl", "no",
               "pt", "se", "zh-cn", "zh-tw"),
  decimal.mark = NULL,
  big.mark = NULL,
  iqr.sep = NULL,
  ci.sep = NULL,
  set_theme = TRUE
)
```

```r
theme_gtsummary_continuous2(
  statistic = "{median} \{p25, {p75}}",
  set_theme = TRUE
)
```

```r
theme_gtsummary_mean_sd(set_theme = TRUE)
```

```r
theme_gtsummary_eda(set_theme = TRUE)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>journal</td>
<td>String indicating the journal theme to follow. One of c(&quot;jama&quot;, &quot;lancet&quot;,</td>
</tr>
<tr>
<td></td>
<td>&quot;nejm&quot;, &quot;qjecon&quot;). Details below.</td>
</tr>
<tr>
<td>set_theme</td>
<td>Logical indicating whether to set the theme. Default is TRUE. When FALSE</td>
</tr>
<tr>
<td></td>
<td>the named list of theme elements is returned invisibly.</td>
</tr>
<tr>
<td>font_size</td>
<td>Numeric font size for compact theme. Default is 13 for gt tables, and 8 for</td>
</tr>
<tr>
<td></td>
<td>all other output types</td>
</tr>
</tbody>
</table>
print_engine String indicating the print method. Must be one of "gt", "kable", "kable_extra", "flextable", "tibble"

language String indicating language. Must be one of "de" (German), "en" (English), "es" (Spanish), "fr" (French), "gu" (Gujarati), "hi" (Hindi), "is" (Icelandic), "ja" (Japanese), "kr" (Korean), "nl" (Dutch), "no" (Norwegian), "pt" (Portuguese), "se" (Swedish), "zh-cn" (Chinese Simplified), "zh-tw" (Chinese Traditional)

If a language is missing a translation for a word or phrase, please feel free to reach out on GitHub with the translated text!

decimal.mark The character to be used to indicate the numeric decimal point. Default is "." or getOption("OutDec")

decimal.mark Character used between every 3 digits to separate hundreds/thousands/millions/etc. Default is ",", except when decimal.mark = "," when the default is a space.

iqr.sep string indicating separator for the default IQR in tbl_summary(). If decimal.mark = NULL, iqr.sep = ",", ". The comma separator, however, can look odd when decimal.mark = ",", ". In this case the argument will default to an en dash

ci.sep string indicating separator for confidence intervals. If decimal.mark = is NULL, ci.sep = ",", ". The comma separator, however, can look odd when decimal.mark = ",", ". In this case the argument will default to an en dash

statistic Default statistic continuous variables

Themes

- theme_gtsummary_journal(journal=
  "jama" The Journal of the American Medical Association
  * Round large p-values to 2 decimal places; separate confidence intervals with "ll to ul".
  * tbl_summary() Doesn’t show percent symbol; use em-dash to separate IQR; run add_stat_label()
  * tbl_regression()/tbl_uvregression() show coefficient and CI in same column

- "lancet" The Lancet
  * Use mid-point as decimal separator; round large p-values to 2 decimal places; separate confidence intervals with "11 to 11".
  * tbl_summary() Doesn’t show percent symbol; use em-dash to separate IQR

- "nejm" The New England Journal of Medicine
  * Round large p-values to 2 decimal places; separate confidence intervals with "11 to 11".
  * tbl_summary() Doesn’t show percent symbol; use em-dash to separate CI

- "qjecon" The Quarterly Journal of Economics
  * tbl_summary() all percentages rounded to one decimal place
  * tbl_regression()/tbl_uvregression() add significance stars with add_significance_stars(); hides CI and p-value from output

- theme_gtsummary_compact()
  - tables printed with gt, flextable, kableExtra, or huxtable will be compact with smaller font size and reduced cell padding

- theme_gtsummary_printer(print_engine=
  - Use this theme to permanently change the default printer.
• theme_gtsummary_continuous2()
  – Set all continuous variables to summary type “continuous2” by default
• theme_gtsummary_mean_sd()
  – Set default summary statistics to mean and standard deviation in tbl_summary()
  – Set default continuous tests in add_p() to t-test and ANOVA
• theme_gtsummary_eda()
  – Set all continuous variables to summary type “continuous2” by default
  – In tbl_summary() show the median, mean, IQR, SD, and Range by default

Use reset_gtsummary_theme() to restore the default settings

Review the themes vignette to create your own themes.

**Example Output**

**See Also**

Themes vignette

set_gtsummary_theme(), reset_gtsummary_theme()

**Examples**

```r
# Setting JAMA theme for gtsummary
theme_gtsummary_journal("jama")
# Themes can be combined by including more than one
theme_gtsummary_compact()

set_gtsummary_theme_ex1 <-
  trial %>%
  select(age, grade, trt) %>%
  tbl_summary(by = trt) %>%
  as_gt()

# reset gtsummary themes
reset_gtsummary_theme()
```

---

**trial**

*Results from a simulated study of two chemotherapy agents*

**Description**

A dataset containing the baseline characteristics of 200 patients who received Drug A or Drug B. Dataset also contains the outcome of tumor response to the treatment.

**Usage**

trial
Format

A data frame with 200 rows–one row per patient

trt  Chemotherapy Treatment
age  Age
marker  Marker Level (ng/mL)
stage  T Stage
grade  Grade
response  Tumor Response
death  Patient Died
ttdeath  Months to Death/Censor