# **R** documentation

# of all in '/home/shaesen2/visR/man'

# April 24, 2022

# ${\sf R}$ topics documented:

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add\_annotation

Add annotations to a visR object

#### **Description**

Wrapper around ggplot2::annotation\_custom for simplified annotation to ggplot2 plots. This function accepts a string, dataframe, data.table, tibble or customized objects of class gtable and places them on the specified location on the ggplot. The layout is fixed: bold columnheaders and plain body. Only the font size and type can be chosen. Both the initial plot as the individual annotation are stored as attribute component in the final object.

# Usage

```
add_annotation(
   gg = NULL,
   label = NULL,
   base_family = "sans",
   base_size = 11,
   xmin = -Inf,
   xmax = Inf,
   ymin = -Inf,
   ymax = Inf
```

#### **Arguments**

gg	Object of class ggplot.
label	String, dataframe, data.table, tibble used to annotate the ggplot.
base_family	character. Base font family
base_size	numeric. Base font size in pt
xmin	x coordinates giving horizontal location of raster in which to fit annotation.
xmax	x coordinates giving horizontal location of raster in which to fit annotation.
ymin	y coordinates giving vertical location of raster in which to fit annotation.
ymax	y coordinates giving vertical location of raster in which to fit annotation.

#### Value

Object of class ggplot with added annotation with an object of class gtable.

#### See Also

```
tableGrob annotation_custom
```

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```
## Estimate survival
surv_object <- visR::estimate_KM(data = adtte, strata = "TRTP")</pre>
## We want to annotate the survival KM plot with a simple string comment
visR::visr(surv_object) %>%
  visR::add_annotation(
    label = "My simple comment",
    base_family = "sans",
    base\_size = 15,
    xmin = 110,
   xmax = 180,
    ymin = 0.80
## Currently, care needs to be taken on the x-y values relative
## to the plot data area. Here we are plotting outside of the data area.
visR::visr(surv_object) %>%
  visR::add_annotation(
 label = "My simple comment",
 base_family = "sans",
 base\_size = 15,
  xmin = 210,
  xmax = 380,
  ymin = 1.0
  )
## We may also want to annotate a KM plot with information
\#\# from additional tests or estimates. This example we annotate
## with p-values contained in a tibble
## we calculate p-values for "Equality across strata"
lbl <- visR::get_pvalue(surv_object,</pre>
             statlist = c("test", "pvalue"),
             type = "All")
## display p-values
## Now annotate survival KM plot with the p-values
visR::visr(surv_object) %>%
 visR::add_annotation(
    label = lbl,
    base_family = "sans",
    base\_size = 9,
    xmin = 100,
    xmax = 180,
    ymin = 0.80
  )
```

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#### **Description**

Method to add pointwise confidence intervals to a an object created by visR through an S3 method. The method is set up to use the pipe %>%. There are two options to display CI's, a "ribbon" or as "step" lines.

No default method is available at the moment.

#### Usage

```
add_CI(gg, ...)
## S3 method for class 'ggsurvfit'
add_CI(gg, alpha = 0.1, style = "ribbon", linetype, ...)
## S3 method for class 'ggtidycuminc'
add_CI(gg, alpha = 0.1, style = "ribbon", linetype, ...)
```

#### **Arguments**

gg	A ggplot created with visR
	other arguments passed on to the method to modify <pre>geom_ribbon</pre>
alpha	aesthetic of ggplot2 geom_ribbon. Default is 0.1.
_	aesthetic of ggplot2 geom_ribbon. Default is "ribbon". An alternative option is "step" that uses a line to display interval bounds.
linetype	aesthetic of ggplot2 geom_ribbon.

#### Value

Pointwise confidence interval overlayed on a visR ggplot

```
library(visR)

# Estimate KM curves by treatment group
survfit_object <- survival::survfit(data = adtte, survival::Surv(AVAL, 1-CNSR) ~ TRTP)

## plot without confidence intervals (CI)
p <- visR::visr(survfit_object)
p

# add CI to plot with default settings
p %>% add_CI()

# change transparency of CI ribbon
p %>% add_CI(alpha = 0.9, style = "ribbon")

# plot CI as a step line instead of ribbon
p %>% add_CI(alpha = 0.1, style = "step")

# change linetype of CI
p %>% add_CI(style = "step", linetype = 1)
```

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add\_CNSR

Add censoring symbols to a visR object

#### **Description**

Add censoring symbols to a visR ggplot through an S3 method. The S3 method is for adding censoring symbols to a visR ggplot. The method is set up to use the pipe %>%.

No default method is available at the moment.

# Usage

```
add_CNSR(gg, ...)
## S3 method for class 'ggsurvfit'
add_CNSR(gg, shape = 3, size = 2, ...)
## S3 method for class 'ggtidycuminc'
add_CNSR(gg, shape = 3, size = 2, ...)
```

#### **Arguments**

```
A ggplot created with visR

other arguments passed on to the method to modify geom_point

shape aesthetic of ggplot2 geom_point. Default is 3.

size aesthetic of ggplot2 geom_point. Default is 2.
```

#### Value

Censoring symbols overlayed on a visR ggplot

```
library(visR)

# Estimate KM curves by treatment group
survfit_object <- survival::survfit(data = adtte, survival::Surv(AVAL, 1-CNSR) ~ TRTP)

## plot without confidence intervals
p <- visR::visr(survfit_object)
p

# add censoring to plot
p %>% visR::add_CNSR()

# change censor symbol shape
p %>% visR::add_CNSR(shape = 1)

# change size and shape
p %>% visR::add_CNSR(size = 4, shape = 2)
```

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add\_highlight

Highlight a specific strata

# Description

S3 method for highlighting a specific strata by lowering the opacity of all other strata.

# Usage

```
add_highlight(gg, ...)
## S3 method for class 'ggsurvfit'
add_highlight(gg = NULL, strata = NULL, bg_alpha = 0.2, ...)
```

# Arguments

gg	A ggplot created with visR other arguments passed on to the method
strata	String representing the name and value of the strata to be highlighted as shown in the legend.
bg_alpha	A numerical value between 0 and 1 that is used to decrease the opacity off all strata not chosen to be highlighted in strata. The other strata's existing alpha values are multiplied by $bg_alpha$ to decrease their opacity, highlighting the target strata. This works on both colour and fill properties, as for example present after applying $visR::add_CI()$ .

# Value

The input ggsurvfit object with adjusted alpha values

```
adtte %>%
  visR::estimate_KM(strata = "SEX") %>%
  visR::visr() %>%
  visR::add_CI(alpha = 0.4) %>%
  visR::add_highlight(strata = "M", bg_alpha = 0.2)

strata = c("Placebo", "Xanomeline Low Dose")

adtte %>%
  visR::estimate_KM(strata = "TRTP") %>%
  visR::visr() %>%
  visR::add_CI(alpha = 0.4) %>%
  visR::add_highlight(strata = strata, bg_alpha = 0.2)
```

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add\_quantiles

Add quantile indicators to visR plot

# Description

Method to add quantile lines to a plot.

# Usage

```
add_quantiles(gg, ...)
## S3 method for class 'ggsurvfit'
add_quantiles(
    gg,
    quantiles = 0.5,
    linetype = "dashed",
    linecolour = "grey50",
    alpha = 1,
    ...
)
```

#### **Arguments**

gg	A ggplot created with visR
	other arguments passed on to the method to modify geom_line
quantiles	vector of quantiles to be displayed on the probability scale, default: 0.5
linetype	string indicating the linetype as described in the aesthetics of ggplot2 geom_line, default: dashed (also supports "mixed" -> horizontal lines are solid, vertical ones are dashed)
linecolour	string indicating the linetype as described in the aesthetics of ggplot2 geom_line, default: grey, (also supports "strata" -> horizontal lines are grey50, vertical ones are the same colour as the respective strata)
alpha	numeric value between 0 and 1 as described in the aesthetics of ggplot2 $\texttt{geom\_line}$ , default: 1

#### Value

Lines indicating the quantiles overlayed on a visR ggplot

```
library(visR)

adtte %>%
   estimate_KM("SEX") %>%
   visr() %>%
   add_quantiles()

adtte %>%
   estimate_KM("SEX") %>%
```

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```
visr() %>%
  add_quantiles(quantiles = c(0.25, 0.50))
adtte %>%
 estimate_KM("SEX") %>%
 visr() %>%
 add_quantiles(
   quantiles = c(0.25, 0.50),
   linetype = "solid",
   linecolour = "grey"
adtte %>%
 estimate_KM("SEX") %>%
 visr() %>%
 add_quantiles(
   quantiles = c(0.25, 0.50),
   linetype = "mixed",
   linecolour = "strata"
```

add\_risktable

Add risk tables to visR plots through an S3 method

# Description

S3 method for adding risk tables to visR plots. The function has following workflow:

- The risktables are calculated using get\_risktable
- The risktables are placed underneath visR plots using plot\_grid
- Both the initial visR plot as the individual risktables are stored as attribute component in the final object to allow post-modification of the individual plots if desired

#### Usage

```
add_risktable(gg, ...)
## S3 method for class 'ggsurvfit'
add_risktable(
   gg,
   times = NULL,
   statlist = "n.risk",
   label = NULL,
   group = "strata",
   collapse = FALSE,
   ...
)

## S3 method for class 'ggtidycuminc'
add_risktable(
   gg,
```

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```
times = NULL,
statlist = "n.risk",
label = NULL,
group = "strata",
collapse = FALSE,
...
)
```

#### **Arguments**

A ggplot created with visR gg other arguments passed on to the method add risktable . . . Numeric vector indicating the times at which the risk set, censored subjects, times events are calculated. statlist Character vector indicating which summary data to present. Current choices are "n.risk" "n.event" "n.censor", "cum.event", "cum.censor". Default is "n.risk". label Character vector with labels for the statlist. Default matches "n.risk" with "At risk", "n.event" with "Events", "n.censor" with "Censored", "cum.event" with "Cum. Event", and "cum.censor" with "Cum. Censor". String indicating the grouping variable for the risk tables. Current options are: group • "strata": groups the risk tables per stratum. The label specifies the label within each risk table. The strata levels are used for the titles of the risk tables. This is the default • "statlist": groups the risk tables per statlist. The label specifies the title for each risk table. The strata levels are used for labeling within each risk table. Default is "strata". collapse Boolean, indicates whether to present the data overall. Default is FALSE.

# Value

Object of class ggplot with added risk table.

#### See Also

```
plot_grid
```

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adtte

adtte - CDISC ADaM compliant time to event data set

# Description

ADTTE data copied from the 2013 CDISC Pilot

#### Usage

adtte

#### **Format**

A data frame with 254 rows and 26 variables:

STUDYID Study Identifier

SITEID Study Site Identifier

USUBJID Unique Subject Identifier

AGE Age

AGEGR1 Pooled Age Group 1

AGEGR1N Pooled Age Group 1 (N)

**RACE** Race

**RACEN** Race (N)

SEX Sex

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRTDUR Duration of treatment (days)

TRTP Planned Treatment

TRTA Actual Treatment

TRTAN Actual Treatment (N)

PARAM Parameter Description

PARAMCD Parameter Code

AVAL Analysis Value

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STARTDT Time to Event Origin Date for Subject

ADT Analysis Date

**CNSR** Censor

**EVNTDESC** Event or Censoring Description

**SRCDOM** Source Domain

**SRCVAR** Source Variable

**SRCSEQ** Source Sequence Number

SAFFL Safety Population Flag

#### Source

```
CDISC SDTM/ADAM Pilot Project. https://bitbucket.cdisc.org/projects/CED
```

#### **Examples**

```
data("adtte")
```

align\_plots

Align multiple ggplot graphs, taking into account the legend

# Description

This function aligns multiple ggplot graphs by making them the same width by taking into account the legend width.

#### Usage

```
align_plots(pltlist)
```

#### **Arguments**

pltlist A list of plots

# Value

List of ggplot with equal width.

# References

https://stackoverflow.com/questions/26159495/align-multiple-ggplot-graphs-with-

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#### **Examples**

apply\_attrition

Apply list of inclusion/exclusion criteria to a patient-level dataframe

#### **Description**

Apply list of inclusion/exclusion criteria to a patient-level dataframe

# Usage

```
apply_attrition(data, criteria_conditions)
```

#### **Arguments**

```
data data.frame. Data set to be filtered criteria_conditions
```

character Dplyr-filter compatible conditions of the filtering criteria. These conditions will be applied to filter the input data set and obtain final analysis data set

#### Value

Filtered data frame

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apply\_theme

Applies a theme to a ggplot object.

# Description

Takes in the styling options defined through visR::define\_theme and applies them to a plot.

# Usage

```
apply_theme(gg, visR_theme_dict = NULL)
```

#### **Arguments**

# Value

object of class ggplot

visR::add\_CI() %>%
visR::apply\_theme(theme)

#### **Examples**

gg

```
library(visR)
theme <- visR::define_theme(strata = list("SEX" = list("F" = "red",
                                                        "M" = "blue"),
                                           "TRTA" = list("Placebo" = "cyan",
                                                         "Xanomeline High Dose" = "purple"
                                                         "Xanomeline Low Dose" = "brown"))
                            fontsizes = list("axis" = 12,
                                              "ticks" = 10,
                                              "legend_title" = 10,
                                              "legend_text" = 8),
                             fontfamily = "Helvetica",
                            grid = FALSE,
                            bg = "transparent",
                            legend_position = "top")
gg <- adtte %>%
 visR::estimate_KM(strata = "SEX") %>%
  visR::visr() %>%
```

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brca\_cohort

Cancer survival data

# Description

Creation script in data-raw

# Usage

```
brca_cohort
```

#### **Format**

An object of class data.frame with 1098 rows and 10 columns.

define\_theme

Provides a simple wrapper for themes

# Description

This function collects several lists if they are present. If absent, reasonable defaults are used.

# Usage

```
define_theme(
   strata = NULL,
   fontsizes = NULL,
   fontfamily = "Helvetica",
   grid = FALSE,
   bg = "transparent",
   legend_position = NULL
)
```

#### **Arguments**

strata list containing the different strata and name:colour value pairs

fontsizes list containing the font sizes for different options

fontfamily string with the name of a supported font

grid boolean that specifies whether the grid should be drawn or not

bg string giving the colour for the background of the plot

legend\_position

string indicating the legend position

#### Value

Nested list with styling preferences for a ggplot object

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#### **Examples**

```
theme <- visR::define_theme(</pre>
  strata = list("SEX" = list("F" = "red",
                              "M" = "blue"
                           ),
               "TRTA" = list("Placebo" = "cyan",
                              "Xanomeline High Dose" = "purple",
                              "Xanomeline Low Dose" = "brown"
            ),
  fontsizes = list("axis" = 12,
                    "ticks" = 10,
                    "legend_title" = 10,
                    "legend_text" = 8),
  fontfamily = "Helvetica",
  grid = list("major" = FALSE,
              "minor" = FALSE
             ),
 bg = "transparent",
  legend_position = "top"
```

estimate\_cuminc

Competing Events Cumulative Incidence

# **Description**

Function creates a cumulative incidence object using the tidycmprsk::cuminc() function.

#### Usage

```
estimate_cuminc(
  data,
  strata = NULL,
  CNSR = "CNSR",
  AVAL = "AVAL",
  conf.int = 0.95,
  ...
)
```

#### Arguments

dat	а

The name of the dataset for Time-to-Event analysis based on the Analysis Data Model (ADaM) principles. The dataset is expected to have one record per subject per analysis parameter. Rows in which the analysis variable (AVAL) or the censor variable (CNSR) contain NA, are removed during analysis.

strata

Character vector, representing the strata for Time-to-Event analysis. When NULL, an overall analysis is performed. Default is NULL.

CNSR

Column name indicating the outcome and censoring statuses. Column must be a factor and the first level indicates censoring, the next level is the outcome of interest, and the remaining levels are the competing events. Default is "CNSR"

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```
AVAL Analysis value for Time-to-Event analysis. Default is "AVAL", as per CDISC ADaM guiding principles.

conf.int Confidence internal level. Default is 0.95.

Additional argument passed to tidycmprsk::cuminc()
```

#### Value

A cumulative incidence object as explained at https://mskcc-epi-bio.github.io/tidycmprsk/reference/cuminc.html

# **Examples**

```
estimate_cuminc(
  tidycmprsk::trial,
  strata = "trt",
  CNSR = "death_cr",
  AVAL = "ttdeath"
) %>%
  visr() %>%
  add_CI() %>%
  add_risktable(statlist = c("n.risk", "cum.event"))
```

estimate\_KM

Wrapper for Kaplan Meier Time-to-Event analysis

#### **Description**

This function is a wrapper around <code>survival::survfit.formula</code> to perform a Kaplan-Meier analysis, assuming right-censored data. The function expects that the data has been filtered on the parameter (PARAM/PARAMCD) of interest. All NA values in the CNSR, AVAL and strata argument are removed. Alternatively, PARAM/PARAMCD can be used in the <code>strata</code> argument. The result is an object of class <code>survfit</code> which can be used in downstream functions and methods that rely on the <code>survfit</code> class. When strata are present, the returned survfit object is supplemented with the a named list of the stratum and associated label, if present. By default:

- The Kaplan Meier estimate is estimated directly (stype = 1).
- The cumulative hazard is estimated using the Nelson-Aalen estimator (ctype = 1): H.tilde = cumsum(x\$n.event/x\$n.risk). The MLE (H.hat(t) = -log(S.hat(t))) can't be requested.
- A two-sided pointwise 0.95 confidence interval is estimated using a log transformation (conf.type = "log").

# Usage

```
estimate KM(data = NULL, strata = NULL, CNSR = "CNSR", AVAL = "AVAL", ...)
```

#### **Arguments**

data

The name of the dataset for Time-to-Event analysis based on the Analysis Data Model (ADaM) principles. The dataset is expected to have one record per subject per analysis parameter. Rows in which the analysis variable (AVAL) or the censor variable (CNSR) contain NA, are removed during analysis.

estimate\_KM

strata	Character vector, representing the strata for Time-to-Event analysis. When NULL, an overall analysis is performed. Default is NULL.
CNSR	Censor for Time-to-Event analysis. Default is "CNSR", as per CDISC ADaM guiding principles.
AVAL	Analysis value for Time-to-Event analysis. Default is "AVAL", as per CDISC ADaM guiding principles.
• • •	additional arguments passed on to the ellipsis of the call survival::survfit.formula(data = data, formula = Surv(AVAL, 1-CNSR) ~ strata),). Use?survival::survfit and?survival::survfitCI for more information.

#### Value

survfit object, extended by elements PARAM/PARAMCD, ready for downstream processing in estimation or visualization functions and methods.

#### References

```
https://github.com/therneau/survival
```

#### See Also

```
survfit.formula survfitCI
```

```
## No stratification
visR::estimate_KM(data = adtte)
## Stratified Kaplan-Meier analysis by `TRTP`
visR::estimate_KM(data = adtte, strata = "TRTP")
## Stratified Kaplan-Meier analysis by `TRTP` and `SEX`
visR::estimate_KM(data = adtte, strata = c("TRTP", "SEX"))
## Stratification with one level
visR::estimate_KM(data = adtte, strata = "PARAMCD")
## Analysis on subset of adtte
visR::estimate_KM(data = adtte[adtte$SEX == "F", ])
## Modify the default analysis by using the ellipsis
visR::estimate_KM(data = adtte, strata = NULL,
  type = "kaplan-meier", conf.int = FALSE, timefix = TRUE)
## Example working with non CDISC data
head(survival::veteran)
# convert time and censoring data to ADaM variables
# convert censoring status to CDISC principles
veteran_adam <- survival::veteran %>%
dplyr::mutate(AVAL = time,
              CNSR = dplyr::if_else(status == 1, 0, 1)
```

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```
visR::estimate_KM(data = veteran_adam, strata = "trt")
```

get\_attrition

Generate cohort attrition table

#### **Description**

[Experimental] This is an experimental function that may be developed over time.

This function calculates the subjects counts excluded and included for each step of the cohort selection process.

#### Usage

```
get_attrition(data, criteria_descriptions, criteria_conditions,
subject_column_name)
```

#### **Arguments**

data

Dataframe. It is used as the input data to count the subjects that meets the criteria of interest

criteria\_descriptions

character It contains the descriptions of the inclusion/exclusion criteria. Each element of the vector corresponds to the description of each criterion.

criteria\_conditions

character It contains the corresponding conditions of the criteria. These conditions will be used in the table to compute the counts of the subjects.

subject\_column\_name

character The column name of the table that contains the subject id.

#### **Details**

criteria\_descriptions and criteria\_conditions need to be of same length

# Value

The counts and percentages of the remaining and excluded subjects for each step of the cohort selection in a table format.

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get\_COX\_HR

Summarize Hazard Ratio from a survival object using S3 method

#### **Description**

S3 method for extracting information regarding Hazard Ratios. The function allows the survival object's formula to be updated. No default method is available at the moment.

#### Usage

```
get_COX_HR(x, ...)
## S3 method for class 'survfit'
get_COX_HR(x, update_formula = NULL, ...)
```

#### **Arguments**

x An object of class survfit... other arguments passed on to the method survival::coxph update formula

Template which specifies how to update the formula of the survfit object update.formula

#### Value

A tidied object of class coxph containing Hazard Ratios

#### See Also

```
coxph update.formula
```

```
## treatment effect
survfit_object_trt <- visR::estimate_KM(data = adtte, strata = c("TRTP"))
visR::get_COX_HR(survfit_object_trt)

## treatment and gender effect
survfit_object_trt_sex <- visR::estimate_KM(data = adtte, strata = c("TRTP", "SEX"))
visR::get_COX_HR(survfit_object_trt_sex)

## update formula of KM estimates by treatment to include "SEX" for HR estimation
visR::get_COX_HR(survfit_object_trt, update_formula = ". ~ . + SEX")

## update formula of KM estimates by treatment to include "AGE" for
## HR estimation with ties considered via the efron method
visR::get_COX_HR(survfit_object_trt,
    update_formula = ". ~ . + survival::strata(AGE)", ties = "efron")</pre>
```

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Summarize the test for equality across strata from a survival object using S3 method

#### **Description**

Wrapper around survival::survdiff that tests the null hypothesis of equality across strata.

#### Usage

```
get_pvalue(
   survfit_object,
   ptype = "All",
   rho = NULL,
   statlist = c("test", "Chisq", "df", "pvalue"),
   ...
)
```

# **Arguments**

survfit\_object

An object of class survfit

ptype

Character vector containing the type of p-value desired. Current options are "Log-Rank" "Wilcoxon" "Tarone-Ware" "Custom" "All". "Custom" allows the user to specify the weights on the Kaplan-Meier estimates using the argument rho. The default is "All" displaying all types possible. When rho is specified in centert of "All" also a custom a value in displayed.

in context of "All", also a custom p-value is displayed.

rho

a scalar parameter that controls the type of test.

statlist

Character vector containing the desired information to be displayed. The order of the arguments determines the order in which they are displayed in the final result. Default is the test name ("test"), Chisquare test statistic ("Chisq"), degrees

of freedom ("df") and p-value ("pvalue").

... other arguments passed on to the method

# Value

A data frame with summary measures for the Test of Equality Across Strata

#### See Also

```
survdiff
```

```
## general examples
survfit_object <- visR::estimate_KM(data = adtte, strata = "TRTP")
visR::get_pvalue(survfit_object)
visR::get_pvalue(survfit_object, ptype = "All")
## examples to obtain specific tests</pre>
```

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```
visR::get_pvalue(survfit_object, ptype = "Log-Rank")
visR::get_pvalue(survfit_object, ptype = "Wilcoxon")
visR::get_pvalue(survfit_object, ptype = "Tarone-Ware")

## Custom example - obtain Harrington and Fleming test
visR::get_pvalue(survfit_object, ptype = "Custom", rho = 1)

## Get specific information and statistics
visR::get_pvalue(survfit_object, ptype = "Log-Rank", statlist = c("test", "Chisq", "df",
visR::get_pvalue(survfit_object, ptype = "Wilcoxon", statlist = c("pvalue"))
```

get\_quantile

Wrapper around quantile methods

### **Description**

S3 method for extracting quantiles. No default method is available at the moment.

# Usage

```
get_quantile(x, ...)
## S3 method for class 'survfit'
get_quantile(
    x,
    ...,
    probs = c(0.25, 0.5, 0.75),
    conf.int = TRUE,
    tolerance = sqrt(.Machine$double.eps)
)
```

#### **Arguments**

X	An object of class survfit
	other arguments passed on to the method
probs	probabilities Default = $c(0.25,0.50,0.75)$
conf.int	should lower and upper confidence limits be returned?
tolerance	tolerance for checking that the survival curve exactly equals one of the quantiles

#### Value

A data frame with quantiles of the object

#### See Also

```
quantile.survfit
```

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#### **Examples**

```
## Kaplan-Meier estimates
survfit_object <- visR::estimate_KM(data = adtte, strata = c("TRTP"))
## visR quantiles
visR::get_quantile(survfit_object)
## survival quantiles
quantile(survfit_object)</pre>
```

get\_risktable

Obtain risk tables for tables and plots

#### **Description**

Create a risk table from an object using an S3 method. Currently, no default method is defined.

# Usage

```
get_risktable(x, ...)
## S3 method for class 'survfit'
get_risktable(
  х,
  times = NULL,
  statlist = "n.risk",
  label = NULL,
  group = c("strata", "statlist"),
  collapse = FALSE,
  . . .
)
## S3 method for class 'tidycuminc'
get_risktable(
  х,
  times = pretty(x$tidy$time, 10),
  statlist = c("n.risk"),
  label = NULL,
  group = c("strata", "statlist"),
  collapse = FALSE,
  . . .
)
```

## **Arguments**

```
    an object of class survfit or tidycuminc
    other arguments passed on to the method
    Numeric vector indicating the times at which the risk set, censored subjects, events are calculated.
```

get\_summary 23

statlist	Character vector indicating which summary data to present. Current choices are "n.risk" "n.event" "n.censor", "cum.event", "cum.censor". Default is "n.risk".
label	Character vector with labels for the statlist. Default matches "n.risk" with "At risk", "n.event" with "Events", "n.censor" with "Censored", "cum.event" with "Cum. Event", and "cum.censor" with "Cum. Censor".
group	String indicating the grouping variable for the risk tables. Current options are:
	• "strata": groups the risk tables per stratum. The label specifies the label within each risk table. The strata levels are used for the titles of the risk tables. This is the default
	• "statlist": groups the risk tables per statlist. The label specifies the title for each risk table. The strata levels are used for labeling within each risk table.
	Default is "strata".
collapse	Boolean, indicates whether to present the data overall. Default is FALSE.

#### Value

return list of attributes the form the risk table i.e. number of patients at risk per strata

### See Also

```
summary.survfit
```

get_summary	Summarize the descriptive statistics across strata from a survival ob-
	ject using S3 method

### **Description**

S3 method for extracting descriptive statistics across strata. No default method is available at the moment.

survfit\_object <- survival::survfit(data = adtte, survival::Surv(AVAL, 1-CNSR) ~ TRTP) get\_summary(survfit\_object)

# Usage

```
get_summary(x, ...)
## S3 method for class 'survfit'
get_summary(
    x,
    statlist = c("strata", "records", "events", "median", "LCL", "UCL", "CI"),
    ...
)
```

24 get\_tableone

#### **Arguments**

x An object of class survfit

other arguments passed on to the method

statlist

Character vector containing the desired information to be displayed. The order of the arguments determines the order in which they are displayed in the final result. Default is the strata ("strata"), number of subjects ("records"), number of events ("events"), the median survival time ("median"), the Confidence Interval ("CI"), the Lower Confidence Limit ("UCL") and the Upper Confidence Limit ("UCL").

#### Value

A data frame with summary measures from a survfit object

get\_tableone

Calculate summary statistics

#### **Description**

S3 method for creating a table of summary statistics. The summary statistics can be used for presentation in tables such as table one or baseline and demography tables.

The summary statistics estimated are conditional on the variable type: continuous, binary, categorical, etc.

By default the following summary stats are calculated:

- Numeric variables: mean, min, 25th-percentile, median, 75th-percentile, maximum, standard deviation
- Factor variables: proportion of each factor level in the overall dataset
- Default: number of unique values and number of missing values

#### Usage

```
get_tableone(
  data,
  strata = NULL,
  overall = TRUE,
  summary_function = summarize_short
)

## Default S3 method:
get_tableone(
  data,
  strata = NULL,
  overall = TRUE,
  summary_function = summarize_short
)
```

get\_tableone 25

# **Arguments**

data	The dataset to summarize as dataframe or tibble	
strata	Stratifying/Grouping variable name(s) as character vector. If NULL, only overall results are returned	
overall	If TRUE, the summary statistics for the overall dataset are also calculated	
summary_function		

A function defining summary statistics for numeric and categorical values

#### **Details**

It is possible to provide your own summary function. Please have a loot at summary for inspiration.

#### Value

object of class tableone. That is a list of data specified summaries for all input variables.

#### Note

All columns in the table will be summarized. If only some columns shall be used, please select only those variables prior to creating the summary table by using dplyr::select()

```
# Example using the ovarian data set
survival::ovarian %>%
  dplyr::select(-fustat) %>%
  dplyr::mutate(
   age_group = factor(
     dplyr::case_when(
        age <= 50 ~ "<= 50 years",
        age <= 60 ~ "<= 60 years",
        age <= 70 ~ "<= 70 years",
        TRUE ~ "> 70 years"
      )
   ),
   rx = factor(rx),
   ecog.ps = factor(ecog.ps)
  dplyr::select(age, age_group, everything()) %>%
  visR::get_tableone()
# Examples using ADaM data
# display patients in an analysis set
adtte %>%
 dplyr::filter(SAFFL == "Y") %>%
  dplyr::select(TRTA) %>%
  visR::get_tableone()
## display overall summaries for demog
adtte %>%
  dplyr::filter(SAFFL == "Y") %>%
  dplyr::select(AGE, AGEGR1, SEX, RACE) %>%
```

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```
visR::get_tableone()

## By actual treatment
adtte %>%
   dplyr::filter(SAFFL == "Y") %>%
   dplyr::select(AGE, AGEGR1, SEX, RACE, TRTA) %>%
   visR::get_tableone(strata = "TRTA")

## By actual treatment, without overall
adtte %>%
   dplyr::filter(SAFFL == "Y") %>%
   dplyr::select(AGE, AGEGR1, SEX, EVNTDESC, TRTA) %>%
   visR::get_tableone(strata = "TRTA", overall = FALSE)
```

legendopts

Translates options for legend into a list that can be passed to ggplot2

# Description

This function takes the legend position and orientation, defined by the user and puts them into a list for ggplot2.

# Usage

```
legendopts(legend_position = "right", legend_orientation = NULL)
```

# **Arguments**

#### Value

List of legend options for ggplot2.

render

Render a data.frame or tibble

#### **Description**

Render a previously created data.frame to html, rtf or latex

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#### Usage

```
render(
  data,
  title,
  datasource,
  footnote = "",
 output_format = "html",
  engine = "gt",
  download_format = c("copy", "csv", "excel")
## S3 method for class 'tableone'
render(
  data,
 title,
 datasource,
 footnote = "",
 output_format = "html",
 engine = "gt",
  download_format = NULL
)
## S3 method for class 'risktable'
render(
 data,
 title,
 datasource,
 footnote = "",
  output_format = "html",
  engine = "gt",
  download_format = NULL
)
## S3 method for class 'data.frame'
render(
 data,
 title,
 datasource,
 footnote = "",
 output_format = "html",
 engine = "gt",
  download_format = c("copy", "csv", "excel")
)
```

#### **Arguments**

data	The data.frame or tibble to visualize
title	Table title to include in the rendered table
datasource	String specifying the data source underlying the data set
footnote	String specifying additional information to be displayed in the table note along- side the data source and specifications of statistical tests.

```
output_format
```

Type of output that is returned, can be html or latex

engine

If html is selected as output format, one can chose between using kable, gt and

DT as engine to create the output table

download\_format

Options generated for downloading the data

#### Value

A table-like data structure, possibly interactive depending on the choice of the engine A table-like data structure, possibly interactive depending on the choice of the engine

summarize\_long

Calculate summary statistics for a vector

# **Description**

Calculates several summary statistics for a vector depending on the vector class

#### Usage

```
summarize_long(x)
```

#### **Arguments**

Х

an object

#### Value

A summarized version of the input.

```
summarize_long.default
```

Create variable summary for all other variable types

# Description

Create variable summary for all other variable types

# Usage

```
## Default S3 method:
summarize_long(x)
```

# **Arguments**

Х

an object of any other class

#### Value

List of counts for unique and missing values in x.

summarize\_long.factor 29

```
summarize_long.factor
```

Create variable summary for factors

# Description

Create variable summary for factors

# Usage

```
## S3 method for class 'factor'
summarize_long(x)
```

# **Arguments**

Х

an object of class "factor"

# Value

Long list of summary statistics for the input factors.

```
summarize_long.integer
```

Create variable summary for numeric variables

# Description

Create variable summary for numeric variables

# Usage

```
## S3 method for class 'integer'
summarize_long(x)
```

#### **Arguments**

Х

an object of class "integer"

# Value

Long list of summary statistics for the input.

30 summarize\_short

```
summarize_long.numeric
```

Create variable summary for numeric variables

# Description

Create variable summary for numeric variables

# Usage

```
## S3 method for class 'numeric'
summarize_long(x)
```

# Arguments

x

an object of class "numeric"

#### Value

Long list of summary statistics for the input.

summarize\_short

Create abbreviated variable summary for table1

# Description

This function creates summaries combines multiple summary measures in a single formatted string.

# Usage

```
summarize_short(x)
```

#### **Arguments**

Х

a vector to be summarized

# Value

A summarized less detailed version of the input.

summarize\_short.default

```
summarize_short.default
```

Create variable summary for all other variable types

# Description

Create variable summary for all other variable types

# Usage

```
## Default S3 method:
summarize_short(x)
```

# **Arguments**

Х

an object of any other class

# Value

List of counts for unique and missing values in x.

```
summarize_short.factor
```

Create variable summary for factors

# Description

Calculates N and % of occurrence for each factor value

# Usage

```
## S3 method for class 'factor'
summarize_short(x)
```

#### **Arguments**

Х

an object of class "factor"

# Value

Short list of summary statistics for the input factors.

```
summarize_short.integer
```

Create variable summary for integer variables

#### **Description**

Calculates mean (standard deviation), median (IQR), min-max range and N/% missing elements for a integer vector.

# Usage

```
## S3 method for class 'integer'
summarize_short(x)
```

# Arguments

Х

an object of class "integer"

#### Value

Short list of summary statistics for the input.

```
summarize_short.numeric
```

Create variable summary for numeric variables

# Description

Calculates mean (standard deviation), median (IQR), min-max range and N/% missing elements for a numeric vector.

# Usage

```
## S3 method for class 'numeric'
summarize_short(x)
```

# **Arguments**

Х

an object of class "numeric"

# Value

Short list of summary statistics for the input.

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tableone

Display a summary Table (i.e. table one)

# Description

Wrapper function to produce a summary table (i.e. Table One). Create and render a summary table for a dataset. A typical example of a summary table are "table one", the first table in an applied medical research manuscript.

Calculate summary statistics and present them in a formatted table

# Usage

```
tableone(
  data,
  title,
  datasource,
  footnote = "",
  strata = NULL,
  overall = TRUE,
  summary_function = summarize_short,
  ...
)
```

# Arguments

data	The dataframe or tibble to visualize		
title	Table title to include in the rendered table. Input is a text string.		
datasource	String specifying the datasource underlying the data set		
footnote	Table footnote to include in the rendered table. Input is a text string.		
strata	Character vector with column names to use for stratification in the summary table. Default: NULL , which indicates no stratification.		
overall	If TRUE, the summary statistics for the overall dataset are also calculated		
summary_function			
	A function defining summary statistics for numeric and categorical values Preimplemented functions are summarize_long and summarize_short		
	Pass options to render_table		

#### Value

A table-like data structure, possibly interactive depending on the choice of the engine

# **Example Output**

34 tableone

```
# metadata for table
t1_title <- "Cohort Summary"</pre>
t1_ds <- "ADaM Interim Dataset for Time-to-Event Analysis"
t1_fn <- "My table one footnote"
## table by treatment - without overall and render with GT
tbl_gt <-
 adtte %>%
 dplyr::filter(SAFFL == "Y") %>%
 dplyr::select(AGE, AGEGR1, SEX, EVNTDESC, TRTA) %>%
 visR::tableone(
    strata = "TRTA",
    overall = FALSE,
    title = t1_title,
    datasource = t1_ds,
    footnote = t1_fn,
    engine = "gt"
  )
## table by treatment - without overall and render with DT
tbl_DT <-
  adtte %>%
  dplyr::filter(SAFFL == "Y") %>%
  dplyr::select(AGE, AGEGR1, SEX, EVNTDESC, TRTA) %>%
  visR::tableone(
     strata = "TRTA",
     overall = FALSE,
     title = t1_title,
     datasource = t1_ds,
     footnote = t1_fn,
     engine = "DT"
  )
## table by treatment - without overall and render with kable
tbl_kable_html <-
  adtte %>%
  dplyr::filter(SAFFL == "Y") %>%
  dplyr::select(AGE, AGEGR1, SEX, EVNTDESC, TRTA) %>%
  visR::tableone(
     strata = "TRTA",
     overall = FALSE,
     title = t1_title,
     datasource = t1_ds,
     footnote = t1_fn,
     engine = "kable"
  )
## table by treatment - without overall and render with kable as
## a latex table format rather than html
tbl_kable_latex <-
  adtte %>%
  dplyr::filter(SAFFL == "Y") %>%
   dplyr::select(AGE, AGEGR1, SEX, EVNTDESC, TRTA) %>%
```

the\_lhs 35

```
visR::tableone(
   strata = "TRTA",
   overall = FALSE,
   title = t1_title,
   datasource = t1_ds,
   footnote = t1_fn,
   output_format = "latex",
   engine = "kable"
)
```

the\_lhs

Find the "lhs" in the pipeline

# Description

This function finds the left-hand sided symbol in a magrittr pipe and returns it as a character.

# Usage

```
the_lhs()
```

# Value

Left-hand sided symbol as string in the magrittr pipe.

#### References

https://github.com/tidyverse/magrittr/issues/115#issuecomment-173894787

#### **Examples**

```
blah <- function(x) the_lhs()
adtte %>%
  blah()
```

tidyme

Extended tidy cleaning of selected objects using S3 method

# Description

S3 method for extended tidying of selected model outputs. Note that the visR method retains the original nomenclature of the objects, and adds the one of broom::tidy to ensure compatibility with tidy workflows. The default method relies on broom::tidy to return a tidied object

#### Usage

```
tidyme(x, ...)
## Default S3 method:
tidyme(x, ...)
## S3 method for class 'survfit'
tidyme(x, ...)
```

# Arguments

x An S3 object

... other arguments passed on to the method

#### Value

Data frame containing all list elements of the S3 object as columns. The column 'strata' is a factor to ensure that the strata are sorted in agreement with the order in the survfit object

#### See Also

tidy

#### **Examples**

```
## Extended tidying for a survfit object
surv_object <- visR::estimate_KM(data = adtte, strata = "TRTA")
tidied <- visR::tidyme(surv_object)

## Tidyme for non-included classes
data <- cars
lm_object <- stats::lm(data = cars, speed ~ dist)
lm_tidied <- visR::tidyme(lm_object)
lm_tidied</pre>
```

visr

Plot a supported S3 object

# Description

S3 method for creating plots directly from objects using ggplot2, similar to the base R plot () function.

# Usage

```
visr(x, ...)
## Default S3 method:
visr(x, ...)
```

```
## S3 method for class 'survfit'
visr(
 x = NULL
  x_{label} = NULL
  y_label = NULL
  x_units = NULL,
  x\_ticks = NULL,
  y_ticks = NULL,
  fun = "surv",
  legend_position = "right",
)
## S3 method for class 'attrition'
visr(
  х,
  description_column_name = "Criteria",
  value_column_name = "Remaining N",
  complement_column_name = "",
 box_width = 50,
  font size = 12,
  fill = "white",
  border = "black",
  . . .
)
## S3 method for class 'tidycuminc'
visr(
  x = NULL
  x_{label} = "Time",
  y_label = "Cumulative Incidence",
  x_units = NULL,
  x_{\text{ticks}} = \text{pretty}(x \approx 10),
  y_{ticks} = pretty(c(0, 1), 5),
  legend_position = "right",
)
```

#### **Arguments**

x\_units

х	Object of class survfit, attritiontable or visr.tidycuminc
	other arguments passed on to the method
x_label	character Label for the x-asis. When not specified, the algorithm will look for "PARAM" information inside the list structure of the survfit object. If no "PARAM" information is available, the algorithm will look for "PARAMCD" information inside the list structure. If both "PARAM" and "PARAMCD" are missing then "time" is used as label. Note that "PARAM"/"PARAMCD" information is automatically added when using visR::estimate_KM if the input data follows ADaM data model structure.
y_label	character Label for the y-axis. When not specified, the default will do a proposal, depending on the ${\tt fun}$ argument.

Unit to be added to the x\_label (x\_label (x\_unit)). Default is NULL.

x ticks Ticks for the x-axis. When not specified, the default will do a proposal. Ticks for the y-axis. When not specified, the default will do a proposal based on y\_ticks the fun argument. fun Function that represents the scale of the estimate. The current options are: • surv is the survival probability. This is the default. • log is log of the survival probability • event is the failure probability • cloglog is log(-log(survival probability)) • pct is survival as a percentage • logpct is log survival as a percentage • cumhaz is the cumulative hazard legend\_position Specifies the legend position in the plot. Character values allowed are "top" "left" "bottom" "right". Numeric coordinates are also allowed. Default is "right". description\_column\_name character Name of the column containing the inclusion descriptions value\_column\_name character Name of the column containing the remaining sample counts complement\_column\_name character Optional: Name of the column containing the exclusion descripcharacter The box width for each box in the flow chart box\_width character The fontsize in pt font size fill The color (string or hexcode) to use to fill the boxes in the flowchart

#### Value

Object of class ggplot and ggsurvplot for survfit objects.

#### See Also

ggplot

border

# **Examples**

```
# fit KM
km_fit <- survival::survfit(survival::Surv(AVAL, 1-CNSR) ~ TRTP, data = adtte)
# plot curves using survival plot function
plot(km_fit)
# plot same curves using visR::visr plotting function
visR::visr(km_fit)
# estimate KM using visR wrapper
survfit_object <- visR::estimate_KM(data = adtte, strata = "TRTP")
# Plot survival probability
visR::visr(survfit_object, fun = "surv")</pre>
```

The color (string or hexcode) to use for the borders of the boxes in the flowchart

```
# Plot survival percentage
visR::visr(survfit_object, fun = "pct")
# Plot cumulative hazard
visR::visr(survfit_object, fun = "cloglog")
## Create attrition
attrition <- visR::get_attrition(adtte,</pre>
    criteria_descriptions = c("1. Not in Placebo Group",
                               "2. Be 75 years of age or older.",
                              "3. White",
                              "4. Female"),
                          = c("TRTP != 'Placebo'",
    criteria_conditions
                              "AGE >= 75",
                              "RACE == 'WHITE'",
                              "SEX=='F'"),
    subject_column_name
                          = "USUBJID")
## Draw a CONSORT attrition chart without specifying extra text for the complement
attrition %>%
  visr("Criteria", "Remaining N")
## Add detailed complement descriptions to the "exclusion" part of the CONSORT diagram
# Step 1. Add new column to attrition dataframe
attrition $Complement <- c("NA",
                          "Placebo Group",
                          "Younger than 75 years",
                          "Non-White",
                          "Male")
# Step 2. Define the name of the column in the call to the plotting function
attrition %>%
 visr("Criteria", "Remaining N", "Complement")
## Styling the CONSORT flowchart
# Change the fill and outline of the boxes in the flowchart
attrition %>%
  visr("Criteria", "Remaining N", "Complement", fill = "lightblue", border="grey")
## Adjust the font size in the boxes
attrition %>%
  visr("Criteria", "Remaining N", font_size = 10)
```

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