Package ‘ggsurvfit’

March 16, 2023

Title Flexible Time-to-Event Figures

Version 0.3.0

Description Ease the creation of time-to-event (i.e. survival) endpoint figures. The modular functions create figures ready for publication. Each of the functions that add to or modify the figure are written as proper 'ggplot2' geoms or stat methods, allowing the functions from this package to be combined with any function or customization from 'ggplot2' and other 'ggplot2' extension packages.

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

Depends R (>= 3.5)

Imports broom (>= 1.0.0), cli (>= 3.0.0), dplyr (>= 1.0.3), ggplot2 (>= 3.4.0), glue (>= 1.6.0), gtable, patchwork (>= 1.1.0), rlang (>= 1.0.0), survival (>= 3.4-0), tidyverse (>= 1.0.0)

Suggests covr, knitr, rmarkdown, scales (>= 1.1.0), spelling, testthat (>= 3.0.0), tidycmprsk (>= 0.2.0), vcd (>= 1.0.0)

VignetteBuilder knitr

Config/testthat/edition 3

Config/Needs/website cowplot, ggeasy, gghighlight

Encoding UTF-8

Language en-US

LazyData true

RoxygenNote 7.2.3

NeedsCompilation no

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add_censor_mark

Description
Add a marking on the figure to represent the time an observation was censored.

Usage
add_censor_mark(...)

Arguments
... arguments passed to ggplot2::geom_point(...) with defaults shape = 3 and size = 2
**add_confidence_interval**

**Value**

a ggplot2 figure

**See Also**

Visit the gallery for examples modifying the default figures

**Examples**

```r
survfit2(Surv(time, status) ~ 1, data = df_lung) %>%
ggsurvfit() +
add_confidence_interval() +
add_censor_mark()
```

---

**Description**

Add a confidence interval represented by either a ribbon or lines.

**Usage**

```
add_confidence_interval(type = c("ribbon", "lines"), ...)
```

**Arguments**

- `type`: string indicating the type of confidence interval to draw. Must be one of c("ribbon", "lines")
- `...`: arguments pass to geom.
  - `type = 'ribbon'`: Defaults are `ggplot2::geom_ribbon(alpha = 0.2, color = NA, ...)`
  - `type = 'lines'`: Defaults are `ggplot2::geom_step(linetype = "dashed", na.rm = TRUE, ...)`

**Value**

a ggplot2 figure

**See Also**

Visit the gallery for examples modifying the default figures
Examples

```r
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit() +
add_confidence_interval()

survfit2(Surv(time, status) ~ 1, data = df_lung) %>%
ggsurvfit() +
add_confidence_interval(type = "lines")
```

---

**Description**

Add a default or custom title to the figure legend.

**Usage**

```r
add_legend_title(title = NULL)
```

**Arguments**

- `title`: a string to override the default legend title. Default is `NULL`

**Value**

a `ggplot2` figure

**See Also**

Visit the `gallery` for examples modifying the default figures

**Examples**

```r
survfit2(Surv(time, status) ~ surg, data = df_colon) %>%
ggsurvfit() +
add_legend_title()
```


**add_pvalue**

### Description

- `add_pvalue("caption")`: Add a p-value to the figure via `ggplot2::labs(caption=)`
- `add_pvalue("annotation")`: Add a p-value text annotation via `ggplot2::annotation("text")`

P-values are calculated with `survival::survdiff()` or `tidycmprsk::glance()`. Examples of custom placement located in the help file for `survfit_p()`.

When a competing risks figure includes multiple outcomes, only the p-value comparing stratum for the *first* outcome can be placed.

### Usage

```r
add_pvalue(
  location = c("caption", "annotation"),
  caption = "{p.value}",
  prepend_p = TRUE,
  pvalue_fun = format_p,
  rho = 0,
  ...
)
```

### Arguments

- `location` string indicating where to place p-value. Must be one of c("caption", "annotation")
- `caption` string to be placed as the caption/annotation. String will be processed with `glue::glue()`, and the default is "{p.value}"
- `prepend_p` prepend "p=" to formatted p-value
- `pvalue_fun` function to round and style p-value with
- `rho` argument passed to `survival::survdiff(rho=)`
- `...` arguments passed to `ggplot2::annotate()`. Commonly used arguments are `x= and y=` to place the p-value at the specified coordinates on the plot.

### Value

A `ggplot2` figure

### See Also

`survfit_p()`
Examples

```r
survfit2(Surv(time, status) ~ surg, df_colon) %>%
ggsurvfit() +
add_pvalue(caption = "Log-rank (p.value)"

survfit2(Surv(time, status) ~ surg, df_colon) %>%
ggsurvfit() +
add_pvalue("annotation", size = 5)
```

---

**add_quantile**

Add Quantile Annotation

### Description

Add quantile information annotated on to the plot.

### Usage

```r
add_quantile(y_value = NULL, x_value = NULL, ...)
```

### Arguments

- `y_value, x_value`  
  Numeric value where the line segment will be drawn. Default is `y_value=0.5` when both `y_value` and `x_value` are unassigned.

- `...`  
  Named arguments passed to `ggplot2::geom_segment()` with default `linetype = 2`

### Value

A ggplot2 figure

### See Also

Visit the gallery for examples modifying the default figures

### Examples

```r
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit() +
add_quantile(linetype = 2) +
scale_ggsurvfit()

survfit2(Surv(time, status) ~ 1, data = df_lung) %>%
ggsurvfit() +
add_quantile(linetype = 2) +
add_quantile(y_value = 0.9, linetype = 3) +
scale_ggsurvfit()
```
```
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit() +
add_quantile(linetype = 2, y_value = NULL, x_value = 10) +
scale_ggsurvfit()
```

---

**Add risk table**

**Description**

Add risk tables below the plot showing the number at risk, events observed, and number of censored observations.

**Usage**

```r
add_risktable(
  times = NULL,
  risktable_stats = c("n.risk", "cum.event"),
  risktable_group = c("auto", "strata", "risktable_stats"),
  risktable_height = NULL,
  stats_label = NULL,
  combine_groups = FALSE,
  theme = theme_risktable_default(),
  size = 3.5,
  ...
)
```

**Arguments**

- `times` numeric vector of times where risk table values will be placed. Default are the times shown on the x-axis. The times passed here will not modify the tick marks shown on the figure. To modify which tick marks are shown, use `ggplot2::scale_x_continuous(breaks=).`

- `risktable_stats` character vector of statistics to show in the risk table. Must be one or more of `c("n.risk", "cum.event", "cum.censor", "n.event", "n.censor")`. Default is `c("n.risk", "cum.event")`.
  - "n.risk" Number of patients at risk
  - "cum.event" Cumulative number of observed events
  - "cum.censor" Cumulative number of censored observations
  - "n.event" Number of events in time interval
  - "n.censor" Number of censored observations in time interval

See additional details below.

- `risktable_group` String indicating the grouping variable for the risk tables. Default is "auto" and will select "strata" or "risktable_stats" based on context.
add_risktable

- "strata" groups the risk tables per stratum when present.
- "risktable_stats" groups the risk tables per risktable_stats.

risktable_height
A numeric value between 0 and 1 indicates the proportion of the final plot the risk table will occupy.

stats_label
named vector or list of custom labels. Names are the statistics from risktable_stats= and the value is the custom label.

combine_groups
logical indicating whether to combine the statistics in the risk table across groups. Default is FALSE

theme
A risk table theme. Default is theme_risktable_default()

size, ... arguments passed to ggplot2::geom_text(...). Pass arguments like, size = 4 to increase the size of the statistics presented in the table.

Value
a ggplot2 figure

Customize Statistics
You can customize how the statistics in the risk table are displayed by utilizing glue-like syntax in the risktable_stats argument.

For example, if you prefer to have the number at risk and the number of events on the same row, you can use risktable_stats = "{n.risk} ({cum.event})".

You can further customize the table to include the risk estimates using elements c("estimate", "conf.low", "conf.high", "std.error"). When using these elements, you’ll likely need to include a function to round the estimates and multiply them by 100.

add_risktable(
  risktable_stats =
    c("{n.risk} {{cum.event}}",
      "{round(estimate*100)}% {{round(conf.low*100)}, {round(conf.high*100)}}"),
  stats_label = c("At Risk (Cum. Events)", "Survival (95% CI)")
)

Competing Risks
The ggcuminc() can plot multiple competing events. The "cum.event" and "n.event" statistics are the sum of all events across outcomes shown on the plot.

See Also
Visit the gallery for examples modifying the default figures
add_risktable_strata_symbol

Examples

```r
p <-
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit() +
add_censor_mark() +
add_confidence_interval()

# using the function defaults
p + add_risktable()

# change the statistics shown and the label
p +
  add_risktable(
    risktable_stats = "n.risk",
    stats_label = list(n.risk = "Number at Risk"),
  )

p +
  add_risktable(
    risktable_stats = "{n.risk} (\{cum.event\})"
  )

p +
  add_risktable(
    risktable_stats = c("n.risk", "cum.event"),
    combine_groups = TRUE
  )
```

add_risktable_strata_symbol

*Use Symbol for Strata in Risk Table*

Description

Replace the stratum level names with a color symbol in the risk tables. Use this function when stratum level names are long.

Usage

```r
add_risktable_strata_symbol(
  symbol = NULL,
  size = 15,
  face = "bold",
  vjust = 0.3,
  ...
)
```
Arguments

symbol UTF-8 code of shape to replace strata level with. Default is a rectangle ("\U25AC"). Other common options are circle ("\U25CF") and diamond ("\U25C6"). While a symbol is the most common string to pass here, any string is acceptable.

size, face, vjust, ...
arguments passed to a function similar to ggtext::element_markdown()

Value

a ggplot2 figure

See Also

Visit the gallery for examples modifying the default figures

Examples

```r
p <- survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit(linewidth = 1) +
  add_confidence_interval() +
  add_risktable(risktable_group = "risktable_stats")

p + add_risktable_strata_symbol()
p + add_risktable_strata_symbol(symbol = "\U25CF", size = 10)
```

Description

Background The example simulated data set is based on large phase III clinical trials in breast cancer such as the ALTTO trial https://ascopubs.org/doi/abs/10.1200/JCO.2015.62.1797. The example “trial” aims to determine if a combination of two therapies tablemab (T) plus vismab (V) improves outcomes for metastatic human epidermal growth factor 2–positive breast cancer and increases the pathologic complete response in the neoadjuvant setting (i.e. treatment given as a first step to shrink a tumor before the main treatment or surgery).

Usage

adtte
Format

The data set contains the following variables:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>STUDYID</td>
<td>The study identifier. A code unique to the clinical trial</td>
</tr>
<tr>
<td>SUBJID</td>
<td>subject identifier. Numeric ID unique to each patient</td>
</tr>
<tr>
<td>USUBJID</td>
<td>unique subject identifier. Text ID combining study and patient IDs</td>
</tr>
<tr>
<td>AGE</td>
<td>age at randomisation (years)</td>
</tr>
<tr>
<td>STR01</td>
<td>Hormone receptor status at randomisation</td>
</tr>
<tr>
<td>STR01N</td>
<td>Hormone receptor positive (Numeric)</td>
</tr>
<tr>
<td>STR01L</td>
<td>Hormone receptor positive (Long format)</td>
</tr>
<tr>
<td>STR02</td>
<td>Prior Radiotherapy at randomisation</td>
</tr>
<tr>
<td>STR02N</td>
<td>Prior Radiotherapy at randomisation (Numeric)</td>
</tr>
<tr>
<td>STR02L</td>
<td>Prior Radiotherapy at randomisation (Long format)</td>
</tr>
<tr>
<td>TRT01P</td>
<td>Planned treatment assigned at randomisation</td>
</tr>
<tr>
<td>TRT01PN</td>
<td>Planned treatment assigned at randomisation (Numeric)</td>
</tr>
<tr>
<td>PARAM</td>
<td>Analysis parameter: Progression free survival</td>
</tr>
<tr>
<td>PARAMCD</td>
<td>Analysis parameter code</td>
</tr>
<tr>
<td>AVAL</td>
<td>Analysis value (time to event (years))</td>
</tr>
<tr>
<td>CNSR</td>
<td>Censoring (0 = Event, 1 = Censored)</td>
</tr>
<tr>
<td>EVNTDESC</td>
<td>Event description</td>
</tr>
<tr>
<td>CNSDTDSC</td>
<td>Censoring description</td>
</tr>
<tr>
<td>DCTREAS</td>
<td>Discontinuation from study reason</td>
</tr>
</tbody>
</table>

Details

The trial has four treatment arms, patients with centrally confirmed human epidermal growth factor 2-positive early breast cancer were randomly assigned to 1 year of adjuvant therapy with V, T, their sequence (T to V), or their combination (T+V) for 52 weeks.

The primary end point was progression-free survival (PFS) as defined by Cancer.gov: “the length of time during and after the treatment of a disease, such as cancer, that a patient lives with the disease but it does not get worse. In a clinical trial, measuring the progression-free survival is one way to see how well a new treatment works”.

A number of baseline measurements (taken at randomization) are also included such as age, hormone receptor status and prior radiotherapy treatment.

Additional details on reasons for study discontinuation and censoring event description are also included.

The data set adopts an abridged version of the CDISC ADaM ADTTE time to event data model. See here for more info on CDISC ADaM data standards https://www.cdisc.org/standards/foundational/adam and specifically the ADTTE time to event data model here https://www.cdisc.org/standards/foundational/adam/adam-basic-data-structure-bds-time-event-tte-analyses-v1-0.

Source

https://github.com/VIS-SIG/Wonderful-Wednesdays/tree/master/data/2020/2020-04-08
Description

This is a copy of the colon data set exported by the survival package. This data set, however, has column labels, numeric categorical variables are now factors with assigned levels, and we only include the recurrence outcome.

Usage

df_colon

Format

An object of class tbl_df (inherits from tbl.data.frame) with 929 rows and 14 columns.

Description

This is a copy of the lung data set exported by the survival package. This data set, however, has column labels and numeric categorical variables are now factors with assigned levels.

Usage

df_lung

Format

An object of class tbl_df (inherits from tbl.data.frame) with 228 rows and 10 columns.
**format_p**  

*Format p-value*

**Description**

Round and format p-values

**Usage**

format_p(x, digits = 1)

**Arguments**

- x: numeric vector of p-values
- digits: number of digits large p-values will be rounded to. Default is 2, and must be one of 1, 2, or 3.

**Value**

a string

**Examples**

```r
p_vec <- c(0.00001, 0.01111, 0.0500000, 0.15, 0.99999)
format_p(p_vec)
format_p(p_vec, 2)
format_p(p_vec, 3)
```

---

**ggcuminc**  

*Plot Cumulative Incidence*

**Description**

Plot a cumulative incidence object created with `tidycmprsk::cuminc()` or a multi-state object created with `survfit2()`. Read more on multi-state models [here](#).

**Usage**

```r
ggcuminc(
  x,
  outcome = NULL,
  linetype_aes = FALSE,
  theme = theme_ggsurvfit_default(),
  ...
)
```
Arguments

- **x**: a `survfit` object created with `survfit2()`
- **outcome**: string indicating which outcome(s) to include in plot. Default is to include the first competing event.
- **linetype_aes**: logical indicating whether to add `ggplot2::aes(linetype = strata)` to the `ggplot2::geom_step()` call. When strata are present, the resulting figure will be a mix of various line types for each stratum.
- **theme**: a survfit theme. Default is `theme_ggsurvfit_default()`
- **...**: arguments passed to `ggplot2::geom_step(...)`, e.g. `size = 2`

Value

A ggplot2 figure

Details

* Why do we not use `cmprsk::cuminc()`?

The implementation of `cmprsk::cuminc()` does not provide the data required to construct the risk table. Moreover, the `tidycmprsk::cuminc()` has a user-friendly interface making it easy to learn and use.

See Also

Visit the gallery for examples modifying the default figures

Examples

```r
library(tidycmprsk)

cuminc(Surv(ttdeath, death_cr) ~ trt, trial) %>%
  ggcuminc(outcome = "death from cancer") +
  add_confidence_interval() +
  add_risktable()

cuminc(Surv(ttdeath, death_cr) ~ trt, trial) %>%
  ggcuminc(outcome = c("death from cancer", "death other causes")) +
  add_risktable()

# using the survival multi-state model
survfit2(Surv(ttdeath, death_cr) ~ trt, trial) %>%
  ggcuminc(outcome = "death from cancer") +
  add_confidence_interval() +
  add_risktable()
```
**ggSurvfit**

*Plot Survival Probability*

**Description**

Plot survival probabilities (and other transformations) using the results from `survfit2()` or `survival::survfit()`; although, the former is recommend to have the best experience with the `ggSurvfit` package.

**Usage**

```r
ggSurvfit(
  x,
  type = "survival",
  linetype_aes = FALSE,
  theme = theme_ggsurvfit_default(),
  ...
)
```

**Arguments**

- `x`: a 'survfit' object created with `survfit2()`
- `type`: type of statistic to report. Available for Kaplan-Meier estimates only. Default is "survival". Must be one of the following:
  
<table>
<thead>
<tr>
<th>type</th>
<th>transformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;survival&quot;</td>
<td>x</td>
</tr>
<tr>
<td>&quot;risk&quot;</td>
<td>1 - x</td>
</tr>
<tr>
<td>&quot;cumhaz&quot;</td>
<td>-log(x)</td>
</tr>
</tbody>
</table>

- `linetype_aes`: logical indicating whether to add `ggplot2::aes(linetype = strata)` to the `ggplot2::geom_step()` call. When strata are present, the resulting figure will be a mix a various line types for each stratum.
- `theme`: a survfit theme. Default is `theme_ggsurvfit_default()`
- `...`: arguments passed to `ggplot2::geom_step(...)`, e.g. `size = 2`

**Value**

a ggplot2 figure

**Details**

This function creates a ggplot figure from the 'survfit' object. To better understand how to modify the figure, review the simplified code used internally:

```r
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
```
tidy_survfit() %>%
ggplot(aes(x = time, y = estimate, y
    min = conf.low, ymax = conf.low,
    color = strata, fill = strata)) +
geom_step()

See Also

Visit the gallery for examples modifying the default figures

Examples

# Default publication ready plot
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit()

# Changing statistic type
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit(type = "cumhaz")

# Configuring KM line type to vary by strata
survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit(linetype_aes = TRUE)

# Customizing the plot to your needs
survfit2(Surv(time, status) ~ 1, data = df_lung) %>%
ggsurvfit() +
add_censor_mark() +
add_confidence_interval() +
add_quantile() +
add_risktable()
scale_ggsurvfit

Arguments

  x
  an object of class 'ggsurvfit' or 'ggcuminc'

  combine_plots
  logical indicating whether to combine the primary plot and the risk tables. When TRUE, plot and risk table(s) are combined with patchwork::wrap_plots(). When FALSE and the plot has risk tables, they are returned in a list of gtable grobs. Default is TRUE.

Value

  a list of ggplot2 objects or a single ggplot2 object

See Also

  Visit the gallery for examples modifying the default figures

Examples

```
# construct plot
p <-
survfit2(Surv(time, status) ~ surg, df_colon) %>%
ggsurvfit() +
  add_risktable() +
  scale_y_continuous(limits = c(0, 1))

# build plots
built_p <- ggsurvfit_build(p, combine_plots = FALSE)

# reconstruct original figure print with risktables
patchwork::wrap_plots(
  built_p[[1]],
  built_p[[2]],
  built_p[[3]],
  ncol = 1,
  heights = c(0.70, 0.15, 0.15)
)

# place plots side-by-side (plots must be built before placement with patchwork)
patchwork::wrap_plots(
  ggsurvfit_build(p),
  ggsurvfit_build(p),
  ncol = 2
)
```
Description

The most common figure created with this package is a survival curve. This scale applies modifications often seen in these figures.

- `scale_y_continuous(expand = c(0.025, 0), limits = c(0, 1), label = scales::label_percent())`.
- `scale_x_continuous(expand = c(0.015, 0), n.breaks = 8)`

If you use this function, you **must** include all scale specifications that would appear in `scale_x_continuous()` or `scale_y_continuous()`. For example, it’s common you’ll need to specify the x-axis break points. `scale_ggsurvfit(x_scales=list(breaks=0:9))`.

To reset any of the above settings to their ggplot2 default, set the value to `NULL`, e.g. `y_scales = list(limits = NULL)`.

Usage

```r
scale_ggsurvfit(x_scales = list(), y_scales = list())
```

Arguments

- `x_scales` a named list of arguments that will be passed to `ggplot2::scale_x_continuous()`.
- `y_scales` a named list of arguments that will be passed to `ggplot2::scale_y_continuous()`.

Value

a ggplot2 figure

See Also

Visit the gallery for examples modifying the default figures

Examples

```r
ggsurvfit <-
survfit2(Surv(time, status) ~ surg, data = df_colon) %>%
ggsurvfit(linewidth = 1) +
  add_confidence_interval()

# use the function defaults
ggsurvfit + scale_ggsurvfit()

# specify additional scales
ggsurvfit +
  scale_ggsurvfit(x_scales = list(breaks = seq(0, 8, by = 2)))
```
stepribbon

Step ribbon statistic

Description

Provides stairstep values for ribbon plots

Usage

```r
stat_stepribbon(
  mapping = NULL,
  data = NULL,
  geom = "ribbon",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  direction = "hv",
  ...)
```

StatStepribbon

Arguments

- **mapping**: Set of aesthetic mappings created by `aes()`. If specified and `inherit.aes = TRUE` (the default), it is combined with the default mapping at the top level of the plot. You must supply `mapping` if there is no plot mapping.

- **data**: The data to be displayed in this layer. There are three options:
  - If `NULL`, the default, the data is inherited from the plot data as specified in the call to `ggplot()`.
  - A `data.frame`, or other object, will override the plot data. All objects will be fortified to produce a data frame. See `fortify()` for which variables will be created.
  - A function will be called with a single argument, the plot data. The return value must be a `data.frame`, and will be used as the layer data. A function can be created from a formula (e.g. `~ head(.x, 10)`).

- **geom**: which geom to use; defaults to "ribbon"

- **position**: Position adjustment, either as a string naming the adjustment (e.g. "jitter" to use `position_jitter`), or the result of a call to a position adjustment function. Use the latter if you need to change the settings of the adjustment.

- **na.rm**: If `FALSE`, the default, missing values are removed with a warning. If `TRUE`, missing values are silently removed.

- **show.legend**: logical. Should this layer be included in the legends? `NA`, the default, includes if any aesthetics are mapped. `FALSE` never includes, and `TRUE` always includes. It can also be a named logical vector to finely select the aesthetics to display.
survfit2

```r
survfit2(formula, ...)  
```

**Description**

Simple wrapper for `survival::survfit()` except the environment is also included in the returned object.

Use this function with all other functions in this package to ensure all elements are calculable.

**Usage**

`survfit2(formula, ...)`
Arguments

- **formula**: a formula object, which must have a Surv object as the response on the left of the ~ operator and, if desired, terms separated by + operators on the right. One of the terms may be a strata object. For a single survival curve the right hand side should be ~ 1.

Arguments passed on to `survival::survfit.formula`

- **data**: a data frame in which to interpret the variables named in the formula, subset and weights arguments.
- **weights**: The weights must be nonnegative and it is strongly recommended that they be strictly positive, since zero weights are ambiguous, compared to use of the subset argument.
- **subset**: expression saying that only a subset of the rows of the data should be used in the fit.
- **na.action**: a missing-data filter function, applied to the model frame, after any subset argument has been used. Default is `options()`$na.action.
- **stype**: the method to be used estimation of the survival curve: 1 = direct, 2 = exp(cumulative hazard).
- **ctype**: the method to be used for estimation of the cumulative hazard: 1 = Nelson-Aalen formula, 2 = Fleming-Harrington correction for tied events.
- **id**: identifies individual subjects, when a given person can have multiple lines of data.
- **cluster**: used to group observations for the infinitesimal jackknife variance estimate, defaults to the value of id.
- **robust**: logical, should the function compute a robust variance. For multi-state survival curves this is true by default. For single state data see details, below.
- **istate**: for multi-state models, identifies the initial state of each subject or observation
- **timefix**: process times through the aeqSurv function to eliminate potential roundoff issues.
- **etype**: a variable giving the type of event. This has been superseded by multi-state Surv objects and is deprecated; see example below.
- **error**: this argument is no longer used

Value

- `survfit2` object

`survfit2()` vs `survfit()`

Both functions have identical inputs, so why do we need `survfit2()`?

The only difference between `survfit2()` and `survival::survfit()` is that the former tracks the environment from which the call to the function was made.

The definition of `survfit2()` is unremarkably simple:
survfit2 <- function(formula, ...) {
  # construct survfit object
  survfit <- survival::survfit(formula, ...)

  # add the environment
  survfit$.Environment = <calling environment>

  # add class and return
  class(survfit) <- c("survfit2", "survfit")
  survfit
}

The environment is needed to ensure the survfit call can be accurately reconstructed or parsed at any
point post estimation. The call is parsed when p-values are reported and when labels are created.
For example, the raw variable names appear in the output of a stratified survfit() result, e.g.
"sex=Female". When using survfit2(), the originating data frame and formula may be parsed
and the raw variable names removed.

Most functions in the package work with both survfit2() and survfit(); however, the output
will be styled in a preferable format with survfit2().

See Also

survival::survfit.formula()

Examples

# With `survfit()`
fit <- survfit(Surv(time, status) ~ sex, data = df_lung)
fit

# With `survfit2()`
fit2 <- survfit2(Surv(time, status) ~ sex, data = df_lung)
fit2

# Consistent behavior with other functions
summary(fit, times = c(10, 20))

summary(fit2, times = c(10, 20))

survfit2_p Calculate p-value

Description

The function survfit2_p() wraps survival::survdiff() and returns a formatted p-value.

Usage

survfit2_p(x, pvalue_fun = format_p, prepend_p = TRUE, rho = 0)
Surv_CNSR

Arguments

- `x` a `survfit2` object
- `pvalue_fun` function to round and style p-value with
- `prepend_p` prepend "p=" to formatted p-value
- `rho` argument passed to `survival::survdiff(rho=)`

Value

- a string

Examples

```r
sf <- survfit2(Surv(time, status) ~ sex, data = df_lung)

sf %>%
  ggsurvfit() +
  add_confidence_interval() +
  add_risktable() +
  labs(caption = glue::glue("Log-rank \{survfit2\_p(sf)\}

sf %>%
  ggsurvfit() +
  add_confidence_interval() +
  add_risktable() +
  annotate("text", x = 2, y = 0.05, label = glue::glue("\{survfit2\_p(sf)\}
```

Surv_CNSR

Create a Survival Outcome from CDISC Data

Description

The aim of `Surv_CNSR()` is to map the inconsistency in data convention between the `survival` package and CDISC ADaM ADTTE data model.

The function creates a survival object (e.g. `survival::Surv()`) that uses CDISC ADaM ADTTE coding conventions and converts the arguments to the status/event variable convention used in the `survival` package.

The AVAL and CNSR arguments are passed to `survival::Surv(time = AVAL, event = 1 - CNSR, type = "right", origin = 0)`.

Usage

`Surv_CNSR(AVAL, CNSR)`
Arguments

AVAL  The follow-up time. The follow-up time is assumed to originate from zero. When no argument is passed, the default value is a column/vector named AVAL.

CNSR  The censoring indicator where 1=censored and 0=death/event. When no argument is passed, the default value is a column/vector named CNSR.

Value

Object of class ‘Surv’

Details

The Surv_CNSR() function creates a survival object utilizing the expected data structure in the CDISC ADaM ADTTE data model, mapping the CDISC ADaM ADTTE coding conventions with the expected status/event variable convention used in the survival package—specifically, the coding convention used for the status/event indicator. The survival package expects the status/event indicator in the following format: 0=alive, 1=dead. Other accepted choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death). A final but risky option is to omit the indicator variable, in which case all subjects are assumed to have an event.

The CDISC ADaM ADTTE data model adopts a different coding convention for the event/status indicator. Using this convention, the event/status variable is named ‘CNSR’ and uses the following coding: censor = 1, status/event = 0.

See Also

survival::Surv()

Examples

# Use the `Surv_CNSR()` function with ggsurvfit functions
survfit2(formula = Surv_CNSR() ~ STR01, data = adtte) %>%
  ggsurvfit() +
  add_confidence_interval()

# Use the `Surv_CNSR()` function with functions from other packages as well
survival::survfit(Surv_CNSR() ~ STR01, data = adtte)
survival::survreg(Surv_CNSR() ~ STR01 + AGE, data = adtte) %>%
  broom::tidy()

theme_ggsurvfit  Survfit Plot Themes

Description

Returns ggplot list of calls defining a theme.

- theme_ggsurvfit_default(): Builds on theme_bw() with increased text sizes.
- theme_ggsurvfit_KMunicate(): Theme to create KMunicate-styled figures. doi:10.1136/bmjopen2019030215
theme_risktable

Usage

theme_ggsurvfit_default()

theme_ggsurvfit_KMunicate()

Value

a ggplot2 theme

Examples

survfit2(Surv(time, status) ~ sex, data = df_lung) %>%
ggsurvfit(theme = theme_ggsurvfit_default())

theme_risktable  Risk Table Themes

Description

Returns ggplot list of calls defining a theme meant to be applied to a risk table.

Usage

theme_risktable_default(axis.text.y.size = 10, plot.title.size = 10.75)

theme_risktable_boxed(axis.text.y.size = 10, plot.title.size = 10.75)

Arguments

axis.text.y.size
text size of the labels on the left of the risk table

plot.title.size
text size of the risk table title

Value

a ggplot2 figure

Examples

p <- survfit2(Surv(time, status) ~ 1, data = df_lung) %>% ggsurvfit()

# default ------------------------------------
p + add_risktable(theme = theme_risktable_default())

# larger text --------------------------------
p +
  add_risktable(size = 4,
tidy_cuminc

Tidy a cuminc object

Description

The tidycmprsk package exports a tidier for "cuminc" objects. This function adds on top of that and returns more information.

Usage

tidy_cuminc(x, times = NULL)

Arguments

x a 'cuminc' object created with tidycmprsk::cuminc()
times numeric vector of times. Default is NULL, which returns all observed times.

Value

a tibble

Examples

library(tidycmprsk)
cuminc(Surv(ttdeath, death_cr) ~ trt, trial) %>% tidy_cuminc()
**tidy_survfit**

**Tidy a survfit object**

**Description**

The broom package exports a tidier for "survfit" objects. This function adds on top of that and returns more information. The function also utilizes additional information stored when the survfit object is created with survfit2(). It's recommended to always use this function with survfit2().

**Usage**

```r
tidy_survfit(x, times = NULL, type = c("survival", "risk", "cumhaz"))
```

**Arguments**

- `x`: a 'survfit' object created with survfit2()
- `times`: numeric vector of times. Default is NULL, which returns all observed times.
- `type`: type of statistic to report. Available for Kaplan-Meier estimates only. Default is "survival". Must be one of the following:

<table>
<thead>
<tr>
<th>type</th>
<th>transformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>&quot;survival&quot;</td>
<td>x</td>
</tr>
<tr>
<td>&quot;risk&quot;</td>
<td>1 - x</td>
</tr>
<tr>
<td>&quot;cumhaz&quot;</td>
<td>-log(x)</td>
</tr>
</tbody>
</table>

**Value**

a tibble

**Examples**

```r
survfit2(Surv(time, status) ~ factor(ph.ecog), data = df_lung) %>%
tidy_survfit()
```
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