

Package ‘TwoTimeScales’

May 7, 2026

Title Analysis of Event Data with Two Time Scales

Type Package

Version 1.2.1

URL <https://github.com/AngelaCar/TwoTimeScales>,
<https://angelacar.github.io/TwoTimeScales/>

Description Analyse time to event data with two time scales by estimating a smooth hazard that varies over two time scales. If covariates are available, estimate a proportional hazards model with such a two-dimensional baseline hazard.
Functions are provided to prepare the raw data for estimation, to fit the model and to plot the two-dimensional smooth hazard.
Extension to a competing risks model are implemented. For details about the method please refer to Carollo et al. (2025) <[doi:10.1002/sim.10297](https://doi.org/10.1002/sim.10297)>.

License GPL-3

Encoding UTF-8

LazyData true

RoxygenNote 7.3.3

Imports colorspace, Epi, fields, JOPS, LMMsolver, popEpi, reshape2,
spam, ucminf, viridis

Depends R (>= 3.5.0)

Suggests knitr, rmarkdown, spelling, testthat (>= 3.0.0)

Config/testthat/edition 3

VignetteBuilder knitr

BugReports <https://github.com/AngelaCar/TwoTimeScales/issues>

Language en-US

NeedsCompilation no

Author Angela Carollo [aut, cre, cph] (ORCID:
<<https://orcid.org/0000-0002-5625-6553>>),
Paul H.C. Eilers [aut],
Jutta Gampe [aut]

Maintainer Angela Carollo <carollo@demogr.mpg.de>

Repository CRAN

Date/Publication 2026-03-30 14:10:35 UTC

Contents

covariates_plot	3
cumhaz2ts	4
cuminc2ts	6
exposures_events_1d	7
exposures_events_2d	8
exposures_events_Lexis	10
fit1ts	11
fit2ts	15
fitpgam	19
fitvcm	22
get_hazard_1d	25
get_hazard_1d_LMM	26
get_hazard_2d	28
get_hazard_2d_LMM	30
get_hr	32
haz2tsLMM_summary	33
haz2tsPGAM_summary	34
haz2tsVCM_summary	35
haz2ts_summary	36
imageplot_2ts	37
imageplot_SE	38
make_bins	40
make_grid	42
plot.haz1ts	43
plot.haz1tsLMM	45
plot.haz2ts	47
plot.haz2tsLMM	51
plot.haz2tsPGAM	54
plot.haz2tsVCM	58
plot_slices	62
predict_cif2ts_pointwise	63
predict_comprisks2ts	64
predict_haz2ts	66
predict_haz2ts_pointwise	68
prepare_data	69
prepare_data_LMMsolver	73
print.data2ts	74
reccolon2ts	75
select_model2ts	76
surv2ts	78

Index

80

covariates_plot *Plot of the covariates' effects*

Description

covariates_plot() produces a plot of the covariates' effects ($\hat{\beta}$) with confidence intervals, or of the Hazard Ratios ($\exp(\hat{\beta})$) with confidence intervals.

Usage

```
covariates_plot(
  fitted_model,
  confidence_lev = 0.95,
  plot_options = list(),
  ...
)
```

Arguments

fitted_model A list returned by the function fit2ts(), fitpgam() or fit1ts().

confidence_lev The level of confidence for the CIs. Default is 0.95 ($\alpha = 0.05$).

plot_options A list of options for the plot:

- **HR** A Boolean. If TRUE the HRs with their CIs will be plotted. Default is FALSE (plot the beta with their CIs).
- **symmetric_ci** A Boolean. Default is TRUE. If a plot of the HRs is required (HR == TRUE), then plot symmetrical Confidence Intervals, based on the SEs for the HRs calculated by delta method. If FALSE, then CIs are obtained by exponentiating the CIs for the betas.
- **main** The title of the plot.
- **ylab** The label for the y-axis.
- **ylim** A vector with two elements defining the limits for the y-axis.
- **col_beta** The color for the plot of the covariates' effects.
- **pch** The symbol for plotting the point estimates.
- **cex_main** The magnification factor for the main of the plot.

... further arguments passed to plot()

Value

A plot of the covariates' effects. The different covariates are plotted on the x-axis, and on the y-axis the effects on the coefficient- or on the HR-scale are plotted. The main estimate is represented by a point and the CIs are added as vertical bars.

Examples

```

# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)
x1 <- c(0, 0, 0, 1, 0, 1, 1, 1, 1, 0, 0, 0, 1, 0, 1, 1, 0, 0, 1, 0)

fakedata <- as.data.frame(cbind(id, u, s, ev, x1))
covs <- subset(fakedata, select = c("x1"))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5,
  individual = TRUE,
  covs = covs
)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)
# Covariates plot with default options
covariates_plot(fakemod)

# Plot the hazard ratios instead
covariates_plot(fakemod,
  plot_options = list(
    HR = TRUE
  )
)

# Change confidence level
covariates_plot(fakemod,
  confidence_lev = .99
)

```

Description

Computes the cumulative hazard surface over two time scales from a fitted model. The function is also called internally from `plot()` if the user wants to plot the cumulative hazard from a fitted model.

Usage

```
cumhaz2ts(
  fitted_model,
  plot_grid = NULL,
  cause = NULL,
  midpoints = FALSE,
  where_slices = NULL,
  direction = c("u", "s", NULL),
  tmax = NULL
)
```

Arguments

<code>fitted_model</code>	(optional) An object of class 'haz2ts', 'haz2tsLMM', 'haz2tsPGAM', or 'haz2tsVCM'.
<code>plot_grid</code>	(optional) A list containing the parameters to build a new finer grid of intervals over <code>u</code> and <code>s</code> for plotting. This must be of the form: <ul style="list-style-type: none"> <code>plot_grid = list(c(umin, umax, du), c(smin, smax, ds))</code> where <code>umin</code>, <code>umax</code> and <code>smin</code>, <code>smax</code> are the minimum and maximum values desired for the grid-points over <code>u</code> and <code>s</code> respectively, and <code>du</code>, <code>ds</code> are distances between two adjacent points over <code>u</code> and <code>s</code> respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of detail. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.
<code>cause</code>	a character string with a short name for the cause (optional).
<code>midpoints</code>	A Boolean. Default is <code>FALSE</code> . If <code>TRUE</code> , the estimated quantities are evaluated at the midpoints of the rectangles (or parallelograms) of the grids, rather than at each grid-point.
<code>where_slices</code>	A vector of values for the cutting points of the desired slices of the surface. This option is included mostly for the plotting function. When using <code>plot.haz2ts()</code> , the user selects <code>which_plot = "cumhaz"</code> and <code>cumhaz_slices = TRUE</code> , then <code>where_slices</code> indicates the location of the cutting points over the <code>u</code> time.
<code>direction</code>	If cross-sectional one-dimensional curves are plotted, this indicates whether the cutting points are located on the <code>u</code> time, or on the <code>s</code> time. For plots of the cumulative hazards, only cutting points over the <code>u</code> time are meaningful.
<code>tmax</code>	The maximum value of <code>t</code> that should be plotted.

Value

A list with the following elements: * Haz a list of estimated hazard and associated SEs (obtained from the function `get_hazard_2d()`); * CumHaz the cumulated hazard estimate over `u` and `s`; * cause (if provided) the short name for the cause.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(u = fakedata$u,
                          s_out = fakedata$s,
                          ev = fakedata$ev,
                          ds = .5)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "grid_search",
                 lrho = list(seq(1, 1.5, .5),
                             seq(1, 1.5, .5)))

# Obtain the fake cumulated hazard
fakecumhaz2ts <- cumhaz2ts(fakemod)
```

cuminc2ts

Cumulative incidence surface over two time scales

Description

Cumulative incidence surface over two time scales

Usage

```
cuminc2ts(haz = list(), ds, cause = NULL)
```

Arguments

haz	a list of cause-specific hazards
ds	the distance between two consecutive intervals over the <code>s</code> time scale. This has to be equal for all cause-specific hazards
cause	is an optional vector of short names for the causes. It should be of the same length as the number of cause-specific cumulated hazards provided.

Value

a list with one cumulative incidence matrix for each cause-specific hazard (named if a vector of short names is passed to cause).

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(u = fakedata$u,
                          s_out = fakedata$s,
                          ev = fakedata$ev,
                          ds = .5)

# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                optim_method = "grid_search",
                lrho = list(seq(1, 1.5, .5),
                           seq(1, 1.5, .5)))

# Obtain the fake cumulated hazard
fakecumhaz2ts <- cumhaz2ts(fakemod)
# Fake cumulative incidence function 2ts
fakecif2ts <- cuminc2ts(haz = list(fakecumhaz2ts$Haz$hazard),
                       ds = .5)
```

exposures_events_1d *Bin data on one time scale*

Description

exposure_events_1d() computes aggregated measures of exposure times and event counts starting from individual records of time at entry, time at exit and event's indicator, over one time scale (s).

Usage

```
exposures_events_1d(s_in = NULL, s_out, ev, bins)
```

Arguments

s_in A vector of (possibly left truncated) times at entry. If this is not provided by the user, the function will consider a value of 0 for all observations.

s_out A vector of times at event or censoring.

ev A vector of events' indicators (possible values 0/1).
 bins A vector of interval breaks for discretization (see also [make_bins\(\)](#)).

Details

The time scale s is divided into bins of equal size, which are provided as input to the function. Then, the time-at-risk for each individual is split according to these bins, and an event indicator is placed in the bin where the exit time is located. Finally, the individual contributions are summed in each bin to provide a vector of total exposure time and total event counts. See also [prepare_data\(\)](#) to conveniently prepare individual data for the analysis with one, or two time scales.

Value

A list with the following elements:

- R A matrix of dimension n by ns containing the exposure times for each individual separately.
- r A vector of exposure times.
- Y A matrix of dimension n by ns containing the event counts for each individual separately
- y A vector of event counts.

If the length of the input vectors do not match, an error message is returned.

Author(s)

Angela Carollo <carollo@demogr.mpg.de> and Paul Eilers <p.eilers@erasmus.nl>

Examples

```
# ---- Bin colon cancer data by time since recurrence ----
# First create vector of bins
bins1ts <- make_bins(s_in = reccolon2ts$entrys, s_out = reccolon2ts$timesr, ds = 30)
bindata <- exposures_events_1d(s_in = reccolon2ts$entrys,
s_out = reccolon2ts$timesr, ev = reccolon2ts$status, bins = bins1ts$bins_s)
```

exposures_events_2d *Bin data on two time scales*

Description

`exposures_events_2d()` computes individual or aggregated matrices of exposure times and event counts starting from individual records of time at entry in the process (measured over the first time scale), duration at entry in the process (measured over the second time scale), duration at exit from the process (measured over the second time scale), and event's indicator.

Usage

```
exposures_events_2d(u, s_in = NULL, s_out, ev, bins_list, individual = FALSE)
```

Arguments

<code>u</code>	A vector of fixed times at entry in the process, measured over the first time scale.
<code>s_in</code>	A vector of (possibly left truncated) times at entry. If this is not provided by the user, the function will consider a value of 0 for all observations.
<code>s_out</code>	A vector of times at event or censoring.
<code>ev</code>	A vector of events' indicators (possible values 0/1).
<code>bins_list</code>	is a list with the following (necessary) elements (usually prepared by <code>make_bins()</code>): <ul style="list-style-type: none"> • <code>bins_u</code> a vector of extreme values for the bins over the <code>u</code> axis • <code>bins_s</code> a vector of extreme values for the bins over the <code>s</code> axis
<code>individual</code>	A Boolean. Default is FALSE: if FALSE computes the matrices R and Y collectively for all observations; if TRUE computes the matrices R and Y separately for each individual record.

Details

The fixed-time variable `u` and the second time scale `s` are divided into n_u and n_s intervals, respectively. The extremes of these intervals are provided as input to the function. First, the fixed-time at entry is located in one of the n_u bins that cover the whole range of `u`. Then, the time-at-risk for each individual is split according to the n_s bins that span the whole range of values for `s`, and an event indicator is placed in the bin where the exit time is located. This is done by calling the function `exposure_events_1d()`. If individual matrices of exposure and events are required, then the function returns two arrays of dimension n_u by n_s by n . If aggregated results are preferred, the individual contributions are summed in each bin to provide a matrix of total exposure times and a matrix of total event counts, both of dimensions n_u by n_s . See also `prepare_data()` to conveniently prepare individual data for the analysis with one, or two time scales.

Value

A list with the following elements:

- R an array of exposure times: if `individual == TRUE`, then R is an array of dimension n_u by n_s by n , otherwise is an array of dimension n_u by n_s
- Y an array of event counts: if `individual == TRUE`, then Y is an array of dimension n_u by n_s by n , otherwise is an array of dimension n_u by n_s

Author(s)

Angela Carollo <carollo@demogr.mpg.de>

Examples

```
# ---- Bin colon cancer data by time at randomization and time since recurrence ----
# First create vectors of bins (using function `make_bins()`)
bins <- make_bins(u = reccolon2ts$timer, s_out = reccolon2ts$timesr,
  du = 30, ds = 30)
# Now bin data (note: the s_in argument is omitted because data are not left truncated)
bindata2d <- exposures_events_2d(u = reccolon2ts$timer,
  s_out = reccolon2ts$timesr, ev = reccolon2ts$status, bins = bins)
```

exposures_events_Lexis

Bin data on the Lexis diagram

Description

exposures_events_Lexis() computes aggregated matrices of exposure times and event counts over two time scales, on the Lexis diagram.

The time scales are t and s . This function uses functions from the package popEpi and from the package Epi, and code shared by Bendix Carstensen on the website bendixcarstensen.com. See also [prepare_data\(\)](#) to conveniently prepare individual data for the analysis with one, or two time scales.

Usage

```
exposures_events_Lexis(t_in = NULL, t_out, s_in = NULL, s_out, ev, bins_list)
```

Arguments

- | | |
|-----------|---|
| t_in | (optional) A vector of entry times on the time scale t . |
| t_out | (optional) A vector of exit times on the time scale t . |
| s_in | (optional) A vector of entry times on the time scale s . |
| s_out | A vector of exit times on the time scale s . |
| ev | A vector of event indicators (possible values 0/1). |
| bins_list | A list with the following (necessary) elements: <ul style="list-style-type: none">• bins_t a vector of extreme values for the bins over the t axis.• nt the number of bins over t.• bins_s a vector of extreme values for the bins over the s axis.• ns the number of bins over s. |

Value

A list with the following elements:

- R an array of exposure times of dimension n_t by n_s
- Y an array of event counts of dimension n_t by n_s

Author(s)

Angela Carollo <carollo@demogr.mpg.de>

References

Carstensen B, Plummer M, Laara E, Hills M (2022). Epi: A Package for Statistical Analysis in Epidemiology. R package version 2.47.1, <https://CRAN.R-project.org/package=Epi>.

Miettinen J, Rantanen M, Seppa K (2023). popEpi: Functions for Epidemiological Analysis using Population Data. R package version 0.4.11, <https://cran.r-project.org/package=popEpi>.

Examples

```
# ---- Bin colon cancer data by time since randomization and time since recurrence ----
# First create vectors of bins (using function `make_bins()`)
bins <- make_bins(t_out = reccolon2ts$timedc, s_out = reccolon2ts$timesr,
dt = 90, ds = 90)
# Now bin data (note: the t_in and s_in arguments are omitted because data are not left truncated)
bindata2d <- exposures_events_Lexis(t_out = reccolon2ts$timedc,
s_out = reccolon2ts$timesr, ev = reccolon2ts$status, bins = bins)
```

fit1ts

Fit a smooth hazard model with one time scale

Description

fit1ts() fits a smooth hazard model with one time scale.

Three methods are implemented for the search of the optimal smoothing parameter (and therefore optimal model): a numerical optimization of the AIC or BIC of the model, a search for the minimum AIC or BIC of the model over a grid of \log_{10} values for the smoothing parameter and the estimation using a sparse mixed model representation of P-splines. Construction of the B-splines basis and of the penalty matrix is incorporated within the function. If a matrix of covariates is provided, the function will estimate a model with covariates.

Usage

```
fit1ts(
  data1ts = NULL,
  y = NULL,
  r = NULL,
  Z = NULL,
  bins = NULL,
  Bbases_spec = list(),
  Wprior = NULL,
  pord = 2,
  optim_method = c("ucminf", "grid_search", "LMMsolver"),
  optim_criterion = c("aic", "bic"),
  lrho = 0,
  ridge = 0,
  control_algorithm = list(),
  par_gridsearch = list()
)
```

Arguments

<code>data1ts</code>	(optional) an object created by the function <code>prepare_data()</code> . Providing this input is the easiest way to use the function <code>fit1ts()</code> . However, the user can also provide the input data together with a list of bins, as explained by the following parameters' descriptions.
<code>y</code>	A vector of event counts of length <code>ns</code> , or an array of dimension <code>ns</code> by <code>n</code> .
<code>r</code>	A vector of exposure times of length <code>ns</code> , or an array of dimension <code>ns</code> by <code>n</code> .
<code>Z</code>	(optional) A regression matrix of covariates of dimension <code>n</code> by <code>p</code> .
<code>bins</code>	a list with the specification for the bins. This is created by the function <code>prepare_data()</code> . Alternatively, a list with the following elements can be provided: * <code>bins_s</code> is a vector of intervals for the time scale <code>s</code> . * <code>mids</code> is a vector with the midpoints of the intervals over <code>s</code> . * <code>ns</code> is the number of bins over <code>s</code> .
<code>Bbases_spec</code>	A list with the specification for the B-splines basis with the following elements: <ul style="list-style-type: none"> • <code>bdeg</code> The degree of the B-splines basis. Default is 3 (for cubic B-splines). • <code>nseg_s</code> The number of segments for the B-splines over <code>s</code>. Default is 10. • <code>min_s</code> (optional) The lower limit of the domain of Bs. Default is <code>min(bins_s)</code>. • <code>max_s</code> (optional) The upper limit of the domain of Bs. Default is <code>max(bins_s)</code>.
<code>Wprior</code>	An optional vector of a-priori weights.
<code>pord</code>	The order of the penalty. Default is 2.
<code>optim_method</code>	The method to be used for optimization: "ucminf" (default) for the numerical optimization of the AIC (or BIC), "grid_search" for a grid search of the minimum AIC (or BIC) over a grid of $\log_{10}(\varrho_s)$ values, and "LMMsolver" to solve the model as sparse linear mixed model using the package LMMsolver.
<code>optim_criterion</code>	The criterion to be used for optimization: "aic" (default) or "bic". BIC penalizes model complexity more strongly than AIC, so that its usage is recommended when a smoother fit is preferable (see also Camarda, 2012).
<code>lrho</code>	A number if <code>optim_method == "ucminf"</code> , default is 0. A vector of values for $\log_{10}(\varrho_s)$ if <code>optim_method == "grid_search"</code> . In the latter case, if a vector is not provided, a default sequence of values is used for $\log_{10}(\varrho_s)$.
<code>ridge</code>	A ridge penalty parameter: default is 0.
<code>control_algorithm</code>	A list with optional values for the parameters of the iterative processes: <ul style="list-style-type: none"> • <code>maxiter</code> The maximum number of iteration for the IWSL algorithm. Default is 20. • <code>conv_crit</code> The convergence criteria, expressed as difference between estimates at iteration <code>i</code> and <code>i+1</code>. Default is <code>1e-5</code>. • <code>verbose</code> A Boolean. Default is FALSE. If TRUE monitors the iteration process. • <code>monitor_ev</code> A Boolean. Default is FALSE. If TRUE monitors the evaluation of the model over the $\log_{10}(\rho_s)$ values. • <code>xtol</code> The relative tolerance to stop the algorithm, only relevant for numerical optimization algorithm. For details see <code>help(ucminf)</code>. Here is it set to <code>1e-5</code>.

par_gridsearch A list of parameters for the grid_search:

- plot_aic A Boolean. Default is FALSE. If TRUE, plot the AIC values over the grid of $\log_{10}(\text{rhos})$ values.
- plot_bic A Boolean. Default is FALSE. If TRUE, plot the BIC values over the grid of $\log_{10}(\text{rhos})$ values.
- return_aic A Boolean. Default is TRUE. Return the AIC values.
- return_bic A Boolean. Default is TRUE. Return the BIC values.
- mark_optimal A Boolean. Default is TRUE. If the plot of the AIC or BIC values is returned, marks the optimal $\log_{10}(\text{rho}_s)$ in the plot.
- main_aic The title of the AIC plot. Default is "AIC grid".
- main_bic The title of the BIC plot. Default is "BIC grid".

Details

Some functions from the R-package LMMsolver are used here. We refer the interested readers to <https://biometris.github.io/LMMsolver/> for more detail on LMMsolver and its usage.

Value

An object of class haz1ts, or of class haz1tsLMM. For objects of class haz1ts this is

- optimal_model A list with:
 - alpha The vector of estimated P-splines coefficients of length c_s .
 - SE_alpha The vector of estimated Standard Errors for the alpha coefficients, of length c_s .
 - beta The vector of estimated covariate coefficients of length p (if model with covariates).
 - se_beta The vector of estimated Standard Errors for the beta coefficients of length p (if model with covariates).
 - eta or eta0. The vector of values of the (baseline) linear predictor (log-hazard) of length n_s .
 - H The hat-matrix.
 - Cov The full variance-covariance matrix.
 - deviance The deviance.
 - ed The effective dimension of the model.
 - aic The value of the AIC.
 - bic The value of the BIC.
 - Bbases a list with the B-spline basis Bs (this is a list for compatibility with functions in 2d).
- optimal_logrho The optimal value of $\log_{10}(\text{rho}_s)$.
- P_optimal The optimal penalty matrix P.
- AIC (if par_gridsearch\$return_aic == TRUE) The vector of AIC values.
- BIC (if par_gridsearch\$return_bic == TRUE) The vector of BIC values.

Objects of class haz1tsLMM have a slight different structure. They are a list with:

- optimal_model an object of class LMMsolve

- AIC_BIC a list with, among other things, the AIC and BIC values and the ED of the model
- n_events the number of events
- ns the number of bins over the s-axis
- cs the number of B-splines over the s-axis
- covariates an indicator for PH model

References

Boer, Martin P. 2023. "Tensor Product P-Splines Using a Sparse Mixed Model Formulation." *Statistical Modelling* 23 (5-6): 465–79. [#](https://doi.org/10.1177/1471082X231178591)

Examples

```
## preparing data - no covariates
dt1ts <- prepare_data(
  data = reccolon2ts,
  s_in = "entrys",
  s_out = "timesr",
  events = "status",
  ds = 180
)

## fitting the model with fit1ts() - default options, that is ucminf optimization
mod1 <- fit1ts(dt1ts)

## fitting with LMMsolver
mod2 <- fit1ts(dt1ts,
  optim_method = "LMMsolver"
)

## preparing the data - covariates
dt1ts_cov <- prepare_data(
  data = reccolon2ts,
  s_in = "entrys",
  s_out = "timesr",
  events = "status",
  ds = 180,
  individual = TRUE,
  covs = c("rx", "node4", "sex")
)

## fitting the model with fit1ts() - grid search over only two log10(rho_s) values
mod3 <- fit1ts(dt1ts_cov,
  optim_method = "grid_search",
  lrho = c(1, 1.5)
)
```

fit2ts

*Fit a smooth hazard model with two time scales***Description**

fit2ts() fits a smooth hazard model with two time scales.

Three methods are implemented for the search of the optimal smoothing parameters (and therefore optimal model): a numerical optimization of the AIC or BIC of the model, a search for the minimum AIC or BIC of the model over a grid of \log_{10} values for the smoothing parameters, and a solution that uses a sparse mixed model representation of the P-spline model to estimate the smoothing parameters. Construction of the B-splines bases and of the penalty matrix is incorporated within the function. If a matrix of covariates is provided, the function will estimate a model with covariates.

Usage

```
fit2ts(
  data2ts = NULL,
  Y = NULL,
  R = NULL,
  Z = NULL,
  bins = NULL,
  Bbases_spec = list(),
  pord = 2,
  optim_method = c("ucminf", "grid_search", "LMMsolver"),
  optim_criterion = c("aic", "bic"),
  lrho = c(0, 0),
  Wprior = NULL,
  ridge = 0,
  control_algorithm = list(),
  par_gridsearch = list()
)
```

Arguments

data2ts	(optional) an object of class "data2ts" created by the function prepare_data(). Providing this input is the easiest way to use the function fit2ts. However, the user can also provide the input data together with a list of bins, as explained by the following parameters' descriptions.
Y	A matrix (or 3d-array) of event counts of dimension nu by ns (or nu by ns by n).
R	A matrix (or 3d-array) of exposure times of dimension nu by ns (or nu by ns by n).
Z	(optional) A regression matrix of covariates values of dimensions n by p.
bins	a list with the specification for the bins. This is created by the function prepare_data. If a list prepared externally from such function if provided, it should contain the following elements: * bins_u A vector of bins extremes for the time scale u. *

	<p>midu A vector with the midpoints of the bins over u. * nu The number of bins over u. * bins_s A vector of bins extremes for the time scale s. * mids A vector with the midpoints of the bins over s. * ns The number of bins over s.</p>
Bbases_spec	<p>A list with the specification for the B-splines basis with the following elements:</p> <ul style="list-style-type: none"> • bdeg The degree of the B-splines basis. Default is 3 (for cubic B-splines). • nseg_u The number of segments for the B-splines over u. Default is 10. • min_u (optional) The lower limit of the domain of Bu. Default is min(bins_u). • max_u (optional) The upper limit of the domain of Bu. Default is max(bins_u). • nseg_s The number of segments for the B-splines over s. Default is 10. • min_s (optional) The lower limit of the domain of Bs. Default is min(bins_s). • max_s (optional) The upper limit of the domain of Bs. Default is max(bins_s).
pord	The order of the penalty. Default is 2.
optim_method	The method to be used for optimization: "ucminf" (default) for the numerical optimization of the AIC (or BIC), "grid_search" for a grid search of the minimum AIC (or BIC) over a grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values, or "LMMsolver" to solve the model as sparse linear mixed model using the package LMMsolver.
optim_criterion	The criterion to be used for optimization: "aic" (default) or "bic". BIC penalizes model complexity more strongly than AIC, so that its usage is recommended when a smoother fit is preferable (see also Camarda, 2012).
lrho	A vector of two elements if optim_method == "ucminf". Default is c(0, 0). A list of two vectors of values for $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ if optim_method == "grid_search". In the latter case, if a list with two vectors is not provided, a default sequence of values is used for both $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$.
Wprior	An optional matrix of a-priori weights.
ridge	A ridge penalty parameter: default is 0. This is useful when, in some cases the algorithm shows convergence problems. In this case, set to a small number, for example 1e-4.
control_algorithm	<p>A list with optional values for the parameters of the iterative processes:</p> <ul style="list-style-type: none"> • maxiter The maximum number of iteration for the IWSL algorithm. Default is 20. • conv_crit The convergence criteria, expressed as difference between estimates at iteration i and i+1. Default is 1e-5. • verbose A Boolean. Default is FALSE. If TRUE monitors the iteration process. • monitor_ev A Boolean. Default is FALSE. If TRUE monitors the evaluation of the model over the $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values. • xtol The relative tolerance to stop the algorithm, only relevant for numerical optimization algorithm. For details see help(ucminf). Here is it set to 1e-5.
par_gridsearch	A list of parameters for the grid_search:

- `plot_aic` A Boolean. Default is FALSE. If TRUE, plot the AIC values over the grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values.
- `plot_bic` A Boolean. Default is FALSE. If TRUE, plot the BIC values over the grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values.
- `return_aic` A Boolean. Default is TRUE. Return the AIC values.
- `return_bic` A Boolean. Default is TRUE. Return the BIC values.
- `col` The color palette to be used for the AIC/BIC plot. Default is `grDevices::gray.colors(n=10)`.
- `plot_contour` A Boolean. Default is TRUE. Adds white contour lines to the AIC/BIC plot.
- `mark_optimal` A Boolean. Default is TRUE. If the plot of the AIC or BIC values is returned, marks the optimal combination of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ in the plot.
- `main_aic` The title of the AIC plot. Default is "AIC grid".
- `main_bic` The title of the BIC plot. Default is "BIC grid".

Details

Some functions from the R-package `LMMsolver` are used here. We refer the interested readers to <https://biometris.github.io/LMMsolver/> for more details on `LMMsolver` and its usage.

For the numerical optimization using ``ucminf`` a tolerance parameter is specified internally: ``control = list(xtol = 1e-5)``. The default in ``ucminf`` is ``xtol = 1e-12``. As `ucminf` is used here to find the minimum AIC or BIC, often this level of precision is not needed. Nevertheless, this value can be adjusted in ``control_algorithm``.

Value

An object of class `haz2ts`, or of class `haz2tsLMM`. For objects of class `haz2ts` this is

- `optimal_model` A list with :
 - `Alpha` The matrix of estimated P-splines coefficients of dimension c_u by c_s .
 - `Cov_alpha` The variance-covariance matrix of the Alpha coefficients, of dimension $c_u c_s$ by $c_u c_s$.
 - `beta` The vector of length p of estimated covariates coefficients (if model with covariates).
 - `Cov_beta` The variance-covariance matrix of the beta coefficients, of dimension p by p (if model with covariates).
 - `SE_beta` The vector of length p of estimated Standard Errors for the beta coefficients (if model with covariates)..
 - `Eta` or `Eta0` The matrix of values of the (baseline) linear predictor (log-hazard) of dimension n_u by n_s .
 - `H` The hat-matrix.
 - `deviance` The deviance.
 - `ed` The effective dimension of the model.
 - `aic` The value of the AIC.
 - `bic` The value of the BIC.

- Bbases a list with the B-spline bases Bu and Bs
- optimal_logrho A vector with the optimal values of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$.
- P_optimal The optimal penalty matrix P.
- AIC (if par_gridsearch\$return_aic == TRUE) The matrix of AIC values.
- BIC (if par_gridsearch\$return_bic == TRUE) The matrix of BIC values.

Objects of class haz2tsLMM have a slight different structure. They are a list with:

- optimal_model an object of class LMMsolve
- AIC_BIC a list with, among other things, the AIC and BIC values and the ED of the model
- n_events the number of events
- nu the number of bins over the u-axis
- ns the number of bins over the s-axis
- cu the number of B-splines over the u-axis
- cs the number of B-splines over the s-axis
- covariates an indicator for PH model

References

Boer, Martin P. 2023. "Tensor Product P-Splines Using a Sparse Mixed Model Formulation." *Statistical Modelling* 23 (5-6): 465–79. doi:10.1177/1471082X231178591. Carollo, A., Eilers, P., Putter, H. and Gampe, J. (2025), "Smooth Hazards With Multiple Time Scales." *Statistics in Medicine*, 44: e10297. doi:10.1002/sim.10297

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5
)
# Fit a fake model - not optimal smoothing
fit2ts(fakedata2ts,
```

```

  optim_method = "grid_search",
  lrho = list(seq(1, 1.5, .5), seq(1, 1.5, .5))
)
# For more examples please check the vignettes! Running more complicated examples
# here would imply longer running times...

```

fitpgam

Fit a log-additive model over two time scales

Description

Fits a log-additive model over two time scales. If covariates are included in the model, `fitpgam()` will fit a proportional hazards model, where the baseline hazard is a surface over the two time scales, where the effect of the time scales is additive on the log-scale. The model is described in Carollo et al. (2025).

Usage

```

fitpgam(
  data2ts = NULL,
  Y = NULL,
  R = NULL,
  Z = NULL,
  bins = NULL,
  Bbases_spec = list(),
  pord = 2,
  ridge = 1e-06,
  lrho = c(0, 0),
  optim_method = c("ucminf", "grid_search"),
  optim_criterion = c("aic", "bic"),
  control_algorithm = list(),
  par_gridsearch = list()
)

```

Arguments

<code>data2ts</code>	(optional) an object of class "data2ts" created by the function <code>prepare_data()</code> . Providing this input is the easiest way to use the function <code>fitpgam()</code> . However, the user can also provide the input data together with a list of bins, as explained by the following parameters' descriptions.
<code>Y</code>	A matrix (or 3d-array) of event counts of dimension <code>nu</code> by <code>ns</code> (or <code>nu</code> by <code>ns</code> by <code>n</code>).
<code>R</code>	A matrix (or 3d-array) of exposure times of dimension <code>nu</code> by <code>ns</code> (or <code>nu</code> by <code>ns</code> by <code>n</code>).
<code>Z</code>	(optional) A regression matrix of covariates values of dimensions <code>n</code> by <code>p</code> .

bins	a list with the specification for the bins. This is created by the function <code>prepare_data()</code> . If a list prepared externally from such function is provided, it should contain the following elements: * <code>bins_u</code> A vector of bins extremes for the time scale <code>u</code> . * <code>midu</code> A vector with the midpoints of the bins over <code>u</code> . * <code>nu</code> The number of bins over <code>u</code> . * <code>bins_s</code> A vector of bins extremes for the time scale <code>s</code> . * <code>mids</code> A vector with the midpoints of the bins over <code>s</code> . * <code>ns</code> The number of bins over <code>s</code> .
Bbases_spec	A list with the specification for the B-splines basis with the following elements: <ul style="list-style-type: none"> • <code>bdeg</code> The degree of the B-splines basis. Default is 3 (for cubic B-splines). • <code>nseg_u</code> The number of segments for the B-splines over <code>u</code>. Default is 10. • <code>min_u</code> (optional) The lower limit of the domain of <code>Bu</code>. Default is <code>min(bins_u)</code>. • <code>max_u</code> (optional) The upper limit of the domain of <code>Bu</code>. Default is <code>max(bins_u)</code>. • <code>nseg_s</code> The number of segments for the B-splines over <code>s</code>. Default is 10. • <code>min_s</code> (optional) The lower limit of the domain of <code>Bs</code>. Default is <code>min(bins_s)</code>. • <code>max_s</code> (optional) The upper limit of the domain of <code>Bs</code>. Default is <code>max(bins_s)</code>.
pord	The order of the penalty. Default is 2.
ridge	A ridge penalty parameter: default is 0. This is useful when, in some cases the algorithm shows convergence problems. In this case, set to a small number, for example <code>1e-4</code> .
lrho	A vector of two elements if <code>optim_method == "ucminf"</code> . Default is <code>c(0, 0)</code> . A list of two vectors of values for $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ if <code>optim_method == "grid_search"</code> . In the latter case, if a list with two vectors is not provided, a default sequence of values is used for both $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$.
optim_method	The method to be used for optimization: <code>"ucminf"</code> (default) for the numerical optimization of the AIC (or BIC), <code>"grid_search"</code> for a grid search of the minimum AIC (or BIC) over a grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values.
optim_criterion	The criterion to be used for optimization: <code>"aic"</code> (default) or <code>"bic"</code> . BIC penalizes model complexity more strongly than AIC, so that its usage is recommended when a smoother fit is preferable (see also Camarda, 2012).
control_algorithm	A list with optional values for the parameters of the iterative processes: <ul style="list-style-type: none"> • <code>maxiter</code> The maximum number of iteration for the IWSL algorithm. Default is 20. • <code>conv_crit</code> The convergence criteria, expressed as difference between estimates at iteration <code>i</code> and <code>i+1</code>. Default is <code>1e-5</code>. • <code>verbose</code> A Boolean. Default is FALSE. If TRUE monitors the iteration process. • <code>monitor_ev</code> A Boolean. Default is FALSE. If TRUE monitors the evaluation of the model over the $\log_{10}(\rho_s)$ values.
par_gridsearch	A list of parameters for the <code>grid_search</code> : <ul style="list-style-type: none"> • <code>plot_aic</code> A Boolean. Default is FALSE. If TRUE, plot the AIC values over the grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values. • <code>plot_bic</code> A Boolean. Default is FALSE. If TRUE, plot the BIC values over the grid of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ values.

- `return_aic` A Boolean. Default is TRUE. Return the AIC values.
- `return_bic` A Boolean. Default is TRUE. Return the BIC values.
- `col` The color palette to be used for the AIC/BIC plot. Default is `grDevices::gray.colors(n=10)`.
- `plot_contour` A Boolean. Default is TRUE. Adds white contour lines to the AIC/BIC plot.
- `mark_optimal` A Boolean. Default is TRUE. If the plot of the AIC or BIC values is returned, marks the optimal combination of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$ in the plot.
- `main_aic` The title of the AIC plot. Default is "AIC grid".
- `main_bic` The title of the BIC plot. Default is "BIC grid".

Details

The name P-GAM follows from the way this class of models is named in Eilers and Marx (2021) Practical Smoothing. The Joys of P-splines. In Carollo et al. (2025) this model is referred to as log-additive.

@references Carollo, A., Putter, H., Eilers, P. H. C., & Gampe, J. (2025). Analysis of Time-to-Event Data With Two Time Scales. An Application to Transitions out of Cohabitation. *Sociological Methods & Research*, 0(0). doi:10.1177/00491241251374193

Value

An object of class `haz2tsPGAM`, that is a list with the following elements:

- `optimal_model` A list with :
 - `alpha` The vector of estimated P-splines coefficients of length $c_u + c_s$.
 - `SE_alpha` The vector of estimated standard errors of the alpha coefficients, of dimension $c_u + c_s$.
 - `beta` (if covariates) The vector of length p of estimated covariates coefficients.
 - `se_beta` The vector of length p of estimated Standard Errors for the beta coefficients.
 - `eta0` (if covariates) The vector of values of the baseline linear predictor (log-hazard).
 - `eta` (if covariates) The vector of values of the baseline linear predictor (log-hazard) of length $n_u * n_s$.
 - `H` The hat-matrix.
 - `deviance` The deviance.
 - `ed` The (total) effective dimension of the model.
 - `aic` The value of the AIC.
 - `bic` The value of the BIC.
 - `Bbases` a list with the B-spline bases `Bu` and `Bs`
- `optimal_logrho` A vector with the optimal values of $\log_{10}(\rho_u)$ and $\log_{10}(\rho_s)$.
- `P_optimal` The optimal penalty matrix `P`.
- `EDu` The effective dimensions along the `u` axis.
- `EDs` The effective dimensions along the `s` axis.
- `AIC` (if `par_gridsearch$return_aic == TRUE`) The matrix of AIC values.
- `BIC` (if `par_gridsearch$return_bic == TRUE`) The matrix of BIC values.

Examples

```

# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5
)
# Fit a fake model - not optimal smoothing
fitpgam(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(seq(1, 1.5, .5), seq(1, 1.5, .5))
)

```

fitvcm

Fit a Varying Coefficient Model (VCM) over two time scales

Description

fitvcm() fits a varying coefficient model over two time scales with P-splines. The model is described in Carollo et al. (2025).

Usage

```

fitvcm(
  data2ts = NULL,
  Y = NULL,
  R = NULL,
  Z = NULL,
  bins = NULL,
  Bbases_spec = list(),
  pord = 2,
  kappa = 1e-10,
  lsmpar = c(0, 0),
  control_algorithm = list()
)

```

Arguments

data2ts	(optional) an object of class "data2ts" created by the function <code>prepare_data()</code> . Providing this input is the easiest way to use the function <code>fitvcm()</code> . However, the user can also provide the input data together with a list of bins, as explained by the following parameters' descriptions.
Y	A matrix (or 3d-array) of event counts of dimension n_u by n_s (or n_u by n_s by n).
R	A matrix (or 3d-array) of exposure times of dimension n_u by n_s (or n_u by n_s by n).
Z	(optional) A regression matrix of covariates values of dimensions n by p .
bins	a list with the specification for the bins. This is created by the function <code>prepare_data()</code> . If a list prepared externally from such function is provided, it should contain the following elements: * <code>bins_u</code> A vector of bins extremes for the time scale u . * <code>midu</code> A vector with the midpoints of the bins over u . * <code>nu</code> The number of bins over u . * <code>bins_s</code> A vector of bins extremes for the time scale s . * <code>mids</code> A vector with the midpoints of the bins over s . * <code>ns</code> The number of bins over s .
Bbases_spec	A list with the specification for the B-splines basis with the following elements: <ul style="list-style-type: none"> • <code>bdeg</code> The degree of the B-splines basis. Default is 3 (for cubic B-splines). • <code>nseg_s</code> The number of segments for the B-splines over s. Default is 10. • <code>min_s</code> (optional) The lower limit of the domain of Bs. Default is <code>min(bins_s)</code>. • <code>max_s</code> (optional) The upper limit of the domain of Bs. Default is <code>max(bins_s)</code>.
pord	The order of the penalty. Default is 2.
kappa	A ridge penalty.
lsmpar	The starting values for the two smoothing parameters. A vector of two elements, default is <code>c(0,0)</code> .
control_algorithm	A list with optional values for the parameters of the iterative processes: <ul style="list-style-type: none"> • <code>maxiter</code> The maximum number of iteration for the IWSL algorithm. Default is 20. • <code>conv_crit</code> The convergence criteria, expressed as difference between estimates at iteration i and $i+1$. Default is $1e-5$. • <code>verbose</code> A Boolean. Default is FALSE. If TRUE monitors the iteration process. • <code>monitor_ev</code> A Boolean. Default is FALSE. If TRUE monitors the evaluation of the model over the <code>log_10(rho_s)</code> values.

Value

An object of class `haz2tsVCM`, that is a list with the following elements:

- `optimal_model` A list with :
 - `Alpha` The matrix of estimated P-splines coefficients of dimension c_s by 2.
 - `Cov_alpha` The variance-covariance matrix of the Alpha coefficients, of dimension $c_u c_s$ by $c_u c_s$.

- Eta The matrix of values of the baseline linear predictor (log-hazard) of dimension n_u by n_s .
- H The hat-matrix.
- deviance The deviance.
- ed The effective dimension of the model.
- aic The value of the AIC.
- bic The value of the BIC.
- Bbases a list with the B-spline bases Bu and Bs
- optimal_logsmpar A vector with the optimal values of $\log_{10}(\theta)$ and $\log_{10}(\phi)$.
- P_optimal The optimal penalty matrix P.
- AIC (if par_gridsearch\$return_aic == TRUE) The matrix of AIC values.
- BIC (if par_gridsearch\$return_bic == TRUE) The matrix of BIC values.

@references Carollo, A., Putter, H., Eilers, P. H. C., & Gampe, J. (2025). Analysis of Time-to-Event Data With Two Time Scales. An Application to Transitions out of Cohabitation. Sociological Methods & Research, 0(0). doi:10.1177/00491241251374193

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5
)
# Fit a fake model - not optimal smoothing
fitvcm(fakedata2ts)
```

`get_hazard_1d`*Get estimated (log-)hazard values with 1 time scale*

Description

`get_hazard_1d()` takes as input the results of a model estimated by `fit1ts()` and it returns the estimated values of the smooth log-hazard and the smooth hazard together with their standard errors.

If the model includes covariates, then only the baseline (log-)hazard is returned. It is possible to provide values that define a new grid for evaluation of the estimated hazard. If not specified, the hazard is evaluated on the same grid used for the binning of the data, and therefore the estimation of the model. The function will check if the parameters for the new grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Usage

```
get_hazard_1d(fitted_model, plot_grid = NULL)
```

Arguments

`fitted_model` is an object of class "haz1ts", the output of the function `fit1ts()`.

`plot_grid` (optional) A named vector containing the parameters to build a new grid of intervals over `s` for plotting the estimated hazard on a finer grid. This must be of the form: `plot_grid = c(smin, smax, ds)`, where `smin`, `smax` are the minimum and maximum values desired for the intervals over `s`, and `ds` is the distance between intervals over `s`. If not specified, the plotting is done using the same B-splines basis as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Value

A list with the following elements:

- `new_plot_grid` A list of parameters that specify the new grid, of the form `list("ints", "smin", "smax", "ds")`
- `hazard` A vector containing the estimated hazard values.
- `loghazard` A vector containing the estimated log-hazard values.
- `log10hazard` A vector containing the estimated log10-hazard values.
- `SE_hazard` A vector containing the estimated SEs for the hazard.
- `SE_loghazard` A vector containing the estimated SEs for the log-hazard.
- `SE_log10hazard` A vector containing the estimated SEs for the log10-hazard.

Examples

```
## preparing data - no covariates
dt1ts <- prepare_data(
  data = reccolon2ts,
  s_in = "entrysr",
  s_out = "timesr",
  events = "status",
  ds = 180
)

## fitting the model with fit1ts() - default options

mod1 <- fit1ts(dt1ts)
# Obtain 1d hazard
get_hazard_1d(mod1)
# Change grid
get_hazard_1d(mod1,
  plot_grid = c(smin = 0, smax = 2730, ds = 30)
)
```

get_hazard_1d_LMM *Get estimated (log-)hazard values with 1 time scale*

Description

get_hazard_1d_LMM() takes as input the results of a model estimated by fit1ts() and it returns the estimated values of the smooth log-hazard and the smooth hazard together with their standard errors. This function is specific for objects of class "haz1tsLMM".

If the model includes covariates, then only the baseline (log-)hazard is returned. It is possible to provide values that define a new grid for evaluation of the estimated hazard. If not specified, the hazard is evaluated on the same grid used for the binning of the data, and therefore the estimation of the model. The function will check if the parameters for the new grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Usage

```
get_hazard_1d_LMM(fitted_model, plot_grid = NULL)
```

Arguments

fitted_model is an object of class "haz1tsLMM", the output of the function fit1ts().

plot_grid (optional) A named vector containing the parameters to build a new grid of intervals over *s* for plotting the estimated hazard on a finer grid. This must be of the form: plot_grid = c(smin, smax, ds), where *smin*, *smax* are the minimum and maximum values desired for the intervals over *s*, and *ds* is the

distance between intervals over s . If not specified, the plotting is done using the same B-splines basis as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Value

A list with the following elements:

- `new_plot_grid` A list of parameters that specify the new grid, of the form `list("ints", "smin", "smax", "ds")`
- `hazard` A vector containing the estimated hazard values.
- `loghazard` A vector containing the estimated log-hazard values.
- `log10hazard` A vector containing the estimated log10-hazard values.
- `SE_hazard` A vector containing the estimated SEs for the hazard.
- `SE_loghazard` A vector containing the estimated SEs for the log-hazard.
- `SE_log10hazard` A vector containing the estimated SEs for the log10-hazard.

Examples

```
## preparing data - no covariates
dt1ts <- prepare_data(
  data = reccolon2ts,
  s_in = "entries",
  s_out = "timesr",
  events = "status",
  ds = 180
)

## fitting the model with fit1ts()

mod1 <- fit1ts(dt1ts,
  optim_method = "LMMsolver"
)
# Obtain 1d hazard
get_hazard_1d_LMM(mod1)
# Change grid
get_hazard_1d_LMM(mod1,
  plot_grid = c(smin = 0, smax = 2730, ds = 30)
)
```

 get_hazard_2d

Get estimated (log-)hazard surface with 2 time scales

Description

get_hazard_2d() takes as input the results of a model estimated by fit2ts(), fitpgam() or fitvcm() and it returns the estimated smooth log-hazard and the smooth hazard together with their standard errors.

It is possible to provide values that define a new grid for evaluation of the estimated hazard. If not specified, the hazard is evaluated on the same grid used for the binning of the data, and therefore the estimation of the model. The function will check if the parameters for the new grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Usage

```
get_hazard_2d(
  fitted_model,
  plot_grid = NULL,
  where_slices = NULL,
  direction = c(NULL, "u", "s"),
  tmax = NULL,
  midpoints = FALSE
)
```

Arguments

- | | |
|--------------|---|
| fitted_model | is an object of class "haz2ts", "haz2tsPGAM", or "haz2tsVCM", the output of the function fit2ts(), fitpgam() or fitvcm(), respectively. |
| plot_grid | (optional) A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: plot_grid = list(c(umin, umax, du), c(smin, smax, ds)), where umin, umax and smin, smax are the minimum and maximum values desired for the intervals over u and s respectively, and du, ds are distances between intervals over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned. |
| where_slices | A vector of values for the cutting points of the desired slices of the surface. If which_plot == "slices", please provide this argument. Please also provide this argument in case which_plot = "survival or which_plot = "cumhaz and surv_slices = TRUE or cumhaz_slices = TRUE, respectively. |

direction	If which_plot == "slices", indicates the direction for cutting the surface. If u, then the surface will be cut at the selected values of u (indicated by where_slices), hence obtaining one-dimensional curves over s. If s, then the surface will be cut at the selected values of s (indicated by where_slices), hence obtaining one-dimensional curves over u.
tmax	The maximum value of t that should be plotted.
midpoints	A Boolean. Default is FALSE. If TRUE, the estimated quantities are evaluated at the midpoints of the rectangles (or parallelograms) of the grids, rather than at each grid-point.

Value

A list with the following elements:

- new_plot_grid A list of parameters that specify the new grid, of the form list("intu", "umin", "umax", "du", "ints", "smin", "smax", "ds")
- nBu The B-spline basis for u, evaluated over the new grid.
- nBs The B-spline basis for s, evaluated over the new grid.
- hazard A matrix containing the estimated hazard values.
- loghazard A matrix containing the estimated log-hazard values.
- log10hazard A matrix containing the estimated log10-hazard values.
- SE_hazard A matrix containing the estimated SEs for the hazard.
- SE_loghazard A matrix containing the estimated SEs for the log-hazard.
- SE_log10haz A matrix containing the estimated SEs for the log10-hazard.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5
)
# Fit a fake model - not optimal smoothing
```

```
fakemod <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)
# Obtain 2d hazard
get_hazard_2d(fakemod)

get_hazard_2d(fakemod,
  plot_grid = list(
    c(umin = 3, umax = 8.5, du = .1),
    c(smin = 0, smax = 7.1, ds = .1)
  )
)
```

get_hazard_2d_LMM *Get estimated (log-)hazard surface with 2 time scales*

Description

get_hazard_2d_LMM() takes as input an object of class 'haz2tsLMM' and it returns the estimated smooth log-hazard, the log10-hazard and the hazard surface together with their standard errors.

It is possible to provide values that define a new grid for evaluation of the estimated hazard. If not specified, the hazard is evaluated on the same grid used for the binning of the data, and therefore the estimation of the model. The function will check if the parameters for the new grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

Usage

```
get_hazard_2d_LMM(
  fitted_model,
  plot_grid = NULL,
  where_slices = NULL,
  direction = c("u", "s", NULL),
  tmax = NULL
)
```

Arguments

fitted_model is an object of class 'haz2tsLMM' the output of the function fit2ts().

plot_grid A list containing the parameters to build a new finer grid of intervals over *u* and *s* for plotting. This must be of the form: plot_grid = list(c(umin, umax, du), c(smin, smax, ds)), where umin, umax and smin, smax are the minimum and maximum values desired for the intervals over *u* and *s* respectively, and du, ds are distances between intervals over *u* and *s* respectively. Specifying

a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned. While for objects of class 'haz2ts' this is an optional input, we strongly recommend to provide the plotting grid for object of class 'haz2tsLMM', given that evaluation of the B-splines works a bit differently in LMMsolver.

where_slices	A vector of values for the cutting points of the desired slices of the surface. If which_plot == "slices", please provide this argument. Please also provide this argument in case which_plot = "survival or which_plot = "cumhaz and surv_slices = TRUE or cumhaz_slices = TRUE, respectively.
direction	If which_plot == "slices", indicates the direction for cutting the surface. If u, then the surface will be cut at the selected values of u (indicated by where_slices), hence obtaining one-dimensional curves over s. If s, then the surface will be cut at the selected values of s (indicated by where_slices), hence obtaining one-dimensional curves over u.
tmax	The maximum value of t that should be plotted.

Value

A list with the following elements:

- new_plot_grid A list of parameters that specify the new grid, of the form list("intu", "umin", "umax", "du", "ints", "smin", "smax", "ds")
- hazard A matrix containing the estimated hazard values.
- loghazard A matrix containing the estimated log-hazard values.
- log10hazard A matrix containing the estimated log10-hazard values.
- SE_hazard A matrix containing the estimated SEs for the hazard.
- SE_loghazard A matrix containing the estimated SEs for the log-hazard.
- SE_log10haz A matrix containing the estimated SEs for the log10-hazard.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'
```

```
fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(data = fakedata,
                           u = "u",
                           s_out = "s",
                           ev = "ev",
                           ds = .5)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "LMMsolver"
                 )

# Get hazard
get_hazard_2d_LMM(fakemod)

# Use a finer grid of points
get_hazard_2d_LMM(fakemod,
                 plot_grid = list(c(umin = 3, umax = 8.5, du = .1),
                                 c(smin = 0, smax = 7.1, ds = .1)))
```

get_hr

Get the Hazard Ratios with their Standard Errors

Description

get_hr() takes as input the results of a model with covariates estimated by fit2ts() or fit1ts() and returns the estimated hazard ratios together with their standard errors.

Usage

```
get_hr(fitted_model)
```

Arguments

fitted_model A list returned by the function fit2ts() or fit1ts().

Value

A list with the following elements:

- HR A vector of hazard ratios (calculated as $\exp(\hat{\beta})$).
- SE_HR A vector of Standard Errors for the hazard ratios calculated via the delta method.
- beta A vector of the estimated $\hat{\beta}$ coefficients.
- SE_beta A vector of the Standard Errors for the beta coefficients.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)
x1 <- c(0, 0, 0, 1, 0, 1, 1, 1, 1, 0, 0, 0, 1, 0, 1, 1, 0, 0, 1, 0)

fakedata <- as.data.frame(cbind(id, u, s, ev, x1))
fakedata2ts <- prepare_data(data = fakedata,
                           u = "u",
                           s_out = "s",
                           ev = "ev",
                           ds = .5,
                           individual = TRUE,
                           covs = "x1")
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "grid_search",
                 lrho = list(seq(1, 1.5, .5),
                             seq(1, 1.5, .5)))

get_hr(fakemod)
```

haz2tsLMM_summary

Summary function for object of class 'haz2tsLMM'

Description

Summary function for object of class 'haz2tsLMM'

Usage

```
haz2tsLMM_summary(x, ...)
```

Arguments

x an object of class 'haz2tsLMM' returned by the function `fit2ts()`
... further arguments

Value

a printed summary of the fitted model, including optimal smoothing paramters, the effective dimension ED and the AIC/BIC. For model with covariates, a regression table is also returned.

Examples

```

# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
  optim_method = "LMMsolver"
)
summary(fakemod)

```

haz2tsPGAM_summary *Summary function for object of class 'haz2tsPGAM'*

Description

Summary function for object of class 'haz2tsPGAM'

Usage

```
haz2tsPGAM_summary(x, ...)
```

Arguments

x an object of class 'haz2tsPGAM' returned by the function `fitpgam()`
 ... further arguments

Value

a printed summary of the fitted model

Examples

```

# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(data = fakedata,
                           u = "u",
                           s_out = "s",
                           ev = "ev",
                           ds = .5)

# Fit a fake model - not optimal smoothing
fakemod <- fitpgam(fakedata2ts,
                  optim_method = "grid_search",
                  lrho = list(seq(1, 1.5, .5),
                              seq(1, 1.5, .5)))

summary(fakemod)

```

haz2tsVCM_summary *Summary function for object of class 'haz2tsVCM'*

Description

Summary function for object of class 'haz2tsVCM'

Usage

```
haz2tsVCM_summary(x, ...)
```

Arguments

x an object of class 'haz2tsVCM' returned by the function `fitvcm()`
 ... further arguments

Value

a printed summary of the fitted model

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(data = fakedata,
                           u = "u",
                           s_out = "s",
                           ev = "ev",
                           ds = .5)

# Fit a fake model - not optimal smoothing
fakemod <- fitvcm(fakedata2ts)
summary(fakemod)
```

haz2ts_summary

Summary function for object of class 'haz2ts'

Description

Summary function for object of class 'haz2ts'

Usage

```
haz2ts_summary(x, ...)
```

Arguments

x an object of class 'haz2ts' returned by the function [fit2ts\(\)](#)
... further arguments

Value

a printed summary of the fitted model

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'
```

```
fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(data = fakedata,
                           u = "u",
                           s_out = "s",
                           ev = "ev",
                           ds = .5)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "grid_search",
                 lrho = list(seq(1, 1.5, .5),
                             seq(1, 1.5, .5)))
summary(fakemod)
```

imageplot_2ts

Image Plot of 2ts hazard

Description

imageplot_2ts() plots an image of the two time scales hazard (or survival or cumulative hazard) with contour lines. This is the default call implemented in plot.haz2ts and other functions.

Usage

```
imageplot_2ts(x, y, z, plot_options = list(), ...)
```

Arguments

- | | |
|--------------|---|
| x | The coordinates for the x-axis. This is a vector of intervals over the u axis (default), a matrix with the corner points of the parallelograms over the t time scale, or a vector of intervals for the t time scale. |
| y | The coordinates for the y-axis. This is a vector of intervals over the s time scale (default), or a matrix with the corner points of the parallelograms over the s time scale. |
| z | The values of the surface to plot, organized in a matrix with dimensions compatible with those of x and y. The default is to plot the hazard. |
| plot_options | A list of options for the plot: <ul style="list-style-type: none"> • loghazard A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard surface, if TRUE the function returns a plot of the log-hazard surface. • log10hazard A Boolean. Default is FALSE. If TRUE, then a log10-hazard surface is plotted. • original A Boolean. Default is TRUE. Plot the (log-)hazard in the (t,s)-plane. If FALSE, the (log-)hazard will be plotted in the (u,s)-plane. |

- `rectangular_grid` A Boolean. Default is FALSE. If TRUE, a rectangular grid is used for plotting also in the (t,s)-plane as opposed to the grid of parallelograms used as default in the (t,s)-plane.
- `col_palette` A function defining the color palette. The default palette is `rev(viridis::plasma())`.
- `n_shades` The number of color shades to plot, default is 50.
- `breaks` The vector of breaks for the color legend. If `n_shades` is provided, this should be of length `n_shades + 1`. Otherwise, `n_shades` will be recalculated accordingly.
- `show_legend` A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid!
- `tmax` The maximum value of `t` that should be plotted.
- `main` The title of the plot.
- `xlab` The label of the first time axis (plotted on the x axis).
- `ylab` The label of the second time axis (plotted on the y axis).
- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of the time scale on the y axis.
- `cex_main` The magnification to be used for the main title, default is 1.2.
- `cex_lab` The magnification to be used for the axis labels, default is 1.
- `contour_lines` A Boolean. Default is FALSE. If TRUE white contour lines are added to the surfaces.
- `contour_col` The color for the contour lines. Default is white.
- `contour_cex` The magnification to be used for the contour lines. Default is .8.
- `contour_nlev` The number of contour levels. Default is 10.

... Further arguments to `image.plot` or `image`

Value

An image plot of an estimated surface.

imageplot_SE

Image Plot of Standard Errors for the 2ts hazard

Description

`imageplot_SE()` plots an image of the SEs of the two time scales hazard with contour lines.

Usage

```
imageplot_SE(x, y, z, plot_options = list(), ...)
```

Arguments

- x** The coordinates for the x-axis. This is a vector of intervals over the *u* axis (default), a matrix with the corner points of the parallelograms over the *t* time scale, or a vector of intervals for the *t* time scale.
- y** The coordinates for the y-axis. This is a vector of intervals over the *s* time scale (default), or a matrix with the corner points of the parallelograms over the *s* time scale.
- z** The values of the surface to plot, organized in a matrix with dimensions compatible with those of *x* and *y*. These can be the SEs for the hazard, the SEs for the log-hazard or the SEs for the log10-hazard.
- plot_options** A list of options for the plot:
- **loghazard** A Boolean. Default is FALSE. If FALSE the function returns a plot of the standard errors of the hazard surface, if TRUE the function returns a plot of the standard errors of the log-hazard surface.
 - **log10hazard** A Boolean. Default is FALSE. If TRUE, the function returns a plot of the standard errors of the log10-hazard surface
 - **original** A Boolean. Default is TRUE. Plot the (log-)hazard in the (t,s)-plane. If FALSE, the (log-)hazard will be plotted in the (u,s)-plane.
 - **rectangular_grid** A Boolean. Default is FALSE. If TRUE, a rectangular grid is used for plotting also in the (t,s)-plane as opposed to the grid of parallelograms used as default in the (t,s)-plane.
 - **col_palette** A function defining the color palette. The default palette is `rev(colorspace::sequential_hcl(n = 50, "Red-Purple"))`.
 - **n_shades** The number of color shades to plot, default is 50.
 - **breaks** The vector of breaks for the color legend. If **n_shades** is provided, this should be of length **n_shades** + 1. Otherwise, **n_shades** will be recalculated accordingly.
 - **show_legend** A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid!
 - **tmax** The maximum value of *t* that should be plotted.
 - **main** The title of the plot.
 - **xlab** The label of the first time axis (plotted on the *x* axis).
 - **ylab** The label of the second time axis (plotted on the *y* axis).
 - **xlim** A vector with two elements defining the limits of the time scale on the *x* axis.
 - **ylim** A vector with two elements defining the limits of the time scale on the *y* axis.
 - **cex_main** The magnification to be used for the main title, default is 1.2.
 - **cex_lab** The magnification to be used for the axis labels, default is 1.
 - **contour_lines** A Boolean. Default is FALSE. If TRUE white contour lines are added to the surfaces.
 - **contour_col** The color for the contour lines. Default is white.
 - **contour_cex** The magnification to be used for the contour lines. Default is .8.

- `contour_nlev` The number of contour levels. Default is 10.

... Further arguments to `image.plot` or `image`

Value

An image plot of the SEs for the (log-) hazard surface.

<code>make_bins</code>	<i>Construct bins over one or more time axes</i>
------------------------	--

Description

`make_bins()` constructs the bins over the time axes and saves the extremes of the bins in a vector.

Usage

```
make_bins(
  t_in = NULL,
  t_out = NULL,
  u = NULL,
  s_in = NULL,
  s_out,
  min_t = NULL,
  max_t = NULL,
  min_u = NULL,
  max_u = NULL,
  min_s = NULL,
  max_s = NULL,
  dt = NULL,
  du = NULL,
  ds
)
```

Arguments

<code>t_in</code>	(optional) A vector of entry times on the time scale <code>t</code> .
<code>t_out</code>	(optional) A vector of exit times on the time scale <code>t</code> .
<code>u</code>	(optional) A vector of fixed-times at entry in the process.
<code>s_in</code>	(optional) A vector of entry times on the time scale <code>s</code> .
<code>s_out</code>	A vector of exit times on the time scale <code>s</code> .
<code>min_t</code>	(optional) A minimum value for the bins over <code>t</code> . If <code>NULL</code> , the minimum of <code>t_in</code> will be used.
<code>max_t</code>	(optional) A maximum value for the bins over <code>t</code> . If <code>NULL</code> , the maximum of <code>t_out</code> will be used.

min_u	(optional) A minimum value for the bins over u. If NULL, the minimum of u will be used.
max_u	(optional) A maximum value for the bins over u. If NULL, the maximum of u will be used.
min_s	(optional) A minimum value for the bins over s. If NULL, the minimum of s_in will be used.
max_s	(optional) A maximum value for the bins over s. If NULL, the maximum of s_out will be used.
dt	(optional) A scalar giving the length of the intervals on the t time scale.
du	(optional) A scalar giving the length of the intervals on the u axis.
ds	A scalar giving the length of the intervals on the s time scale.

Details

It allows construction of bins over the time scales t and s and/or over the fixed-time axis u. The time scale s is always required. See also [prepare_data\(\)](#) to conveniently prepare individual data for the analysis with one, or two time scales.

A few words about constructing the grid of bins. There is no 'golden rule' or optimal strategy for setting the number of bins over each time axis, or deciding on the bins' width. It very much depends on the data structure, however, we try to give some directions here. First, in most cases, more bins is better than less bins. A good number is about 30 bins. However, if data are scarce, the user might want to find a compromise between having a larger number of bins, and having many bins empty. Second, the chosen width of the bins (that is du and ds) does depend on the time unit over which the time scales are measured. For example, if the time is recorded in days, as in the example below, and several years of follow-up are available, the user can split the data in bins of width 30 (corresponding to about one month), 60 (about two months), 90 (about three months), etc. If the time scale is measured in years, then appropriate width could be 0.25 (corresponding to a quarter of a year), or 0.5 (that is half year). However, in some cases, time might be measure in completed years, as is often the case for age. In this scenario, an appropriate bin width is 1.

Finally, it is always a good idea to plot your data first, and explore the range of values over which the time scale(s) are recorded. This will give insight about reasonable values for the arguments min_s, min_u, max_s and max_u (that in any case are optional).

Value

A list with the following elements:

- bins_t if t_out is provided, this is a vector of bins extremes for the time scale t
- midt if t_out is provided, this is a vector with the midpoints of the bins over t
- nt if t_out is provided, this is the number of bins over t
- bins_u if u is provided, this is a vector of bins extremes for u axis
- midu if u is provided, this is a vector with the midpoints of the bins over u
- nu if u is provided, this is the number of bins over u
- bins_s is a vector of bins extremes for the time scale s
- mids is a vector with the midpoints of the bins over s
- ns is the number of bins over s

Examples

```
# Make bins for colon cancer data by time at randomization and time since recurrence
bins <- make_bins(u = reccolon2ts$timer, s_out = reccolon2ts$timesr,
                 du = 30, ds = 30)

# Make bins for colon cancer data only over time since recurrence
bins <- make_bins(s_out = reccolon2ts$timesr, ds = 60)
```

make_grid

*Make a grid of points to evaluate B-splines***Description**

Make a grid of points to evaluate B-splines

Usage

```
make_grid(
  plot_grid,
  model_specifications,
  class_fitmodel,
  where_slices = NULL,
  direction = c(NULL, "u", "s"),
  tmax = NULL,
  midpoints = FALSE
)
```

Arguments

- plot_grid** (optional) A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: `plot_grid = list(c(umin, umax, du), c(smin, smax, ds))`, where `umin`, `umax` and `smin`, `smax` are the minimum and maximum values desired for the intervals over u and s respectively, and `du`, `ds` are distances between intervals over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.
- model_specifications** A list with 'umin', 'umax', 'smin', and 'smax' used for fitting.
- class_fitmodel** The class of the fitted model. Can be 'haz2ts', 'haz2tsPGAM', or 'haz2tsVCM'.
- where_slices** A vector of values for the cutting points of the desired slices of the surface. If `which_plot == "slices"`, please provide this argument. Please also provide this argument in case `which_plot = "survival"` or `which_plot = "cumhaz"` and `surv_slices = TRUE` or `cumhaz_slices = TRUE`, respectively.

direction	If which_plot == "slices", indicates the direction for cutting the surface. If u, then the surface will be cut at the selected values of u (indicated by where_slices), hence obtaining one-dimensional curves over s. If s, then the surface will be cut at the selected values of s (indicated by where_slices), hence obtaining one-dimensional curves over u.
tmax	The maximum value of t that should be plotted.
midpoints	A Boolean. Default is FALSE. If TRUE, the estimated quantities are evaluated at the midpoints of the rectangles (or parallelograms) of the grids, rather than at each grid-point.

Value

A list of specification for the grid

Examples

```
make_grid(plot_grid = list(
  c(umin = 3, umax = 8.5, du = .1),
  c(smin = 0, smax = 7.1, ds = .1)
),
model_specification = list(umin = 2, umax = 8.5, smin = 0, smax = 7.5),
class_fitmodel = "haz2ts")
```

plot.haz1ts

Plot method for a haz1ts object.

Description

plot.haz1ts() is a plot method for objects of class haz1ts.

Usage

```
## S3 method for class 'haz1ts'
plot(
  x,
  which_plot = c("hazard", "covariates"),
  plot_grid = NULL,
  plot_options = list(),
  ...
)
```

Arguments

x	The output of the function fit1ts.
which_plot	The type of plot required. Can be one of "hazard" (default) or "covariates".

plot_grid	(optional) A named vector containing the parameters to build a new grid of intervals over s for plotting the estimated hazard on a finer grid. This must be of the form: <code>plot_grid = c(smin, smax, ds)</code> , where <code>smin</code> , <code>smax</code> are the minimum and maximum values desired for the intervals over s , and <code>ds</code> is the distance between intervals over s . If not specified, the plotting is done using the same B-splines basis as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.
plot_options	<p>A list with all possible options for any of the plots:</p> <ul style="list-style-type: none"> • <code>loghazard</code> A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard curve, if TRUE the function returns a plot of the log-hazard curve. • <code>log10hazard</code> A Boolean. Default is FALSE. If TRUE it returns a plot of the log10-hazard curve. • <code>col</code> The color of the curve plotted. Default is "black". • <code>add_CI</code> A Boolean. If TRUE (default) the confidence bands will be added. • <code>col_CI</code> The color for the confidence bands. The default is the same color of the curve, with a 50% transparency level. • <code>main</code> The title of the plot. • <code>xlab</code> The label of the time axis (plotted on the x axis). • <code>ylab</code> The label of the y-axis (hazard, log-hazard or log10-hazard). • <code>xlim</code> A vector with two elements defining the limits of the time scale on the x axis. • <code>ylim</code> A vector with two elements defining the limits of function plotted on the y axis (hazard, log-hazard or log10-hazard). • <code>xmin</code> The minimum value on the x-axis. • <code>ymin</code> The minimum value on the y-axis. • <code>cex_main</code> The magnification to be used for the main title, default is 1.2 . • <code>cex_lab</code> The magnification to be used for the axis labels, default is 1 . • <code>HR</code> A Boolean. If TRUE the HRs with their CIs will be plotted. Default is FALSE (plot the beta with their CIs). • <code>symmetric_CI</code> A Boolean. Default is TRUE. If a plot of the HRs is required (<code>HR == TRUE</code>), then plot symmetrical Confidence Intervals, based on the SEs for the HRs calculated by delta method. If FALSE, then CIs are obtained by exponentiating the CIs for the betas. • <code>confidence</code> The level of confidence for the CIs. Default is .95 (alpha = 0.05). • <code>col_beta</code> The color for the plot of the covariates' effects. • <code>pch</code> The symbol for plotting the point estimates. • <code>lwd</code> The line width. • <code>lty</code> The line type. <p>... Further arguments to plot.</p>

Value

A plot of the type required.

Examples

```
## preparing data - no covariates
dt1ts <- prepare_data(
  data = reccolon2ts,
  s_in = "entrysr",
  s_out = "timesr",
  events = "status",
  ds = 180
)

## fitting the model with fit1ts() - default options

mod1 <- fit1ts(dt1ts)

plot(mod1)
```

plot.haz1tsLMM *Plot method for a haz1ts object.*

Description

plot.haz1tsLMM() is a plot method for objects of class haz1tsLMM.

Usage

```
## S3 method for class 'haz1tsLMM'
plot(
  x,
  which_plot = c("hazard", "covariates"),
  plot_grid = NULL,
  plot_options = list(),
  ...
)
```

Arguments

x	The output of the function fit1ts.
which_plot	The type of plot required. Can be one of "hazard" (default) or "covariates".
plot_grid	(optional) A named vector containing the parameters to build a new grid of intervals over s for plotting the estimated hazard on a finer grid. This must be of the form: plot_grid = c(smin, smax, ds), where smin, smax are the minimum and maximum values desired for the intervals over s, and ds is the

distance between intervals over s . If not specified, the plotting is done using the same B-splines basis as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

`plot_options` A list with all possible options for any of the plots:

- `loghazard` A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard curve, if TRUE the function returns a plot of the log-hazard curve.
- `log10hazard` A Boolean. Default is FALSE. If TRUE it returns a plot of the log10-hazard curve.
- `col` The color of the curve plotted. Default is "black".
- `add_CI` A Boolean. If TRUE (default) the confidence bands will be added.
- `col_CI` The color for the confidence bands. The default is the same color of the curve, with a 50% transparency level.
- `main` The title of the plot.
- `xlab` The label of the time axis (plotted on the x axis).
- `ylab` The label of the y-axis (hazard, log-hazard or log10-hazard).
- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of function plotted on the y axis (hazard, log-hazard or log10-hazard).
- `xmin` The minimum value on the x-axis.
- `ymin` The minimum value on the y-axis.
- `cex_main` The magnification to be used for the main title, default is 1.2 .
- `cex_lab` The magnification to be used for the axis labels, default is 1 .
- `HR` A Boolean. If TRUE the HRs with their CIs will be plotted. Default is FALSE (plot the beta with their CIs).
- `symmetric_CI` A Boolean. Default is TRUE. If a plot of the HRs is required (`HR == TRUE`), then plot symmetrical Confidence Intervals, based on the SEs for the HRs calculated by delta method. If FALSE, then CIs are obtained by exponentiating the CIs for the betas.
- `confidence` The level of confidence for the CIs. Default is .95 (alpha = 0.05).
- `col_beta` The color for the plot of the covariates' effects.
- `pch` The symbol for plotting the point estimates.
- `lwd` The line width.
- `lty` The line type.

... Further arguments to plot.

Details

The function `obtainSmoothTrend` from the R-package `LMMsolver` is used here. We refer the interested readers to <https://biometris.github.io/LMMsolver/> for more details on `LMMsolver` and its usage.

Value

A plot of the type required.

Examples

```
## preparing data - no covariates
dt1ts <- prepare_data(data = reccolon2ts,
                      s_in = "entrysr",
                      s_out = "timesr",
                      events = "status",
                      ds = 180)

## fitting the model with fit1ts() - default options

mod1 <- fit1ts(dt1ts,
              optim_method = "LMMsolver")
plot(mod1)
```

plot.haz2ts

Plot method for a haz2ts object.

Description

plot.haz2ts() is the plot method for objects of class haz2ts. It produces several kinds of plots of the fitted model with two time scales (see [fit2ts\(\)](#)), either in the original (t,s) plane, while respecting the constraint imposed by the relation of the two time scales, or in the transformed (u,s) plane.

Usage

```
## S3 method for class 'haz2ts'
plot(
  x,
  plot_grid = NULL,
  which_plot = c("hazard", "covariates", "SE", "slices", "survival", "cumhaz"),
  where_slices = NULL,
  direction = c(NULL, "u", "s"),
  plot_options = list(),
  ...
)
```

Arguments

x The output of the function fit2ts. This is an object of class "haz2ts".

plot_grid (optional) A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: plot_grid = list(c(umin, umax, du), c(smin, smax, ds)), where umin, umax and smin, smax are the

minimum and maximum values desired for the intervals over u and s respectively, and du , ds are distances between intervals over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

which_plot	The type of plot required. Can be one of "hazard" (default), "covariates", "SE", "slices", "survival" or "cumhaz" (see details section).
where_slices	A vector of values for the cutting points of the desired slices of the surface. If which_plot == "slices", please provide this argument. Please also provide this argument in case which_plot = "survival" or which_plot = "cumhaz" and surv_slices = TRUE or cumhaz_slices = TRUE, respectively.
direction	If which_plot == "slices", indicates the direction for cutting the surface. If u , then the surface will be cut at the selected values of u (indicated by where_slices), hence obtaining one-dimensional curves over s . If s , then the surface will be cut at the selected values of s (indicated by where_slices), hence obtaining one-dimensional curves over u .
plot_options	A list with all possible options for any of the plots: <ul style="list-style-type: none"> • loghazard A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard surface, if TRUE the function returns a plot of the log-hazard surface. • log10hazard A Boolean. Default is FALSE. If TRUE, then a log₁₀ hazard surface is plotted. • cut_extrapolated A Boolean. Default is TRUE. Cuts away the extrapolated area of the (log-)hazard surface before plotting. • rectangular_grid A Boolean. Default is FALSE. If TRUE, a rectangular grid is used for plotting also in the (t,s)-plane as opposed to the grid of parallelograms used as default in the (t,s)-plane. • original A Boolean. Default is TRUE. Plot the (log-)hazard (and/or the SEs) in the (t,s)-plane. If FALSE, the (log-)hazard (and/or the SEs) will be plotted in the (u,s)-plane. • tmax The maximum value of t that should be plotted. • surv_slices A Boolean. Default is FALSE. If TRUE and which_plot == "survival", plot survival curves over the time s for selected values of u, that are cross-sections of the 2D survival surface. • cumhaz_slices A Boolean. Default is FALSE. If TRUE and which_plot == "cumhaz", plot cumulative hazards curves over the time s for selected values of u, that are cross-sections of the 2D cumulative hazard surface. • midpoints A Boolean. Default is FALSE. If TRUE, the estimated quantities (hazard, survival, etc.) will be evaluated in the mid-points of the bins rather than at the extremes. Set to TRUE if plotting estimated number of events. • col_palette A function defining the color palette. The default palette is viridis::rev(plasma()). Specifying the color palette as a function allows for greater flexibility than passing the palette as a vector.

- `n_shades` The number of color shades to plot, default is 50.
- `breaks` The vector of breaks for the color legend. If `n_shades` is provided, this should be of length `n_shades + 1`.
- `show_legend` A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid (i.e. transformed (u,s) plane)
- `main` The title of the plot.
- `xlab` The label of the first time axis (plotted on the x axis).
- `ylab` The label of the second time axis (plotted on the y axis).
- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of the time scale on the y axis.
- `contour_lines` A Boolean. Default is FALSE. If TRUE white contour lines are added to the surfaces.
- `contour_col` The color for the contour lines. Default is white.
- `contour_cex` The magnification to be used for the contour lines. Default is .8.
- `contour_nlev` The number of contour levels desired. Default is 10.
- `cex_main` The magnification to be used for the main title, default is 1.2 .
- `cex_lab` The magnification to be used for the axis labels, default is 1 .
- `HR` A Boolean. If TRUE the HRs with their CIs will be plotted. Default is FALSE (plot the beta with their CIs).
- `symmetric_CI` A Boolean. Default is TRUE. If a plot of the HRs is required (`HR == TRUE`), then plot symmetrical Confidence Intervals, based on the SEs for the HRs calculated by delta method. If FALSE, then CIs are obtained by exponentiating the CIs for the betas.
- `confidence` The level of confidence for the CIs. Default is .95 (alpha = 0.05).
- `col_beta` The color for the plot of the covariates' effects.
- `pch` The symbol for plotting the point estimates.
- `lwd` The line width.

... Further arguments to `image.plot` or `image`

Details

The vignette "visualization" presents and discusses all the different plotting options for the fitted model over two time scales. In most of the cases, the user will want to visualize the hazard surface over the two time scales. This can be plotted on the hazard scale, the log-hazard scale or the log10-hazard scale, by switching to TRUE the corresponding argument in `plot_options`. The survival and cumulative hazard functions can be plotted as two-dimensional surfaces over `u` and `s` or `t` and `s`. However, it is also very informative to plot them as one-dimensional curves over `s` (cross-sections or slices). This is done by selecting `which_plot = "survival"` and `surv_slices = TRUE` in `plot_options`. Additionally, a vector of values for the cutting points over the `u`-axis should be passed to the argument `where_slices`, together with setting `direction = u`. Similar plot is obtained for the cumulative hazard by selecting `which_plot = "cumhaz"`, `cumhaz_slices = TRUE`,

see examples section. Please, notice that for the survival function and the cumulative hazard, only cross-sections of the surface for selected values of u (over the s time) can be plotted.

Value

A plot of the fitted model.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5
)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)

# plot the hazard surface
plot(fakemod)

# plot the survival function as one-dimension curves over `s`
plot(fakemod,
  which_plot = "survival",
  direction = "u",
  where_slices = c(4, 6, 8),
  plot_options = list(
    surv_slices = TRUE
  )
)

# Plot cross-sections of the hazard over `s` for selected values of `u`
plot(fakemod,
```

```

which_plot = "slices",
where_slices = c(4, 6, 8),
direction = "u",
plot_options = list(
  main = "Cross-sections of the hazard",
  xlab = "Time",
  ylab = "Hazard"
)
)

```

plot.haz2tsLMM

Plot method for a haz2tsLMM object.

Description

plot.haz2tsLMM() is the plot method for objects of class haz2tsLMM. It produces plots of the fitted model with two time scales (see [fit2ts\(\)](#)), fitted via LMMsolver. The two-dimensional plots are limited to the transformed plane, that is only plots over u and s axes are produced.

Usage

```

## S3 method for class 'haz2tsLMM'
plot(
  x,
  plot_grid = NULL,
  which_plot = c("hazard", "covariates", "SE", "slices", "survival", "cumhaz"),
  where_slices = NULL,
  direction = c(NULL, "u", "s"),
  plot_options = list(),
  ...
)

```

Arguments

x The output of the function fit2ts. This is an object of class "haz2tsLMM".

plot_grid A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: plot_grid = list(c(umin, umax, du), c(smin, smax, ds)), where umin, umax and smin, smax are the minimum and maximum values desired for the intervals over u and s respectively, and du, ds are distances between intervals over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned. While for objects of class 'haz2ts' this is an optional

input, we strongly recommend to provide the plotting grid for object of class 'haz2tsLMM', given that evaluation of the B-splines works a bit differently in LMMsolver.

which_plot	The type of plot required. Can be one of "hazard" (default), "covariates", "SE", "slices", "survival" or "cumhaz" (see details section).
where_slices	A vector of values for the cutting points of the desired slices of the surface. If which_plot == "slices", please provide this argument. Please also provide this argument in case which_plot = "survival" or which_plot = "cumhaz" and surv_slices = TRUE or cumhaz_slices = TRUE, respectively.
direction	If which_plot == "slices", indicates the direction for cutting the surface. If u, then the surface will be cut at the selected values of u (indicated by where_slices), hence obtaining one-dimensional curves over s. If s, then the surface will be cut at the selected values of s (indicated by where_slices), hence obtaining one-dimensional curves over u.
plot_options	<p>A list with all possible options for any of the plots:</p> <ul style="list-style-type: none"> • loghazard A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard surface, if TRUE the function returns a plot of the log-hazard surface. • log10hazard A Boolean. Default is FALSE. If TRUE, then a log10 hazard surface is plotted. • cut_extrapolated A Boolean. Default is TRUE. Cuts away the extrapolated area of the (log-)hazard surface before plotting. • tmax The maximum value of t that should be plotted. • surv_slices A Boolean. Default is FALSE. If TRUE and which_plot == "survival", plot survival curves over the time s for selected values of u, that are cross-sections of the 2D survival surface. • cumhaz_slices A Boolean. Default is FALSE. If TRUE and which_plot == "cumhaz", plot cumulative hazards curves over the time s for selected values of u, that are cross-sections of the 2D cumulative hazard surface. • midpoints A Boolean. Default is FALSE. If TRUE, the estimated quantities (hazard, survival, etc.) will be evaluated in the mid-points of the bins rather than at the extremes. Set to TRUE if plotting estimated number of events. • col_palette A function defining the color palette. The default palette is viridis::rev(plasma()). Specifying the color palette as a function allows for greater flexibility than passing the palette as a vector. We provide an example on how to create a function from any color palette below. • n_shades The number of color shades to plot, default is 50. • breaks The vector of breaks for the color legend. If n_shades is provided, this should be of length n_shades + 1. • show_legend A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid (i.e. transformed (u,s) plane). • main The title of the plot. • xlab The label of the first time axis (plotted on the x axis). • ylab The label of the second time axis (plotted on the y axis).

- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of the time scale on the y axis.
- `contour_lines` A Boolean. Default is FALSE. If TRUE contour lines are added to the surfaces.
- `contour_col` The color for the contour lines. Default is white.
- `contour_cex` The magnification to be used for the contour lines. Default is .8.
- `contour_nlev` The number of contour levels desired. Default is 10.
- `cex_main` The magnification to be used for the main title, default is 1.2 .
- `cex_lab` The magnification to be used for the axis labels, default is 1 .
- `HR` A Boolean. If TRUE the HRs with their CIs will be plotted. Default is FALSE (plot the beta with their CIs).
- `symmetric_CI` A Boolean. Default is TRUE. If a plot of the HRs is required (`HR == TRUE`), then plot symmetrical Confidence Intervals, based on the SEs for the HRs calculated by delta method. If FALSE, then CIs are obtained by exponentiating the CIs for the betas.
- `confidence` The level of confidence for the CIs. Default is .95 (alpha = 0.05).
- `col_beta` The color for the plot of the covariates' effects.
- `pch` The symbol for plotting the point estimates.
- `lwd` The line width.

... Further arguments to `image.plot` or `image`

Details

The vignette "visualization" presents and discusses all the different plotting options for the fitted model over two time scales. In most of the cases, the user will want to visualize the hazard surface over the two time scales. This can be plotted on the hazard scale, the log-hazard scale or the log10-hazard scale, by switching to TRUE the corresponding argument in `plot_options`. The survival and cumulative hazard functions can be plotted as two-dimensional surfaces over `u` and `s` or `t` and `s`. However, it is also very informative to plot them as one-dimensional curves over `s` (cross-sections or slices). This is done by selecting `which_plot = "survival"` and `surv_slices = TRUE` in `plot_options`. Additionally, a vector of values for the cutting points over the `u`-axis should be passed to the argument `where_slices`, together with setting `direction = u`. Similar plot is obtained for the cumulative hazard by selecting `which_plot = "cumhaz"`, `cumhaz_slices = TRUE`, see examples section. Please, notice that for the survival function and the cumulative hazard, only cross-sections of the surface for selected values of `u` (over the `s` time) can be plotted.

The function `obtainSmoothTrend` from the R-package `LMMsolver` is used here. We refer the interested readers to <https://biometris.github.io/LMMsolver/> for more details on `LMMsolver` and its usage.

Value

A plot of the fitted model.

Examples

```

# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)#'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(u = fakedata$u,
                          s_out = fakedata$s,
                          ev = fakedata$ev,
                          ds = .5)

# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "LMMsolver")

# plot the hazard surface
plot(fakemod)
# plot the survival function as one-dimension curves over `s`
plot(fakemod,
     which_plot = "survival",
     direction = "u",
     where_slices = c(4, 6, 8),
     plot_options = list(
       surv_slices = TRUE
     ))

# Plot cross-sections of the hazard over `s` for selected values of `u`
plot(fakemod,
     which_plot = "slices",
     where_slices = c(4, 6, 8),
     direction = "u",
     plot_options = list(
       main = "Cross-sections of the hazard",
       xlab = "Time",
       ylab = "Hazard"
     ))
)
)

```

Description

plot.haz2tsPGAM() is the plot method for objects of class haz2tsPGAM. It produces several kinds of plots of the fitted log-additive model with two time scales (see [fitpgam\(\)](#)), either on the original (t,s) plane, while respecting the constraint imposed by the relation of the two time scales, or on the transformed (u,s) plane.

Usage

```
## S3 method for class 'haz2tsPGAM'
plot(
  x,
  plot_grid = NULL,
  which_plot = c("hazard", "SE", "slices", "survival", "cumhaz"),
  where_slices = NULL,
  direction = c(NULL, "u", "s"),
  plot_options = list(),
  ...
)
```

Arguments

x	The output of the function <code>fitpgam()</code> . This is an object of class "haz2tsPGAM".
plot_grid	(optional) A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: <code>plot_grid = list(c(umin, umax, du), c(smin, smax, ds))</code> , where <code>umin</code> , <code>umax</code> and <code>smin</code> , <code>smax</code> are the minimum and maximum values desired for the intervals over u and s respectively, and <code>du</code> , <code>ds</code> are distances between intervals over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.
which_plot	The type of plot required. Can be one of "hazard" (default), "SE", "slices", "survival" or "cumhaz" (see details section).
where_slices	A vector of values for the cutting points of the desired slices of the surface. If <code>which_plot == "slices"</code> , please provide this argument. Please also provide this argument in case <code>which_plot = "survival"</code> or <code>which_plot = "cumhaz"</code> and <code>surv_slices = TRUE</code> or <code>cumhaz_slices = TRUE</code> , respectively.
direction	If <code>which_plot == "slices"</code> , indicates the direction for cutting the surface. If u, then the surface will be cut at the selected values of u (indicated by <code>where_slices</code>), hence obtaining one-dimensional curves over s. If s, then the surface will be cut at the selected values of s (indicated by <code>where_slices</code>), hence obtaining one-dimensional curves over u.
plot_options	A list with all possible options for any of the plots:

- `loghazard` A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard surface, if TRUE the function returns a plot of the log-hazard surface.
- `log10hazard` A Boolean. Default is FALSE. If TRUE, then a log₁₀ hazard surface is plotted.
- `cut_extrapolated` A Boolean. Default is TRUE. Cuts away the extrapolated area of the (log-)hazard surface before plotting.
- `rectangular_grid` A Boolean. Default is FALSE. If TRUE, a rectangular grid is used for plotting also in the (t,s)-plane as opposed to the grid of parallelograms used as default in the (t,s)-plane.
- `original` A Boolean. Default is TRUE. Plot the (log-)hazard (and/or the SEs) in the (t,s)-plane. If FALSE, the (log-)hazard (and/or the SEs) will be plotted in the (u,s)-plane.
- `tmax` The maximum value of t that should be plotted.
- `surv_slices` A Boolean. Default is FALSE. If TRUE and `which_plot == "survival"`, plot survival curves over the time s for selected values of u, that are cross-sections of the 2D survival surface.
- `cumhaz_slices` A Boolean. Default is FALSE. If TRUE and `which_plot == "cumhaz"`, plot cumulative hazards curves over the time s for selected values of u, that are cross-sections of the 2D cumulative hazard surface.
- `midpoints` A Boolean. Default is FALSE. If TRUE, the estimated quantities (hazard, survival, etc.) will be evaluated in the mid-points of the bins rather than at the extremes. Set to TRUE if plotting estimated number of events.
- `col_palette` A function defining the color palette. The default palette is `viridis::rev(plasma())`. Specifying the color palette as a function allows for greater flexibility than passing the palette as a vector.
- `n_shades` The number of color shades to plot, default is 50.
- `breaks` The vector of breaks for the color legend. If `n_shades` is provided, this should be of length `n_shades + 1`.
- `show_legend` A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid (i.e. transformed (u,s) plane)
- `main` The title of the plot.
- `xlab` The label of the first time axis (plotted on the x axis).
- `ylab` The label of the second time axis (plotted on the y axis).
- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of the time scale on the y axis.
- `contour_lines` A Boolean. Default is FALSE. If TRUE white contour lines are added to the surfaces.
- `contour_col` The color for the contour lines. Default is white.
- `contour_cex` The magnification to be used for the contour lines. Default is .8.
- `contour_nlev` The number of contour levels desired. Default is 10.

- `cex_main` The magnification to be used for the main title, default is 1.2 .
 - `cex_lab` The magnification to be used for the axis labels, default is 1 .
 - `lwd` The line width.
 - `lty` The line type.
- ... Further arguments to `image.plot` or `image`

Details

The vignette "visualization" presents and discusses all the different plotting options for the fitted model over two time scales. In most of the cases, the user will want to visualize the hazard surface over the two time scales. This can be plotted on the hazard scale, the log-hazard scale or the log10-hazard scale, by switching to TRUE the corresponding argument in `plot_options`. The survival and cumulative hazard functions can be plotted as two-dimensional surfaces over `u` and `s` or `t` and `s`. However, it is also very informative to plot them as one-dimensional curves over `s` (cross-sections or slices). This is done by selecting `which_plot = "survival"` and `surv_slices = TRUE` in `plot_options`. Additionally, a vector of values for the cutting points over the `u`-axis should be passed to the argument `where_slices`, together with setting `direction = u`. Similar plot is obtained for the cumulative hazard by selecting `which_plot = "cumhaz"`, `cumhaz_slices = TRUE`, see examples section. Please, notice that for the survival function and the cumulative hazard, only cross-sections of the surface for selected values of `u` (over the `s` time) can be plotted.

Value

A plot of the fitted model.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5
)
# Fit a fake model - not optimal smoothing
fakemod <- fitpgam(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
```

```

    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)

# plot the hazard surface
plot(fakemod)

# plot the survival function as one-dimension curves over `s`
plot(fakemod,
     which_plot = "survival",
     direction = "u",
     where_slices = c(4, 6, 8),
     plot_options = list(
       surv_slices = TRUE
     )
)

# Plot cross-sections of the hazard over `s` for selected values of `u`

plot(fakemod,
     which_plot = "slices",
     where_slices = c(4, 6, 8),
     direction = "u",
     plot_options = list(
       main = "Cross-sections of the hazard",
       xlab = "Time",
       ylab = "Hazard"
     )
)

```

plot.haz2tsVCM

Plot method for a haz2tsVCM object.

Description

plot.haz2tsVCM() is the plot method for objects of class haz2tsVCM. It produces several kinds of plots of the fitted varying coefficients model with two time scales (see `fitvcm()`), either on the original (t,s) plane, while respecting the constraint imposed by the relation of the two time scales, or on the transformed (u,s) plane.

Usage

```

## S3 method for class 'haz2tsVCM'
plot(
  x,
  plot_grid = NULL,
  which_plot = c("hazard", "SE", "slices", "survival", "cumhaz"),

```

```

    where_slices = NULL,
    direction = c(NULL, "u", "s"),
    plot_options = list(),
    ...
)

```

Arguments

- | | |
|--------------|--|
| x | The output of the function <code>fitvcm()</code> . This is an object of class "haz2tsVCM". |
| plot_grid | (optional) A list containing the parameters to build a new finer grid of intervals over <i>u</i> and <i>s</i> for plotting. This must be of the form: <code>plot_grid = list(c(umin, umax, du), c(smin, smax, ds))</code> , where <i>umin</i> , <i>umax</i> and <i>smin</i> , <i>smax</i> are the minimum and maximum values desired for the intervals over <i>u</i> and <i>s</i> respectively, and <i>du</i> , <i>ds</i> are distances between intervals over <i>u</i> and <i>s</i> respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned. |
| which_plot | The type of plot required. Can be one of "hazard" (default), "SE", "slices", "survival" or "cumhaz" (see details section). |
| where_slices | A vector of values for the cutting points of the desired slices of the surface. If <code>which_plot == "slices"</code> , please provide this argument. Please also provide this argument in case <code>which_plot = "survival"</code> or <code>which_plot = "cumhaz"</code> and <code>surv_slices = TRUE</code> or <code>cumhaz_slices = TRUE</code> , respectively. |
| direction | If <code>which_plot == "slices"</code> , indicates the direction for cutting the surface. If <i>u</i> , then the surface will be cut at the selected values of <i>u</i> (indicated by <code>where_slices</code>), hence obtaining one-dimensional curves over <i>s</i> . If <i>s</i> , then the surface will be cut at the selected values of <i>s</i> (indicated by <code>where_slices</code>), hence obtaining one-dimensional curves over <i>u</i> . |
| plot_options | A list with all possible options for any of the plots: <ul style="list-style-type: none"> • <code>loghazard</code> A Boolean. Default is FALSE. If FALSE the function returns a plot of the hazard surface, if TRUE the function returns a plot of the log-hazard surface. • <code>log10hazard</code> A Boolean. Default is FALSE. If TRUE, then a log₁₀ hazard surface is plotted. • <code>cut_extrapolated</code> A Boolean. Default is TRUE. Cuts away the extrapolated area of the (log-)hazard surface before plotting. • <code>rectangular_grid</code> A Boolean. Default is FALSE. If TRUE, a rectangular grid is used for plotting also in the (t,s)-plane as opposed to the grid of parallelograms used as default in the (t,s)-plane. • <code>original</code> A Boolean. Default is TRUE. Plot the (log-)hazard (and/or the SEs) in the (t,s)-plane. If FALSE, the (log-)hazard (and/or the SEs) will be plotted in the (u,s)-plane. |

- `tmax` The maximum value of `t` that should be plotted.
- `surv_slices` A Boolean. Default is FALSE. If TRUE and `which_plot == "survival"`, plot survival curves over the time `s` for selected values of `u`, that are cross-sections of the 2D survival surface.
- `cumhaz_slices` A Boolean. Default is FALSE. If TRUE and `which_plot == "cumhaz"`, plot cumulative hazards curves over the time `s` for selected values of `u`, that are cross-sections of the 2D cumulative hazard surface.
- `midpoints` A Boolean. Default is FALSE. If TRUE, the estimated quantities (hazard, survival, etc.) will be evaluated in the mid-points of the bins rather than at the extremes. Set to TRUE if plotting estimated number of events.
- `col_palette` A function defining the color palette. The default palette is `viridis::rev(plasma())`. Specifying the color palette as a function allows for greater flexibility than passing the palette as a vector.
- `n_shades` The number of color shades to plot, default is 50.
- `breaks` The vector of breaks for the color legend. If `n_shades` is provided, this should be of length `n_shades + 1`.
- `show_legend` A Boolean. Default is TRUE. If FALSE no legend will be plotted, useful for multi-panel figures with common legend. Works only for plots on rectangular grid (i.e. transformed (u,s) plane)
- `main` The title of the plot.
- `xlab` The label of the first time axis (plotted on the x axis).
- `ylab` The label of the second time axis (plotted on the y axis).
- `xlim` A vector with two elements defining the limits of the time scale on the x axis.
- `ylim` A vector with two elements defining the limits of the time scale on the y axis.
- `contour_lines` A Boolean. Default is FALSE. If TRUE white contour lines are added to the surfaces.
- `contour_col` The color for the contour lines. Default is white.
- `contour_cex` The magnification to be used for the contour lines. Default is .8.
- `contour_nlev` The number of contour levels desired. Default is 10.
- `cex_main` The magnification to be used for the main title, default is 1.2 .
- `cex_lab` The magnification to be used for the axis labels, default is 1 .
- `lwd` The line width.
- `lty` The line type.

... Further arguments to `image.plot` or `image`

Details

The vignette "visualization" presents and discusses all the different plotting options for the fitted model over two time scales. In most of the cases, the user will want to visualize the hazard surface over the two time scales. This can be plotted on the hazard scale, the log-hazard scale or the log10-hazard scale, by switching to TRUE the corresponding argument in `plot_options`. The survival and cumulative hazard functions can be plotted as two-dimensional surfaces over `u` and `s` or `t`

and s . However, it is also very informative to plot them as one-dimensional curves over s (cross-sections or slices). This is done by selecting `which_plot = "survival"` and `surv_slices = TRUE` in `plot_options`. Additionally, a vector of values for the cutting points over the u -axis should be passed to the argument `where_slices`, together with setting `direction = u`. Similar plot is obtained for the cumulative hazard by selecting `which_plot = "cumhaz"`, `cumhaz_slices = TRUE`, see examples section. Please, notice that for the survival function and the cumulative hazard, only cross-sections of the surface for selected values of u (over the s time) can be plotted.

Value

A plot of the fitted model.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1) #'

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5
)
# Fit a fake model - not optimal smoothing
fakemod <- fitvcm(fakedata2ts)

# plot the hazard surface
plot(fakemod)

# plot the survival function as one-dimension curves over `s`
plot(fakemod,
  which_plot = "survival",
  direction = "u",
  where_slices = c(4, 6, 8),
  plot_options = list(
    surv_slices = TRUE
  )
)

# Plot cross-sections of the hazard over `s` for selected values of `u`

plot(fakemod,
```

```

which_plot = "slices",
where_slices = c(4, 6, 8),
direction = "u",
plot_options = list(
  main = "Cross-sections of the hazard",
  xlab = "Time",
  ylab = "Hazard"
)
)

```

plot_slices

Plot slices of the (log-) hazard

Description

plot_slices() plots slices of the (log-)hazard with two time scales, at selected values of one of the two time dimensions.

Usage

```
plot_slices(x, y, direction, plot_options = list())
```

Arguments

x	A vector of values for the x-axis. This is a vector of values over the axis opposite to the one where the sliced are cut.
y	A matrix of (log-)hazard values.
direction	Either "u" or "s".
plot_options	A list of options for the plot: <ul style="list-style-type: none"> loghazard A Boolean. Default is FALSE. If FALSE the function returns a plot of cross-sections from the hazard surface, if TRUE the function returns a plot of cross-sections from the log-hazard surface. log10hazard A Boolean. Default is FALSE. If TRUE returns a plot of cross-sections from the log10-hazard surface. col_palette A function defining the color palette. The default palette is grDevices::gray.colors(). n_shades The number of color shades to plot, default is 50. main The title of the plot. xlab The label of the first time axis (plotted on the x axis). ylab The label of the second time axis (plotted on the y axis). xlim A vector with two elements defining the limits of the time scale on the x axis. ylim A vector with two elements defining the limits of the time scale on the y axis.

- `cex_main` The magnification to be used for the main title, default is 1.2.
- `cex_lab` The magnification to be used for the axis labels, default is 1.
- `lwd` The line width. Default is 2.
- `lty` The line type.

Value

A plot of the slices of the hazard cut at selected points.

predict_cif2ts_pointwise

Point-wise prediction of cumulative incidence over 2 time scale

Description

Point-wise prediction of cumulative incidence over 2 time scale

Usage

```
predict_cif2ts_pointwise(fitted_models = list(), u, s, ds = NULL)
```

Arguments

`fitted_models` a list with cause-specific hazard models

`u` The value(s) of `u` where prediction is required

`s` The value(s) of `s` where prediction is required

`ds` (optional) The distance between two consecutive points on the `s` axis. If not provided, an optimal minimum value will be chosen automatically and a warning is returned.

Value

A data.frame with one row containing: the values of `u` and `s` for which predictions of the overall survival (`surv`) probability, and the values of the cumulative incidence functions, one for each cause, are obtained.

Examples

```
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
```

```

ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)

fakedata <- as.data.frame(cbind(id, u, s, ev))
# cause 1
fakedata2ts1 <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  min_u = 2, min_s = 0,
  ds = .5
)
# Fit a fake model - not optimal smoothing for cause type 1
fakemod1 <- fit2ts(fakedata2ts1,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)
# cause 2
fakedata2ts2 <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = 1 - (fakedata$ev),
  min_u = 2, min_s = 0,
  ds = .5
)
# Fit a fake model - not optimal smoothing for cause 2
fakemod2 <- fit2ts(fakedata2ts2,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)
predict_cif2ts_pointwise(
  fitted_models = list(fakemod1, fakemod2),
  u = 5.2, s = 4.4
)

```

predict_comprisks2ts *Predict overall survival and cumulative incidence functions*

Description

Predict overall survival and cumulative incidence functions

Usage

```
predict_comprisks2ts(models = list(), newdata, u, s, ds = NULL)
```

Arguments

models	A list with cause-specific hazard models of class 'haz2ts', fitted via fit2ts(). The models have to be fitted on the same grid of points over the time axes.
newdata	A dataframe with columns containing the values of the variable u and the time scale s for which predictions are to be obtained.
u	The name of the variable in newdata containing values for the variable u.
s	The name of the variable in newdata containing values for the variable s.
ds	(optional) The distance between two consecutive points on the s axis. If not provided, an optimal minimum value will be chosen automatically and a warning is returned.

Details

Predictions of cumulated quantities can be provided only within intervals of values on the s time scale.

Value

A dataframe containing the values of u and s in newdata, the predicted survival probability and the values of the cumulative incidence functions (cif), for each combination of u and s. There will be as many values of the cif as the number of cause-specific hazard models.

Examples

```
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)

fakedata <- as.data.frame(cbind(id, u, s, ev))
# cause 1
fakedata2ts1 <- prepare_data(u = fakedata$u,
                           s_out = fakedata$s,
                           ev = fakedata$ev,
                           ds = .5)

# Fit a fake model - not optimal smoothing for cause type 1
fakemod1 <- fit2ts(fakedata2ts1,
                  optim_method = "grid_search",
                  lrho = list(seq(1, 1.5, .5),
                             seq(1, 1.5, .5)))

# cause 2
fakedata2ts2 <- prepare_data(u = fakedata$u,
                           s_out = fakedata$s,
                           ev = 1-(fakedata$ev),
                           ds = .5)

# Fit a fake model - not optimal smoothing for cause 2
fakemod2 <- fit2ts(fakedata2ts2,
                  optim_method = "grid_search",
```

```

lrho = list(seq(1 ,1.5 ,.5),
            seq(1 ,1.5 ,.5))

newdata <- as.data.frame(cbind("u" = c(2.5,3.4,6), "s" = c(.2,.5,1.3)))

predict_comprisks2ts(models = list(fakemod1, fakemod2),
                    newdata,
                    u = "u", s = "s")

```

predict_haz2ts *Prediction method for objects of class 'haz2ts'*

Description

Prediction method for objects of class 'haz2ts'

Usage

```

predict_haz2ts(
  x,
  newdata = NULL,
  originaldata = NULL,
  u,
  s,
  z = NULL,
  id = NULL,
  ds = NULL
)

```

Arguments

x	an object of class 'haz2ts', the output of the function fit2ts().
newdata	(optional) A dataframe with columns containing the values of the variable u and the time scale s for which predictions are to be obtained.
originaldata	(optional) The original dataset. Provide it to obtain individual predictions for each observation in the data.
u	The name of the variable in newdata, or in originaldata containing values for the variable u.
s	The name of the variable in newdata, or in originaldata containing values for the variable s. Note that over the s axis predictions are provided only within intervals of values, as it is necessary to approximate cumulated quantities.
z	Covariates value
id	(optional) The name of the variable in newdata, or in originaldata containing the identification of each observation. It is not required for predictions on a new dataset.
ds	(optional) The distance between two consecutive points on the s axis. If not provided, an optimal minimum value will be chosen automatically and a warning is returned.

Details

Predictions of cumulated quantities can be provided only within intervals of values on the *s* time scale.

Value

A dataframe. This can be the original dataframe (*originaldata*), where only the variables *id*, *u* and *s* are selected, or the new data frame (*newdata*), together with the predicted values for the hazard hazard and its standard errors *se_hazard*, the cumulative hazard *cumhazard* and the survival probability *survival*.

Examples

```
# Create the same fake data as in other examples
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)
z1 <- rbinom(n = 20, size = 20, prob = .5)
z2 <- rnorm(n = 20)

fakedata <- as.data.frame(cbind(id, u, s, ev, z1, z2))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5, individual = TRUE,
  covs = subset(fakedata, select = c("z1", "z2"))
)

# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)

# Create a new dataset for prediction
newdata <- as.data.frame(cbind("u" = c(2.5, 3.4, 6),
  "s" = c(.2, .5, 1.3)))

# First - predict on original data
predict(object = fakemod,
  originaldata = fakedata, u = "u", s = "s", id = "id"
)
```

```

# Now - predict on new dataset
predict(object = fakemod,
  newdata = newdata, u = "u", s = "s"
)

# Now - predict including covariates
predict(object = fakemod,
  originaldata = fakedata,
  u = "u", s = "s", id = "id",
  z = c("z1", "z2")
)

# If one wants to predict with only one of the covariates at a different
# value than the baseline, the other one(s) should be fixed at their
# baseline levels

newdata2 <- as.data.frame(cbind("u" = c(2.5, 3.4, 6),
  "s" = c(.2, .5, 1.3),
  "z1" = c(1, 2, 3),
  "z2" = c(0, 0, 0)))

predict(object = fakemod,
  newdata = newdata2,
  u = "u", s = "s", id = "id",
  z = c("z1", "z2")
)

```

predict_haz2ts_pointwise

Point-wise prediction hazard 2 time scale

Description

Point-wise prediction hazard 2 time scale

Usage

```
predict_haz2ts_pointwise(fitted_model, upred, spread, zpred = NULL, ds = NULL)
```

Arguments

fitted_model	An object of class 'haz2ts' fitted via fit2ts().
upred	The value(s) of u where prediction is required
spread	The value(s) of s where prediction is required
zpred	(optional) is a vector of values for the covariates in the model
ds	(optional) The distance between two consecutive points on the s axis. If not provided, an optimal minimum value will be chosen automatically and a warning is returned.

Value

A data.frame with one row and 6 variable: the values of `u` and `s` for which predictions of hazard, `se_hazard`, the cumulative hazard `cumhaz` and the survival probability are obtained

Examples

```
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)
fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  u = fakedata$u,
  s_out = fakedata$s,
  ev = fakedata$ev,
  ds = .5
)
# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(
    seq(1, 1.5, .5),
    seq(1, 1.5, .5)
  )
)
predict_haz2ts_pointwise(fakemod, upred = 5, spread = 4.44)
```

```
prepare_data
```

Prepare raw data by binning them in 1d or 2d

Description

`prepare_data()` prepares the raw individual time-to-event data for hazard estimation in 1d or 2d.

Given the raw data, this function first constructs the bins over one or two time axes and then computes the aggregated (or individual) vectors, or matrices, of exposure times and events indicators.

Usage

```
prepare_data(
  data = NULL,
  t_in = NULL,
  t_out = NULL,
```

```

u = NULL,
s_in = NULL,
s_out,
events,
min_t = NULL,
max_t = NULL,
min_u = NULL,
max_u = NULL,
min_s = NULL,
max_s = NULL,
dt = NULL,
du = NULL,
ds,
individual = FALSE,
covs = NULL
)

```

Arguments

<code>data</code>	A data frame. If the object provided is a tibble, or a data.table, it will be converted to data.frame.
<code>t_in</code>	(optional) A vector of entry times on the time scale t.
<code>t_out</code>	(optional) A vector of exit times on the time scale t.
<code>u</code>	(optional) A vector of fixed-times at entry in the process.
<code>s_in</code>	(optional) A vector of entry times on the time scale s.
<code>s_out</code>	A vector of exit times on the time scale s.
<code>events</code>	A vector of event's indicators (possible values 0/1).
<code>min_t</code>	(optional) A minimum value for the bins over t. If NULL, the minimum of <code>t_in</code> will be used.
<code>max_t</code>	(optional) A maximum value for the bins over t. If NULL, the maximum of <code>t_out</code> will be used.
<code>min_u</code>	(optional) A minimum value for the bins over u. If NULL, the minimum of u will be used.
<code>max_u</code>	(optional) A maximum value for the bins over u. If NULL, the maximum of u will be used.
<code>min_s</code>	(optional) A minimum value for the bins over s. If NULL, the minimum of <code>s_in</code> will be used.
<code>max_s</code>	(optional) A maximum value for the bins over s. If NULL, the maximum of <code>s_out</code> will be used.
<code>dt</code>	(optional) A scalar giving the length of the intervals on the t time scale.
<code>du</code>	(optional) A scalar giving the length of the intervals on the u axis.
<code>ds</code>	A scalar giving the length of the intervals on the s time scale.
<code>individual</code>	A Boolean. Default is FALSE: if FALSE computes the matrices R and Y collectively for all observations; if TRUE computes the matrices R and Y separately for each individual record.

covs A data.frame with the variables to be used as covariates, or a vector with the names of the covariates to be included. The function will create dummy variables for any factor variable passed as argument in covs. If a variable of class character is passed as argument, it will be converted to factor.

Details

A few words about constructing the grid of bins. Bins are containers for the individual data. There is no 'golden rule' or optimal strategy for setting the number of bins over each time axis, or deciding on the bins' width. It very much depends on the data structure, however, we try to give some directions here. First, in most cases, more bins is better than less bins. A good number is about 30 bins. It also depends on the user's expectation about how rapidly the hazard of the event changes over the time scales. However, if data are scarce, the user might want to find a compromise between having a larger number of bins, and having many bins empty. Second, the chosen width of the bins (that is du and ds) does depend on the time unit over which the time scales are measured. For example, if the time is recorded in days, as in the example below, and several years of follow-up are available, the user can split the data in bins of width 30 (corresponding to about one month), 60 (about two months), 90 (about three months), etc. If the time scale is measured in years, then appropriate width could be 0.25 (corresponding to a quarter of a year), or 0.5 (that is half year). However, in some cases, time might be measure in completed years, as is often the case for age. In this scenario, an appropriate bin width is 1.

Finally, it is always a good idea to plot the data first, and explore the range of values over which the time scale(s) are recorded. This will give insight about reasonable values for the arguments `min_s`, `min_u`, `max_s` and `max_u` (that in any case are optional).

Regarding the names of covariates or levels of categorical covariates/factors: When using "LMM-solver" to fit a model with covariates that have names (or factor labels) including a symbol such as "+", "-", "<" or ">", these will result in an error. To avoid this, the responsible names (labels) will be rewritten without mathematical symbols. For example: "Lev+5FU" (in the colon cancer data) is replaced by "Lev&5FU".

Value

A list with the following elements:

- bins a list:
 - bins_t if `t_out` is provided, this is a vector of bins extremes for the time scale `t`.
 - mid_t if `t_out` is provided, this is a vector with the midpoints of the bins over `t`.
 - nt if `t_out` is provided, this is the number of bins over `t`.
 - bins_u if `u` is provided, this is a vector of bins extremes for `u` axis.
 - midu if `u` is provided, this is a vector with the midpoints of the bins over `u`.
 - nu if `u` is provided, this is the number of bins over `u`.
 - bins_s is a vector of bins extremes for the time scale `s`.
 - mids is a vector with the midpoints of the bins over `s`.
 - ns is the number of bins over `s`.
- bindata:

- r or R an array of exposure times: if binning the data over one time scale only this is a vector. If binning the data over two time scales and if `individual == TRUE` then R is an array of dimension nu by ns by n , otherwise it is an array of dimension nu by ns
- y or Y an array of event counts: if binning the data over one time scale only this is a vector. If binning the data over two time scales and if `individual == TRUE` then Y is an array of dimension nu by ns by n , otherwise it is an array of dimension nu by ns
- Z A matrix of covariates' values to be used in the model, of dimension n by p

Author(s)

Angela Carollo <carollo@demogr.mpg.de>

Examples

```
# Bin data over s = time since recurrence only, with intervals of length 30 days
# aggregated data (no covariates)
# The following example provide the vectors of data directly from the dataset
binned_data <- prepare_data(s_out = reccolon2ts$timesr, events = reccolon2ts$status, ds = 30)
# Visualize vector of event counts
print(binned_data$bindata$y)
# Visualize midpoints of the bins
print(binned_data$bins$mids)
# Visualize number of bins
print(binned_data$bins$ns)

# Now, the same thing is done by providing a dataset and the name of all relevant variables
binned_data <- prepare_data(data = reccolon2ts, s_out = "timesr", events = "status", ds = 30)
# Visualize vector of event counts
print(binned_data$bindata$y)

# Now using ds = .3 and the same variable measured in years
binned_data <- prepare_data(s_out = reccolon2ts$timesr_y, events = reccolon2ts$status, ds = .3)
# Visualize vector of exposure times
print(binned_data$bindata$r)

# Bin data over u = time at recurrence and s = time since recurrence, measured in days
# aggregated data (no covariates)
# Note that if we do not provide du this is taken to be equal to ds
binned_data <- prepare_data(
  u = reccolon2ts$timer, s_out = reccolon2ts$timesr,
  events = reccolon2ts$status, ds = 30
)

# Visualize matrix of event counts
print(binned_data$bindata$Y)

# Visualize midpoints of bins over u
print(binned_data$bins$midu)

# Bin data over u = time at recurrence and s = time since recurrence, measured in day
```

```

# individual-level data required
# we provide two covariates: nodes (numerical) and rx (factor)
covs <- subset(reccolon2ts, select = c("nodes", "rx"))
binned_data <- prepare_data(
  u = reccolon2ts$timer, s_out = reccolon2ts$timesr,
  events = reccolon2ts$status, ds = 30, individual = TRUE, covs = covs
)

# Visualize structure of binned data
print(str(binned_data$bindata))

# Alternately:
binned_data <- prepare_data(
  data = reccolon2ts,
  u = "timer", s_out = "timesr",
  events = "status", ds = 30, individual = TRUE, covs = c("nodes", "rx")
)

```

```
prepare_data_LMMsolver
```

Process data to fit model with LMMsolver

Description

Process data to fit model with LMMsolver

Usage

```
prepare_data_LMMsolver(Y = Y, R = R, Z = NULL, bins = bins)
```

Arguments

Y	A matrix (or 3d-array) of event counts of dimension nu by ns (or nu by ns by n).
R	A matrix (or 3d-array) of exposure times of dimension nu by ns (or nu by ns by n).
Z	(optional) A regression matrix of covariates values of dimensions n by p.
bins	a list with the specification for the bins. This is created by the function prepare_data. If a list prepared externally from such function if provided, it should contain the following elements: * bins_u A vector of bins extremes for the time scale u. * midu A vector with the midpoints of the bins over u. * nu The number of bins over u. * bins_s A vector of bins extremes for the time scale s. * mids A vector with the midpoints of the bins over s. * ns The number of bins over s.

Value

A dataset in long form to fit the model with LMMsolver

print.data2ts	<i>Print method for a data2ts object</i>
---------------	--

Description

Print method for an object of class data2ts

Usage

```
## S3 method for class 'data2ts'  
print(x, ...)
```

Arguments

x	of class data2ts, as prepared by prepare_data
...	Further arguments to print

Value

No return value

Author(s)

Angela Carollo <carollo@demogr.mpg.de>

Examples

```
# Bin the colon cancer data over s (time since recurrence)  
  
dt1ts <- prepare_data(data = reccolon2ts,  
                     s_in = "entrys",  
                     s_out = "timesr",  
                     events = "status",  
                     ds = 180)  
  
print(dt1ts)  
  
# Bin the colon cancer data over u (time at recurrence) and s (time since recurrence)  
dt2ts <- prepare_data(data = reccolon2ts,  
                     u = "timer",  
                     s_in = "entrys",  
                     s_out = "timesr",  
                     events = "status",  
                     ds = 180)  
  
print(dt2ts)
```

reccolon2ts

*Data from the chemotherapy for stage B/C colon cancer study***Description**

This dataset is a reduced version of the dataset colon from the package survival (Therneau, T., 2023). Each observation is a transition from recurrence of colon cancer to death or censoring. The time scales are time from randomization to recurrence, time from randomization to death or censoring and time from recurrence of the cancer to death or censoring. Only observations about individuals with a recurrence of the cancer are selected. Additionally, 7 individuals with exit times from the risk set equal to entry times in the recurrence state (0 exposure time) were dropped from the sample. In the original dataset, all times of recurrence are known, so that after recurrence all individuals are followed from entry in the state, without left truncation. To be able to illustrate how to include left truncated times in the model, artificial left truncated entry in the 'recurrence' state are randomly introduced for 40 individuals.

Usage

```
data(reccolon2ts)
```

Format

reccolon2ts A data.table with 461 rows and 25 columns::

id patient's id

study 1 for all patients

rx Treatment - Obs(ervation), Lev(amisole), Lev(amisole)+5-FU

sex 1=male, 0=female

age Age at transplant in years

obstruct obstruction of colon by tumor: 1=yes

perfor perforation of colon: 1=yes

adhere adherence to nearby organs: 1=yes

nodes number of lymph nodes with detectable cancer

status censoring status: 0=censoring, 1=event

differ differentiation of tumour: 1=well, 2=moderate, 3=poor

extent extent of local spread: 1=submucosa, 2=muscle, 3=serosa, 4=contiguous structures

surg time from surgery to registration: 0=short, 1=long

node4 more than 4 positive lymph nodes

etype 2 for all patients (2=death)

timedc time in days from randomization to death or censoring

timer time in days from randomization to recurrence

timesr time in days from recurrence to death or censoring

entrys artificial entry time on the time since recurrence scale. For most of the individual this is 0 (no left truncation). For 40 individuals a random number between 1 and the exit time on the time since recurrence scale (timesr) is simulated.

entryt time in days from randomization to observation in the recurrence state. If the individual is observed from entry in the recurrence state this is equal to the time at recurrence. If the entry in the recurrence state is not observed from the beginning, left truncation is observed. This is not present in the original data, but has been here introduced artificially for 40 individuals. This is done by first increasing the time at recurrence by a random number between 1 and the exit time on the time since recurrence scale. Then, the time at recurrence is added to the artificial entry time.

timedc_y time in years from randomization to death or censoring

timer_y time in years from randomization to recurrence

entrys_y left truncated entry in the recurrence state measured in years since recurrence

entryt_y left truncated entry in the recurrence state measured in years since randomization

timesr_y time in years from recurrence to death or censoring

Source

Therneau, T. (2023). A Package for Survival Analysis in R. R package version 3.5-3, <https://CRAN.R-project.org/package=survival>

References

Moertel, C.G., et al. (1995). Fluorouracil plus Levamisole as Effective Adjuvant Therapy after Resection of Stage III Colon Carcinoma: A Final Report. *Annals of Internal Medicine*, 122:321-326

Moerel, C.G., et al. (1990). Levamisole and Fluorouracil for Adjuvant Therapy of Resected Colon Carcinoma. *The New England Journal of Medicine*, 322:352-8

Examples

```
data(reccolon2ts)
rm(reccolon2ts)
```

<code>select_model2ts</code>	<i>Selects best model among a group of fitted models</i>
------------------------------	--

Description

`select_model2ts()` takes as input a list of hazard models with two time scales, and returns a table with the best fitting model indicated. The selection is based on minimization of the AIC or BIC, so the models do not need to be nested. It can be used, for example, to identify the best model among the log-additive model, the varying coefficient model, and the full 2D model.

Usage

```
select_model2ts(model_list, sel_criteria = c("aic", "bic"))
```

Arguments

`model_list` is a list of fitted objects, of class `haz2ts`, `hat2tsLMM`, `haz2tsPGAM`, or `haz2tsVCM`.
`sel_criteria` to determine the best model - it can be `'aic'` or `'bic'`.

Details

In the model list, it is possible to provide a character name for each model. In such case, these names will also be reported in the results' table. An example is provided (See examples).

Value

A data.frame with as many rows as the number of models being compared, and the following columns: * Model: the model name (see details and examples) * Type: the class of the model * AIC * BIC * Best: indicates which of the model is the best fitting one with respect to the criteria indicated

Examples

```
id <- 1:20
u <- c(
  5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
  4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96
)
s <- c(
  0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
  7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00
)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(
  data = fakedata,
  u = "u",
  s_out = "s",
  ev = "ev",
  ds = .5
)
# Fit a fake model - not optimal smoothing
full2d <- fit2ts(fakedata2ts,
  optim_method = "grid_search",
  lrho = list(seq(1, 1.5, .5), seq(1, 1.5, .5))
)
pgam <- fitpgam(fakedata2ts,
  optim_method = "grid_search",
  optim_criterion = "aic",
  lrho = list(seq(1, 1.5, .5), seq(1, 1.5, .5))
)
vcm <- fitvcm(fakedata2ts)

select_model2ts(model_list = list(
  "full interaction" = full2d,
```

```

    "additive" = pgam,
    "varying coeff" = vcm
  ))

```

surv2ts

Survival function with two time scales

Description

Computes the survival matrix containing the probability of not experiencing an event (of any cause) by time s and fixed entry time u . The survival function can be obtained from one fitted model with only one event type, or combining information from several cause-specific hazard in a competing risks model. In the first case, a fitted object of class 'haz2ts', 'haz2tsLMM', 'haz2tsPGAM', or 'haz2tsVCM' can be passed directly as argument to the function. In the competing risks framework, the user should provide a list of cause-specific cumulative hazard matrices. The function is also called internally from `plot()` if the user wants to plot the cumulative hazard from a fitted model.

Usage

```

surv2ts(
  cumhaz = NULL,
  fitted_model = NULL,
  plot_grid = NULL,
  cause = NULL,
  midpoints = FALSE,
  where_slices = NULL,
  direction = c("u", "s", NULL),
  tmax = NULL
)

```

Arguments

- | | |
|---------------------------|--|
| <code>cumhaz</code> | (optional) a list with all the cause-specific cumulated hazard matrices (minimum one element needs to be supplied). If more than one cause-specific cumulated hazard is provided, then they should all be matrices of the same dimension. |
| <code>fitted_model</code> | (optional) An object of class 'haz2ts', 'haz2tsLMM', 'haz2tsPGAM', or 'haz2tsVCM'. |
| <code>plot_grid</code> | (optional) A list containing the parameters to build a new finer grid of intervals over u and s for plotting. This must be of the form: <ul style="list-style-type: none"> <code>plot_grid = list(c(umin, umax, du), c(smin, smax, ds))</code> where <code>umin</code>, <code>umax</code> and <code>smin</code>, <code>smax</code> are the minimum and maximum values desired for the grid-points over u and s respectively, and <code>du</code>, <code>ds</code> are distances between points over u and s respectively. Specifying a new denser grid is used to evaluate the B-spline bases used for estimation on such grid and plot the estimated surfaces with a greater level of details. If not specified, the plotting is done using the same B-splines bases as for the estimation. The function |

will check if the parameters for the grid provided by the user are compatible with those originally used to construct the B-splines for estimating the model. If not, the grid will be adjusted accordingly and a warning will be returned.

cause	a character string with a short name for the cause (optional).
midpoints	A Boolean. Default is FALSE. If TRUE, the estimated quantities are evaluated at the midpoints of the rectangles (or parallelograms) of the grids, rather than at each grid-point.
where_slices	A vector of values for the cutting points of the desired slices of the surface. This option is included mostly for the plotting function. When using <code>plot.haz2ts()</code> , the user selects <code>which_plot = "survival"</code> and <code>surv_slices = TRUE</code> , then <code>where_slices</code> indicates the location of the cutting points over the <code>u</code> time.
direction	If cross-sectional one-dimensional curves are plotted, this indicates whether the cutting points are located on the <code>u</code> time, or on the <code>s</code> time. For plots of the survival function, only cutting points over the <code>u</code> time are meaningful.
tmax	The maximum value of <code>t</code> that should be plotted.

Value

a matrix containing the values of the survival function over `s` and `u`.

Examples

```
# Create some fake data - the bare minimum
id <- 1:20
u <- c(5.43, 3.25, 8.15, 5.53, 7.28, 6.61, 5.91, 4.94, 4.25, 3.86, 4.05, 6.86,
      4.94, 4.46, 2.14, 7.56, 5.55, 7.60, 6.46, 4.96)
s <- c(0.44, 4.89, 0.92, 1.81, 2.02, 1.55, 3.16, 6.36, 0.66, 2.02, 1.22, 3.96,
      7.07, 2.91, 3.38, 2.36, 1.74, 0.06, 5.76, 3.00)
ev <- c(1, 0, 0, 1, 0, 1, 0, 1, 1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 1)

fakedata <- as.data.frame(cbind(id, u, s, ev))
fakedata2ts <- prepare_data(u = fakedata$u,
                          s_out = fakedata$s,
                          ev = fakedata$ev,
                          ds = .5)

# Fit a fake model - not optimal smoothing
fakemod <- fit2ts(fakedata2ts,
                 optim_method = "grid_search",
                 lrho = list(seq(1, 1.5, .5),
                             seq(1, 1.5, .5)))

# Obtain the fake cumulated hazard
fakecumhaz2ts <- cumhaz2ts(fakemod)
# Fake survival curve
fakesurv2ts <- surv2ts(fitted_model = fakemod)
```

Index

* datasets

reccolon2ts, 75

covariates_plot, 3

cumhaz2ts, 4

cuminc2ts, 6

exposures_events_1d, 7

exposures_events_2d, 8

exposures_events_Lexis, 10

fit1ts, 11

fit2ts, 15

fit2ts(), 33, 36, 47, 51

fitpgam, 19

fitpgam(), 34, 55

fitvcm, 22

fitvcm(), 35, 58

get_hazard_1d, 25

get_hazard_1d_LMM, 26

get_hazard_2d, 28

get_hazard_2d_LMM, 30

get_hr, 32

haz2ts_summary, 36

haz2tsLMM_summary, 33

haz2tsPGAM_summary, 34

haz2tsVCM_summary, 35

imageplot_2ts, 37

imageplot_SE, 38

make_bins, 40

make_bins(), 8, 9

make_grid, 42

plot.haz1ts, 43

plot.haz1tsLMM, 45

plot.haz2ts, 47

plot.haz2tsLMM, 51

plot.haz2tsPGAM, 54

plot.haz2tsVCM, 58

plot_slices, 62

predict_cif2ts_pointwise, 63

predict_comprisks2ts, 64

predict_haz2ts, 66

predict_haz2ts_pointwise, 68

prepare_data, 69, 74

prepare_data(), 8–10, 41

prepare_data_LMMsolver, 73

print.data2ts, 74

reccolon2ts, 75

select_model2ts, 76

surv2ts, 78