

Package: **weightedsurv** (via r-universe)

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Title Survival Analysis with Subject-Specific (Case Weights) and Time-Dependent Weighting

Version 0.1.0

Description Provides survival analysis functions with support for time-dependent and subject-specific (e.g., propensity score) weighting. Implements weighted estimation for Cox models, Kaplan-Meier survival curves, and treatment differences with point-wise and simultaneous confidence bands. Includes restricted mean survival time (RMST) comparisons evaluated across all potential truncation times with both point-wise and simultaneous confidence bands. See Cole, S. R. & Hernán, M. A. (2004) <[doi:10.1016/j.cmpb.2003.10.004](https://doi.org/10.1016/j.cmpb.2003.10.004)> for methodological background.

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URL <https://larry-leon.github.io/weightedsurv/>,
<https://github.com/larry-leon/weightedsurv>

BugReports <https://github.com/larry-leon/weightedsurv/issues>

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add_legends	<i>Add legends to KM plot</i>
-------------	-------------------------------

Description

Adds legends for Cox model, log-rank test, and arms to a Kaplan-Meier plot.

Usage

```
add_legends(
  dfcount,
  show.cox,
  cox.cex,
  put.legend.cox,
  show.logrank,
  logrank.cex,
  put.legend.lr,
  show_arm_legend,
  arms,
  col.1,
  col.0,
  ltys,
  lwds,
  arm.cex,
  put.legend.arms
)
```

Arguments

dfcount	List with results, typically output from a survival analysis function. Should contain elements such as <code>cox_results</code> , <code>z.score</code> , and <code>zlogrank_text</code> .
show.cox	Logical; show Cox legend.
cox.cex	Numeric; Cox legend size.
put.legend.cox	Character; Cox legend position (e.g., "topright").
show.logrank	Logical; show logrank legend.
logrank.cex	Numeric; logrank legend size.
put.legend.lr	Character; logrank legend position (e.g., "topleft").
show_arm_legend	Logical; show arm legend.
arms	Character vector of arm labels.
col.1	Color for group 1.
col.0	Color for group 0.
ltys	Line types.

lwds	Line widths.
arm.cex	Numeric; arm legend size.
put.legend.arms	Character; arm legend position (e.g., "left").

Value

Invisibly returns NULL. Used for plotting side effects.

add_median_annotation *Add median annotation to KM plot*

Description

Adds median survival annotation to a Kaplan-Meier plot.

Usage

```
add_median_annotation(
  medians_df,
  med.digits,
  med.cex,
  med.font,
  xmed.fraction,
  ymed.offset
)
```

Arguments

medians_df	Data frame with quantile results. Should contain columns quantile, lower, upper, and group.
med.digits	Integer; number of digits to display for median and confidence interval.
med.cex	Numeric; text size for median annotation.
med.font	Integer; font for median annotation.
xmed.fraction	Numeric; fraction of the x-axis for annotation placement (e.g., 0.8 for 80% to the right).
ymed.offset	Numeric; offset from the top of the plot for annotation placement.

Value

Invisibly returns NULL. Used for plotting side effects.

add_risk_table	<i>Add risk table annotation to KM plot</i>
----------------	---

Description

Adds risk set counts for two groups to a Kaplan-Meier plot.

Usage

```
add_risk_table(  
  risk.points,  
  rpoints0,  
  rpoints1,  
  col.0,  
  col.1,  
  risk.cex,  
  ymin,  
  risk_offset,  
  risk_delta,  
  y.risk0,  
  y.risk1  
)
```

Arguments

risk.points	Numeric vector of risk time points.
rpoints0	Numeric vector of risk set counts for group 0.
rpoints1	Numeric vector of risk set counts for group 1.
col.0	Color for group 0.
col.1	Color for group 1.
risk.cex	Numeric; text size for risk table.
ymin	Numeric; minimum y value.
risk_offset	Numeric; offset for risk table.
risk_delta	Numeric; delta for risk table.
y.risk0	Numeric; y position for group 0 risk table.
y.risk1	Numeric; y position for group 1 risk table.

Value

Invisibly returns NULL. Used for plotting side effects.

calculate_risk_event_counts

Calculate risk set and event counts at time points

Description

Calculates risk set and event counts for a group at specified time points, with variance estimation.

Usage

```
calculate_risk_event_counts(U, D, W, at_points, draws = 0, seedstart = 816951)
```

Arguments

U	Numeric vector of times for group.
D	Numeric vector of event indicators for group.
W	Numeric vector of weights for group.
at_points	Numeric vector of time points.
draws	Number of draws for variance estimation (default 0).
seedstart	Random seed for draws (default 816951).

Value

List with ybar (risk set counts), nbar (event counts), sig2w_multiplier (variance term).

check_km_curve

Check KM curve for validity

Description

Checks that a Kaplan-Meier curve is valid (values in [0,1], non-increasing).

Usage

```
check_km_curve(S.KM, group_name = "Group", stop_on_error = TRUE)
```

Arguments

S.KM	Numeric vector of survival probabilities.
group_name	Character; name of the group.
stop_on_error	Logical; whether to stop on error (default TRUE).

Value

None. Stops or warns if invalid.

check_results	<i>Check and Compare Statistical Test Results</i>
---------------	---

Description

Calculates and compares three equivalent test statistics from weighted survival analysis: the squared standardized weighted log-rank statistic, the log-rank chi-squared statistic, and the squared Cox model z-score. These should be approximately equal under correct implementation.

Usage

```
check_results(dfcount, verbose = TRUE)
```

Arguments

dfcount	A list or data frame from <code>df_counting</code> containing: lr Weighted log-rank statistic sig2_lr Variance of the weighted log-rank statistic z.score Standardized z-score from Cox model logrank_results List containing chi sq, the log-rank chi-squared statistic
verbose	Logical; if TRUE, prints the comparison table to the console. Default: TRUE.

Details

This function serves as a diagnostic tool to verify computational consistency. The three statistics should be numerically equivalent (within rounding error):

$$(lr/\sqrt{sig2_lr})^2 \approx logrank_chisq \approx z.score^2$$

Discrepancies between these values may indicate:

- Numerical instability in variance estimation
- Incorrect weighting scheme application
- Data processing errors

Value

A data frame with one row and three columns, returned invisibly:

zlr_sq Squared standardized weighted log-rank: $(lr/\sqrt{sig2_lr})^2$

logrank_chisq Chi-squared statistic from log-rank test

zCox_sq Squared z-score from Cox model: $z.score^2$

Note

This function is primarily used for package development and validation. End users typically don't need to call it directly.

See Also

[df_counting](#) for generating the input object

Examples

```
# After running df_counting
library(survival)
data(veteran)
veteran$treat <- as.numeric(veteran$trt) - 1

result <- df_counting(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat"
)

# Check consistency of test statistics
check_results(result)

# Store results without printing
stats_comparison <- check_results(result, verbose = FALSE)
print(stats_comparison)

# Simple example with constructed data
dfcount_example <- list(
  lr = 2.5,
  sig2_lr = 1.0,
  z.score = 2.5,
  logrank_results = list(chisq = 6.25)
)
check_results(dfcount_example)
```

ci_cox

Confidence interval for Cox model estimate

Description

Calculates the confidence interval for a Cox model log hazard ratio estimate.

Usage

```
ci_cox(bhat, sig_bhat, alpha = 0.05, verbose = FALSE)
```

Arguments

bhat	Estimated log hazard ratio.
sig_bhat	Standard error of the estimate.
alpha	Significance level (default: 0.05).
verbose	Logical; if TRUE, prints interval.

Value

Data frame with log hazard ratio, standard error, hazard ratio, lower and upper confidence limits.

count_weighted	<i>Weighted counting process</i>
----------------	----------------------------------

Description

Computes the weighted count of events up to a specified time.

Usage

```
count_weighted(x, y, w = rep(1, length(y)))
```

Arguments

x	Time point.
y	Vector of event/censoring times.
w	Weights (default 1).

Value

Weighted count of events up to time x.

cox_rhogamma	<i>Weighted Cox Model with Rho-Gamma Weights</i>
--------------	--

Description

Fits a weighted Cox proportional hazards model using flexible time-dependent weights (e.g., Fleming-Harrington, Magirr-Burman). Supports resampling-based inference for variance estimation and bias correction.

Usage

```
cox_rhogamma(
  dfcount,
  scheme = "fh",
  scheme_params = list(rho = 0, gamma = 0.5),
  draws = 0,
  alpha = 0.05,
  verbose = FALSE,
  lr.digits = 4
)
```

Arguments

dfcount	List; output from <code>df_counting</code> containing counting process data including time, delta, z, w_hat, survP_all, survG_all, etc.
scheme	Character; weighting scheme. See <code>wt.rg.S</code> for options. Default: "fh".
scheme_params	List; parameters for the selected scheme. Default: <code>list(rho = 0, gamma = 0.5)</code> . Required parameters depend on scheme: <ul style="list-style-type: none"> • "fh": rho, gamma • "MB": mb_tstar • "custom_time": t.tau, w0.tau, w1.tau
draws	Integer; number of resampling draws for variance estimation and bias correction. If 0, only asymptotic inference is performed. Default: 0.
alpha	Numeric; significance level for confidence intervals. Default: 0.05.
verbose	Logical; whether to print detailed output. Default: FALSE.
lr.digits	Integer; number of decimal places for formatted output. Default: 4.

Details

This function solves the weighted Cox partial likelihood score equation: $U(\beta) = \sum_i w_i K_i (dN_0/Y_0 - dN_1/Y_1) = 0$

where K_i are time-dependent weights and Y_j , dN_j are risk sets and event counts for group j .

When $draws > 0$, the function performs resampling to:

- Estimate finite-sample variance (more accurate than asymptotic)
- Compute bias correction for $\hat{\beta}$
- Provide improved confidence intervals for small samples

The score test at $\beta = 0$ corresponds to the weighted log-rank test.

Value

A list containing:

fit List with fitted model components:

- bhat: Estimated log hazard ratio

- `sig_bhat_asy`: Asymptotic standard error
- `u.zero`: Score statistic at $\beta=0$ (log-rank)
- `z.score`: Standardized score statistic
- `sig2_score`: Variance of score statistic
- `wt_rg`: Vector of time-dependent weights
- `bhat_debiased`: Bias-corrected estimate (if draws > 0)
- `sig_bhat_star`: Resampling-based standard error (if draws > 0)

hr_ci_asy Data frame with asymptotic HR and CI

hr_ci_star Data frame with resampling-based HR and CI (if draws > 0)

cox_text_asy Formatted string with HR and asymptotic CI

cox_text_star Formatted string with HR and resampling CI (if draws > 0)

z.score_debiased Bias-corrected score statistic (if draws > 0)

zlogrank_text Formatted log-rank test result

Note

The treatment variable in `dfcount` must be coded as 0=control, 1=experimental.

References

Magirr, D. and Burman, C. F. (2019). Modestly weighted logrank tests. *Statistics in Medicine*, 38(20), 3782-3790.

See Also

[df_counting](#) for preprocessing `wt_rg.S` for weighting schemes [cox_score_rhogamma](#) for score function

Other survival_analysis: [KM_diff\(\)](#), [df_counting\(\)](#), [wt_rg.S\(\)](#)

Other weighted_tests: [df_counting\(\)](#), [wt_rg.S\(\)](#)

Examples

```
# First get counting process data
library(survival)
str(veteran)
veteran$treat <- as.numeric(veteran$trt) - 1

dfcount <- df_counting(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat"
)

# Fit weighted Cox model with FH(0,0.5) weights
fit <- cox_rhogamma(
  dfcount = dfcount,
```

```

    scheme = "fh",
    scheme_params = list(rho = 0, gamma = 0.5),
    draws = 1000,
    verbose = TRUE
  )

  print(fit$cox_text_star) # Resampling-based CI
  print(fit$zlogrank_text) # Weighted log-rank test

  # Compare asymptotic and resampling CIs
  print(fit$hr_ci_asy)
  print(fit$hr_ci_star)

```

cox_rhogamma_resample *Resampling for Weighted Cox Model (rho, gamma)*

Description

Performs resampling to estimate uncertainty for the weighted Cox model (rho, gamma).

Usage

```

cox_rhogamma_resample(
  fit_rhogamma,
  i_bhat,
  K_wt_rg,
  i_zero,
  K_zero,
  G1.draws = NULL,
  G0.draws = NULL,
  draws = 100,
  seedstart = 8316951
)

```

Arguments

fit_rhogamma	List with fitted Cox model results.
i_bhat	Information at estimated beta.
K_wt_rg	Weights at estimated beta.
i_zero	Information at beta=0.
K_zero	Weights at beta=0.
G1.draws	Optional: pre-generated random draws for groups.
G0.draws	Optional: pre-generated random draws for groups.
draws	Number of resampling iterations (default: 100).
seedstart	Random seed for reproducibility (default: 8316951).

Value

List with resampling results (score, beta, standard error, etc.).

cox_score_rhogamma	<i>Cox score with rho-gamma weights</i>
--------------------	---

Description

Calculates the Cox score statistic for weighted log-rank tests.

Usage

```
cox_score_rhogamma(
  beta,
  time,
  delta,
  z,
  w_hat = rep(1, length(time)),
  wt_rg = rep(1, length(time)),
  score_only = TRUE
)
```

Arguments

beta	Log hazard ratio parameter.
time	Numeric vector of event times.
delta	Numeric vector of event indicators.
z	Numeric vector of group indicators.
w_hat	Numeric vector of weights.
wt_rg	Numeric vector of rho-gamma weights.
score_only	Logical; if TRUE, returns only the score.

Value

Numeric value of the score or a list with additional results.

create_baseline_table *Create Baseline Characteristics Table by Treatment Arm*

Description

This function generates publication-ready baseline characteristic tables commonly used in clinical trials and observational studies. It calculates summary statistics, p-values, and standardized mean differences for continuous, categorical, and binary variables.

Usage

```
create_baseline_table(  
  data,  
  treat_var = "treat",  
  vars_continuous = NULL,  
  vars_categorical = NULL,  
  vars_binary = NULL,  
  var_labels = NULL,  
  digits = 1,  
  show_pvalue = TRUE,  
  show_smd = TRUE,  
  show_missing = TRUE  
)
```

Arguments

data	Data frame containing baseline variables
treat_var	Name of treatment variable (default: "treat")
vars_continuous	Character vector of continuous variable names
vars_categorical	Character vector of categorical variable names
vars_binary	Character vector of binary variable names
var_labels	Named vector for variable labels (e.g., c(age = "Age (years)"))
digits	Number of decimal places for continuous variables (default: 1)
show_pvalue	Logical, whether to show p-values (default: TRUE)
show_smd	Logical, whether to show standardized mean differences (default: TRUE)
show_missing	Logical, whether to show missing data counts (default: TRUE)

Value

A gt table object (if gt package is available) or data frame

Examples

```
# Create sample data
set.seed(123)
n <- 500
sample_data <- data.frame(
  treat = rbinom(n, 1, 0.5),
  age = rnorm(n, mean = 55, sd = 10),
  stage = sample(c("I", "II", "III", "IV"), n, replace = TRUE),
  sex = rbinom(n, 1, 0.45),
  smoking = rbinom(n, 1, 0.3)
)

# Create table
table <- create_baseline_table(
  data = sample_data,
  treat_var = "treat",
  vars_continuous = "age",
  vars_categorical = "stage",
  vars_binary = c("sex", "smoking"),
  var_labels = c(
    age = "Age (years)",
    stage = "Disease Stage",
    sex = "Female",
    smoking = "Current Smoker"
  )
)
```

cumulative_rmst_bands *Cumulative RMST bands for survival curves*

Description

Calculates cumulative Restricted Mean Survival Time (RMST) and confidence bands for survival curves using resampling. Optionally plots the cumulative RMST curve, pointwise confidence intervals, and simultaneous confidence bands.

Usage

```
cumulative_rmst_bands(
  df,
  fit,
  tte.name,
  event.name,
  treat.name,
  weight.name = NULL,
  draws_sb = 1000,
```

```

xlab = "months",
ylim_pad = 0.5,
rmst_max_legend = "left",
rmst_max_cex = 0.7,
plot = TRUE
)

```

Arguments

<code>df</code>	Data frame containing survival data.
<code>fit</code>	Survival fit object (output from <code>KM_diff</code>).
<code>tte.name</code>	Name of time-to-event variable in <code>df</code> .
<code>event.name</code>	Name of event indicator variable in <code>df</code> .
<code>treat.name</code>	Name of treatment group variable in <code>df</code> .
<code>weight.name</code>	Optional name of weights variable in <code>df</code> .
<code>draws_sb</code>	Number of resampling draws for simultaneous bands (default: 1000).
<code>xlab</code>	Label for x-axis (default: "months").
<code>ylim_pad</code>	Padding for y-axis limits (default: 0.5).
<code>rmst_max_legend</code>	Position for RMST legend (default: "left").
<code>rmst_max_cex</code>	Text size for RMST legend (default: 0.7).
<code>plot</code>	Logical; if TRUE, plot the results. Default is TRUE.

Value

A list with elements:

<code>at_points</code>	Time points used for RMST calculation
<code>rmst_time</code>	Cumulative RMST estimates
<code>sig2_rmst_time</code>	Variance of RMST estimates
<code>rmst_time_lower</code>	Pointwise lower confidence interval
<code>rmst_time_upper</code>	Pointwise upper confidence interval
<code>rmst_maxtau_ci</code>	RMST and CI at maximum time
<code>rmst_text</code>	Text summary for legend
<code>c_alpha_band</code>	Critical value for simultaneous band
<code>rmst_time_sb_lower</code>	Simultaneous band lower bound
<code>rmst_time_sb_upper</code>	Simultaneous band upper bound

Description

Performs comprehensive weighted and/or stratified survival analysis, including Cox proportional hazards model, logrank/Fleming-Harrington tests, and calculation of risk/event sets, Kaplan-Meier curves, quantiles, and variance estimates.

Usage

```
df_counting(  
  df,  
  tte.name = "tte",  
  event.name = "event",  
  treat.name = "treat",  
  weight.name = NULL,  
  strata.name = NULL,  
  arms = c("treat", "control"),  
  time.zero = 0,  
  tpoints.add = c(0),  
  by.risk = 6,  
  time.zero.label = 0,  
  risk.add = NULL,  
  get.cox = TRUE,  
  cox.digits = 2,  
  lr.digits = 2,  
  cox.eps = 0.001,  
  lr.eps = 0.001,  
  verbose = FALSE,  
  qprob = 0.5,  
  scheme = "fh",  
  scheme_params = list(rho = 0, gamma = 0),  
  conf_level = 0.95,  
  check.KM = TRUE,  
  check.seKM = FALSE,  
  draws = 0,  
  seedstart = 8316951,  
  stop.onerror = FALSE,  
  censoring_allmarks = TRUE  
)
```

Arguments

df	Data frame containing survival data.
tte.name	Character; name of the time-to-event column in df.

event.name	Character; name of the event indicator column in df (1=event, 0=censored).
treat.name	Character; name of the treatment/group column in df (must be coded as 0=control, 1=experimental).
weight.name	Character or NULL; name of the weights column in df. If NULL, equal weights are used.
strata.name	Character or NULL; name of the strata column in df for stratified analysis.
arms	Character vector of length 2; group labels. Default: c("treat", "control").
time.zero	Numeric; time value to use as zero. Default: 0.
tpoints.add	Numeric vector; additional time points to include in calculations. Default: c(0).
by.risk	Numeric; interval for risk set time points. Default: 6.
time.zero.label	Numeric; label for time zero in output. Default: 0.0.
risk.add	Numeric vector or NULL; additional specific risk points to include.
get.cox	Logical; whether to fit Cox proportional hazards model. Default: TRUE.
cox.digits	Integer; number of decimal places for Cox output formatting. Default: 2.
lr.digits	Integer; number of decimal places for logrank output formatting. Default: 2.
cox.eps	Numeric; threshold for Cox p-value formatting (values below shown as "<eps"). Default: 0.001.
lr.eps	Numeric; threshold for logrank p-value formatting. Default: 0.001.
verbose	Logical; whether to print warnings and diagnostic messages. Default: FALSE.
qprob	Numeric in (0,1); quantile probability for KM quantile table. Default: 0.5 (median).
scheme	Character; weighting scheme for logrank/Fleming-Harrington test. Options: "fh", "schemper", "XO", "MB", "custom_time", "fh_exp1", "fh_exp2". Default: "fh".
scheme_params	List; parameters for the selected weighting scheme. Default: list(rho = 0, gamma = 0). <ul style="list-style-type: none"> • For "fh": rho and gamma (Fleming-Harrington parameters) • For "MB": mb_tstar (cutoff time) • For "custom_time": t.tau, w0.tau, w1.tau
conf_level	Numeric in (0,1); confidence level for quantile intervals. Default: 0.95.
check.KM	Logical; whether to check KM curve validity against survival::survfit. Default: TRUE.
check.seKM	Logical; whether to check KM standard error estimates. Default: FALSE.
draws	Integer; number of draws for resampling-based variance estimation. Default: 0 (no resampling).
seedstart	Integer; random seed for reproducible resampling. Default: 8316951.
stop.onerror	Logical; whether to stop execution on errors (TRUE) or issue warnings (FALSE). Default: FALSE.
censoring_allmarks	Logical; if FALSE, removes events from censored time points. Default: TRUE.

Details

This function implements a comprehensive survival analysis framework supporting:

- Weighted observations via `weight.name`
- Stratified analysis via `strata.name`
- Multiple weighting schemes for log-rank tests
- Resampling-based variance estimation
- Automatic validation against survival package results

The function performs time-fixing using `survival::aeqSurv` to handle tied event times. For stratified analyses, stratum-specific estimates are computed and combined appropriately.

Value

A list with the following components:

cox_results List with Cox model results including hazard ratio, confidence interval, p-value, and formatted text

logrank_results List with log-rank test chi-square statistic, p-value, and formatted text

z.score Standardized weighted log-rank test statistic

at_points Vector of all time points used in calculations

surv0, surv1 Kaplan-Meier survival estimates for control and treatment groups

sig2_surv0, sig2_surv1 Variance estimates for survival curves

survP Pooled survival estimates

survG Censoring distribution estimates

quantile_results Data frame with median survival and confidence intervals by group

lr, sig2_lr Weighted log-rank statistic and its variance

riskpoints0, riskpoints1 Risk set counts at specified time points

z.score_stratified Stratified z-score (if stratified analysis)

Weighting Schemes

fh Fleming-Harrington: $w(t) = S(t)^\rho * (1-S(t))^\gamma$

MB Magirr-Burman: $w(t) = 1/\max(S(t), S(t^*))$

schemper Schemper: $w(t) = S(t)/G(t)$ where G is the censoring distribution

XO Xu-O'Quigley: $w(t) = S(t)/Y(t)$ where Y is risk set size

References

Fleming, T. R. and Harrington, D. P. (1991). Counting Processes and Survival Analysis. Wiley.

Magirr, D. and Burman, C. F. (2019). Modestly weighted logrank tests. *Statistics in Medicine*, 38(20), 3782-3790.

See Also

[coxph](#), [survdif](#), [survfit](#) [cox_rhogamma](#) for weighted Cox models [KM_diff](#) for Kaplan-Meier differences

Other survival_analysis: [KM_diff\(\)](#), [cox_rhogamma\(\)](#), [wt.rg.S\(\)](#)

Other weighted_tests: [cox_rhogamma\(\)](#), [wt.rg.S\(\)](#)

Examples

```
# Basic survival analysis
library(survival)
str(veteran)
veteran$treat <- as.numeric(veteran$strtrt) - 1

result <- df_counting(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat",
  arms = c("Treatment", "Control")
)

# Print results
print(result$cox_results$cox_text)
print(result$zlogrank_text)

# Fleming-Harrington (0,1) weights (emphasizing late differences)
result_fh <- df_counting(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat",
  scheme = "fh",
  scheme_params = list(rho = 0, gamma = 1)
)

# Stratified analysis
result_strat <- df_counting(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat",
  strata.name = "celltype"
)
```

extract_and_calc_weights

Extract and calculate weights for multiple schemes

Description

Extracts and calculates weights for multiple schemes and returns a combined data frame.

Usage

```
extract_and_calc_weights(atpoints, S.pool, weights_spec_list)
```

Arguments

atpoints	Numeric vector of time points.
S.pool	Numeric vector of pooled survival probabilities.
weights_spec_list	List of weighting scheme specifications.

Value

Data frame with weights for each scheme.

extract_group_data	<i>Extract time, event, and weight data for a group</i>
--------------------	---

Description

Extracts time, event, and weight vectors for a specified group.

Usage

```
extract_group_data(time, delta, wgt, z, group = 1)
```

Arguments

time	Numeric vector of times.
delta	Numeric vector of event indicators (1=event, 0=censored).
wgt	Numeric vector of weights.
z	Numeric vector of group indicators.
group	Value of group to extract (default 1).

Value

List with U (times), D (events), W (weights).

find_cox_root	<i>Root-finding for Cox score function</i>
---------------	--

Description

Finds the root of the Cox model score equation for weighted log-rank statistics.

Usage

```
find_cox_root(time, delta, z, w_hat, wt_rg)
```

Arguments

time	Numeric vector of event times.
delta	Numeric vector of event indicators.
z	Numeric vector of group indicators.
w_hat	Numeric vector of weights.
wt_rg	Numeric vector of rho-gamma weights.

Value

List with root and additional information, or NA if not found.

format_pval	<i>Format p-value for display</i>
-------------	-----------------------------------

Description

Formats a p-value for display, showing "<eps" for small values.

Usage

```
format_pval(pval, eps = 0.001, digits = 3)
```

Arguments

pval	Numeric p-value.
eps	Threshold for small p-values.
digits	Number of digits to display.

Value

Formatted p-value as character.

 get_censoring_and_events

Get censoring and event times and their indices

Description

Extracts censoring and event times and their indices for a group at specified time points.

Usage

```
get_censoring_and_events(time, delta, z, group, censoring_allmarks, at_points)
```

Arguments

time	Numeric vector of times.
delta	Numeric vector of event indicators.
z	Numeric vector of group indicators.
group	Value of group to extract.
censoring_allmarks	Logical; if FALSE, remove events from censored.
at_points	Numeric vector of time points.

Value

List with cens (censored times), ev (event times), idx_cens, idx_ev, idx_ev_full.

 get_dfcounting

Get df_counting results

Description

Wrapper for df_counting, safely executes and returns results for survival analysis.

Usage

```
get_dfcounting(...)
```

Arguments

... Arguments passed to [df_counting](#).

Value

Result from df_counting or NULL if error.

See Also[df_counting](#)

`get_event_risk_matrices`*Event and Risk Matrices for Survival Analysis*

Description

Constructs matrices indicating event and risk status for each subject at specified time points.

Usage

```
get_event_risk_matrices(U, at.points)
```

Arguments

`U` Vector of observed times (e.g., time-to-event).
`at.points` Vector of time points at which to evaluate events and risk.

Value

A list with event and risk matrices.

`get_riskpoints`*Get risk set counts at specified risk points*

Description

Returns risk set counts at specified risk points.

Usage

```
get_riskpoints(ybar, risk_points, at_points)
```

Arguments

`ybar` Numeric vector of risk set counts.
`risk_points` Numeric vector of risk points.
`at_points` Numeric vector of time points.

Value

Numeric vector of risk set counts at risk points.

get_validated_weights *Get validated weights for a data frame*

Description

Validates and returns weights for a data frame according to specified schemes.

Usage

```
get_validated_weights(
  df_weights,
  scheme = "fh",
  scheme_params = list(rho = 0, gamma = 0),
  details = FALSE
)
```

Arguments

df_weights	Data frame containing weights and related data.
scheme	Character string specifying the weighting scheme.
scheme_params	List of parameters for the scheme.
details	Logical; if TRUE, returns detailed output.

Value

Numeric vector or list of validated weights.

get_weights *Get weights for a weighting scheme*

Description

Calculates weights for a specified scheme at given time points.

Usage

```
get_weights(scheme, scheme_params, S.pool, tpoints)
```

Arguments

scheme	Character string specifying the weighting scheme.
scheme_params	List of parameters for the scheme.
S.pool	Numeric vector of pooled survival probabilities.
tpoints	Numeric vector of time points.

Value

Numeric vector of weights.

 KM_diff

Kaplan-Meier Difference Between Groups

Description

Calculates the difference in Kaplan-Meier curves between two groups, with confidence intervals and optional resampling-based simultaneous confidence bands.

Usage

```

KM_diff(
  df,
  tte.name = "tte",
  event.name = "event",
  treat.name = "treat",
  weight.name = NULL,
  at_points = sort(df[[tte.name]]),
  alpha = 0.05,
  seedstart = 8316951,
  draws = 0,
  risk.points = NULL,
  draws.band = 0,
  tau.seq = 0.25,
  qtau = 0.025,
  show_resamples = TRUE,
  modify_tau = FALSE
)

```

Arguments

df	Data frame containing survival data.
tte.name	Character; name of time-to-event variable in df.
event.name	Character; name of event indicator variable in df (1=event, 0=censored).
treat.name	Character; name of treatment group variable in df (0=control, 1=treatment).
weight.name	Character or NULL; name of weights variable in df.
at_points	Numeric vector; time points for calculation. Default: sorted unique event times.
alpha	Numeric; significance level for confidence intervals. Default: 0.05.
seedstart	Integer; random seed for reproducibility. Default: 8316951.
draws	Integer; number of draws for pointwise variance estimation. Default: 0.
risk.points	Numeric vector; time points for risk table display.

<code>draws.band</code>	Integer; number of draws for simultaneous confidence bands. Default: 0.
<code>tau.seq</code>	Numeric; step size for tau sequence when <code>modify_tau=TRUE</code> . Default: 0.25.
<code>qtau</code>	Numeric; quantile for tau range restriction. Default: 0.025.
<code>show_resamples</code>	Logical; whether to plot resampled curves. Default: TRUE.
<code>modify_tau</code>	Logical; whether to restrict time range for simultaneous bands. Default: FALSE.

Details

This function computes the difference in Kaplan-Meier survival curves, $\delta(t) = S_1(t) - S_0(t)$, along with variance estimates and confidence intervals.

When `draws.band > 0`, simultaneous confidence bands are constructed using the supremum distribution of the standardized difference process. These bands maintain the specified coverage probability across all time points simultaneously.

The variance is estimated using Greenwood's formula for unweighted data, or resampling-based methods when `draws > 0`.

Value

A list containing:

- at_points** Time points used in calculations
- surv0, surv1** Survival estimates for control and treatment groups
- sig2_surv0, sig2_surv1** Variance estimates for survival curves
- dhat** Survival difference ($S_1 - S_0$) at each time point
- sig2_dhat** Variance of survival difference
- lower, upper** Pointwise confidence limits ($1 - \alpha/2$)
- sb_lower, sb_upper** Simultaneous band limits (if `draws.band > 0`)
- c_alpha_band** Critical value for simultaneous band (if `draws.band > 0`)
- dhat_star** Matrix of resampled differences (if `draws.band > 0`)
- Zdhat_star** Standardized resampled differences (if `draws.band > 0`)

Confidence Intervals vs Bands

- Pointwise CIs (`lower, upper`): Cover the true difference at each time point with probability $1 - \alpha$
- Simultaneous bands (`sb_lower, sb_upper`): Cover the entire difference curve with probability $1 - \alpha$

Note

Treatment must be coded as 0=control, 1=experimental. Event must be binary (0/1).

See Also

[df_counting](#) for full survival analysis [plotKM.band_subgroups](#) for visualization [cumulative_rmst_bands](#) for RMST analysis

Other survival_analysis: [cox_rhogamma\(\)](#), [df_counting\(\)](#), [wt.rg.S\(\)](#)

Examples

```
library(survival)
str(veteran)
veteran$treat <- as.numeric(veteran$trt) - 1

# Basic KM difference
result <- KM_diff(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat"
)

# Plot the difference
plot(result$at_points, result$dhat, type = "s",
      xlab = "Time", ylab = "Survival Difference")

# With simultaneous confidence bands
result_band <- KM_diff(
  df = veteran,
  tte.name = "time",
  event.name = "status",
  treat.name = "treat",
  draws.band = 1000,
  modify_tau = TRUE
)
```

KM_estimates

Kaplan-Meier Survival Estimates and Variance

Description

Computes Kaplan-Meier survival estimates and their variances given risk and event counts.

Usage

```
KM_estimates(ybar, nbar, sig2w_multiplier = NULL)
```

Arguments

ybar Vector of risk set sizes at each time point.
nbar Vector of event counts at each time point.
sig2w_multiplier Optional vector for variance calculation. If NULL, calculated internally.

Value

List with survival estimates and variances.

KM_plot_2sample_weighted_counting

Plot Weighted Kaplan-Meier Curves for Two Samples (Counting Process Format)

Description

Plots Kaplan-Meier survival curves for two groups using precomputed risk/event counts and survival estimates. Optionally displays confidence intervals, risk tables, median survival annotations, and statistical test results.

Usage

```
KM_plot_2sample_weighted_counting(
  dfcount,
  show.cox = TRUE,
  cox.cex = 0.725,
  show.logrank = FALSE,
  logrank.cex = 0.725,
  cox.eps = 0.001,
  lr.eps = 0.001,
  show_arm_legend = TRUE,
  arms = c("treat", "control"),
  put.legend.arms = "left",
  stop.onerror = TRUE,
  check.KM = TRUE,
  put.legend.cox = "topright",
  put.legend.lr = "topleft",
  lr.digits = 2,
  cox.digits = 2,
  tpoints.add = c(0),
  by.risk = NULL,
  Xlab = "time",
  Ylab = "proportion surviving",
  col.0 = "black",
  col.1 = "blue",
```

```

show.med = TRUE,
med.digits = 2,
med.font = 4,
conf.int = FALSE,
conf.level = 0.95,
choose.ylim = FALSE,
arm.cex = 0.7,
quant = 0.5,
med.cex = 0.725,
ymed.offset = 0.1,
xmed.fraction = 0.8,
risk.cex = 0.725,
ltys = c(1, 1),
lwds = c(1, 1),
censor.mark.all = TRUE,
censor.cex = 0.5,
show.ticks = TRUE,
risk.set = TRUE,
ymin = 0,
ymax = 1,
ymin.del = 0.035,
ymin2 = NULL,
risk_offset = 0.125,
risk_delta = 0.05,
y.risk0 = NULL,
show.Y.axis = TRUE,
cex_Yaxis = 1,
y.risk1 = NULL,
add.segment = FALSE,
risk.add = NULL,
xmin = 0,
xmax = NULL,
x.truncate = NULL,
time.zero = 0,
prob.points = NULL
)

```

Arguments

<code>dfcount</code>	List containing precomputed survival data.
<code>show.cox</code>	Logical; show Cox model results.
<code>cox.cex</code>	Numeric; text size for Cox annotation.
<code>show.logrank</code>	Logical; show log-rank test results.
<code>logrank.cex</code>	Numeric; text size for log-rank annotation.
<code>cox.eps</code>	Numeric; small values for Cox calculations.
<code>lr.eps</code>	Numeric; small values for log-rank calculations.

show_arm_legend	Logical; show arm legend.
arms	Character vector of arm labels.
put.legend.arms	Character; legend positions.
stop.onerror	Logical; stop on KM curve errors.
check.KM	Logical; check KM curve validity.
put.legend.cox	Character; legend positions.
put.legend.lr	Character; legend positions.
lr.digits	Integer; digits for test results.
cox.digits	Integer; digits for test results.
tpoints.add	Numeric vector; additional time points to include (default: c(0)).
by.risk	Numeric; interval for risk table time points.
Xlab	Character; axis labels.
Ylab	Character; axis labels.
col.0	Color for control curve.
col.1	Color for treatment curve.
show.med	Logical; annotate median survival.
med.digits	Median annotation settings.
med.font	Median annotation settings.
conf.int	Logical; plot confidence intervals.
conf_level	Numeric; confidence level for intervals.
choose_ylim	Logical; auto-select y-axis limits.
arm.cex	Numeric; text size for arm legend.
quant	Numeric; quantile for annotation.
med.cex	Median annotation settings.
ymed.offset	Median annotation settings.
xmed.fraction	Median annotation settings.
risk.cex	Numeric; text size for risk table.
ltys	Integer; line types for curves.
lwds	Integer; line widths for curves.
censor.mark.all	Logical; mark all censored times.
censor.cex	Numeric; size of censor marks.
show.ticks	Logical; show axis ticks.
risk.set	Logical; display risk table.
ymin	Additional graphical and calculation parameters.
ymax	Additional graphical and calculation parameters.

ymin.del	Additional graphical and calculation parameters.
ymin2	Additional graphical and calculation parameters.
risk_offset	Additional graphical and calculation parameters.
risk_delta	Additional graphical and calculation parameters.
y.risk0	Additional graphical and calculation parameters.
show.Y.axis	Additional graphical and calculation parameters.
cex.Yaxis	Additional graphical and calculation parameters.
y.risk1	Additional graphical and calculation parameters.
add.segment	Additional graphical and calculation parameters.
risk.add	Additional graphical and calculation parameters.
xmin	Additional graphical and calculation parameters.
xmax	Additional graphical and calculation parameters.
x.truncate	Additional graphical and calculation parameters.
time.zero	Numeric; time zero value for risk table alignment (default: 0).
prob.points	Numeric vector; probability points for additional annotations (default: NULL).

Value

Invisibly returns NULL. Used for plotting side effects.

km_quantile	<i>Kaplan-Meier quantile and confidence interval</i>
-------------	--

Description

Calculates the quantile and confidence interval for a Kaplan-Meier curve.

Usage

```
km_quantile(
  time_points,
  survival_probs,
  se_probs = NULL,
  qprob = 0.5,
  type = c("midpoint", "min"),
  conf_level = 0.95
)
```

Arguments

time_points	Vector of time points.
survival_probs	Vector of survival probabilities.
se_probs	Standard errors of survival probabilities.
qprob	Quantile probability (default 0.5).
type	Calculation type (midpoint or min).
conf_level	Confidence level (default 0.95).

Value

List with quantile and confidence interval.

km_quantile_table	<i>Table of KM quantiles for two groups</i>
-------------------	---

Description

Returns a data frame of quantiles and confidence intervals for two groups.

Usage

```
km_quantile_table(
  time_points,
  surv0,
  se0,
  surv1,
  se1,
  arms = c("treat", "control"),
  qprob = 0.5,
  type = c("midpoint", "min"),
  conf_level = 0.95
)
```

Arguments

time_points	Vector of time points.
surv0	Survival probabilities for group 0.
se0	Standard errors for group 0.
surv1	Survival probabilities for group 1.
se1	Standard errors for group 1.
arms	Group labels.
qprob	Quantile probability.
type	Calculation type.
conf_level	Confidence level.

Value

Data frame of quantiles and CIs for each group.

kmq_calculations	<i>Kaplan-Meier quantile calculation</i>
------------------	--

Description

Calculates the quantile time for a Kaplan-Meier curve.

Usage

```
kmq_calculations(time_points, survival_probs, qprob = 0.5, type = "midpoint")
```

Arguments

time_points	Vector of time points.
survival_probs	Vector of survival probabilities.
qprob	Quantile probability (default 0.5).
type	Calculation type (midpoint or min).

Value

Estimated quantile time.

N_rhogamma	<i>Weighted event count for Cox model</i>
------------	---

Description

Calculates the weighted event count for the Cox model with rho-gamma weights.

Usage

```
N_rhogamma(x, error, delta, weight = 1)
```

Arguments

x	Numeric vector of time points.
error	Numeric vector of error terms.
delta	Numeric vector of event indicators.
weight	Numeric vector of weights (default: 1).

Value

Numeric value of weighted event count.

plot_km	<i>Plot Kaplan-Meier curves</i>
---------	---------------------------------

Description

Plots Kaplan-Meier survival curves for groups in the data.

Usage

```
plot_km(df, tte.name, event.name, treat.name, weights = NULL, ...)
```

Arguments

df	Data frame containing survival data.
tte.name	Name of time-to-event column.
event.name	Name of event indicator column.
treat.name	Name of treatment/group column.
weights	Optional; name of weights column.
...	Additional arguments passed to plot().

Value

Kaplan-Meier fit object (invisible).

plot_km_confint_polygon	<i>Plot confidence interval polygon for KM curve</i>
-------------------------	--

Description

Plots a shaded polygon representing the confidence interval for a Kaplan-Meier survival curve.

Usage

```
plot_km_confint_polygon(x, surv, se, conf_level, col)
```

Arguments

x	Numeric vector of time points.
surv	Numeric vector of survival probabilities.
se	Numeric vector of standard errors of survival probabilities.
conf_level	Numeric; confidence level for interval (default 0.95).
col	Color for the polygon.

Value

Invisibly returns NULL. Used for plotting side effects.

plot_km_curves_counting

Plot KM curves for two groups with optional confidence intervals and censoring marks

Description

Plots Kaplan-Meier survival curves for two groups, with options for confidence intervals and censoring marks.

Usage

```
plot_km_curves_counting(  
  at_points,  
  S0.KM,  
  idx0,  
  idv0,  
  S1.KM,  
  idx1,  
  idv1,  
  col.0,  
  col.1,  
  ltys,  
  lwds,  
  Xlab,  
  Ylab,  
  ylim,  
  xlim,  
  show.ticks = FALSE,  
  cens0 = NULL,  
  risk.points,  
  risk.points.label,  
  cens1 = NULL,  
  se0.KM = NULL,  
  se1.KM = NULL,  
  conf.int = FALSE,  
  conf_level = 0.95,  
  censor.cex = 1,  
  time.zero = 0,  
  tpoints.add = c(0),  
  ...  
)
```

Arguments

at_points	Numeric vector of time points for plotting.
S0.KM	Numeric vector of survival probabilities for group 0.
idx0	Indices for censoring in group 0.
idv0	Indices for events in group 0.
S1.KM	Numeric vector of survival probabilities for group 1.
idx1	Indices for censoring in group 1.
idv1	Indices for events in group 1.
col.0	Color for group 0.
col.1	Color for group 1.
ltys	Line types for groups.
lwds	Line widths for groups.
Xlab	X-axis label.
Ylab	Y-axis label.
ylim	Y-axis limits.
xlim	X-axis limits.
show.ticks	Logical; show censoring marks (default FALSE).
cens0	Numeric vector of censoring times for group 0.
risk.points	Numeric vector of risk time points.
risk.points.label	Character vector of labels for risk time points.
cens1	Numeric vector of censoring times for group 1.
se0.KM	Numeric vector of standard errors for group 0.
se1.KM	Numeric vector of standard errors for group 1.
conf.int	Logical; show confidence intervals (default FALSE).
conf_level	Confidence level (default 0.95).
sensor.cex	Numeric; censoring mark size (default 1.0).
time.zero	Numeric; time zero value for risk table alignment (default 0).
tpoints.add	Numeric vector; additional time points to include (default c(0)).
...	Additional arguments to plot.

Value

Invisibly returns NULL. Used for plotting side effects.

plot_weight_schemes *Plot weight schemes for survival analysis*

Description

Plots the weights for different schemes over time using ggplot2.

Usage

```
plot_weight_schemes(
  dfcount,
  tte.name = "time_months",
  event.name = "status",
  treat.name = "treat",
  arms = c("treat", "control"),
  weights_spec_list = list(`MB(12)` = list(scheme = "MB", mb_tstar = 12), `MB(6)` =
    list(scheme = "MB", mb_tstar = 6), `FH(0,1)` = list(scheme = "fh", rho = 0, gamma =
      1), `FH(0.5,0.5)` = list(scheme = "fh", rho = 0.5, gamma = 0.5), custom_time =
      list(scheme = "custom_time", t.tau = 25, w0.tau = 1, w1.tau = 1.5), FHexp2 =
      list(scheme = "fh_exp2")),
  custom_colors = c(`FH(0,1)` = "grey", FHexp2 = "black", `MB(12)` = "#1b9e77", `MB(6)` =
    "#d95f02", `FH(0.5,0.5)` = "#7570b3", custom_time = "green"),
  custom_sizes = c(`FH(0,1)` = 2, FHexp2 = 1, `MB(12)` = 1, `MB(6)` = 1, `FH(0.5,0.5)` =
    1, custom_time = 1),
  transform_fh = FALSE,
  rescheme_fhexp2 = TRUE
)
```

Arguments

dfcount	Data frame containing counting process results, including time points and survival probabilities.
tte.name	Name of the time-to-event variable (default: 'time_months').
event.name	Name of the event indicator variable (default: 'status').
treat.name	Name of the treatment group variable (default: 'treat').
arms	Character vector of group names (default: c('treat', 'control')).
weights_spec_list	List of weighting scheme specifications.
custom_colors	Named character vector of colors for each scheme.
custom_sizes	Named numeric vector of line sizes for each scheme.
transform_fh	Logical; whether to transform FH weights (default: FALSE).
rescheme_fhexp2	Logical; whether to rescheme FHexp2 and custom_time (default: TRUE).

Details

This function visualizes the weights used in various survival analysis schemes (e.g., FH, MB, custom) using ggplot2. Facets and colors are customizable.

Value

A ggplot object showing the weight schemes over time.

plot_weighted_km	<i>Plot weighted Kaplan-Meier curves</i>
------------------	--

Description

Plots weighted Kaplan-Meier curves using a custom function.

Usage

```
plot_weighted_km(dfcount, ...)
```

Arguments

dfcount	Result object from df_counting.
...	Additional arguments passed to KM_plot_2sample_weighted_counting.

Value

None. Plots the curves.

plotKM.band_subgroups	<i>Plot Kaplan-Meier Survival Difference Curves with Subgroups and Confidence Bands</i>
-----------------------	---

Description

Plots the difference in Kaplan-Meier survival curves between two groups, optionally including simultaneous confidence bands and subgroup curves. Also displays risk tables for the overall population and specified subgroups.

Usage

```
plotKM.band_subgroups(  
  df,  
  tte.name,  
  event.name,  
  treat.name,  
  weight.name = NULL,  
  sg_labels = NULL,  
  ltype = "s",  
  lty = 1,  
  draws = 20,  
  lwd = 2,  
  sg_colors = NULL,  
  color = "lightgrey",  
  ymax.pad = 0.01,  
  ymin.pad = -0.01,  
  taus = c(-Inf, Inf),  
  yseq_length = 5,  
  cex_Yaxis = 0.8,  
  risk_cex = 0.8,  
  by.risk = 6,  
  risk.add = NULL,  
  xmax = NULL,  
  ymin = NULL,  
  ymax = NULL,  
  ymin.del = 0.035,  
  y.risk1 = NULL,  
  y.risk2 = NULL,  
  ymin2 = NULL,  
  risk_offset = NULL,  
  risk.pad = 0.01,  
  risk_delta = 0.0275,  
  tau_add = NULL,  
  time.zero.pad = 0,  
  time.zero.label = 0,  
  xlabel = NULL,  
  ylabel = NULL,  
  Maxtau = NULL,  
  seedstart = 8316951,  
  ylim = NULL,  
  draws.band = 20,  
  qtau = 0.025,  
  show_resamples = FALSE,  
  modify_tau = FALSE  
)
```

Arguments

<code>df</code>	Data frame containing survival data.
<code>tte.name</code>	Name of the time-to-event column.
<code>event.name</code>	Name of the event indicator column (0/1).
<code>treat.name</code>	Name of the treatment group column (0/1).
<code>weight.name</code>	Optional name of the weights column.
<code>sg_labels</code>	Character vector of subgroup definitions (as logical expressions).
<code>ltype</code>	Line type for curves (default: "s").
<code>lty</code>	Line style for curves (default: 1).
<code>draws</code>	Number of draws for resampling (default: 20).
<code>lwd</code>	Line width for curves (default: 2).
<code>sg_colors</code>	Colors for subgroup curves.
<code>color</code>	Color for confidence band polygon (default: "lightgrey").
<code>ymin.pad</code>	Padding for y-axis limits.
<code>ymin.pad</code>	Padding for y-axis limits.
<code>taus</code>	Vector for time truncation (default: c(-Inf, Inf)).
<code>yseq_length</code>	Number of y-axis ticks (default: 5).
<code>cex_Yaxis</code>	Text size for axis.
<code>risk_cex</code>	Text size for risk table.
<code>by.risk</code>	Interval for risk table time points (default: 6).
<code>risk.add</code>	Additional time points for risk table.
<code>xmax</code>	Additional graphical and calculation parameters.
<code>ymin</code>	Additional graphical and calculation parameters.
<code>ymin</code>	Additional graphical and calculation parameters.
<code>ymin.del</code>	Additional graphical and calculation parameters.
<code>y.risk1</code>	Additional graphical and calculation parameters.
<code>y.risk2</code>	Additional graphical and calculation parameters.
<code>ymin2</code>	Additional graphical and calculation parameters.
<code>risk_offset</code>	Additional graphical and calculation parameters.
<code>risk.pad</code>	Additional graphical and calculation parameters.
<code>risk_delta</code>	Additional graphical and calculation parameters.
<code>tau_add</code>	Additional graphical and calculation parameters.
<code>time.zero.pad</code>	Additional graphical and calculation parameters.
<code>time.zero.label</code>	Additional graphical and calculation parameters.
<code>xlabel</code>	Additional graphical and calculation parameters.
<code>ylabel</code>	Additional graphical and calculation parameters.

Maxtau	Additional graphical and calculation parameters.
seedstart	Additional graphical and calculation parameters.
ylim	Additional graphical and calculation parameters.
draws.band	Number of draws for simultaneous confidence bands (default: 20).
qtau	Quantile for time range in simultaneous bands (default: 0.025).
show_resamples	Logical; whether to plot resampled curves (default: FALSE).
modify_tau	Logical; restrict time range for bands.

Value

(Invisible) list containing KM_diff results, time points, subgroup curves, risk tables, and confidence intervals.

resampling_survival *Resampling Survival Curves for Confidence Bands*

Description

Performs resampling to generate survival curves for a group, used for constructing confidence bands.

Usage

```
resampling_survival(U, W, D, at.points, draws.band, surv, G_draws)
```

Arguments

U	Vector of observed times.
W	Vector of weights.
D	Vector of event indicators (0/1).
at.points	Vector of time points for evaluation.
draws.band	Number of resampling draws.
surv	Vector of survival estimates.
G_draws	Matrix of random draws for resampling.

Value

Matrix of resampled survival curves.

risk_weighted	<i>Weighted risk set</i>
---------------	--------------------------

Description

Computes the weighted number at risk at a specified time.

Usage

```
risk_weighted(x, y, w = rep(1, length(y)))
```

Arguments

x	Time point.
y	Vector of event/censoring times.
w	Weights (default 1).

Value

Weighted number at risk at time x.

safe_run	<i>Safe execution wrapper</i>
----------	-------------------------------

Description

Executes an R expression safely, returning NULL and printing an error message if an error occurs.

Usage

```
safe_run(expr)
```

Arguments

expr	An R expression to evaluate.
------	------------------------------

Value

The result of expr, or NULL if an error occurs.

score_calculation	<i>Score calculation for weighted Cox model</i>
-------------------	---

Description

Calculates the score and variance for the weighted Cox model.

Usage

```
score_calculation(ybar1, ybar0, dN1, dN0, wt_rg)
```

Arguments

ybar1	Numeric vector of event counts for group 1.
ybar0	Numeric vector of event counts for group 0.
dN1	Numeric vector of event increments for group 1.
dN0	Numeric vector of event increments for group 0.
wt_rg	Numeric vector of rho-gamma weights.

Value

List with score, variance, information, and weights.

validate_input	<i>Validate required columns in a data frame</i>
----------------	--

Description

Checks that all required columns are present in a data frame.

Usage

```
validate_input(df, required_cols)
```

Arguments

df	Data frame to check.
required_cols	Character vector of required column names.

Value

NULL if all columns present, otherwise error.

validate_scheme_params
Validate weighting scheme parameters

Description

Checks and validates the parameters for a given weighting scheme.

Usage

```
validate_scheme_params(scheme, scheme_params, S.pool)
```

Arguments

scheme Character string specifying the weighting scheme.
scheme_params List of parameters for the scheme.
S.pool Numeric vector of pooled survival probabilities.

Value

Logical indicating if parameters are valid, or stops with error.

wlr_cumulative *Weighted log-rank cumulative statistics*

Description

Calculates cumulative weighted log-rank statistics for survival data.

Usage

```
wlr_cumulative(  
  df_weights,  
  scheme,  
  scheme_params = list(rho = 0, gamma = 0),  
  return_cumulative = FALSE  
)
```

Arguments

df_weights Data frame with weights and risk/event counts.
scheme Weighting scheme.
scheme_params List of scheme parameters.
return_cumulative Logical; whether to return cumulative statistics.

Value

List with score and variance.

wlr_dhat_estimates	<i>Weighted Log-Rank and Difference Estimate at a Specified Time</i>
--------------------	--

Description

Computes the weighted log-rank statistic, its variance, the difference in survival at a specified time (`tzero`), the variance of the difference, their covariance, and correlation, using flexible time-dependent weights. The weighting scheme is selected via the `scheme` argument and is calculated using `wt.rg.S`.

Usage

```
wlr_dhat_estimates(
  dfcounting,
  scheme = "fh",
  scheme_params = list(rho = 0, gamma = 0),
  tzero = NULL
)
```

Arguments

<code>dfcounting</code>	List output from <code>df_counting</code> containing risk sets, event counts, and survival estimates.
<code>scheme</code>	Character string specifying weighting scheme. One of: "fh" (Fleming-Harrington), "schemper", "XO", "MB", "custom_time", or "custom_code".
<code>scheme_params</code>	Named list with numeric weighting parameters <code>rho</code> and <code>gamma</code> (used for "fh" and "custom_code" schemes).
<code>tzero</code>	Time point at which to evaluate the difference in survival (default: 24).

Details

The weighting scheme is selected via the `scheme` argument and calculated using `wt.rg.S`. Supports standard Fleming-Harrington, Schemper, XO (Xu & O'Quigley), MB (Maggir-Burman), custom time-based, and custom code weights.

Value

A list with elements:

<code>lr</code>	Weighted log-rank test statistic.
<code>sig2_lr</code>	Variance of the log-rank statistic.
<code>dhat</code>	Difference in survival at <code>tzero</code> .
<code>cov_wlr_dhat</code>	Covariance between log-rank and difference at <code>tzero</code> .

sig2_dhat Variance of the difference at tzero.
 cor_wlr_dhat Correlation between log-rank and difference at tzero.

wt.rg.S

*Compute Time-Dependent Weights for Survival Analysis***Description**

Calculates time-dependent weights for survival analysis according to various schemes (Fleming-Harrington, Schemper, XO, MB, custom).

Usage

```
wt.rg.S(  
  S,  
  scheme = c("fh", "schemper", "XO", "MB", "custom_time", "fh_exp1", "fh_exp2"),  
  rho = NULL,  
  gamma = NULL,  
  Scensor = NULL,  
  Ybar = NULL,  
  tpoints = NULL,  
  t.tau = NULL,  
  w0.tau = 0,  
  w1.tau = 1,  
  mb_tstar = NULL,  
  details = FALSE  
)
```

Arguments

S Numeric vector of survival probabilities at each time point.
 scheme Character; weighting scheme. One of: 'fh', 'schemper', 'XO', 'MB', 'custom_time', 'fh_exp1', 'fh_exp2'.
 rho Numeric; rho parameter for Fleming-Harrington weights. Required if scheme='fh'.
 gamma Numeric; gamma parameter for Fleming-Harrington weights. Required if scheme='fh'.
 Scensor Numeric vector; censoring KM curve for Schemper weights. Must have same length as S.
 Ybar Numeric vector; risk set sizes for XO weights. Must have same length as S.
 tpoints Numeric vector; time points corresponding to S. Required for MB and custom_time schemes.
 t.tau Numeric; cutoff time for custom_time weights.
 w0.tau Numeric; weight before t.tau for custom_time weights.
 w1.tau Numeric; weight after t.tau for custom_time weights.
 mb_tstar Numeric; cutoff time for Magirr-Burman weights.
 details Logical; if TRUE, returns list with weights and diagnostic info. Default: FALSE.

Details

This function implements several weighting schemes for weighted log-rank tests: See vignette for details This function implements several weighting schemes for weighted log-rank tests:

Fleming-Harrington (fh) $w(t) = S(t)^{\rho} * (1-S(t))^{\gamma}$. See vignette for common choices.

Schemper $w(t) = S(t)/G(t)$ where G is the censoring distribution.

Xu-O'Quigley (XO) $w(t) = S(t)/Y(t)$ where Y is risk set size.

Magirr-Burman (MB) $w(t) = 1/\max(S(t), S(t^*))$.

Custom time Step function with weight w_0 before t^* and w_1 after t^* .

Value

If `details=FALSE` (default): Numeric vector of weights with length equal to S.

If `details=TRUE`: List containing:

weights Numeric vector of calculated weights

S Input survival probabilities

S_left Left-continuous survival probabilities

scheme Scheme name

Note

All weights are calculated using left-continuous survival probabilities $S(t-)$ to ensure consistency with counting process notation.

References

Fleming, T. R. and Harrington, D. P. (1991). Counting Processes and Survival Analysis. Wiley.

Magirr, D. and Burman, C. F. (2019). Modestly weighted logrank tests. *Statistics in Medicine*, 38(20), 3782-3790.

Schemper, M., Wakounig, S., and Heinze, G. (2009). The estimation of average hazard ratios by weighted Cox regression. *Statistics in Medicine*, 28(19), 2473-2489.

See Also

Other survival_analysis: [KM_diff\(\)](#), [cox_rhogamma\(\)](#), [df_counting\(\)](#)

Other weighted_tests: [cox_rhogamma\(\)](#), [df_counting\(\)](#)

Examples

```
# Generate example survival curve
time <- seq(0, 24, by = 0.5)
surv <- exp(-0.05 * time) # Exponential survival

# Fleming-Harrington (0,1) weights
w_fh01 <- wt.rg.S(surv, scheme = "fh", rho = 0, gamma = 1, tpoints = time)
```

```
# Magirr-Burman weights
w_mb <- wt.rg.S(surv, scheme = "MB", mb_tstar = 12, tpoints = time)

# Standard log-rank (equal weights)
w_lr <- wt.rg.S(surv, scheme = "fh", rho = 0, gamma = 0)

# Plotting examples
plot(time, w_fh01, type = "l", main = "FH(0,1) Weights")
plot(time, w_mb, type = "l", main = "MB(12) Weights")

# Compare multiple schemes
plot(time, w_lr, type = "l", ylim = c(0, 2))
lines(time, w_fh01, col = "blue")
legend("topleft", c("Log-rank", "FH(0,1)"), col = 1:2, lty = 1)
```

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